

1 Difference between Cohen's d and Hedges' g for effect size metrics

To my understanding, Hedges's g is a somewhat more accurate version of Cohen's d (with pooled SD) in that we add a correction factor for small sample. Both measures generally agree when the homoscedasticity assumption is not violated, but we may find situations where this is not the case, see e.g. McGrath & Meyer, *Psychological Methods* 2006, **11**(4): 386-401 ([pdf](#)). Other papers are listed at the end of my reply.

I generally found that in almost every psychological or biomedical studies, this is the Cohen's d that is reported; this probably stands from the well-known rule of thumb for interpreting its magnitude (Cohen, 1988). I don't know about any recent paper considering Hedges's g (or Cliff delta as a non-parametric alternative). Bruce Thompson has a [revised version](#) of the APA section on effect size.

Googling about Monte Carlo studies around effect size measures, I found this paper which might be interesting (I only read the abstract and the simulation setup): [Robust Confidence Intervals for Effect Sizes: A Comparative Study of Cohen's d and Cliff's Delta Under Non-normality and Heterogeneous Variances](#) ([pdf](#)).

About your 2nd comment, the [MBESS](#) R package includes various utilities for ES calculation (e.g., [smd](#) and related functions).

Other references

1. Zakzanis, K.K. (2001). Statistics to tell the truth, the whole truth, and nothing but the truth: Formulae, illustrative numerical examples, and heuristic interpretation of effect size analyses for neuropsychological researchers. *Archives of Clinical Neuropsychology*, 16(7), 653-667. ([pdf](#))
2. Durlak, J.A. (2009). How to Select, Calculate, and Interpret Effect Sizes. *Journal of Pediatric Psychology* ([pdf](#))

2 Under what conditions should Likert scales be used as ordinal or interval data?

Maybe too late but I add my answer anyway...

It depends on what you intend to do with your data: If you are interested in showing that scores differ when considering different group of participants (gender, country, etc.), you may treat your scores as numeric values, provided they fulfill usual assumptions about variance (or shape) and sample size. If you are rather interested in highlighting how response patterns vary across subgroups, then you should consider item scores as discrete choice among a set of answer options and look for log-linear modeling, ordinal logistic regression, item-response models or any other statistical model that allows to cope with polytomous items.

As a rule of thumb, one generally considers that having 12 distinct points on a scale is sufficient to approximate an interval scale (for interpretation purpose). Lickert items may be regarded as true ordinal scale, but they are often used as numeric and we can compute their mean or SD. This is often done in attitude surveys, although it is wise to report both mean/SD and % of response in, e.g. the two highest categories.

When using summated scale scores (i.e., we add up score on each item to compute a "total score"), usual statistics may be applied, but you have to keep in mind that you are now working with a latent variable so the underlying construct should make sense! In psychometrics, we generally check that (1) unidimensionality of the scale holds, (2) scale reliability is sufficient. When comparing two such scale scores (for two different instruments), we might even consider using attenuated correlation measures instead of classical Pearson correlation coefficient.

Classical textbooks include:

1. Nunnally, J.C. and Bernstein, I.H. (1994). *Psychometric Theory* (3rd ed.). McGraw-Hill Series in Psychology.
2. Streiner, D.L. and Norman, G.R. (2008). *Health Measurement Scales. A practical guide to their development and use* (4th ed.). Oxford.
3. Rao, C.R. and Sinharay, S., Eds. (2007). *Handbook of Statistics, Vol. 26: Psychometrics*. Elsevier Science B.V.
4. Dunn, G. (2000). *Statistics in Psychiatry*. Hodder Arnold.

You may also have a look at [Applications of latent trait and latent class models in the social sciences](#), from Rost & Langeheine, and W. Revelle's website on [personality research](#).

When validating a psychometric scale, it is important to look at so-called ceiling/floor effects (large

asymmetry resulting from participants scoring at the lowest/highest response category), which may seriously impact on any statistics computed when treating them as numeric variable (e.g., country aggregation, t-test). This raises specific issues in cross-cultural studies since it is known that overall response distribution in attitude or health surveys differ from one country to the other (e.g. chinese people vs. those coming from western countries tend to highlight specific response pattern, the former having generally more extreme scores at the item level, see e.g. Song, X.-Y. (2007) Analysis of multisample structural equation models with applications to Quality of Life data, in *Handbook of Latent Variable and Related Models*, Lee, S.-Y. (Ed.), pp 279-302, North-Holland).

More generally, you should look at the psychometric-related literature which makes extensive use of Likert items if you are interested with measurement issue. Various statistical models have been developed and are currently headed under the Item Response Theory framework.

3 Group differences on a five point Likert item

Clason & Dormody discussed the issue of statistical testing for Likert items ([Analyzing data measured by individual Likert-type items](#)). I think that a bootstrapped test is ok when the two distributions look similar (bell shaped and equal variance). However, a test for categorical data (e.g. trend or Fisher test, or ordinal logistic regression) would be interesting too since it allows to check for response distribution across the item categories, see Agresti's book on *Categorical Data Analysis* (Chapter 7 on *Logit models for multinomial responses*).

Aside from this, you can imagine situations where the t-test or any other non-parametric tests would fail if the response distribution is strongly imbalanced between the two groups. For example, if all people from group A answer 1 or 5 (in equally proportion) whereas all people in group B answer 3, then you end up with identical within-group mean and the test is not meaningful at all, though in this case the homoscedasticity assumption is largely violated.

4 What is a statistical journal with quick turnaround?

Maybe [Statistics Surveys](#) (but I think they are seeking review more than short note), [Statistica Sinica](#), or the [Electronic Journal of Statistics](#). They are not as quoted as SPL, but I hope this may help.

5 Quantitative methods and statistics conferences in psychology?

The [European Association of Methodology](#) has a meeting turning around statistics and psychometrics for applied research in social, educational and psychological science every two years. The latest was held in [Postdam](#) two months ago.

6 A survey of data-mining software tools.

Have a look at

- [Weka](#) (java)
- [Orange](#)
- the open-source [R](#) statistical software (Check the [Machine Learning](#) taskview)

and the [UCI Machine Learning Repository](#).

7 Interpretation of biplots in principal components analysis in R

PCA is one of the many ways to analyse the structure of a given correlation matrix. By construction, the first principal axis is the one which maximizes the variance (reflected by its eigenvalue) when data are projected onto a line (which stands for a direction in the p-space, assuming you have p variables) and the second one is orthogonal to it, and still maximizes the remaining variance. This is the reason why using the first two axes should yield the better approximation of the original variables space (say, a matrix X of dim n x p) when it is projected onto a plane.

Principal components are just linear combinations of the original variables. Therefore, plotting indi-

vidual factor scores (defined as Xu , where u is the vector of loadings of any principal component) may help to highlight groups of homogeneous individuals, for example, or to interpret one's overall scoring when considering all variables at the same time. In other words, this is a way to summarize one's location with respect to his value on the p variables, or a combination thereof. In your case, Fig. 13.3 in HSAUR shows that Joyner-Kersee (Jy-K) has a high (negative) score on the 1st axis, suggesting he performed overall quite good on all events. The same line of reasoning applies for interpreting the second axis. I take a very short look at the figure so I will not go into details and my interpretation is certainly superficial. I assume that you will find further information in the HSAUR textbook. Here it is worth noting that both variables and individuals are shown on the same diagram (this is called a *biplot*), which helps to interpret the factorial axes while looking at individuals' location. Usually, we plot the variables into a so-called correlation circle (where the angle formed by any two variables, represented here as vectors, reflects their actual pairwise correlation, since $r(x_1, x_2) = \cos(\angle(x_1, x_2))$).

I think, however, you'd better start reading some introductory book on multivariate analysis to get deep insight into PCA-based methods. For example, B.S. Everitt wrote an excellent textbook on this topic, *An R and S-Plus® Companion to Multivariate Analysis*, and you can check the [companion website](#) for illustration. There are other great R packages for applied multivariate data analysis, like [ade4](#) and [FactoMineR](#).

8 What is the best way to identify outliers in multivariate data?

Have a look at the [mvoutlier](#) package which relies on ordered robust mahalanobis distances, as suggested by @drknexus.

9 FA: Choosing Rotation matrix, based on "Simple Structure Criteria"

The R [psych](#) package includes various routines to apply Factor Analysis (whether it be PCA-, ML- or FA-based), but see my short review on [crantastic](#). Most of the usual rotation techniques are available, as well as algorithm relying on simple structure criteria; you might want to have a look at W. Rev-

elle's paper on this topic, [Very Simple Structure: An Alternative Procedure For Estimating The Optimal Number Of Interpretable Factors](#) (MBR 1979 (14)) and the [VSS\(\)](#) function.

Many authors are using orthogonal rotation (VARIMAX), considering loadings higher than, say 0.3 or 0.4 (which amounts to 9 or 16% of variance explained by the factor), as it provides simpler structures for interpretation and scoring purpose (e.g., in quality of life research); others (e.g. Cattell, 1978; Kline, 1979) would recommend oblique rotations since "in the real world, it is not unreasonable to think that factors, as important determiners of behavior, would be correlated" (I'm quoting Kline, *Intelligence. The Psychometric View*, 1991, p. 19).

To my knowledge, researchers generally start with FA (or PCA), using a scree-plot together with simulated data (parallel analysis) to help choosing the right number of factors. I often found that item cluster analysis and VSS nicely complement such an approach. When one is interested in second-order factors, or to carry on with SEM-based methods, then obviously you need to use oblique rotation and factor out the resulting correlation matrix.

Other packages/software:

- [lavaan](#), for latent variable analysis in R;
- [OpenMx](#) based on [Mx](#), a general purpose software including a matrix algebra interpreter and numerical optimizer for structural equation modeling.

References

1. Cattell, R.B. (1978). The scientific use of factor analysis in behavioural and life sciences. New York, Plenum.
2. Kline, P. (1979). Psychometrics and Psychology. London, Academic Press.

10 Machine Learning conferences?

[Artificial Intelligence In Medicine](#) (AIME), odd years starting from 1985.

11 How to get Sphericity in R for a nested within subject design?

Did you try the [car](#) package, from John Fox? It

includes the function `Anova()` which is very useful when working with experimental designs. It should give you corrected p-value following Greenhouse-Geisser correction and Huynh-Feldt correction. I can post a quick R example if you wonder how to use it.

Also, there is a nice tutorial on the use of R with repeated measurements and mixed-effects model for [psychology experiments and questionnaires](#); see Section 6.10 about sphericity.

As a sidenote, the Mauchly's Test of Sphericity is available in `mauchly.test()`, but it doesn't work with `aov` object if I remembered correctly. The [R Newsletter](#) from October 2007 includes a brief description of this topic.

12 Checking assumptions for random effects in nested mixed-effects models in R / S-Plus

It seems you are using the `nlme` package. Maybe it would be worth trying R and the `lme4` instead, although it is not fully comparable wrt. syntax or function call.

In your case, I would suggest to specify the `level` when you called `ranef()`, see `?ranef.lme`:

This is also present in the official [documentation](#) for NLME 3.0 (e.g., p. 17).

Check out Douglas Bates's neat handouts on GLMM. He is also writing a textbook entitled *lme4: Mixed-effects modeling with R*. All are available on [R-forge](#).

13 Flowcharts to help selecting the proper analysis technique and test?

These are not really interactive flowcharts, but maybe this could be useful: (1) <http://j.mp/cmakYq>, (2) <http://j.mp/aaxUusz>, and (3) <http://j.mp/bDMyAR>.

14 For a classification problem if class variable has unequal distribution which technique we should use?

Your class sample sizes do not seem so unbalanced since you have 30% of observations in your minority class. Logistic regression should be well performing in your case. Depending on the number of predictors that enter your model, you may consider some kind of penalization for parameters estimation, like ridge (L2) or lasso (L1). For an overview of problems with very unbalanced class, see Cramer (1999), *The Statistician*, 48: 85-94 ([PDF](#)).

I am not familiar with credit scoring techniques, but I found some papers that suggest that you could use SVM with weighted classes, e.g. [Support Vector Machines for Credit Scoring: Extension to Non Standard Cases](#). As an alternative, you can look at [boosting](#) methods with CART, or Random Forests (in the latter case, it is possible to adapt the sampling strategy so that each class is represented when constructing the classification trees). The paper by Novak and LaDue discuss the pros and cons of [GLM vs Recursive partitioning](#). I also found this article, [Scorecard construction with unbalanced class sizes](#) by Hand and Vinciotti.

15 R package for fixed-effect logistic regression

Conditional logistic regression (I assume that this is what you referred to when talking about Chamberlain's estimator) is available through `clogit()` in the [survival](#) package. I also found this page which contains R code to estimate [conditional logit parameters](#). The [survey](#) package also includes a lot of wrapper function for GLM and Survival model in the case of complex sampling, but I didn't look at.

Try also to look at `logit.mixed` in the [Zelig](#) package, or directly use the `lme4` package which provide methods for mixed-effects models with binomial link (see `lmer` or `glmer`).

Did you take a look at [Econometrics in R](#), from Grant V. Farnsworth? It seems to provide a gentle overview of applied econometrics in R (with which I am not familiar).

16 How can I estimate coefficient standard errors when using ridge regression?

I think bootstrap would be the best option to obtain robust SEs. This was done in some applied work using shrinkage methods, e.g. [Analysis of North American Rheumatoid Arthritis Consortium data using a penalized logistic regression approach](#) (BMC Proceedings 2009). There is also a nice paper from Casella on SE computation with penalized model, [Penalized Regression, Standard Errors, and Bayesian Lasso](#) (Bayesian Analysis 2010 5(2)). But they are more concerned with *lasso* and *elasticnet* penalization.

I always thought of ridge regression as a way to get better predictions than standard OLS, where the model is generally not parsimonious. For variable selection, the *lasso* or *elasticnet* criteria are more appropriate, but then it is difficult to apply a bootstrap procedure (since selected variables would change from one sample to the other, and even in the inner k -fold loop used to optimize the ℓ_1/ℓ_2 parameters); this is not the case with ridge regression, since you always consider all variables.

I have no idea about R packages that would give this information. It doesn't seem to be available in the [glmnet](#) package (see Friedman's paper in JSS, [Regularization Paths for Generalized Linear Models via Coordinate Descent](#)). However, Jelle Goeman who authored the [penalized](#) package discuss this point too. Cannot find the original PDF on the web, so I simply quote his words:

It is a very natural question to ask for standard errors of regression coefficients or other estimated quantities. In principle such standard errors can easily be calculated, e.g. using the bootstrap.

Still, this package deliberately does not provide them. The reason for this is that standard errors are not very meaningful for strongly biased estimates such as arise from penalized estimation methods. Penalized estimation is a procedure that reduces the variance of estimators by introducing substantial bias. The bias of each estimator is therefore a major component of its mean squared error, whereas its variance may contribute only a small part.

Unfortunately, in most applications of penal-

ized regression it is impossible to obtain a sufficiently precise estimate of the bias. Any bootstrap-based calculations can only give an assessment of the variance of the estimates. Reliable estimates of the bias are only available if reliable unbiased estimates are available, which is typically not the case in situations in which penalized estimates are used.

Reporting a standard error of a penalized estimate therefore tells only part of the story. It can give a mistaken impression of great precision, completely ignoring the inaccuracy caused by the bias. It is certainly a mistake to make confidence statements that are only based on an assessment of the variance of the estimates, such as bootstrap-based confidence intervals do.

17 How to plot ROC curves in multiclass classification?

It seems you are looking for multi-class ROC analysis, which is a kind of multi-objective optimization covered in a [tutorial](#) at ICML'04. As in several multi-class problem, the idea is generally to carry out pairwise comparison (one class vs. all other classes, one class vs. another class, see (1) or the *Elements of Statistical Learning*), and there is a recent paper by Landgrebe and Duin on that topic, [Approximating the multiclass ROC by pairwise analysis](#), Pattern Recognition Letters 2007 28: 1747-1758. Now, for visualization purpose, I've seen some papers some time ago, most of them turning around [volume under the ROC surface](#) (VUS) or [Cobweb diagram](#).

I don't know, however, if there exists an R implementation of these methods, although I think the [stars\(\)](#) function might be used for cobweb plot. I just ran across a Matlab toolbox that seems to offer multi-class ROC analysis, [PRSD Studio](#).

Other papers that may also be useful as a first start for visualization/computation:

- [Visualisation of multi-class ROC surfaces](#)
- [A simplified extension of the Area under the ROC to the multiclass domain](#)

References:

1. Allwein, E.L., Schapire, R.E. and Singer, Y.

(2000). Reducing multiclass to binary: A unifying approach for margin classifiers. *Journal of Machine Learning Research*, 1:113–141.

18 What do ROC curves tell you that traditional inference wouldn't?

In case you're interested in further references, an extensive list of papers is available on K.H. Zou's website, [Receiver Operating Characteristic \(ROC\) Literature Research](#).

ROC curves are also used when one is interested in comparing different classifiers performance, with wide applications in biomedical research and bioinformatics.

19 Recommended books on experiment design?

Not really a book but a gentle introduction on DoE in R: [An R companion to Experimental Design](#).

20 PCA on Correlation or Covariance?

A late reply, but you may find VERY useful handouts on multivariate data analysis "à la française" on the [Bioinformatics department](#) of Lyon. These come from the authors of the R [ade4](#) package. It is in french, though.

21 Applying the "kernel trick" to linear methods?

Two further references from [B. Schölkopf](#):

- Schölkopf, B. and Smola, A.J. (2002). [Learning with kernels](#). The MIT Press.
- Schölkopf, B., Tsuda, K., and Vert, J.-P. (2004). [Kernel methods in computational biology](#). The MIT Press.

and a website dedicated to [kernel machines](#).

22 Book recommendations for multivariate analysis

Off the top of my head, I would say that the following general purpose books are rather interesting as a first start:

- Izenman, J. [Modern Multivariate Statistical Techniques: Regression, Classification, and Manifold Learning](#). Springer. [companion website](#)
- Tinsley, H. and Brown, S. (2000). [Handbook of Applied Multivariate Statistics and Mathematical Modeling](#). Academic Press.

There is also many applied textbook, like

- Everitt, B.S. (2005). [An R and S-Plus Companion to Multivariate Analysis](#). Springer. [companion website](#)

It is difficult to suggest you specific books as there are many ones that are domain-specific (e.g. social sciences, machine learning, categorical data, biomedical data).

23 Variable selection procedure for binary classification

I have a slight preference for [Random Forests](#) by Leo Breiman & Adele Cutler for several reasons:

- it allows to cope with categorical and continuous predictors, as well as unbalanced class sample size;
- as an ensemble/embedded method, cross-validation is embedded and allows to estimate a generalization error;
- it is relatively insensible to its tuning parameters (% of variables selected for growing a tree, # of trees built);
- it provides an original measure of variable importance and is able to uncover complex interactions between variables (although this may lead to hard to read results).

Some authors argued that it performed as well as penalized SVM or Gradient Boosting Machines (see, e.g. Cutler et al., 2009, for the latter point).

A complete coverage of its applications or advantages may be off the topic, so I suggest the [Elements of Statistical Learning](#) from Hastie et al. (chap. 15) and Sayes et al. (2007) for further readings.

Last but not least, it has a nice implementation in R, with the [randomForest](#) package. Other R packages also extend or use it, e.g. [party](#) and [caret](#).

References:

Cutler, A., Cutler, D.R., and Stevens, J.R. (2009). Tree-Based Methods, in *High-Dimensional Data Analysis in Cancer Research*, Li, X. and Xu, R. (eds.), pp. 83-101, Springer.

Saeys, Y., Inza, I., and Larrañaga, P. (2007). A review of feature selection techniques in bioinformatics. *Bioinformatics*, **23**(19): 2507-2517.

24 Working through a clustering problem

You can try *Latent Semantic Analysis*, which basically provides a way to represent in a reduced space your news feeds and any term (in your case, keyword appearing in the title). As it relies on Singular Value Decomposition, I suppose you may then be able to check if there exists a particular association between those two attributes. I know this is used to find documents matching a specific set of criteria, as in information retrieval, or to construct a tree reflecting terms similarity (like a dictionary) based on a large corpus (which here plays the role of the concept space).

See for a gentle introduction [An Introduction to Latent Semantic Analysis](#), by Landauer et al.

Moreover, there is an R package that implements this technique, namely [lsa](#).

25 How to calculate the “exact confidence interval” for relative risk?

Check out the R [Epi](#) and [epitools](#) packages, which include many functions for computing exact and approximate CIs/p-values for various measures of association found in epidemiological studies, including relative risk (RR). I know there is also [PropCIs](#), but I never tried it. Bootstrapping is also an option, but

generally these are exact or approximated CIs that are provided in epidemiological papers, although most of the explanatory studies rely on GLM, and thus make use of odds-ratio (OR) instead of RR (although, wrongly it is often the RR that is interpreted because it is easier to understand, but this is another story).

You can also check your results with online calculator, like on [statpages.org](#), or [Relative Risk and Risk Difference Confidence Intervals](#). The latter explains how computations are done.

By “exact” tests, we generally mean tests/CIs not relying on an asymptotic distribution, like the chi-square or standard normal; e.g. in the case of an RR, an 95% CI may be approximated as $\exp \left[\log(\text{rr}) - 1.96 \sqrt{\text{Var}(\log(\text{rr}))} \right], \exp \left[\log(\text{rr}) + 1.96 \sqrt{\text{Var}(\log(\text{rr}))} \right]$, where $\text{Var}(\log(\text{rr})) = 1/a - 1/(a+b) + 1/c - 1/(c+d)$ (assuming a 2-way cross-classification table, with a , b , c , and d denoting cell frequencies). The explanations given by @Keith are, however, very insightful.

For more details on the calculation of CIs in epidemiology, I would suggest to look at Rothman and Greenland’s textbook, [Modern Epidemiology](#) (now in its 3rd edition), [Statistical Methods for Rates and Proportions](#), from Fleiss et al., or [Statistical analyses of the relative risk](#), from J.J. Gart (1979).

You will generally get similar results with [fisher.test\(\)](#), as pointed by @gd047, although in this case this function will provide you with a 95% CI for the odds-ratio (which in the case of a disease with low prevalence will be very close to the RR).

Notes:

1. I didn’t check your Excel file, for the reason advocated by @csgillespie.
2. Michael E Dewey provides an interesting summary of [confidence intervals for risk ratios](#), from a digest of posts on the R mailing-list.

26 What is the proper way to analyze discrete data?

What you are looking for seems to be a test for comparing two groups where observations are kind of ordinal data. In this case, I would suggest to apply a trend test to see if there are any differences between the CTL and TRT group.

Using a t-test would not acknowledge the fact your data are discrete, and the Gaussian assumption may be seriously violated if scores distribution isn't symmetric as is often the case with Likert scores (such as the ones you seem to report). Don't know if these data come from a case-control study or not, but you might also apply rank-based method as suggested by @propfol: If it is not a matched design, the Wilcoxon-Mann-Whitney test (`wilcox.test()` in R) is fine, and ask for an exact p-value although you may encounter problem with tied observations. The efficiency of the WMW test is $3/\pi$ with respect to the t-test if normality holds but it may even be better otherwise, I seem to remember.

Given your sample size, you may also consider applying a permutation test (see the `perm` or `coin` R packages).

Check also those related questions:

- [Group differences on a five point Likert item](#)
- [Under what conditions should Likert scales be used as ordinal or interval data?](#)

27 Adding labels to points using mds and scatter3d package with R

Basically, what you need is to store your `scatterplot3d` in a variable and reuse it like this:

```
x <- replicate(10, rnorm(100))
x.mds <- cmdscale(dist(x), eig=TRUE, k=3)
s3d <- scatterplot3d(x.mds$points[,1:3])
text(s3d$xyz.convert(0,0,0), labels="Origin")
```

Replace the coordinates and text by whatever you want to draw. You can also use a color vector to highlight the groups of interest.

The `R.basic` package, from Henrik Bengtsson, seems to provide additional facilities to customize 3D plots, but I never tried it.

28 When is it acceptable to collapse across groups when performing a factor analysis?

There seems to be two cases to consider, depending

on whether your scale was already validated using standard psychometric methods (from classical test or item response theory). In what follows, I will consider the first case where I assume preliminary studies have demonstrated construct validity and scores reliability for your scale.

In this case, there is no formal need to apply exploratory factor analysis, unless you want to examine the pattern matrix within each group (but I generally do it, just to ensure that there are no items that unexpectedly highlight low factor loading or cross-load onto different factors); in order to be able to pool all your data, you need to use a multi-group factor analysis (hence, a confirmatory approach as you suggest), which basically amount to add extra parameters for testing a group effect on factor loading (1st order model) or factor correlation (2nd order model, if this makes sense) which would impact measurement invariance across subgroups of respondents. This can be done using `Mplus` (see the discussion about CFA [there](#)) or `Mx` (e.g. [Conor et al.](#), 2009), not sure about `Amos` as it seems to be restricted to simple factor structure. The `Mx` software has been redesigned to work within the R environment, `OpenMx`. The wiki is well responding so you can ask questions if you encounter difficulties with it. There is also a more recent package, `lavaan`, which appears to be a promising package for SEMs.

Alternatives models coming from IRT may also be considered, including a Latent Regression Rasch Model (for each scale separately, see De Boeck and Wilson, 2004), or a Multivariate Mixture Rasch Model (von Davier and Carstensen, 2007). You can take a look at [Volume 20](#) of the [Journal of Statistical Software](#), entirely devoted to psychometrics in R, for further information about IRT modeling with R. You may be able to reach similar tests using Structural Equation Modeling, though.

If factor structure proves to be equivalent across the two groups, then you can aggregate the scores (on your four summated scales) and report your statistics as usual. However, it is always a challenging task to use CFA since not rejecting H_0 does by no mean allow you to check that your postulated theoretical model is correct in the true world, but just that there is no reason to reject it on statistical grounds; on the other hand, rejecting the null would lead to accept the alternative, which is generally left unspecified, unless you apply sequential testing of nested models. Anyway, this is the way we go in cross-cultural settings, especially when we want to assess whether a given questionnaire (e.g.,

on Patients Reported Outcomes) measures what it purports to do whatever the population it is administered to.

Now, regarding the apparent differences between the two groups – one is drawn from a population of students, the other is a clinical sample, assessed at a later date – it depends very much on your own considerations: Does mixing of these two samples makes sense from the literature surrounding the questionnaire used (esp., it should have shown temporal stability and applicability in a wide population), do you plan to generalize your findings over a larger population (obviously, you gain power by increasing sample size). At first sight, I would say that you need to ensure that both groups are comparable with respect to the characteristics thought to influence one's score on this questionnaire (e.g., gender, age, SES, biomedical history, etc.), and this can be done using classical statistics for two-groups comparison (on raw scores). It is worth noting that in clinical studies, we face the reverse situation: We usually want to show that scores differ between different clinical subgroups (or between treated and naive patients), which is often referred to as *known-group validity*.

Reference:

1. De Boeck, P. and Wilson, M. (2004). *Explanatory Item Response Models. A Generalized Linear and Nonlinear Approach*. Springer.
2. von Davier, M. and Carstensen, C.H. (2007). *Multivariate and Mixture Distribution Rasch Models*. Springer.

29 I just installed the latest version of R. What packages should I obtain?

You can also take a look at [Task views](#) on CRAN and see if something suit your needs. I agree with @Jeromy for these must-have packages (for data manipulation and plotting).

30 How to plot a violin scatter box-plot (in R) ?

Try the [vioplot](#) package:

```
library(vioplot)
vioplot(rnorm(100))
```

(with awful default color ;-)

There is also `wvioplot()` in the [wvioplot](#) package, for weighted violin plot, and [beanplot](#), which combines violin and rug plots. They are also available through the [lattice](#) package, see `?panel.violin`.

31 Area Under Curve (AUC) - given peak mean and standard deviation (SD)

Given how your plot looks like, I would suggest rather to fit a mixture of gaussians and get their respective densities. Look at the [mclust](#) package; basically this is referred to model-based clustering (you are seeking groups of points belonging to a given distribution, that is to be estimated, whose location parameter – but also shape – varies along a common dimension). A full explanation of MClust is available [here](#).

It seems the [delt](#) package offers an alternative way to fit 1D data with a mixture of gaussians, but I didn't get into details.

Anyway, I think this is the best way to get automatic estimates and avoid cutting your x-scale at arbitrary locations.

32 Recommended books or articles as introduction to Cluster Analysis?

It may be worth looking at M.W. Berry's books:

1. *Survey of Text Mining I: Clustering, Classification, and Retrieval* (2003)
2. *Survey of Text Mining II: Clustering, Classification, and Retrieval* (2008)

They consist of series of applied and review papers. The latest seems to be available as PDF at the following address: <http://bit.ly/deNeiy>.

Here are few links related to CA as applied to text

mining:

- [Document Topic Generation in Text Mining by Using Cluster Analysis with EROCK](#)
- [An Approach to Text Mining using Information Extraction](#)

You can also look at *Latent Semantic Analysis*, but see my response there: [Working through a clustering problem](#).

Gene selection and classification of microarray data using random forest. *BMC Bioinformatics*, **7**:3.

4. Diaz-Uriarte, R. (2007). GeneSrF and varSelRF: a web-based tool and R package for gene selection and classification using random forest. *BMC Bioinformatics*, **8**: 328

Since you are talking of SVM, you can look for *penalized SVM*.

33 Feature selection for “final” model when performing cross-validation in machine learning

This is a very good question that I faced myself when working with SNPs data... And I didn't find any obvious answer through the literature.

Whether you use LOO or K-fold CV, you'll end up with different features since the cross-validation iteration must be the most outer loop, as you said. You can think of some kind of voting scheme which would rate the n-vectors of features you got from your LOO-CV (can't remember the paper but it is worth checking the work of [Harald Binder](#) or [Antoine Cornuéjols](#)). In the absence of a new test sample, what is usually done is to re-apply the ML algorithm to the whole sample once you have found its optimal cross-validated parameters. But proceeding this way, you cannot ensure that there is no overfitting (since the sample was already used for model optimization).

Or, alternatively, you can use embedded methods which provide you with features ranking through a measure of variable importance, e.g. like in [Random Forests](#) (RF). As cross-validation is included in RFs, you don't have to worry about the $n \ll p$ case or curse of dimensionality. Here are nice papers of their applications in gene expression studies:

1. Cutler, A., Cutler, D.R., and Stevens, J.R. (2009). Tree-Based Methods, in *High-Dimensional Data Analysis in Cancer Research*, Li, X. and Xu, R. (eds.), pp. 83-101, Springer.
2. Saeys, Y., Inza, I., and Larrañaga, P. (2007). A review of feature selection techniques in bioinformatics. *Bioinformatics*, **23**(19): 2507-2517.
3. Díaz-Uriarte, R., Alvarez de Andrés, S. (2006).

34 What are some valuable Statistical Analysis open source projects?

There are also those projects initiated by the FSF or redistributed under GNU General Public License, like:

- [PSPP](#), which aims to be a free alternative to SPSS
- [GRET](#), mostly dedicated to regression and econometrics

There is even applications that were released just as a companion software for a textbook, like [JMulTi](#), but are still in use by few people.

I am still playing with [xliostat](#), from time to time, although Lisp has been largely superseded by R (see Jan de Leeuw's overview on [Lisp vs. R](#) in the *Journal of Statistical Software*). Interestingly, one of the cofounders of the R language, Ross Ihaka, argued on the contrary that the future of statistical software is... Lisp: [Back to the Future: Lisp as a Base for a Statistical Computing System](#). @Alex already pointed to the Clojure-based statistical environment [Incanter](#), so maybe we will see a revival of Lisp-based software in the near future? :-)

35 Explain the difference between multiple regression and multivariate regression, with minimal use of symbols/math.

Very quickly, I would say: 'multiple' applies to the number of predictors that enter the model (or equivalently the design matrix) with a single outcome (Y response), while 'multivariate' refers to a matrix of response vectors. Cannot remember the author

who starts its introductory section on multivariate modeling with that consideration, but I think it is Brian Everitt in his textbook [An R and S-Plus Companion to Multivariate Analysis](#). For a thorough discussion about this, I would suggest to look at his latest book, [Multivariable Modeling and Multivariate Analysis for the Behavioral Sciences](#).

For ‘variate’, I would say this is a common way to refer to any random variable that follows a known or hypothesized distribution, e.g. we speak of gaussian variates X_i as a series of observations drawn from a normal distribution (with parameters μ and σ^2). In probabilistic terms, we said that these are some random *realizations* of X , with mathematical expectation μ , and about 95% of them are expected to lie on the range $[\mu - 2\sigma; \mu + 2\sigma]$.

36 What books provide an overview of computational statistics as it applies to computer science?

Here is a very nice book from James E. Gentle, [Computational Statistics](#) (Springer, 2009), which covers both computational and statistical aspects of data analysis. Gentle also authored other great books, check his publications.

Another great book is the [Handbook of Computational Statistics](#), from Gentle et al. (Springer, 2004); it is circulating as PDF somewhere on the web, so just try looking at it on Google.

37 Is there a biglm equivalent for coxph?

Maybe take a look at the [DatABEL](#) package. I know it is used in genomic studies with large data that may be stored on the HD instead of RAM. From what I read in the help file, you can then apply different kind of model, including survival model.

38 Likert scales analysis

From what I’ve seen so far, FA is used for attitude items as it is for other kind of rating scales. The problem arising from the metric used (that is, “are Likert scales really to be treated as numeric scales?” is a long-standing debate, but providing you check for the bell-shaped response distribution

you may handle them as continuous measurements, otherwise check for non-linear FA models or *optimal scaling*) may be handled by polytomous IRT models, like the Graded Response, Rating Scale, or Partial Credit Model. The latter two may be used as a rough check of whether the threshold distances, as used in Likert-type items, are a characteristic of the response format (RSM) or of the particular item (PCM).

Regarding your second point, it is known, for example, that response distributions in attitude or health surveys differ from one country to the other (e.g. chinese people tend to highlight ‘extreme’ response patterns compared to those coming from western countries, see e.g. Song, X.-Y. (2007) Analysis of multisample structural equation models with applications to Quality of Life data, in *Handbook of Latent Variable and Related Models*, Lee, S.-Y. (Ed.), pp 279-302, North-Holland). Some methods to handle such situation off the top of my head:

- use of log-linear models (marginal approach) to highlight strong between-groups imbalance at the item level (coefficients are then interpreted as relative risks instead of odds);
- the multi-sample SEM method from Song cited above (Don’t know if they do further work on that approach, though).

Now, the point is that most of these approaches focus at the item level (ceiling/floor effect, decreased reliability, bad item fit statistics, etc.), but when one is interested in how people deviate from what would be expected from an ideal set of observers/respondents, I think we must focus on person fit indices instead.

Such χ^2 statistics are readily available for IRT models, like INFIT or OUTFIT mean square, but generally they apply on the whole questionnaire. Moreover, since estimation of items parameters rely in part on persons parameters (e.g., in the marginal likelihood framework, we assume a gaussian distribution), the presence of outlying individuals may lead to potentially biased estimates and poor model fit.

As proposed by Eid and Zickar (2007), combining a latent class model (to isolate group of respondents, e.g. those always answering on the extreme categories vs. the others) and an IRT model (to estimate item parameters and persons locations on the latent trait in both groups) appears a nice solution.

Other modeling strategies are described in their paper (e.g. HYBRID model, see also Holden and Book, 2009).

Likewise, **unfolding models** may be used to cope with *response style*, which is defined as a consistent and content-independent pattern of response category (e.g. tendency to agree with all statements). In the social sciences or psychological literature, this is known as Extreme Response Style (ERS). References (1–3) may be useful to get an idea on how it manifests and how it may be measured.

Here is a short list of papers that may help to progress on this subject:

1. Hamilton, D.L. (1968). **Personality attributes associated with extreme response style**. *Psychological Bulletin*, **69**(3): 192–203.
2. Greanleaf, E.A. (1992). Measuring extreme response style. *Public Opinion Quarterly*, **56**(3): 328–351.
3. de Jong, M.G., Steenkamp, J.-B.E.M., Fox, J.-P., and Baumgartner, H. (2008). Using Item Response Theory to Measure Extreme Response Style in Marketing Research: A Global Investigation. *Journal of marketing research*, **45**(1): 104–115.
4. Morren, M., Gelissen, J., and Vermunt, J.K. (2009). **Dealing with extreme response style in cross-cultural research: A restricted latent class factor analysis approach**
5. Moors, G. (2003). Diagnosing Response Style Behavior by Means of a Latent-Class Factor Approach. Socio-Demographic Correlates of Gender Role Attitudes and Perceptions of Ethnic Discrimination Reexamined. *Quality & Quantity*, **37**(3), 277–302.
6. de Jong, M.G. Steenkamp J.B., Fox, J.-P., and Baumgartner, H. (2008). Item Response Theory to Measure Extreme Response Style in Marketing Research: A Global Investigation. *Journal of Marketing Research*, **45**(1), 104–115.
7. Javaras, K.N. and Ripley, B.D. (2007). An “Unfolding” Latent Variable Model for Likert Attitude Data. *JASA*, **102**(478): 454–463.
8. slides from Moustaki, Knott and Mavridis, **Methods for detecting outliers in latent variable**

models

9. Eid, M. and Zickar, M.J. (2007). Detecting response styles and faking in personality and organizational assessments by Mixed Rasch Models. In von Davier, M. and Carstensen, C.H. (Eds.), *Multivariate and Mixture Distribution Rasch Models*, pp. 255–270, Springer.
10. Holden, R.R. and Book, A.S. (2009). Using hybrid Rasch-latent class modeling to improve the detection of fakers on a personality inventory. *Personality and Individual Differences*, **47**(3): 185–190.

39 Is it appropriate to treat n-point Likert scale data as n trials from a binomial process?

I don’t know of any articles related to your question in the psychometric literature. It seems to me that ordered logistic models allowing for random effect components can handle this situation pretty well.

I agree with @Srikant and think that a proportional odds model or an ordered probit model (depending on the link function you choose) might better reflect the intrinsic coding of Likert items, and their typical use as rating scales in opinion/attitude survey or questionnaire.

Other alternatives are: (1) use of adjacent instead of proportional or cumulative categories (where there is a connection with log-linear models); (2) use of item-response models like the partial-credit model or the rating scale model (as was mentioned in my response on **Likert scales analysis**). The latter case is comparable to a mixed-effects approach, with subjects treated as random effects, and is readily available in the SAS system (e.g., **Fitting mixed-effects models for repeated ordinal outcomes with the NLMIXED procedure**) or R (see vol. 20 of the *Journal of Statistical Software*). You might also be interested in the discussion provided by John Linacre about **Optimizing Rating Scale Category Effectiveness**.

The following papers may also be useful:

1. Wu, C-H (2007). **An Empirical Study on the Transformation of Likert-scale Data to Numerical Scores**. *Applied Mathematical Sciences*, **1**(58): 2851–2862.

2. Rost, J and Luo, G (1997). [An Application of a Rasch-Based Unfolding Model to a Questionnaire on Adolescent Centrism](#). In Rost, J and Langeheine, R (Eds.), *Applications of latent trait and latent class models in the social sciences*, New York: Waxmann.
3. Lubke, G and Muthen, B (2004). [Factor-analyzing Likert-scale data under the assumption of multivariate normality complicates a meaningful comparison of observed groups or latent classes](#). *Structural Equation Modeling*, **11**: 514-534.
4. Nering, ML and Ostini, R (2010). *Handbook of Polytomous Item Response Theory Models*. Routledge Academic
5. Bender R and Grouven U (1998). Using binary logistic regression models for ordinal data with non-proportional odds. *Journal of Clinical Epidemiology*, **51(10)**: 809-816. (Cannot find the pdf but this one is available, [Ordinal logistic regression in medical research](#))

test (relying on the hypergeometric distribution).

The `prop.test()` function in R allows to test whether proportions are comparable between groups or does not differ from theoretical probabilities. It is referred to as a z-test because the statistics looks like

$$z = (f_1 - f_2) / \sqrt{\hat{p} \left(\frac{1}{n_1} + \frac{1}{n_2} \right)}$$

where $\hat{p} = (p_1 + p_2) / (n_1 + n_2)$, and indices (1, 2) refers to the first and second line of your table. In a two-way contingency table where $H_0 : p_1 = p_2$, this should yield comparable results to the ordinary χ^2 test:

```
> tab <- matrix(c(100, 80, 20, 10), ncol = 2)
> chisq.test(tab)

Pearson's Chi-squared test with Yates' continuity correction

data:  tab
X-squared = 0.8823, df = 1, p-value = 0.3476

> prop.test(tab)

2-sample test for equality of proportions with continuity
correction

data:  tab
X-squared = 0.8823, df = 1, p-value = 0.3476
alternative hypothesis: two.sided
95 percent confidence interval:
 -0.15834617  0.04723506
sample estimates:
 prop 1    prop 2 
0.8333333 0.8888889
```

40 Boosted Decision Trees in python?

My first look would be at [Orange](#), which is a fully-featured app for ML, with a backend in Python. See e.g. [orngEnsemble](#).

Other promising projects are [mlpy](#) and the [scikit.learn](#).

I know that [PyCV](#) include several boosting procedures, but apparently not for CART. Take also a look at [MLboost](#)

41 What is the relationship between a chi square test and test of equal proportions?

Very short answer:

The chi-Square test (`chisq.test()` in R) compares the observed frequencies in each category of a contingency table with the expected frequencies (computed as the product of the marginal frequencies). It is used to determine whether the deviations between the observed and the expected counts are too large to be attributed to chance. Departure from independence is easily checked by inspecting residuals (try `?mosaicplot` or `?assocplot`, but also look at the `vcd` package). Use `fisher.test()` for an exact

For analysis of discrete data with R, I highly recommend [R \(and S-PLUS\) Manual to Accompany Agresti's Categorical Data Analysis \(2002\)](#), from Laura Thompson.

42 Post hoc tests in ANCOVA

Multiple testing following ANCOVA, or more generally any GLM, but the comparisons now focus on the adjusted group/treatment or marginal means (i.e. what the scores would be if groups did not differ on the covariate of interest). To my knowledge, Tukey HSD and Scheffé tests are used. Both are quite conservative and will tend to bound type I error rate. The latter is preferred in case of unequal sample size in each group. I seem to remember that

some people also use Sidak correction on specific contrasts (when it is of interest of course) as it is less conservative than the Bonferroni correction.

Such tests are readily available in the R `multcomp` package (see `?glht`). The accompanying vignette include example of use in the case of a simple linear model (section 2), but it can be extended to any other model form. Other examples can be found in the `HH` packages (see `?MMC`). Several MCP and resampling procedures (recommended for strong inferences, but it relies on a different approach to the correction for Type I error rate inflation) are also available in the `multtest` package, through `Bioconductor`, see refs (3–4). The definitive reference to multiple comparison is the book from the same authors: Dudoit, S. and van der Laan, M.J., *Multiple Testing Procedures with Applications to Genomics* (Springer, 2008).

Reference 2 explained the difference between MCP in the general case (ANOVA, working with unadjusted means) vs. ANCOVA. There are also several papers that I can't remember actually, but I will look at them.

Other useful references:

1. Westfall, P.H. (1997). Multiple Testing of General Contrasts Using Logical Constraints and Correlations. *JASA* **92**: 299-306.
2. Westfall, P.H. and Young, S.S. (1993) *Resampling Based Multiple Testing, Examples and Methods for p-Value Adjustment*. John Wiley and Sons: New York.
3. Pollard, K.S., Dudoit, S., and van der Laan, M.J. (2004). *Multiple Testing Procedures: R multtest Package and Applications to Genomics*.
4. Taylor, S.L. Lang, D.T., and Pollard, K.S. (2007). *Improvements to the multiple testing package multtest*. *R News* **7(3)**: 52-55.
5. Bretz, F., Genz, A., and Hothorn, L.A. (2001). On the numerical availability of multiple comparison procedures. *Biometrical Journal*, **43(5)**: 645–656.
6. Hothorn, T., Bretz, F., and Westfall, P. (2008). *Simultaneous Inference in General Parametric Models*. Department of Statistics: Technical Reports, Nr. 19.

The first two are referenced in SAS PROC related to MCP.

43 Under what conditions does correlation imply causation?

I'll just add some additional comments about causality as viewed from an *epidemiological perspective*. Most of these arguments are taken from *Practical Psychiatric Epidemiology*, by Prince et al. (2003).

Causation, or *causality interpretation*, are by far the most difficult aspects of epidemiological research. *Cohort* and *cross-sectional* studies might both lead to confounding effects for example. Quoting S. Menard (*Longitudinal Research*, Sage University Paper 76, 1991), H.B. Asher in *Causal Modeling* (Sage, 1976) initially proposed the following set of criteria to be fulfilled:

- The phenomena or variables in question must covary, as indicated for example by differences between experimental and control groups or by nonzero correlation between the two variables.
- The relationship must not be attributable to any other variable or set of variables, i.e., it must not be spurious, but must persist even when other variables are controlled, as indicated for example by successful randomization in an experimental design (no difference between experimental and control groups prior to treatment) or by a nonzero partial correlation between two variables with other variable held constant.
- The supposed cause must precede or be simultaneous with the supposed effect in time, as indicated by the change in the cause occurring no later than the associated change in the effect.

While the first two criteria can easily be checked using a cross-sectional or time-ordered cross-sectional study, the latter can only be assessed with longitudinal data, except for biological or genetic characteristics for which temporal order can be assume without longitudinal data. Of course, the situation becomes more complex in case of a non-recursive causal relationship.

I also like the following illustration (Chapter 13, in the aforementioned reference) which summarizes the approach promulgated by Hill (1965) which includes 9 different criteria related to causation effect,

as also cited by @James. The original article was indeed entitled “The environment and disease: association or causation?” (PDF version)

Finally, Chapter 2 of Rothman’s most famous book, *Modern Epidemiology* (1998, Lippincott Williams & Wilkins, 2nd Edition), offers a very complete discussion around causation and causal inference, both from a statistical and philosophical perspective.

I’d like to add the following references (roughly taken from an online course in epidemiology) are also very interesting:

- Swaen, G and van Amelsvoort, L (2009). *A weight of evidence approach to causal inference. Journal of Clinical Epidemiology*, 62, 270-277.
- Botti, C, Comba, P, Forastiere, F, and Settimi, L (1996). *Causal inference in environmental epidemiology: the role of implicit values. The Science of the Total Environment*, 184, 97-101.
- Weed, DL (2002). *Environmental epidemiology. Basics and proof of cause-effect. Toxicology*, 181, 399-403.
- Franco, EL, Correa, P, Santella, RM, Wu, X, Goodman, SN, and Petersen, GM (2004). *Role and limitations of epidemiology in establishing a causal association. Seminars in Cancer Biology*, 14, 413-426.

Finally, this review offers a larger perspective on causal modeling, *Causal inference in statistics: An overview* (J Pearl, SS 2009 (3)).

44 Confidence intervals on differences in choices in a GEE framework: methods and alternatives?

Well, the `gee` package has facilities for fitting GEE and `gee()` return asymptotic and robust SE. I never used the `geepack` package. From what I saw in the online example, output seems to resemble more or less that of `gee`. To compute 100(1- α) CIs for your main effects (e.g. gender), why not use the robust SE (in the following I will assume it is extracted from, say `summary(gee.fit)`, and stored in a variable `rob.se`)? I suppose that

```
exp(coef(gee.fit)[“gender”]+c(-1,1)*rob.se*qnrm(0.975))
```

should yield 95% CIs expressed on the odds scale.

Criterion	Description	Limitations
1. Strength	A strong effect size for an association	This reduces the chance of minor unmeasured confounding, but assumes that major confounding factors have been accounted for. Weak associations may also be causal.
2. Consistency	Repeated observations of an association in different populations/circumstances	This assumes that all necessary causal factors are evenly distributed between populations. If a factor-outcome association were present in all populations, would this imply non-causality?
3. Specificity	A risk factor leads to a single outcome	There is no reason why a risk factor should be associated with a single disorder (e.g. multiple disorders associated with alcohol misuse).
4. Temporality	The cause should precede the effect	A study should ideally demonstrate this. However, the fact that one event follows another does not rule out the opposite direction of causation or other circumstances. For example, depression may cause physical ill-health but the opposite may also occur.
5. Biological gradient	A 'dose-response' relationship	This assumes that the 'ceiling' of risk has been reached. A single life-event may be sufficient to cause depression with no influence of further events. Of little use for cross-sectional associations since a 'dose-response' pattern would be predicted with either direction of causation.
6. Plausibility	That the hypothesis is biologically plausible	Frequently a highly subjective judgement based on the volume of the biological literature. There are many historical examples of important findings rejected on the ground of implausibility at the time (e.g. Darwin's theory of the Origin of Species).
7. Coherence	That the interpretation does not conflict with the known biology of the disease	This depends heavily on the quality of the information. It also is not entirely consistent with the principle of hypothesis refutation.
8. Experimental evidence	Evidence from interventional research	Interventional research may not be ethical or feasible for many cause effect investigations. An intervention may not be discrete enough to establish causation.
9. Analogy	Similar associations in other fields	A highly subjective judgement.

Hi

Now, in fact I rarely use GEE except when I am working with binary endpoints in longitudinal studies, because it's easy to pass or estimate a given working correlation matrix. In the case you summarize here, I would rather rely on an IRT model for dichotomous items (see the [psychometrics](#) task view), or (it is quite the same in fact) a mixed-effects GLM such as the one that is proposed in the [lme4](#) package, from Doug Bates. For study like yours, as you said, subjects will be considered as random effects, and your other covariates enter the model as fixed effects; the response is the 0/1 rating on each item (which enter the model as well). Then you will get 95% CI for fixed effects, either from the SE computed as `sqrtdiag(vcov(glm.fit))` or as read in `summary(glm.fit)`, or using `confint()` together with an `lmList` object. Doug Bates gave nice illustrations in the following two paper/handout:

- [Estimating the Multilevel Rasch Model: With the lme4 Package](#) (JSS, 2007 20(2))
- [Item Response Models as GLMMs](#) (from a workshop at UseR 2008)

There is also a discussion about profiling `lmer` fits (based on *profile deviance*) to investigate variability in fixed effects, but I didn't investigate that point. I think it is still in section 1.5 of Doug's [draft on mixed models](#). There are a lot of discussion about computing SE and CI for GLMM as implemented in the `lme4` package (whose interface differs from the previous `nlme` package), so that you will easily find other interesting threads after googling about that.

It's not clear to me why GEE would have to be preferred in this particular case. Maybe, look at the R translation of Agresti's book by Laura Thompson, [R \(and S-PLUS\) Manual to Accompany Agresti's Categorical Data](#).

Update:

I just realized that the above solution would only work if you're interested in getting a confidence interval for the gender effect alone. If it is the interaction `item*gender` that is of concern, you have to model it explicitly in the GLMM (my second reference on Bates's has an example on how to do it with `lmer`).

Another solution is to use an explanatory IRT model, where you explicitly acknowledge the potential effect of person covariates, like gender or age, and consider fitting them within a Rasch model, for example. This is called a Latent Regression

Rasch Model, and is fully described in de Boeck and Wilson's book, *Explanatory item response models: a generalized linear and nonlinear approach* (Springer, 2004), which you can read online on [Google books](#) (section 2.4). There are some facilities to fit this kind of model in Stata (see [there](#)). In R, we can mimic such model with a mixed-effects approach; a toy example would look something like

```
lmer(response ~ 0 + Age + Sex + item + (Sex|id), data=df,
      binomial)
```

if I remember correctly. I'm not sure whether the `eRm` allows to easily incorporate person covariates (because we need to construct a specific design matrix), but it may be worth checking out since it provides 95% CIs too.

45 How to rank the results of questions with categorical answers?

Recoding your data with numerical values seems ok, provided the assumption of an ordinal scale holds. This is often the case for Likert-type item, but see these related questions:

- [Is it appropriate to treat n-point Likert scale data as n trials from a binomial process?](#)
- [Under what conditions should Likert scales be used as ordinal or interval data?](#)

When validating a questionnaire, we often provide usual numerical summaries (mean \pm sd, range, quartiles) to highlight ceiling/floor effect, that is higher response rate in the extreme range of the scale. Dotplots are also great tool to summarize such data.

This is just for visualization/summary purpose. If you want to get into more statistical stuff, you can use proportional odds model or ordinal logistic regression, for ordinal items, and multinomial regression, for discrete ones.

46 When to use (non)parametric test of homoscedasticity assumption?

It seems that the FK test is to be preferred in case of strong departure from the normality (to which the Bartlett test is sensible). Quoting the on-line help,

The Fligner-Killeen (median) test has been determined in a simulation study as one of the many tests for homogeneity of variances which is most robust against departures from normality, see Conover, Johnson & Johnson (1981).

Generally speaking, the Levene test works well in the ANOVA framework, providing there are small to moderate deviations from the normality. In this case, it outperforms the Bartlett test. If the distribution are nearly normal, however, the Bartlett test is better. I've also heard of the Brown-Forsythe test as a non-parametric alternative to the Levene test. Basically, it relies on either the median or the trimmed mean (as compared to the mean in the Levene test). According to Brown and Forsythe (1974), a test based on the mean provided the best power for symmetric distributions with moderate tails.

In conclusion, I would say that if there is strong evidence of departure from the normality (as seen e.g., with the help of a Q-Q plot), then use a non-parametric test (FK or BF test); otherwise, use Levene or Bartlett test.

There was also a small discussion about this test for small and large samples in the R Journal, last year, [asymptTest: A Simple R Package for Classical Parametric Statistical Tests and Confidence Intervals in Large Samples](#). It seems that the FK test is also available through the [coin](#) interface for permutation tests, see the [vignette](#).

References

Brown, M. B. and Forsythe, A. B. (1974). Robust Tests for Equality of Variances. *JASA*, 69, 364-367.

47 How to project a vector onto matrix of rotation in PCA

Well, @Srikant already gave you the right answer since the rotation (or loadings) matrix contains eigenvectors arranged column-wise, so that you just have to multiply (using `%*%`) your vector or matrix of new data with e.g. `prcomp(X)$rotation`. Be careful, however, with any extra centering or scaling parameters that were applied when computing PCA EVs.

In R, you may also find useful the `predict()` function, see `?predict.prcomp`. BTW, you can check how projection of new data is implemented by sim-

ply entering:

```
getS3method("predict", "prcomp")
```

48 What stop-criteria for agglomerative hierarchical clustering are used in practice?

It is rather difficult to provide a clear-cut solution about how to choose the “best” number of clusters in your data, whatever the clustering method you use, because Cluster Analysis seeks to isolate groups of statistical units (whether it be individuals or variables) for exploratory or descriptive purpose, essentially. Hence, you also have to interpret the output of your clustering scheme and several cluster solutions may be equally interesting.

Now, regarding usual statistical criteria used to decide when to stop to aggregate data, as pointed by @ars most are *visual-guided criteria*, including the analysis of the dendrogram or the inspection of clusters profiles, also called *silhouette* plots (Rousseeuw, 1987). Several *numerical criteria*, also known as validity indices, were also proposed, e.g. Dunn's validity index, Davies-Bouldin validity index, C index, Hubert's gamma, to name a few. Hierarchical clustering is often run together with k-means (in fact, several instances of k-means since it is a stochastic algorithm), so that it add support to the clustering solutions found. I don't know if all of this stuff is readily available in Python, but a huge amount of methods is available in R (see the [Cluster](#) task view, already cited by @mbq for a related question, [What tools could be used for applying clustering algorithms on MovieLens?](#)). Other approaches include [fuzzy clustering](#) and [model-based clustering](#) (also called *latent trait analysis*, in the psychometric community) if you seek more robust way to choose the number of clusters in your data.

BTW, I just came across this webpage, [scipy-cluster](#), which is *an extension to Scipy for generating, visualizing, and analyzing hierarchical clusters*. Maybe it includes other functionalities? I've also heard of [PyChem](#) which offers pretty good stuff for multivariate analysis.

The following reference may also be helpful:

Steinley, D., & Brusco, M. J. (2008). Selection of variables in cluster analysis: An empirical comparison of eight procedures. *Psychometrika*, 73, 125-

144.

49 How to combine confidence intervals for a variance component of a mixed-effects model when using multiple imputation

This is a great question! Not sure this is a full answer, however, I drop these few lines in case it helps.

It seems that Yucel and Demirtas (2010) refer to an older paper published in the JCGS, [Computational strategies for multivariate linear mixed-effects models with missing values](#), which uses an hybrid EM/Fisher scoring approach for producing likelihood-based estimates of the VCs. It has been implemented in the R package [mlmmm](#). I don't know, however, if it produces CIs.

Otherwise, I would definitely check the [WinBUGS](#) program, which is largely used for multilevel models, including those with missing data. I seem to remember it will only works if your MV are in the response variable, not in the covariates because we generally have to specify the full conditional distributions (if MV are present in the independent variables, it means that we must give a prior to the missing Xs, and that will be considered as a parameter to be estimated by WinBUGS...). It seems to apply to R as well, if I refer to the following thread on [r-sig-mixed](#), [missing data in lme, lmer, PROC MIXED](#). Also, it may be worth looking at the [MLwiN](#) software.

50 Mathematical Statistics Videos

I just came across this website, [CensusAtSchool – Informal inference](#). Maybe worth looking at the videos and handouts...

51 Swamy Random Coefficient Model and time fixed effects

I think you use Stata, given your other post about [Panel data and selection models issue](#). Did you look at the following paper, [From the help desk: Swamy's random-coefficients model](#) from the Stata Journal (2003 3(3))? It seems that the command [xtcrhh2](#) (available through [findit xtcrhh](#) in Stata com-

mand line) includes an option about time, but I'm afraid it only allow to estimate the panel-specific coefficients. Looking around, I only found this article, [Estimation and testing of fixed-effect panel-data systems](#) (SJ 2005 5(2)), but it doesn't seem to address your question. So maybe it is better to use the [xtreg](#) command directly. If you have more than one random coefficient, then it may be better to [gllamm](#).

Otherwise, I would suggest trying the [plm](#) R package (it has a lot of dependencies, but it mainly relies on the [nlme](#) and [survival](#) packages). The [effect](#) parameter that is passed to [plm\(\)](#) seems to return individual, time or both (for balanced design) kind of effects; there's also a function names [plstest\(\)](#). I'm not a specialist of econometrics, I only used it for clinical trials in the past, but quoting the online help, it seems you will be able to get fixed effects for your time covariate (expressed as deviations from the overall mean or as deviations from the first value of the index):

```
library(plm)
data("Grunfeld", package = "plm")
gi <- plm(inv ~ value + capital, data = Grunfeld,
          model = "within", effect = "twoways")
summary(gi)
fixef(gi, effect = "time")
```

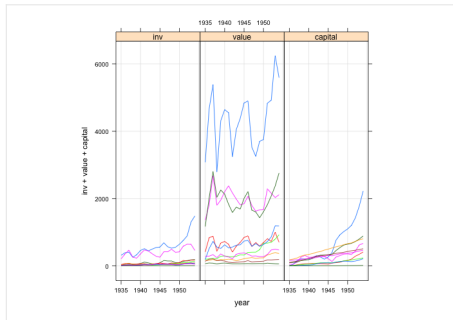
where the data looks like (or see the plot below to get a rough idea):

	firm	year	inv	value	capital
1	1	1935	317.6	3078.5	2.8
2	1	1936	391.8	4661.7	52.6
3	1	1937	410.6	5387.1	156.9
...					
198	10	1952	6.00	74.42	9.93
199	10	1953	6.53	63.51	11.68
200	10	1954	5.12	58.12	14.33

For more information, check the accompanying vignette or this paper, [Panel Data Econometrics in R: The plm Package](#), published in the JSS (2008 27(2)).

52 Difference between Norm of Residuals and what is a “good” Norm of Residual

So, I would recommend using standard method for comparing nested models. In your case, you consider two alternative models, the cubic fit being the more “complex” one. An F- or χ^2 -test tells you



whether the residual sum of squares or deviance significantly decrease when you add further terms. It is very like comparing a model including only the intercept (in this case, you have residual variance only) vs. another one which include one meaningful predictor: does this added predictor account for a sufficient part of the variance in the response? In your case, it amounts to say: Modeling a cubic relationship between X and Y decreases the unexplained variance (equivalently, the R^2 will increase), and thus provide a better fit to the data compared to a linear fit.

It is often used as a *test of linearity* between the response variable and the predictor, and this is the reason why Frank Harrell advocates the use of **restricted cubic spline** instead of assuming a strict linear relationship between Y and the continuous X s (e.g. age).

The following example comes from a book I was reading some months ago (**High-dimensional data analysis in cancer research**, Chap. 3, p. 45), but it may well serves as an illustration. The idea is just to fit different kind of models to a simulated data set, which clearly highlights a non-linear relationship between the response variable and the predictor. The true generative model is shown in black. The other colors are for different models (restricted cubic spline, B-spline close to yours, and CV smoothed spline).

```
library(rms)
library(splines)
set.seed(101)
f <- function(x) sin(sqrt(2*pi*x))
n <- 1000
x <- runif(n, 0, 2*pi)
sigma <- rnorm(n, 0, 0.25)
y <- f(x) + sigma
plot(x, y, cex=.4)
curve(f, 0, 6, lty=2, add=TRUE)
# linear fit
```

```
lm00 <- lm(y~x)
# restricted cubic spline, 3 knots (2 Df)
lm0 <- lm(y~rcs(x,3))
lines(seq(0,6,length=1000),
      predict(lm0,data.frame(x=seq(0,6,length=1000))),
      col="red")
# use B-spline and a single knot at x=1.13 (4 Df)
lm1 <- lm(y~bs(x, knots=1.13))
lines(seq(0,6,length=1000),
      predict(lm1,data.frame(x=seq(0,6,length=1000))),
      col="green")
# cross-validated smoothed spline (approx. 20 Df)
xy.spl <- smooth.spline(x, y, cv=TRUE)
lines(xy.spl, col="blue")
legend("bottomleft", c("f(x)", "RCS {rms}", "BS {splines}", "SS
{stats}"),
      col=1:4, lty=c(2,rep(1,3)),bty="n", cex=.6)
```

Now, suppose you want to compare the linear fit (**lm00**) and model relying on B-spline (**lm1**), you just have to do an F-test to see that the latter provides a better fit:

```
> anova(lm00, lm1)
Analysis of Variance Table

Model 1: y ~ x
Model 2: y ~ bs(x, knots = 1.13)
  Res.Df    RSS Df Sum of Sq    F    Pr(>F)
1     998 309.248
2     995  63.926   3    245.32 1272.8 < 2.2e-16 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

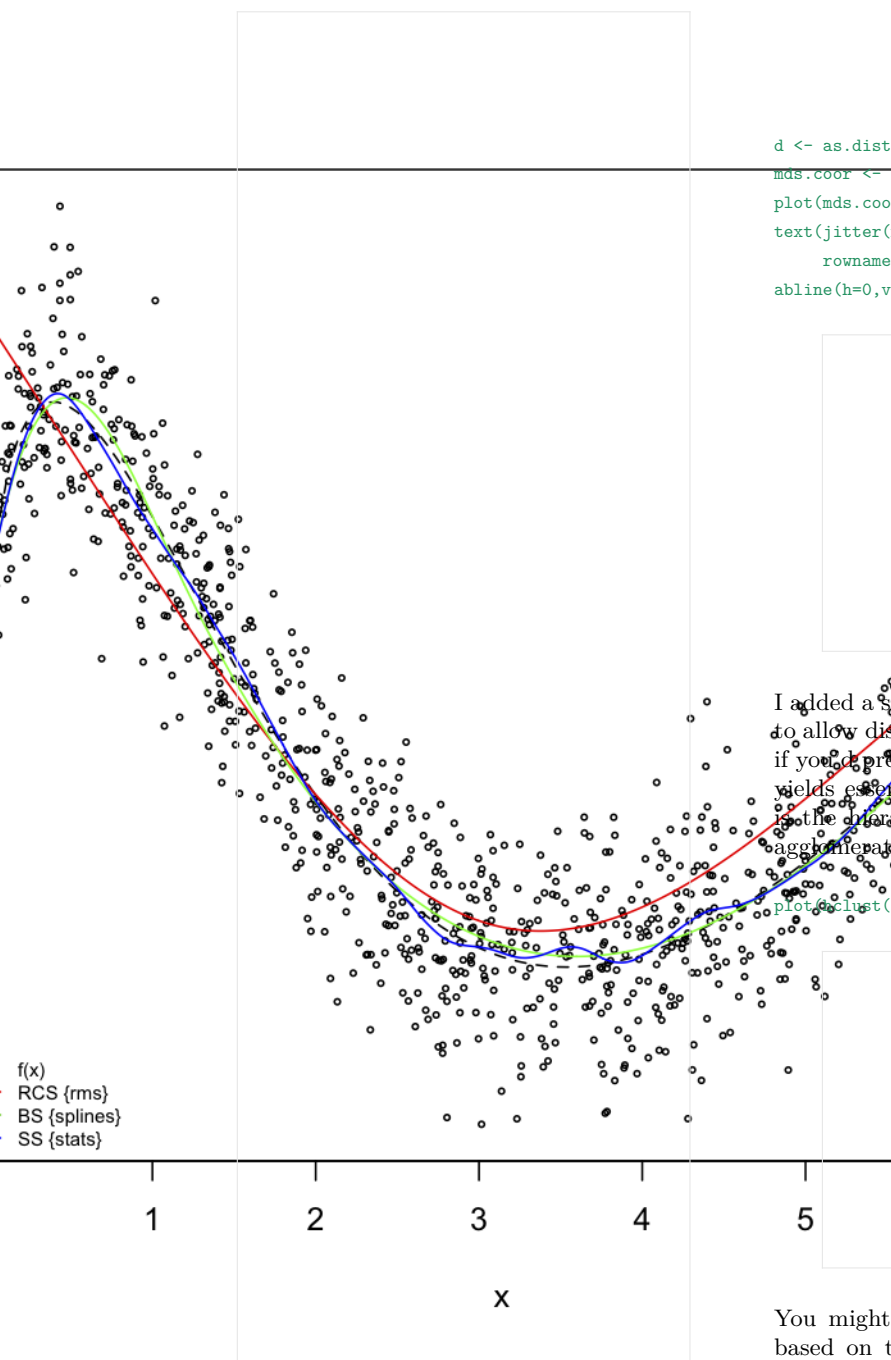
Likewise, it is quite usual to compare **GLM** with **GAM** based on the results of a χ^2 -test.

53 Clustering with a distance matrix

One way to highlight clusters on your distance matrix is by way of **Multidimensional scaling**. When projecting individuals (here what you call your nodes) in an 2D-space, it provides a comparable solution to PCA. This is unsupervised, so you won't be able to specify a priori the number of clusters, but I think it may help to quickly summarize a given distance or similarity matrix.

Here is what you would get with your data:

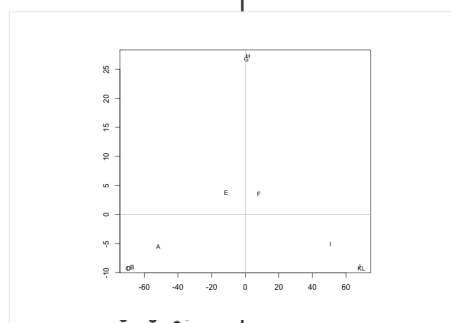
```
tmp <- matrix(c(0,20,20,20,40,60,60,60,100,120,120,120,
```

```

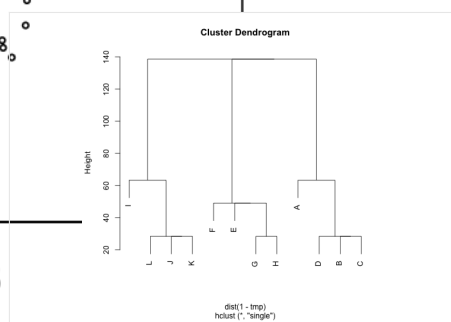
120,140,140,140,80,60,80,80,20,0,20,20,
120,140,140,140,80,60,80,80,20,0,20,
120,140,140,140,80,60,80,80,20,0,20,
nr=12, dimnames=list(LETTERS[1:12], LETTERS[1:12]))
d <- as.dist(tmp)
mds.coor <- cmdscale(d)
plot(mds.coor[,1], mds.coor[,2], type="n", xlab="", ylab="")
text(jitter(mds.coor[,1]), jitter(mds.coor[,2]),
     rownames(mds.coor), cex=0.8)
abline(h=0,v=0,col="gray75")

```



I added a small jittering on the x and y coordinates to allow distinguishing cases. Replace `tmp` by `1-tmp` if you'd prefer working with dissimilarities, but this yields essentially the same picture. However, here is the hierarchical clustering solution, with *single* agglomeration criteria:

```
plot(hclust(dist(1-tmp), method="single"))
```



You might further refine the selection of clusters based on the dendrogram, or more robust methods, see e.g. this related question: [What stop-criteria for agglomerative hierarchical clustering are used in practice?](#)

```

20,0,20,20,60,80,80,80,120,140,140,140,
20,20,0,20,60,80,80,80,120,140,140,140,
20,20,20,0,60,80,80,80,120,140,140,140,
40,60,60,60,0,20,20,20,60,80,80,80,
60,80,80,80,20,0,20,20,40,60,60,60,
60,80,80,80,20,20,0,20,60,80,80,80,
60,80,80,80,20,20,20,0,60,80,80,80,
100,120,120,120,60,40,60,60,0,20,20,20,

```


54 Good resource to understand anova and ancova?

So, in addition to this paper, [Misunderstanding Analysis of Covariance](#), which enumerates common pitfalls when using ANCOVA, I would recommend starting with:

- Frank Harrell’s [homepage](#), especially his hand-out on [Regression Modeling Strategies](#) and [Biostatistical Modeling](#)
- John Fox’s [homepage](#) includes great material on [Linear Model](#)
- [Practical Regression and Anova using R](#)

This is mostly R-oriented material, but I feel you might better catch the idea if you start playing a little bit with these models on toy examples or real datasets (and R is great for that).

As for a good book, I would recommend [Design and Analysis of Experiments](#) by Montgomery (now in its 7th ed.); ANCOVA is described in chapter 15. [Plane Answers to Complex Questions](#) by Christensen is an excellent book on the theory of linear model (ANCOVA in chapter 9); it assumes a good mathematical background. Any biostatistical textbook should cover both topics, but I like [Biostatistical Analysis](#) by Zar (ANCOVA in chapter 12), mainly because this was one of my first textbook.

And finally, H. Baayen’s textbook is very complete, [Practical Data Analysis for the Language Sciences with R](#). Although it focus on linguistic data, it includes a very comprehensive treatment of the Linear Model and mixed-effects models.

55 On the use of oblique rotation after PCA

I think there are different opinions or views about PCA, but basically we often think of it as either a *reduction technique* (you reduce your features space to a smaller one, often much more “readable” providing you take care of properly centering/standardizing the data when it is needed) or a way to construct *latent factors* or dimensions that account for a significant part of the inter-individual dispersion (here, the “individuals” stand for the statistical units on which data are collected; this may be country, people, etc.). In both case, we construct

linear combinations of the original variables that account for the maximum of variance (when projected on the principal axis), subject to a constraint of orthogonality between any two principal components. Now, what has been described is purely algebraical or mathematical and we don’t think of it as a (generating) model, contrary to what is done in the factor analysis tradition where we include an error term to account for some kind of measurement error. I also like the introduction given by William Revelle in his forthcoming handbook on [applied psychometrics using R](#) (Chapter 6), if we want to analyze the structure of a correlation matrix, then

The first [approach, PCA] is a model that approximates the correlation matrix in terms of the product of components where each component is a weighted linear sum of the variables, the second model [factor analysis] is also an approximation of the correlation matrix by the product of two factors, but the factors in this are seen as causes rather than as consequences of the variables.

In other words, with PCA you are expressing each component (factor) as a linear combination of the variables whereas in FA these are the variables that are expressed as a linear combination of the factors. It is well acknowledged that both methods will generally yield quite similar results (see e.g. Harman, 1976 or Catell, 1978), especially in the “ideal” case where we have a large number of individuals and a good ratio factor:variables (typically varying between 2 and 10 depending on the authors you consider!). This is because, by estimating the diagonals in the correlation matrix (as is done in FA, and these elements are known as the communalities), the error variance is eliminated from the factor matrix. This is the reason why PCA is often used as a way to uncover latent factors or psychological constructs in place of FA developed in the last century. But, as we go on this way, we often want to reach an easier interpretation of the resulting factor structure (or the so-called pattern matrix). And then comes the useful trick of rotating the factorial axis so that we maximize loadings of variables on specific factor, or equivalently reach a “simple structure”. Using orthogonal rotation (e.g. VARIMAX), we preserve the independence of the factors. With oblique rotation (e.g. OBLIMIN, PROMAX), we break it and factors are allowed to correlate. This has been largely debated in the literature, and has lead some authors (not psychometricians, but statisticians in the early 1960’s) to conclude that FA is an unfair approach due to the fact that researchers might seek the factor solution that is the more convenient to interpret.

But the point is that rotation methods were originally developed in the context of the FA approach and are now routinely used with PCA. I don't think this contradicts the algorithmic computation of the principal components: You can rotate your factorial axes the way you want, provided you keep in mind that once correlated (by oblique rotation) the interpretation of the factorial space becomes less obvious.

PCA is routinely used when developing new questionnaires, although FA is probably a better approach in this case because we are trying to extract meaningful factors that take into account measurement errors and whose relationships might be studied on their own (e.g. by factoring out the resulting pattern matrix, we get a second-order factor model). But PCA is also used for checking the factorial structure of already validated ones. Researchers don't really matter about FA vs. PCA when they have, say 500 representative subjects who are asked to rate a 60-item questionnaire tackling five dimensions (this is the case of the **NEO-FFI**, for example), and I think they are right because in this case we aren't very much interested in identifying a generating or conceptual model (the term "representative" is used here to alleviate the issue of *measurement invariance*).

Now, about the choice of rotation method and why some authors argue against the strict use of orthogonal rotation, I would like to quote Paul Kline, as I did in response to the following question, **FA: Choosing Rotation matrix, based on "Simple Structure Criteria"**,

(...) in the real world, it is not unreasonable to think that factors, as important determiners of behavior, would be correlated. – P. Kline, *Intelligence. The Psychometric View*, 1991, p. 19

I would thus conclude that, depending on the objective of your study (do you want to highlight the main patterns of your correlation matrix or do you seek to provide a sensible interpretation of the underlying mechanisms that may have caused you to observe such a correlation matrix), you are up to choose the method that is the most appropriate: This doesn't have to do with the construction of linear combinations, but merely on the way you want to interpret the resulting factorial space.

References

1. Harman, H.H. (1976). *Modern Factor Analysis*.

Chicago, University of Chicago Press.

2. Cattell, R.B. (1978). *The Scientific Use of Factor Analysis*. New York, Plenum.
3. Kline, P. (1991). *Intelligence. The Psychometric View*. Routledge.

56 How to conduct conditional Cox regression for matched case-control study?

If you are using Stata, you can just look at the **stcox** command. Examples are available from **Stata** or **UCLA** website. Also, take a look at **Analysis of matched cohort data** from the *Stata Journal* (2004 4(3)).

Under R, you can use the **coxph()** function from the **survival** library.

57 How to use STATA to pool Cohen's d?

For SPSS, look at :

- www.mathkb.com/Uwe/Forum.aspx/stat-consult/1201/Effect-Size-in-SPSS
- www.spsstools.net/Syntax/T-Test/StandardizedEffectsSize.txt (maybe better organized)

For Stata, I used **SIZEFX: Stata module to compute effect size correlations** (**findit sizefx** at Stata command prompt), but **metan** as suggested by @onestop is probably more featured.

58 Good text on Clinical Trials?

I would definitively recommend **Design and Analysis of Clinical Trials: Concepts and Methodologies** which seems actually the most complete one given your request.

Statistical Issues in Drug Development also covers a broad range of concepts but is less oriented toward design of experiment. **Statistics Applied to Clinical Trials** includes more technical stuff, but mainly focus on crossover trials and applications in epidemiology. And there is always the most famous Rothman, **Modern Epidemiology**, which provides valuable

help for interpretation and examples of application of clinical biostatistics. Finally, [The Design and Analysis of Clinical Experiments](#) is more centered onto the analysis of specific experimental settings, but it does not address other points and is a bit older.

59 What is a consistency check?

I suppose this has to do with some form of Quality Control about *data integrity*, and more specifically that you regularly check that your working database isn't corrupted (due to error during transfer, copy, or after an update or a sanity check). This may also mean ensuring that your intermediate computation are double-checked (either manually or through additional code or macros in your statistical software).

Other information may be found here: the ICH E6 (R1) reference guide about [Guideline for Good Clinical Practice](#) from the EMEA, [Guidelines on Good Clinical Laboratory Practice](#), or [Clinical Research Study Investigator's Toolbox](#).

60 Calculating Orwin's (1983) modified Fail-safe N in a meta-analysis with Odds Ratio as summary statistic?

So, perhaps check these additional resources: <http://j.mp/d8znoP> for SPSS. Don't know about Stata. There is some R code about fail-safe N in the following handout: [Tests for funnel plot asymmetry and failsafe N](#), but I didn't check on the www.metaanalysis.com website.

Otherwise, [ClinTools Software](#) may be an option (I hope the demo version let you do some computation on real data), or better the [MIX](#) software.

61 Modelling longitudinal data where the effect of time varies in functional form between individuals

I would suggest to look at the following three directions:

- *longitudinal clustering*: this is unsupervised, but you use k-means approach relying on the Calinsky criterion for assessing quality of the partitioning (package [kml](#), and references included in the online help); so basically, it won't

help identifying specific shape for individual time course, but just separate homogeneous evolution profile

- some kind of *latent growth curve* accounting for heteroscedasticity: my best guess would be to look at the extensive references around [MPlus](#) software, especially the FAQ and mailing. I've also heard of random effect multiplicative heteroscedastic model (try googling around those keywords). I find these papers ([1](#), [2](#)) interesting, but I didn't look at them in details. I will update with references on neuropsychological assessment once back to my office.
- *functional PCA* ([fpca](#) package) but it may be worth looking at [functional data analysis](#)

Other references (just browsed on the fly):

- Willett & Bull (2004), [Latent Growth Curve Analysis](#) – the authors use LGC on non-linear reading trajectories
- Welch (2007), [Model Fit and Interpretation of Non-Linear Latent Growth Curve Models](#) – a PhD on modeling non-linear change in the context of latent growth modeling
- Berkey CS, Laird NM (1986). [Nonlinear growth curve analysis: estimating the population parameters](#). Ann Hum Biol. 1986 Mar-Apr;13(2):111-28
- Rice (2003), [Functional and Longitudinal Data Analysis: Perspectives on Smoothing](#)
- Wu, Fan and Müller (2007). [Varying-Coefficient Functional Linear Regression](#)

62 Correcting p values for multiple tests where tests are correlated (genetics)

This is actually a hot topic in Genomewide analysis studies (GWAS)! I am not sure the method you are thinking of is the most appropriate in this context. Pooling of p-values was described by some authors, but in a different context (replication studies or meta-analysis, see e.g. ([1](#)) for a recent review). Combining SNP p-values by Fisher's method is generally used when one wants to derive an unique p-value for a given gene; this allows to work at the

gene level, and reduce the amount of dimensionality of subsequent testing, but as you said the non-independence between markers (arising from spatial colocation or linkage disequilibrium, LD) introduce a bias. More powerful alternatives rely on resampling procedures, for example the use of maxT statistics for combining p-value and working at the gene level or when one is interested in pathway-based approaches, see e.g. (2) (§2.4 p. 93 provides details on their approach).

My main concerns with bootstrapping (with replacement) would be that you are introducing an artificial form of relatedness, or in other words you create virtual twins, hence altering Hardy-Weinberg equilibrium (but also minimum allele frequency and call rate). This would not be the case with a permutation approach where you permute individual labels and keep the genotyping data as is. Usually, the [plink](#) software can give you raw and permuted p-values, although it uses (by default) an adaptive testing strategy with a sliding window that allows to stop running all permutations (say 1000 per SNP) if it appears that the SNP under consideration is not “interesting”; it also has option for computing maxT, see the [online help](#).

But given the low number of SNPs you are considering, I would suggest relying on FDR-based or maxT tests as implemented in the [multtest](#) R package (see [mt.maxT](#)), but the definitive guide to resampling strategies for genomic application is [Multiple Testing Procedures with Applications to Genomics](#), from Dudoit & van der Laan (Springer, 2008). See also Andrea Foulkes’s book on [genetics with R](#), which is reviewed in the JSS. She has great material on multiple testing procedures.

Further Notes

Many authors have pointed to the fact that simple multiple testing correcting methods such as the Bonferroni or Sidak are too stringent for adjusting the results for the individual SNPs. Moreover, neither of these methods take into account the correlation that exists between SNPs due to LD which tags the genetic variation across gene regions. Other alternative have been proposed, like a derivative of Holm’s method for multiple comparison (3), Hidden Markov Model (4), conditional or positive FDR (5) or derivative thereof (6), to name a few. So-called gap statistics or sliding window have been proved successful in some case, but you’ll find a good review in (7) and (8).

I’ve also heard of methods that make effective use of the haplotype structure or LD, e.g. (9), but I never used them. They seem, however, more related to estimating the correlation between markers, not p-value as you meant. But in fact, you might better think in terms of the dependency structure between successive test statistics, than between correlated p-values.

References

1. Cantor, RM, Lange, K and Sinsheimer, JS. [Prioritizing GWAS Results: A Review of Statistical Methods and Recommendations for Their Application](#). Am J Hum Genet. 2010 86(1): 6–22.
2. Corley, RP, Zeiger, JS, Crowley, T et al. [Association of candidate genes with antisocial drug dependence in adolescents](#). Drug and Alcohol Dependence 2008 96: 90–98.
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4. Wei, Z, Sun, W, Wang, K, and Hakonarson, H. [Multiple Testing in Genome-Wide Association Studies via Hidden Markov Models](#). Bioinformatics 2009 25(21): 2802–2808.
5. Broberg, P. [A comparative review of estimates of the proportion unchanged genes and the false discovery rate](#). BMC Bioinformatics 2005 6: 199.
6. Need, AC, Ge, D, Weale, ME, et al. [A Genome-Wide Investigation of SNPs and CNVs in Schizophrenia](#). PLoS Genet. 2009 5(2): e1000373.
7. Han, B, Kang, HM, and Eskin, E. [Rapid and Accurate Multiple Testing Correction and Power Estimation for Millions of Correlated Markers](#). PLoS Genetics 2009
8. Liang, Y and Kelemen, A. [Statistical advances and challenges for analyzing correlated high dimensional snp data in genomic study for complex diseases](#). Statistics Surveys 2008 2 :43–60. – the best recent review ever
9. Nyholt, DR. [A Simple Correction for Multiple Testing for Single-Nucleotide Polymorphisms in Linkage Disequilibrium with Each Other](#). Am J

Hum Genet. 2004 74(4): 765–769.

10. Nicodemus, KK, Liu, W, Chase, GA, Tsai, Y-Y, and Fallin, MD. **Comparison of type I error for multiple test corrections in large single-nucleotide polymorphism studies using principal components versus haplotype blocking algorithms**. BMC Genetics 2005; 6(Suppl 1): S78.
11. Peng, Q, Zhao, J, and Xue, F. **PCA-based bootstrap confidence interval tests for gene-disease association involving multiple SNPs**. BMC Genetics 2010, 11:6
12. Li, M, Romero, R, Fu, WJ, and Cui, Y (2010). **Mapping Haplotype-haplotype Interactions with Adaptive LASSO**. BMC Genetics 2010, 11:79 – although not directly related to the question, it covers haplotype-based analysis/epistatic effect

63 What ways are there to show two analytical methods are equivalent ?

The simple correlation approach isn't the right way to analyze results from method comparison studies. There are (at least) two highly recommended books on this topic that I referenced at the end (1,2). Briefly stated, when comparing measurement methods we usually expect that (a) our conclusions should not depend on the particular sample used for the comparison, and (b) measurement error associated to the particular measurement instrument should be accounted for. This precludes any method based on correlations, and we shall turn our attention to variance components or mixed-effects models that allow to reflect the systematic effect of item (here, item stands for individual or sample on which data are collected), which results from (a).

In your case, you have single measurements collected using two different methods (I assume that none of them might be considered as a gold standard) and the very basic thing to do is to plot the differences ($X_1 - X_2$) versus the means $((X_1 + X_2)/2)$; this is called a **Bland-Altman plot**. It will allow you to check if (1) the variations between the two set of measurements are constant and (2) the variance of the difference is constant across the range of observed values. Basically, this is just a 45° rotation of a simple scatterplot of X_1 vs. X_2 , and its interpretation is close to a plot of fitted vs.

residuals values used in linear regression. Then,

- if the difference is constant (*constant bias*), you can compute the limit of agreement (see (3))
- if the difference is not constant across the range of measurement, you can fit a linear regression model between the two methods (choose the one you want as predictor)
- if the variance of the differences is not constant, try to find a suitable transformation that makes the relationship linear with constant variance

Other details may be found in (2), chapter 4.

References

1. Dunn, G (2004). *Design and Analysis of Reliability Studies*. Arnold. See the review in the *International Journal of Epidemiology*.
2. Carstensen, B (2010). *Comparing clinical measurement methods*. Wiley. See the **companion website**, including R code.
3. The original article from Bland and Altman, **Statistical methods for assessing agreement between two methods of clinical measurement**.
4. Carstensen, B (2004). **Comparing and predicting between several methods of measurement**. *Biostatistics*, 5(3), 399–413.

64 What is the difference between summary and loadings for princomp?

The first output is the correct and most useful one. Calling `loadings()` on your object just returns a summary where the SS are always equal to 1, hence the % variance is just the SS loadings divided by the number of variables. It makes sense only when using Factor Analysis (like in `factanal`). I never use `princomp` or its SVD-based alternative (`prcomp`), and I prefer the `FactoMineR` or `ade4` package which are by far more powerful!

About your second question, the `summary()` function just returns the SD for each component (`pc.cr$sdev` in your case), and the rest of the table seems to be computed afterwards (through the

print or show method, I didn't investigate this in details).

```
> getS3method("summary","princomp")
function (object, loadings = FALSE, cutoff = 0.1, ...)
{
  object$cutoff <- cutoff
  object$print.loadings <- loadings
  class(object) <- "summary.princomp"
  object
}
<environment: namespace:stats>
```

What `princomp()` itself does may be viewed using `getAnywhere("princomp.default")`.

65 Variation in PCA weights

It looks like you are referring to eigenanalysis for SNPs data and the article from Nick Patterson, **Population Structure and Eigenanalysis** (PLoS Genetics 2006), where the first component explains the largest variance on allele frequency wrt. potential stratification in the sample (due to ethnicity or, more generally, ancestry). So I wonder why you want to consider all three first components, unless they appear to be significant from their expected distribution according to **TW distribution**. Anyway, in R you can isolate the most informative SNPs (i.e. those that are at the extreme of the successive principal axes) with the `apply()` function, working on row, e.g.

```
apply(snp.df, 1, function(x) any(abs(x)>threshold))
```

where `snp.df` stands for the data you show and which is stored either as a `data.frame` or `matrix` under R, and `threshold` is the value you want to consider (this can be $\text{Mean} \pm 6 \text{ SD}$, as in Price et al. *Nature Genetics* 2007 38(8): 904, or whatever value you want). You may also implement the iterative PCA yourself.

Finally, the TW test can be implemented as follows:

```
##' Test for the largest eigenvalue in a gaussian covariance
matrix
##'
##' This function computes the test statistic and associated
p-value
##' for a Wishart matrix focused on individuals (Tracy-Widom
distribution).
##'
##' @param C a rectangular matrix of bi-allelic markers (columns)
```

```
indexed
##' on m individuals. Caution: genotype should be in
{0,1,2}.
##' @return test statistic and tabulated p-value
##' @reference \cite{Johnstone:2001}
##' @seealso The RMTstat package provides other interesting
functions to
##' deal with Wishart matrices.
##' @example
##' X <- replicate(100,sample(0:2,20,rep=T))
tw.test <- function(C) {
  m <- nrow(C) # individuals
  n <- ncol(C) # markers
  # compute M
  C <- scale(C, scale=F)
  pj <- attr(C,"scaled:center")/2
  M <- C/sqrt(pj*(1-pj))
  # compute X=MM'
  X <- M %*% t(M)
  ev <- sort(svd(X)$d, decr=T)[1:(m-1)]
  nprime <- ((m+1)*sum(ev^2)/(((m-1)*sum(ev^2))-sum(ev)^2)
  l <- (m-1)*ev[1]/sum(ev)
  # normalize l and compute test statistic
  num <- (sqrt(nprime-1)+sqrt(m))
  mu <- num^2/nprime
  sigma <- num/nprime*(1/sqrt(nprime-1)+1/sqrt(m))^(1/3)
  l <- (l-mu)/sigma
  # estimate associated p-value
  if (require(RMTstat)) pv <- ptw(l, lower.tail=F)
  else pv <- NA
  return(list(stat=l, pval=pv))
}
```

66 An easy explanation for the parallel coordinates plot

It seems to me that the main function of PCP is to highlight homogeneous groups of individuals, or conversely (in the dual space, by analogy with PCA) specific patterns of association on different variables. It produces an effective graphical summary of a multivariate data set, when there are not too much variables. Variables are automatically scaled to a fixed range (typically, 0–1) which is equivalent to working with standardized variables (to prevent the influence of one variable onto the others due to scaling issue), but for very high-dimensional data set (# of variables > 10), you definitely have to look at other displays, like **fluctuation plot** or **heatmap** as used in microarray studies.

It helps answering questions like:

- are there any consistent pattern of individual

scores that measure membership (i.e. their

- are there any scores observed (e.g. low scores) ways associated with

In the following plot, we can see that species within each cluster show very distinct patterns of petal length and sepal width, and are more homogeneous in sepal length (i.e. their

You can even use dimension reduction techniques, when performing the features of clusters of individuals who systematically vary in some kind of biological scores and highlighting membership on the face of the package).

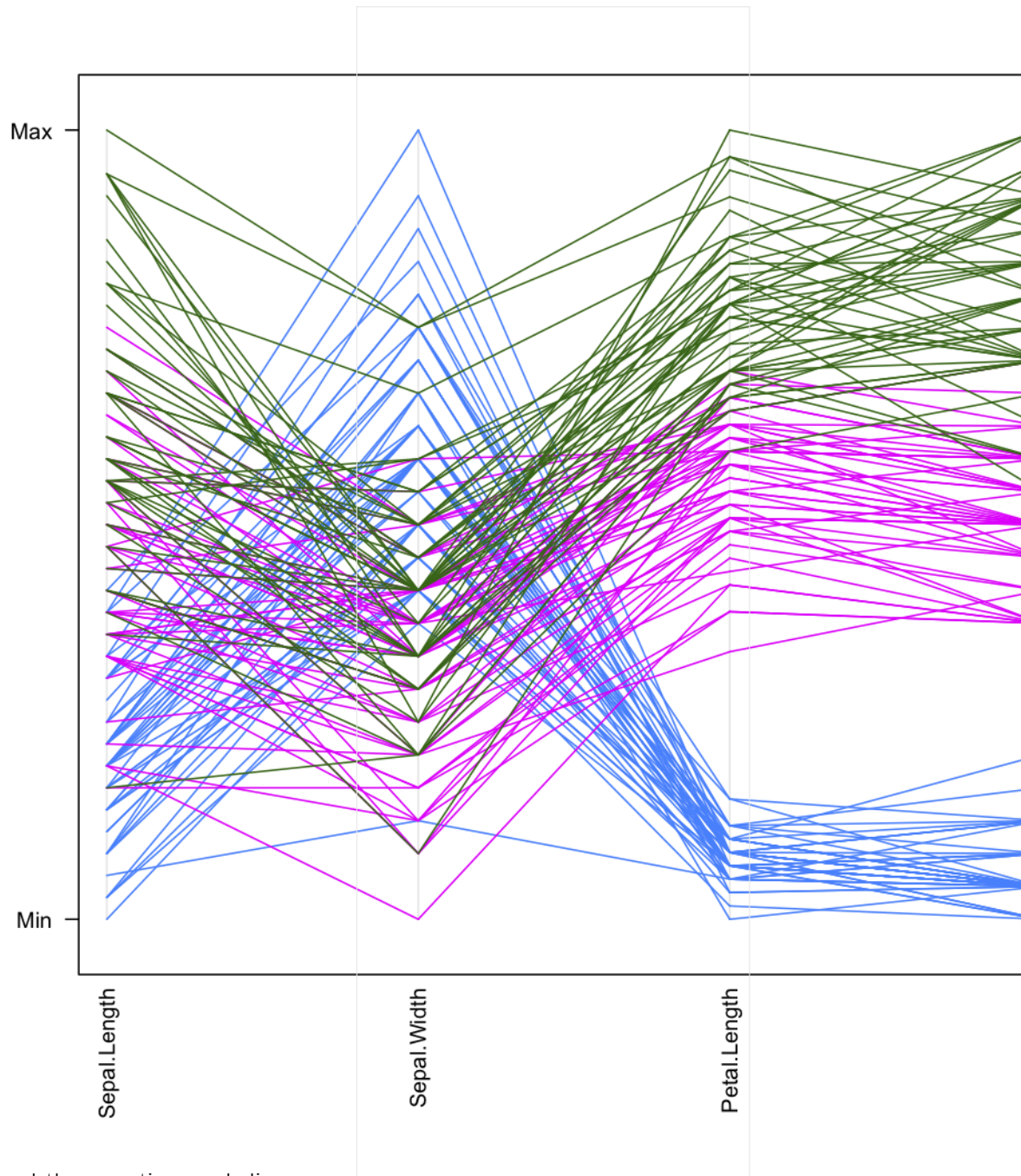
It is also used in clustering and hierarchical clustering, including how cluster membership, the number of clusters, for agglomerative practice?).

Such displays are scatterplots (which show a 2D-relationship) is available in the `ggplot2` or the `Mondrian` s

67 How to

For most of your data, the distributions have identical shapes, which is thrown by R:

for all observations, and there are ties, precluding the computation of an exact p-value. For your variable named `V2`, there are in fact only two distinct values (out of 7), so I am very puzzled by the approach you took to analysis your data. With such a high number of tied data, I would not trust any Wilcoxon test. Moreover, in most non-parametric tests we assume that the sampled populations are symmetric and have the same dispersion or shape,



which is hardly verifiable in your case.

Thus, I think a permutation test would be more appropriate in your case, see e.g. `permTS` (`perm`), `pperm` (`exactRankTests`), or the `coin` package.

68 How will you deal with “don’t know” and “missing data” in survey data?

Well, you should also considered that “don’t know” is at least some kind of answer, whereas non-response is a purely missing value. Now, we often allow for “don’t know” response in survey just to avoid forcing people to provide a response anyway (which might bias the results). For example, in the National Health and Nutrition Examination Survey, they are coded differently but subsequently discarded from the analysis.

You could try analyzing the data both ways: (1) treating “don’t know response” as specific response category and handling all responses set with some kind of multivariate data analysis (e.g. [multiple correspondence analysis](#) or multiple factor analysis for mixed data, see the [FactoMineR](#) package), and (2) if it doesn’t bring any evidence of distortion on items distribution, just merge it with missing values.

For (2), I would also suggest you to check that “don’t know” and MV are at least missing at random (MAR), or that they are not specific of one respondents group (e.g. male/female, age class, SES, etc.).

69 How to do community detection in a weighted social network/graph?

I know that [Gephi](#) can process undirected weighted graph, but I seem to remember it has to be stored in [GDF](#), which is pretty close to CSV, or Ucinet [DL](#). Be aware that it’s still an alpha release. Now, about clustering your graph, Gephi seems to lack clustering pipelines, except for the MCL algorithm that is now available in the latest version. There was a [Google Code Project](#) in 2009, [Gephi Network Statistics](#) (featuring e.g. Newman’s modularity metric), but I don’t know if something has been released in this direction. Anyway, it seems to allow some kind of modularity/clustering computations, but see also [Social Network Analysis using R and Gephi](#) and [Data preparation for Social Network Analysis using R and Gephi](#) (Many thanks to @Tal).

If you are used to Python, it is worth trying [NetworkX](#) (Here is an example of a [weighted graph](#)

with the corresponding code). Then you have many ways to carry out your analysis.

You should also look at [INSNA - Social Network Analysis Software](#) or Tim Evans’s webpage about [Complex Networks and Complexity](#).

70 kNN randomization test in R?

I am not sure to understand your question since you talk about k-means, which is basically an *unsupervised method* (i.e. where classes are not known a priori), while at the same time you are saying that you already identified groups of individuals. So I would suggest to look at classification methods, or other *supervised methods* where class membership is known and the objective is to find a weighted combination of your variables that minimize your classification error rate (this is just an example). For instance, [LDA](#) does a good job (see the CRAN task view on [Multivariate Statistics](#)), but look also at the machine learning community (widely represented on the [stats.stackexchange](#)) for other methods.

Now since you also talked of k-nearest neighbor, I wonder if you are not introducing a confusion between k-means and [kNN](#). In this case, the corresponding R function is [knn\(\)](#) in the [class](#) package (it includes cross-validation), or see the [kkn](#) package.

71 Factor analysis of dyadic data

Structural equation models are better suited for this kind of data, e.g. by introducing an extra factor for couple which allows to account for the dependence structure (paired responses). David A. Kenny reviewed the main points for [analysis dyadic data](#); although it doesn’t focus on questionnaire analysis, it may help.

A couple of references :

1. Olsen, JA and Kenny, DA (2006). [Structural Equation Modeling With Interchangeable Dyads](#). *Psychological Methods*, 11(2), 127–141.
2. McMahon,, JM, Pouget, ER, and Tortu, S (2006). [A guide for multilevel modeling of dyadic data with binary outcomes using SAS PROC NL MIXED](#). *Comput Stat Data Anal.*, 50(12), 3663–3680.

3. Thompson, L and Walker, AJ (1982). **The Dyad as the Unit of Analysis: Conceptual and Methodological Issues**. *Journal of Marriage and the Family*, 889-900.
4. Newsom, JT (2002). **A multilevel structural equation model for dyadic data**. *Structural Equation Modeling*, 9(3), 441-447.
5. González, J, Tuerlinckx, F, and De Boeck, P (2009). **Analyzing structural relations in multivariate dyadic binary data**. *Applied Multivariate Research*, 13, 77-92.
6. Gill, PS (2005). **Bayesian Analysis of Dyadic Data**.

For more thorough description of the models for dyadic data (although not restrained to item analysis), I would suggest

- Kenny, DA, Kashy, DA, and Cook, WL (2006). **Dyadic Data Analysis**. Guilford Press.
- Card, NA, Selig, JP, and Little, TD (2008). **Modeling Dyadic and Interdependent Data in the Developmental and Behavioral Sciences**. Mahwah, NJ: Lawrence Erlbaum Associates.

72 Regression analysis and parameter estimates with populations

What you describe seems to refer to a particular case of multilevel modeling, where data are organized into a hierarchical structure; in your case, forests (1st level unit) nested in counties nested in states, but see **Under what conditions should one use multilevel/hierarchical analysis?**.

Now, the “particular” case comes from the fact that you want to account for the spatial proximity which might be a vector for the propagation of **Lyme disease** (correct me if I am wrong), as is done in epidemiology where one is interested in studying the geography of infectious disease. In the usual case, we can use so-called *spatial models* like the *multiple membership model* or the *conditional autoregressive model*, among others. I enclose a couple of references about these approaches at the end, but I think you will find more references by looking at related studies in ecology or epidemiology.

Now, I think that you may pay a particular attention at the following paper of Langford et al. which features multilevel modeling with spatially correlated data:

Langford, IH, Leyland, AHL, Rasbash, and Goldstein, H (1999). **Multilevel modelling of the geographical distributions of diseases**. *Journal of Royal Statistical Society C*, 48, 253-268.

Harvey Goldstein is the author of an excellent book on multilevel modeling, **Multilevel Statistical Models** (the 2nd edition is available for free). Finally, the book of Andrew Gelman, **Data Analysis Using Regression and Multilevel/Hierarchical Models**, may provide additional clues about hierarchical/multilevel modeling.

About software, I know there is the R **spdep** package for modeling spatially correlated outcomes, but there are some examples of analysis of spatial hierarchical data with WinBUGS on the **BUGS Project**.

References

1. Browne, W.J., Goldstein, H. and Rasbash, J. (2001) **Multiple membership multiple classification (MMMC) models**. *Statistical Modelling*, 1, 103-124.
2. Lichstein, JW, Simons, TR, Shiner, SA, and Franzreb, KE (2002). **Spatial autocorrelation and autoregressive models in ecology**. *Ecological Monographs*, 72(3), 445-463.
3. Feldkircher, M (2007). **A Spatial CAR Model applied to a Cross-Country Growth Regression**.
4. Lawson, AB, Browne, WJ, and Vidal Rodeiro, CL (2003). *Disease mapping with WinBUGS and MLwiN*. John Wiley & Sons.

73 How to look for valleys in a graph?

Some of the **Bioconductor**’s packages (e.g., **ShortRead**, **Biostrings**, **BSgenome**, **IRanges**, **genomIntervals**) offer facilities for dealing with genome positions or coverage vectors, e.g. for **ChIP-seq** and identifying enriched regions. As for the other answers, I agree that any method relying on ordered observations with some threshold-based filter would allow to iso-

late low signal within a specific bandwidth.

Maybe you can also look at the methods used to identify so-called “islands”

Zang, C, Schones, DE, Zeng, C, Cui, K, Zhao, K, and Peng, W (2009). [A clustering approach for identification of enriched domains from histone modification ChIP-Seq data](#). *Bioinformatics*, 25(15), 1952-1958.

74 Calculation of Relative Risk Confidence Interval

The three options that are proposed in `riskratio()` refer to an asymptotic or large sample approach, an approximation for small sample, a resampling approach (asymptotic bootstrap, i.e. not based on percentile or bias-corrected). The former is described in Rothman’s book (as referenced in the online help), chap. 14, pp. 241-244. The latter is relatively trivial so I will skip it. The small sample approach is just an adjustment on the calculation of the estimated relative risk.

If we consider the following table of counts for subjects cross-classified according to their exposure and disease status,

	Exposed	Non-exposed	Total
Cases	a1	a0	m1
Non-case	b1	b0	m0
Total	n1	n0	N

the MLE of the risk ratio (RR), $RR = R_1/R_0$, is $RR = \frac{a_1/n_1}{a_0/n_0}$. In the *large sample approach*, a score statistic (for testing $R_1 = R_0$, or equivalently, $RR = 1$) is used, $\chi_S = \frac{a_1 - \tilde{a}_1}{\sqrt{V^{1/2}}}$, where the numerator reflects the difference between the observed and expected counts for exposed cases and $V = (m_1 n_1 m_0 n_0) / (n^2 (n - 1))$ is the variance of a_1 . Now, that’s all for computing the p -value because we know that χ_S follow a chi-square distribution. In fact, the three p -values (mid- p , Fisher exact test, and χ^2 -test) that are returned by `riskratio()` are computed in the `tab2by2.test()` function. For more information on mid- p , you can refer to

Berry and Armitage (1995). [Mid-P confidence intervals: a brief review](#). *The Statistician*, 44(4), 417-423.

Now, for computing the $100(1 - \alpha)$ CIs, this asymp-

totic approach yields an approximate SD estimate for $\ln(RR)$ of $(\frac{1}{a_1} - \frac{1}{n_1} + \frac{1}{a_0} - \frac{1}{n_0})^{1/2}$, and the Wald limits are found to be $\exp(\ln(RR)) \pm Z_c \text{SD}(\ln(RR))$, where Z_c is the corresponding quantile for the standard normal distribution.

The *small sample approach* makes use of an adjusted RR estimator: we just replace the denominator a_0/n_0 by $(a_0 + 1)/(n_0 + 1)$.

As to how to decide whether we should rely on the large or small sample approach, it is mainly by checking expected cell frequencies; for the χ_S to be valid, \tilde{a}_1 , $m_1 - \tilde{a}_1$, $n_1 - \tilde{a}_1$ and $m_0 - n_1 + \tilde{a}_1$ should be > 5 .

Working through the example of Rothman (p. 243),

```
sel <- matrix(c(2,9,12,7), 2, 2)
riskratio(sel, rev="row")
```

which yields

```
$data
      Outcome
Predictor Disease1 Disease2 Total
Exposed2      9         7     16
Exposed1      2        12     14
Total        11        19     30

$measure
      risk ratio with 95% C.I.
Predictor estimate lower upper
Exposed2 1.000000    NA     NA
Exposed1 1.959184 1.080254 3.553240

$.value
      two-sided
Predictor midp.exact fisher.exact chi.square
Exposed2      NA         NA         NA
Exposed1 0.02332167 0.02588706 0.01733469

$correction
[1] FALSE

attr(,"method")
[1] "Unconditional MLE & normal approximation (Wald) CI"
```

By hand, we would get $RR = (12/14)/(7/16) = 1.96$, $\tilde{a}_1 = 19 \times 14/30 = 8.87$, $V = (8.87 \times 11 \times 16)/(30 \times (30 - 1)) = 1.79$, $\chi_S = (12 - 8.87)/\sqrt{1.79} = 2.34$, $\text{SD}(\ln(RR)) = (1/12 - 1/14 + 1/7 - 1/16)^{1/2} = 0.304$, $95\% \text{CIs} = \exp(\ln(1.96) \pm 1.645 \times 0.304) = [1.2; 3.2]$ (rounded).

The following partition of the test statistic

1. Miettinen and *analysis of two* 213-226.
2. Becker (1989) *likelihood and relative risk in Medicine*, 8
3. Tian, Tang, and *intervals for the* *Statistics in Medicine*
4. Walter and Cheng *point estimation of contingency tables*

Notes

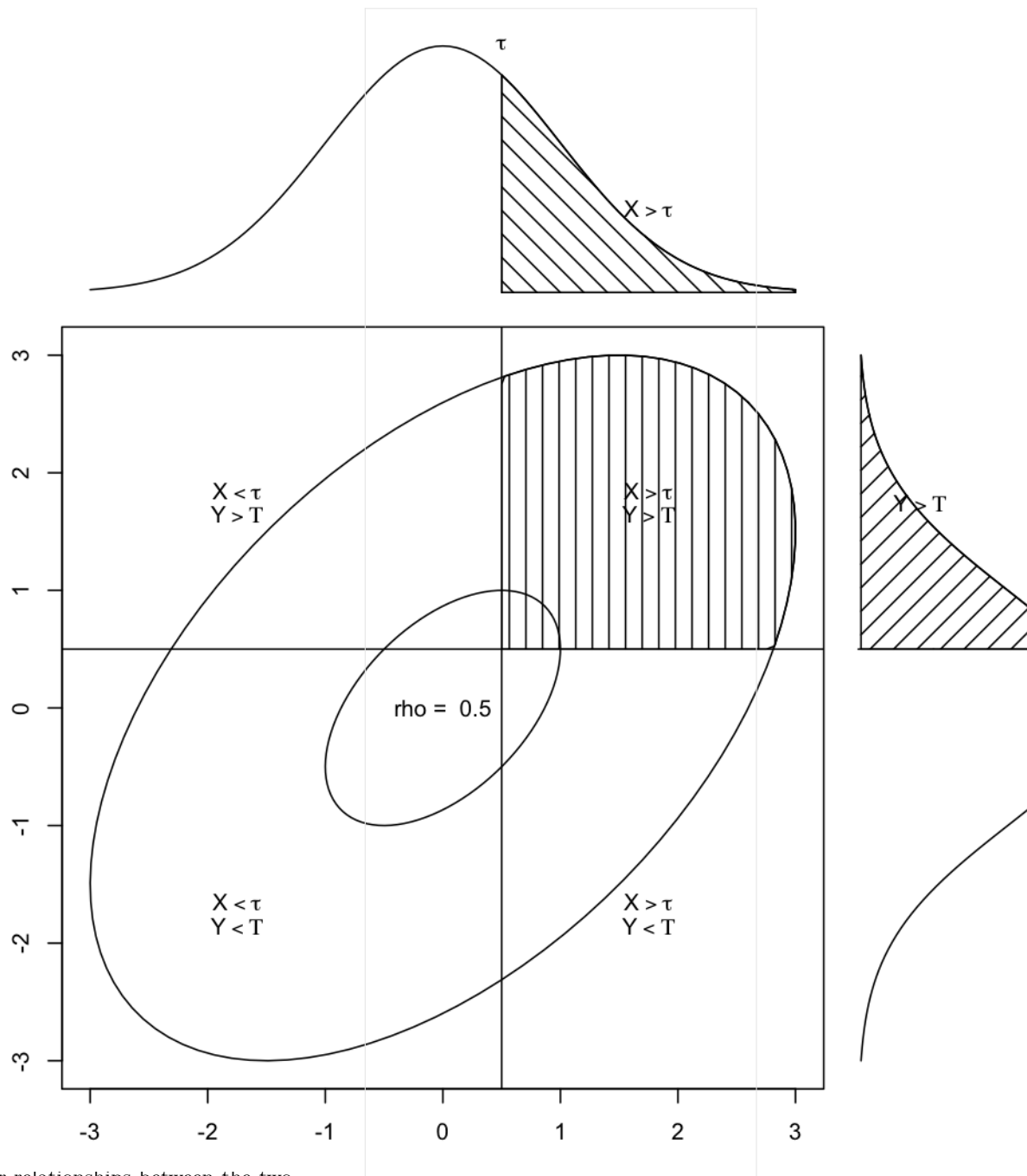
1. As far as I know, there is no online help).
2. Alan Agresti (1996)

75 Different and Pearson

My best bet is that between your categories.

If you assume that the individual locations (continuous) trait, then ok, provided the bivariate density, are set symmetric (0.5):

In this case, the estimate the true linear relationships between the two latent traits, especially in the mid-range of the correlation metric. On the other hand, when the cut-offs are clearly asymmetrical on both continuous variables, the tetrachoric correlation will generally overestimate the true relationship. The following picture illustrates the ideal case.



```
library(polycor)
set.seed(101)
n <- 500
rho <- seq(0,1,length=500)
pc1 <- pc2 <- tc <- numeric(500)

for (i in 1:500) {
```



```

data <- rmvnorm(n
1), 2, 2))
x <- data[,1]; y
xb <- ifelse(x>=m
1, 0)
pc1[i] <- cor(x,
pc2[i] <- cor(xb,
tc[i] <- polychor
}

```

```

plot(pc1, pc2, cex=
ylab="Observed
lines(lowess(pc1, p
abline(0, 1, col="l
points(pc1, tc, cex
lines(lowess(pc1, t
legend("topleft", c
lty=1, bty="n")

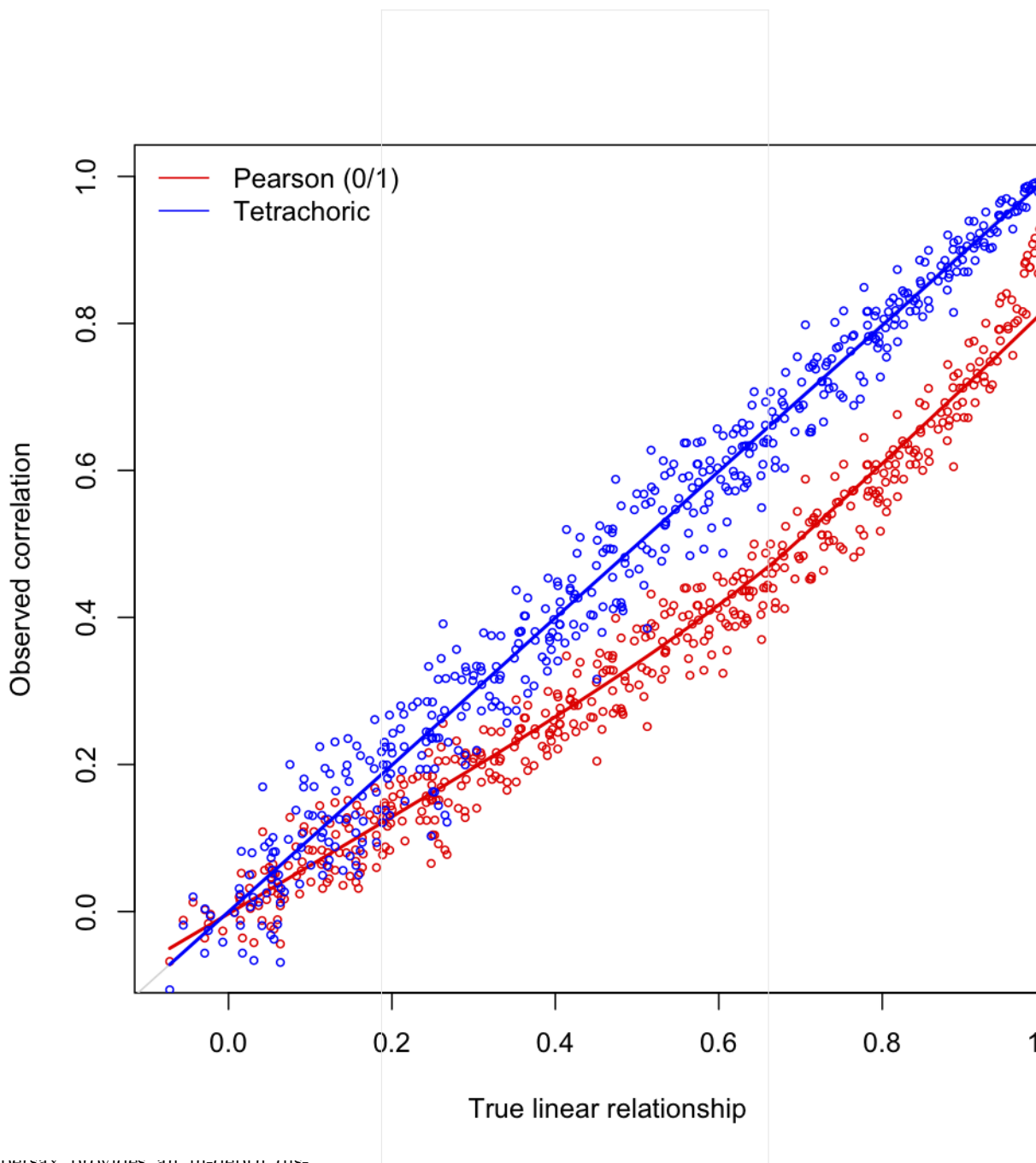
```

Now, you can pla
and see what ha
largely departs fr
of x and y .

To complement @
efficient is genera
variables (no hyp
erating process a
correlation in thi
then to factor out
in such a way bec
ingless.

To avoid this pro
item response mo
model (in this cas
off, since it is esti
like Mokken scali
sume monotonicit
tional form relatir
and the outcome
the item). Howe
pain and would r
ture in your corre
afterwards.

Finally, John Uebersax provides an in-depen dis-
cussion of tetrachoric correlation in relation of
latent trait modeling, see [Introduction to the
Tetrachoric and Polychoric Correlation Coefficients](#).
Also, Nunnally discussed a long ago of the advan-
tages/disadvantages of relying on Pearson vs. Tetra-
choric correlation coefficients in Factor Analysis, see
e.g. pp. 570-573 (3rd ed.).



References

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categorization. *Psychological Bulletin*, 105, 467-477.

3. Edwards, J.H. and Edwards, A.W.F. (1984). Approximating the tetrachoric correlation coefficient. *Biometrics*, 40, 563.
4. Castellan, N.J. (1966). On the estimation of the tetrachoric correlation coefficient. *Psychometrika*, 31(1), 67-73.
5. Fitzgerald, P., Knuiman, M.W., Divitini, M.L., and Bartholomew, H.C. (1999). Effect of dichotomising a continuous variable on the assessment of familial aggregation: an empirical study using body mass index data from the Busselton Health Study. *J. Epidemiol. Biostat.*, 4(4), 321-327.
6. Nunnally, J.C. and Bernstein, I.H. (1994). *Psychometric Theory* (Third ed.). McGraw-Hill.

76 How to capture R text+image output into one file (html, doc, pdf etc)?

Well, I just remind that I was using [AsciiDoc](#) for short reporting or editing webpage. Now there's an [R plugin](#) (`ascii` on CRAN), which allows to embed R code into an asciidoc document. The syntax is quite similar to Markdown or Textile, so you'll learn it very fast.

Output are (X)HTML, Docbook, LaTeX, and of course PDF through one of the last two backends.

Unfortunately, I don't think you can wrap all your code into a single statement. However, it supports a large number of R objects, see below.

```
> methods(ascii)
[1] ascii.anova*          ascii.aov*
ascii.cast_df*
[5] ascii.character*      ascii.coxph*
ascii.data.frame*
[9] ascii.default*        ascii.density*
ascii.describe.single*
[13] ascii.factor*         ascii.freqtable*
ascii.glm*
[17] ascii.htest*          ascii.integer*
ascii.lm*
[21] ascii.matrix*         ascii.meanscomp*
ascii.packageDescription*
```

```
[25] ascii.prcomp*          ascii.sessionInfo*    ascii.simple.list*
ascii.smooth.spline*
[29] ascii.summary.aov*     ascii.summary.aovlist* ascii.summary.glm*
ascii.summary.lm*
[33] ascii.summary.prcomp*  ascii.summary.survfit* ascii.summary.table*
ascii.survdiff*
[37] ascii.survfit*         ascii.table*          ascii.ts*
ascii.zoo*
```

Non-visible functions are asterisked

77 Mixed effects log-linear models

Log-linear or Poisson model are part of generalized linear models. Look at the [lme4](#) package which allows for mixed-effects modeling, with `family=poisson()`.

Here is an example of use:

```
> data(homerun, package="Zelig")
> with(homerun, table(homeruns, month))
      month
homeruns April August July June March May September
0      36      36  40  26      1  33      30
1      13      17  11  21      1  14      14
2       0       3   3   3      0   3       6
3       1       0   0   1      0   1       0

> library(lme4)
> mod <- glmer(homeruns ~ player + (player - 1 | month), data=homerun,
family=poisson())
> summary(mod)

Generalized linear mixed model fit by the Laplace approximation
Formula: homeruns ~ player + (player - 1 | month)

Data: homerun
      AIC      BIC logLik deviance
305.8 324.6 -147.9   295.8

Random effects:
Groups Name          Variance Std.Dev.  Corr
month  playerMcGwire 7.9688e-10 2.8229e-05
      playerSosa    6.6633e-02 2.5813e-01 0.000

Number of obs: 314, groups: month, 7
ascii.aovlist*

Fixed effects:
ascii.CrossTable*
      Estimate Std. Error z value Pr(>|z|)
(Intercept) -0.7949      0.1195  -6.651 2.91e-11 ***
playerSosa   -0.1252      0.2020  -0.620 0.535
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

ascii.list*

Correlation of Fixed Effects:
ascii.numeric*(Intr)
playerSosa -0.592
```

The scale parameter (useful to check for possible overdispersion) is available through the following slot:

```
summary(mod)$sigma
```

(The equivalent for usual GLM would be `summary(glm(...))$dispersion`). good discussion on Medstats, [Documenting analyses and data edits Options](#)

More information about mixed-effects modeling as implemented in `lme4` can be found on R-forge, in the [Mixed-effects models project](#).

The `gamm4` package may also be of interest as it allows to fit generalized additive mixed models.

78 A good way to show lots of data graphically

[Mondrian](#) provides interactive features and handles quite large data sets (it's in Java, though).

[Paraview](#) includes 2D/3D viz. features.

79 How to efficiently manage a statistical analysis project?

I am compiling a quick series of guidelines I found on [SO](#) (as suggested by @Shane), [Biostar](#) (hereafter, BS), and this SE. I tried my best to acknowledge ownership for each item, and to select first or highly upvoted answer. I also added things of my own, and flagged items that are specific to the [R] environment.

Data management

- create a project structure for keeping all things at the right place (data, code, figures, etc., [giovanni](#) /BS)
- never modify raw data files (ideally, they should be read-only), copy/rename to new ones when making transformations, cleaning, etc.
- check data consistency ([whuber](#) /SE)

Coding

- organize source code in logical units or building blocks ([Josh Reich/hadley/ars](#) /SO; [giovanni/Khader Shameer](#) /BS)

- separate source code from editing stuff, especially for large project – partly overlapping with previous item and reporting
- document everything, with e.g. [R]oxygen ([Shane](#) /SO) or consistent self-annotation in the [source files](#). good discussion on Medstats, [Documenting analyses and data edits Options](#)
- [R] custom functions can be put in a dedicated file (that can be sourced when necessary), in a new environment (so as to avoid populating the top-level namespace, [Brendan OConnor](#) /SO), or a package ([Dirk Eddelbuettel/Shane](#) /SO)

Analysis

- don't forget to set/record the seed you used when calling RNG or stochastic algorithms (e.g. k-means)
- for Monte Carlo studies, it may be interesting to store specs/parameters in a separate file ([sumatra](#) may be a good candidate, [giovanni](#) /BS)
- don't limit yourself to one plot per variable, use multivariate (Trellis) displays and interactive visualization tools (e.g. GGobi)

Versioning

- use some kind of CVS for easy tracking/export, e.g. Git ([Sharpie/VonC/JD Long](#) /SO) – this follows from nice questions asked by @Jeromy and @Tal
- backup everything, on a regular basis ([Sharpie/JD Long](#) /SO)
- keep a log of your ideas, or rely on an issue tracker, like [ditz](#) ([giovanni](#) /BS) – partly redundant with the previous item since it is available in Git

Editing/Reporting

- [R] Sweave ([Matt Parker](#) /SO)
- [R] brew ([Shane](#) /SO)
- [R] [R2HTML](#) or [ascii](#)

As a side note, Hadley Wickham offers a comprehensive overview of [R project management](#), including

reproducible exemplification and an unified philosophy of data.

80 Is adjusting p-values in a multiple regression for multiple comparisons a good idea?

It seems your question more generally addresses the problem of identifying good predictors. In this case, you should consider using some kind of penalized regression (methods dealing with variable or **feature selection** are relevant too), with e.g. L1, L2 (or a combination thereof, the so-called **elasticnet**) penalties (look for related questions on this site, or the **R penalized** and **elasticnet** package, among others).

Now, about correcting p-values for your regression coefficients (or equivalently your partial correlation coefficients) to protect against over-optimism (e.g. with Bonferroni or, better, step-down methods), it seems this would only be relevant if you are considering one model and seek those predictors that contribute a significant part of explained variance, that is if you don't perform model selection (with stepwise selection, or hierarchical testing). This article may be a good start: **Bonferroni Adjustments in Tests for Regression Coefficients**. Be aware that such correction won't protect you against multicollinearity issue, which affects the reported p-values.

Given your data, I would recommend using some kind of iterative model selection techniques. In R for instance, the **stepAIC** function allows to perform stepwise model selection by exact AIC. You can also estimate the relative importance of your predictors based on their contribution to R^2 using bootstrap (see the **relaimpo** package). I think that reporting effect size measure or % of explained variance are more informative than p-value, especially in a confirmatory model.

It should be noted that stepwise approaches have also their drawbacks (e.g., Wald tests are not adapted to conditional hypothesis as induced by the stepwise procedure), or as indicated by Frank Harrell on **R mailing**, "stepwise variable selection based on AIC has all the problems of stepwise variable selection based on P-values. AIC is just a restatement of the P-Value" (but AIC remains useful if the set of predictors is already defined); a related question – **Is a variable significant in a linear regression model?** – raised interesting comments (**@Rob**, among others) about the use of AIC for variable selection. I

append a couple of references at the end (including papers kindly provided by **@Stephan**); there is also a lot of other references on **P.Mean**.

Frank Harrell authored a book on **Regression Modeling Strategy** which includes a lot of discussion and advices around this problem (§4.3, pp. 56-60). He also developed efficient R routines to deal with generalized linear models (See the **Design** or **rms** packages). So, I think you definitely have to take a look at it (his **handouts** are available on his homepage).

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models. *International Conference on Multiple Comparison Procedures*. – see `step.adj()` in the R `someMTP` package.

9. Wiegand, RE (2010). **Performance of using multiple stepwise algorithms for variable selection.** *Statistics in Medicine*, 29(15), 1647–1659.
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13. Flom, PL and Cassell, DL (2007). **Stopping Stepwise: Why stepwise and similar selection methods are bad, and what you should use.** *NE-SUG 2007 Proceedings*.
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81 Do some of you use Google Docs spreadsheet to conduct and share your statistical work with others?

As an enthusiast user of R, bash, Python, asciidoc, (La)TeX, open source software or any un*x tools, I cannot provide an objective answer. Moreover, as I often argue against the use of MS Excel or spreadsheet of any kind (well, you see your data, or part of it, but what else?), I would not contribute positively to the debate. I’m not the only one, e.g.

- **Spreadsheet Addiction**, from P. Burns.
- **MS Excel’s precision and accuracy**, a post on the 2004 R mailing-list

- L. Knusel, **On the accuracy of statistical distributions in Microsoft Excel 97**, *Computational Statistics & Data Analysis*, 26: 375–377, 1998. ([pdf](#))
- B.D. McCullough & B. Wilson, **On the accuracy of statistical procedures in Microsoft Excel 2000 and Excel XP**, *Computational Statistics & Data Analysis*, 40: 713–721, 2002.
- M. Altman, J. Gill & M.P. McDonald, *Numerical Issues in Statistical Computing for the Social Scientist*, Wiley, 2004. [e.g., pp. 12–14]

A colleague of mine lost all his macros because of the lack of backward compatibility, etc. Another colleague tried to import genetics data (around 700 subjects genotyped on 800,000 markers, 120 Mo), just to “look at them”. Excel failed, Notepad gave up too... I am able to “look at them” with vi, and quickly reformat the data with some sed/awk or perl script. So I think there are different levels to consider when discussing about the usefulness of spreadsheets. Either you work on small data sets, and only want to apply elementary statistical stuff and maybe it’s fine. Then, it’s up to you to trust the results, or you can always ask for the source code, but maybe it would be simpler to do a quick test of all inline procedures with the **NIST benchmark**. I don’t think it corresponds to a good way of doing statistics simply because this is not a *true* statistical software (IMHO), although as an update of the aforementioned list, newer versions of MS Excel seem to have demonstrated improvements in its accuracy for statistical analyses, see Keeling and Pavur, **A comparative study of the reliability of nine statistical software packages** (*CSDA* 2007 51: 3811).

Still, about one paper out of 10 or 20 (in biomedicine, psychology, psychiatry) includes graphics made with Excel, sometimes without removing the gray background, the horizontal black line or the automatic legend (Andrew Gelman and Hadley Wickham are certainly as happy as me when seeing it). But more generally, it tends to be the most used “software” according to a **recent poll** on FlowingData, which reminds me of an old talk of Brian Ripley (who co-authored the MASS R package, and wrote an excellent book on pattern recognition, among others):

Let’s not kid ourselves: the most widely used piece of software for statistics is Excel (B. Ripley via Jan De Leeuw), <http://bit.ly/dB5K6r>

Now, if you feel it provides you with a quick and eas-

ier way to get your statistics done, why not? The problem is that there are still things that cannot be done (or at least, it's rather tricky) in such an environment. I think of bootstrap, permutation, multivariate exploratory data analysis, to name a few. Unless you are very proficient in VBA (which is neither a scripting nor a programming language), I am inclined to think that even minor operations on data are better handled under R (or Matlab, or Python, providing you get the right tool for dealing with e.g. so-called data.frame). Above all, I think Excel does not promote very good practices for the data analyst (but it also applies to any “cliquodrome”, see the discussion on Medstats about the need to maintain a record of data processing, [Documenting analyses and data edits](#)), and I found this post on [Practical Stats](#) relatively illustrative of some of Excel pitfalls. Still, it applies to Excel, I don't know how it translates to GDocs.

About sharing your work, I tend to think that [Github](#) (or [Gist](#) for source code) or [Dropbox](#) (although EULA might discourage some people) are very good options (revision history, grant management if needed, etc.). I cannot encourage the use of a software which basically store your data in a binary format. I know it can be imported in R, Matlab, Stata, SPSS, but to my opinion:

- data should definitively be in a text format, that can be read by another statistical software;
- analysis should be reproducible, meaning you should provide a complete script for your analysis and it should run (we approach the ideal case near here. . .) on another operating system at any time;
- your own statistical software should implement acknowledged algorithms and there should be an easy way to update it to reflect current best practices in statistical modeling;
- the sharing system you choose should include versioning and collaborative facilities.

That's it.

82 How to teach students who fear statistics?

Not very much about how to deal with students' fear, but Andrew Gelman wrote an excellent book, [Teaching Statistics, a bag of tricks](#) (there's also some

[slides](#)).

I like introducing a course by talking about randomness, elementary probability as found in games, causal association, permutation tests (because parametric tests provide good approximation to them :).

I just put an example that I like to show to students. This is from Phillip Good, in his book [Permutation, Parametric, and Bootstrap Tests of Hypotheses](#) (Springer, 2005 3rd ed.), where he introduces the general strategy of testing or decision making about statistical hypothesis and how to carry out a very simple and exact permutation test to solve the following problem.

Shortly after I received my doctorate in statistics, I decided that if I really wanted to help bench scientists apply statistics I ought to become a scientist myself. So I went back to school to learn physiology and aging in cells raised in petri dishes.

I soon learned there was a great deal more to an experiment than the random assignment of subjects to treatments. In general, 90% of experimental effort was spent mastering various arcane laboratory techniques, another 9% in developing new techniques to span the gap between what had been done and what I really wanted to do, and a mere 1% on the experiment itself. But the moment of truth came finally—it had to if I were to publish and not perish—and I succeeded in cloning human diploid fibroblasts in eight culture dishes: Four of these dishes were filled with a conventional nutrient solution and four held an experimental “life-extending” solution to which vitamin E had been added.

I waited three weeks with fingers crossed that there was no contamination of the cell cultures, but at the end of this test period three dishes of each type had survived. My technician and I transplanted the cells, let them grow for 24 hours in contact with a radioactive label, and then fixed and stained them before covering them with a photographic emulsion.

Ten days passed and we were ready to examine the autoradiographs. Two years had elapsed since I first envisioned this experiment and now the results were in: I had the six numbers I needed.

“I’ve lost the labels,” my technician said as she handed me the results. This was a dire situation. Without the labels, I had no way of knowing which cell cultures had been treated with vitamin E and which had not.

83 Interpreting 2D Correspondence Analysis Plots

First, there are different ways to construct so-called **biplots** in the case of correspondence analysis. In all cases, the basic idea is to find a way to show the best 2D approximation of the “distances” between row cells and column cells. In other words, we seek a hierarchy (we also speak of “ordination”) of the relationships between rows and columns of a contingency table.

Very briefly, CA decomposes the chi-square statistic associated with the two-way table into orthogonal factors that maximize the separation between row and column scores (i.e. the frequencies computed from the table of profiles). Here, you see that there is some connection with PCA but the measure of variance (or the metric) retained in CA is the χ^2 , which only depends on column profiles (As it tends to give more importance to modalities that have large marginal values, we can also re-weight the initial data, but this is another story).

Here is a more detailed answer. The implementation that is proposed in the `corresp()` function (in **MASS**) follows from a view of CA as an SVD decomposition of dummy coded matrices representing the rows and columns (such that $R^t C = N$, with N the total sample). This is in light with canonical correlation analysis. In contrast, the French school of data analysis considers CA as a variant of the PCA, where you seek the directions that maximize the “inertia” in the data cloud. This is done by diagonalizing the inertia matrix computed from the centered and scaled (by marginals frequencies) two-way table, and expressing row and column profiles in this new coordinate system.

If you consider a table with $i = 1, \dots, I$ rows, and $j = 1, \dots, J$ columns, each row is weighted by its corresponding marginal sum which yields a series of conditional frequencies associated to each row: $f_{j|i} = n_{ij}/n_{i.}$. The marginal column is called the *mean profile* (for rows). This gives us a vector of coordinates, also called a *profile* (by row). For the column, we have $f_{i|j} = n_{ij}/n_{.j}$. In both cases, we will consider the I row profiles (associated to their

weight $f_{i.}$) as individuals in the column space, and the J column profiles (associated to their weight $f_{.j}$) as individuals in the row space. The metric used to compute the proximity between any two individuals is the χ^2 distance. For instance, between two rows i and i' , we have

$$d_{\chi^2}^2(i, i') = \sum_{j=1}^J \frac{n_{.j}}{n} \left(\frac{n_{ij}}{n_{i.}} - \frac{n_{i'j}}{n_{i'.}} \right)^2$$

You may also see the link with the χ^2 statistic by noting that it is simply the distance between observed and expected counts, where expected counts (under H_0 , independence of the two variables) are computed as $n_{i.} \times n_{.j} / n$ for each cell (i, j) . If the two variables were to be independent, the row profiles would be all equal, and identical to the corresponding marginal profile. In other words, when there is independence, your contingency table is entirely determined by its margins.

If you realize an PCA on the row profiles (viewed as individuals), replacing the euclidean distance by the χ^2 distance, then you get your CA. The first principal axis is the line that is the closest to all points, and the corresponding eigenvalue is the inertia explained by this dimension. You can do the same with the column profiles. It can be shown that there is a symmetry between the two approaches, and more specifically that the principal components (PC) for the column profiles are associated to the same eigenvalues than the PCs for the row profiles. What is shown on a biplot is the coordinates of the individuals in this new coordinate system, although the individuals are represented in a separate factorial space. Provided each individual/modality is well represented in its factorial space (you can look at the \cos^2 of the modality with the 1st principal axis, which is a measure of the correlation/association), you can even interpret the proximity between elements i and j of your contingency table (as can be done by looking at the residuals of your χ^2 test of independence, e.g. `chisq.test(tab)$expected - chisq.test(tab)$observed`).

The total inertia of your CA (= the sum of eigenvalues) is the χ^2 statistic divided by n (which is Pearson’s ϕ^2).

Actually, there are several packages that may provide you with enhanced CAs compared to the function available in the **MASS** package: `ade4`, `FactoMineR`, `anacor`, and `ca`.

The latest is the one that was used for your partic-

ular illustration, and a paper was published in the Journal of Statistical Software that explains most of its functionalities: [Correspondence Analysis in R, with Two- and Three-dimensional Graphics: The ca Package](#).

So, your example on eye/hair colors can be reproduced in many ways:

```
data(HairEyeColor)
tab <- apply(HairEyeColor, c(1, 2), sum) # aggregate on gender
tab

library(MASS)
plot(corresp(tab, nf=2))
corresp(tab, nf=2)

library(ca)
plot(ca(tab))
summary(ca(tab, nd=2))

library(FactoMineR)
CA(tab)
CA(tab, graph=FALSE)$eig # == summary(ca(tab))$scree[, "values"]
CA(tab, graph=FALSE)$row$contrib

library(ade4)
scatter(dudi.coa(tab, scannf=FALSE, nf=2))
```

In all cases, what we read in the resulting biplot is basically (I limit my interpretation to the 1st axis which explained most of the inertia):

- the first axis highlights the clear opposition between light and dark hair color, and between blue and brown eyes;
- people with blond hair tend to also have blue eyes, and people with black hair tend to have brown eyes.

There is a lot of additional resources on data analysis on the [bioinformatics lab](#) from Lyon, in France. This is mostly in French, but I think it would not be too much of a problem for you. The following two handouts should be interesting as a first start:

- [Initiation à l'analyse factorielle des correspondances](#)
- [Pratique de l'analyse des correspondances](#)

Finally, when you consider a full disjunctive (dummy) coding of k variables, you get the *multiple correspondence analysis*.

84 Regularization and Mean Estimation

Ridge regression (Hoerl and Kennard, 1988) was initially developed to overcome singularities when inverting $X^t X$ (by adding λ to its diagonal elements). Thus, the *regularization* in this case consists in working with a vc matrix $(X^t X - \lambda I)^{-1}$. This L2 penalization leads to “better” predictions than with usual OLS by optimizing the compromise between bias and variance (shrinkage), but it suffers from considering all coefficients in the model. The regression coefficients are found to be

$$\hat{\beta} = \underset{\beta}{\operatorname{argmin}} \|Y - X\beta\|^2 + \lambda \|\beta\|^2$$

with $\|\beta\|^2 = \sum_{j=1}^p \beta_j^2$ (L2-norm).

From a bayesian perspective, you can consider that the β 's must be small and plug them into a prior distribution. The likelihood $\ell(y, X, \beta, \sigma^2)$ can thus be weighted by the prior probability for β (assumed i.i.d. with zero mean and variance τ^2), and the posterior is found to be

$$f(\beta|y, X, \sigma^2, \tau^2) = (y - \hat{\beta}^t X)^t (y - \hat{\beta}^t X) + \frac{\sigma^2}{\tau^2} \hat{\beta}^t \hat{\beta}$$

where σ^2 is the variance of your y 's. It follows that this density is the opposite of the residual sum of squares that is to be minimized in the Ridge framework, after setting $\lambda = \sigma^2/\tau^2$.

The bayesian estimator for $\hat{\beta}$ is thus the same as the OLS one when considering the Ridge loss function with a prior variance τ^2 . More details can be found in *The Elements of Statistical Learning* from Hastie, Tibshirani, and Friedman (§3.4.3, p.60 in the 1st ed.). The [second edition](#) is also available for free.

85 Clustering genes in a time course experiment

In complement to @mbq's response ([Mfuzz](#) looks fine), I'll just put some references (PDFs) about clustering of time-course gene expression data:

1. Futschik, ME and Charlisle, B (2005). [Noise robust clustering of gene expression time-course data](#). *Journal of Bioinformatics and Computational Biology*, 3(4), 965-988.

2. Luan, Y and Li, H (2003). [Clustering of time-course gene expression data using a mixed-effects model with B-splines](#). *Bioinformatics*, 19(4), 474-482.
3. Tai YC and Speed, TP (2006). [A multivariate empirical Bayes statistic for replicated microarray time course data](#). *The Annals of Statistics*, 34, 2387–2412.
4. Schliep, A, Steinhoff, C, and Schönhuth, A (2004). [Robust inference of groups in gene expression time-courses using mixtures of HMMs](#). *Bioinformatics*, 20(1), i283-i228.
5. Costa, IG, de Carvalho, F, and de Souto, MCP (2004). [Comparative analysis of clustering methods for gene expression time course data](#). *Genetics and Molecular Biology*, 27(4), 623-631.
6. Inoue, LYT, Neira, M, Nelson, C, Gleave, M, and Etzioni, R (2006). [Cluster-based network model for time-course gene expression data](#). *Biostatistics*, 8(3), 507-525.
7. Phang, TL, Neville, MC, Rudolph, M, and Hunter, L (2003). [Trajectory Clustering: A Non-Parametric Method for Grouping Gene Expression Time Courses with Applications to Mammary Development](#). *Pacific Symposium on Biocomputing*, 8, 351-362.

Did you try the [timecourse](#) package (as suggested by @csgillespie in his [handout](#))?

86 Building a Statistics Library, with Knapsack Constraint

1. Harrell, FE. [Regression Modeling Strategies](#) (Springer, 2010, 2nd ed.)
2. Izenman, AJ. [Modern Multivariate Statistical Techniques: Regression, Classification, and Manifold Learning](#) (Springer, 2008)

You should have money left to print part of *The Handbook of Computational Statistics* (Gentle et al., Springer 2004) and *The Elements of Statistical Learning* (Hastie et al., Springer 2009 2nd ed.) that are circulating on the web. As the latter mostly covers the same topics than Izenman’s book (as pointed by @kwak), either may be replaced by one of the *Handbook of Statistics* published by North-Holland,

depending on your field of interests.

87 Resources for Learning Markov Chain and Hidden Markov Models

Here are some tutorials (available as PDFs):

1. Dugad and Desai, [A tutorial on hidden markov models](#)
2. Valeria De Fonzo¹, Filippo Aluffi-Pentini² and Valerio Parisi (2007). [Hidden Markov Models in Bioinformatics](#). *Current Bioinformatics*, 2, 49-61.
3. Smith, K. [Hidden Markov Models in Bioinformatics with Application to Gene Finding in Human DNA](#)

Also take a look at [Bioconductor](#) tutorials.

I assume you want free resources; otherwise, [Bioinformatics](#) from Polanski and Kimmel (Springer, 2007) provides a nice overview (§2.8-2.9) and applications (Part II).

88 How to cope with exploratory data analysis and data dredging in small-sample studies?

I just drop some references about **data dredging** and **clinical studies** for the interested reader. This is intended to extend @onestop’s fine answer. I tried to avoid articles focusing only on multiple comparisons or design issues, although studies with multiple endpoints continue to present challenging and controversial discussions (long after Rothman’s claims about [useless adjustments](#), *Epidemiology* 1990, 1: 43-46; or see Feise’s review in *BMC Medical Research Methodology* 2002, 2:8).

My understanding is that, although I talked about *exploratory data analysis*, my question more generally addresses the use of data mining, with its potential pitfalls, in parallel to hypothesis-driven testing.

1. Koh, HC and Tan, G (2005). [Data Mining Applications in Healthcare](#). *Journal of Healthcare Information Management*, 19(2), 64-72.
2. Ioannidis, JPA (2005). [Why most published research findings are false](#). *PLoS Medicine*, 2(8),

e124.

3. Anderson, DR, Link, WA, Johnson, DH, and Burnham, KP (2001). [Suggestions for Presenting the Results of Data Analysis](#). *The Journal of Wildlife Management*, 65(3), 373-378. – this echoes @onestop’s comment about the fact that we have to acknowledge the data-driven exploration/modeling beyond the initial set of hypotheses
4. Michels, KB and Rosner, BA (1996). [Data trawling: to fish or not to fish](#). *Lancet*, 348, 1152-1153.
5. Lord, SJ, Gebiski, VJ, and Keech, AC (2004). [Multiple analyses in clinical trials: sound science or data dredging?](#). *The Medical Journal of Australia*, 181(8), 452-454.
6. Smith, GD and Ebrahim, S (2002). [Data dredging, bias, or confounding](#). *BMJ*, 325, 1437-1438.
7. Afshartous, D and Wolf, M (2007). [Avoiding ‘data snooping’ in multilevel and mixed effects models](#). *Journal of the Royal Statistical Society A*, 170(4), 1035–1059
8. Anderson, DR, Burnham, KP, Gould, WR, and Cherry, S (2001). [Concerns about finding effects that are actually spurious](#). *Wildlife Society Bulletin*, 29(1), 311-316.

89 Is it possible to do time-series clustering based on curve shape?

Several directions for analyzing longitudinal data were discussed in the link provided by @Jeromy, so I would suggest you to read them carefully, especially those on functional data analysis. Try googling for “Functional Clustering of Longitudinal Data”, or the PACE Matlab toolbox which is specifically concerned with model-based clustering of irregularly sampled trajectories (Peng and Müller, [Distance-based clustering of sparsely observed stochastic processes, with applications to online auctions](#), *Annals of Applied Statistics* 2008 2: 1056). I can imagine that there may be a good statistical framework for financial time series, but I don’t know about that.

The `kml` package basically relies on k-means, work-

ing (by default) on euclidean distances between the t measurements observed on n individuals. What is called a *trajectory* is just the series of observed values for individual i , $y_i = (y_{i1}, y_{i2}, \dots, y_{it})$, and $d(y_i, y_j) = \sqrt{t^{-1} \sum_{k=1}^t (y_{ik} - y_{jk})^2}$. Missing data are handled through a slight modification of the preceding distance measure (Gower adjustment) associated to a nearest neighbor-like imputation scheme (for computing Calinski criterion). As I don’t represent myself what you real data would look like, I cannot say if it will work. At least, it work with longitudinal growth curves, “polynomial” shape, but I doubt it will allow you to detect very specific patterns (like local minima/maxima at specific time-points with time-points differing between clusters, by a translation for example). If you are interested in clustering possibly misaligned curves, then you definitively have to look at other solutions; [Functional clustering and alignment](#), from Sangalli et al., and references therein may provide a good starting point.

Below, I show you some code that may help to experiment with it (my seed is generally set at 101, if you want to reproduce the results). Basically, for using `kml` you just have to construct a `clusterizLongData` object (an `id` number for the first column, and the t measurements in the next columns).

```
library(lattice)
xyplot(var0 ~ date, data=test.data, groups=store, type=c("l","g"))

tw <- reshape(test.data, timevar="date", idvar="store", direction="wide")
parallel(tw[, -1], horizontal.axis=F,
  scales=list(x=list(rot=45,
    at=seq(1, ncol(tw)-1, by=2),
    labels=substr(names(tw[, -1])[seq(1, ncol(tw)-1, by=2)], 6, 100),
    cex=.5)))

library(kml)
names(tw) <- c("id", paste("t", 1:(ncol(tw)-1)))
tw.cld <- as.cld(tw)
cld.res <- kml(tw.cld, nbRedrawing=5)
plot(tw.cld)
```

The next two figures are the raw simulated data and the five-cluster solution (according to Calinski criterion, also used in the `fpc` package). I don’t show the [scaled version](#).

90 R package to simulate titration data

Try using the argument `items=1` e.g.

```
dat <- Science[c(1,
fit1 <- grm(dat)
plot(fit1, items=1,
")])
```

See `help(plot.grm)`

Also, you can enable the figure by using `par(mfrow=c(1,4))`

```
opar <- par(mfrow=c(1,4))
for (i in 1:4) {
  plot(fit1, items=i,
  par(opar)
}
```

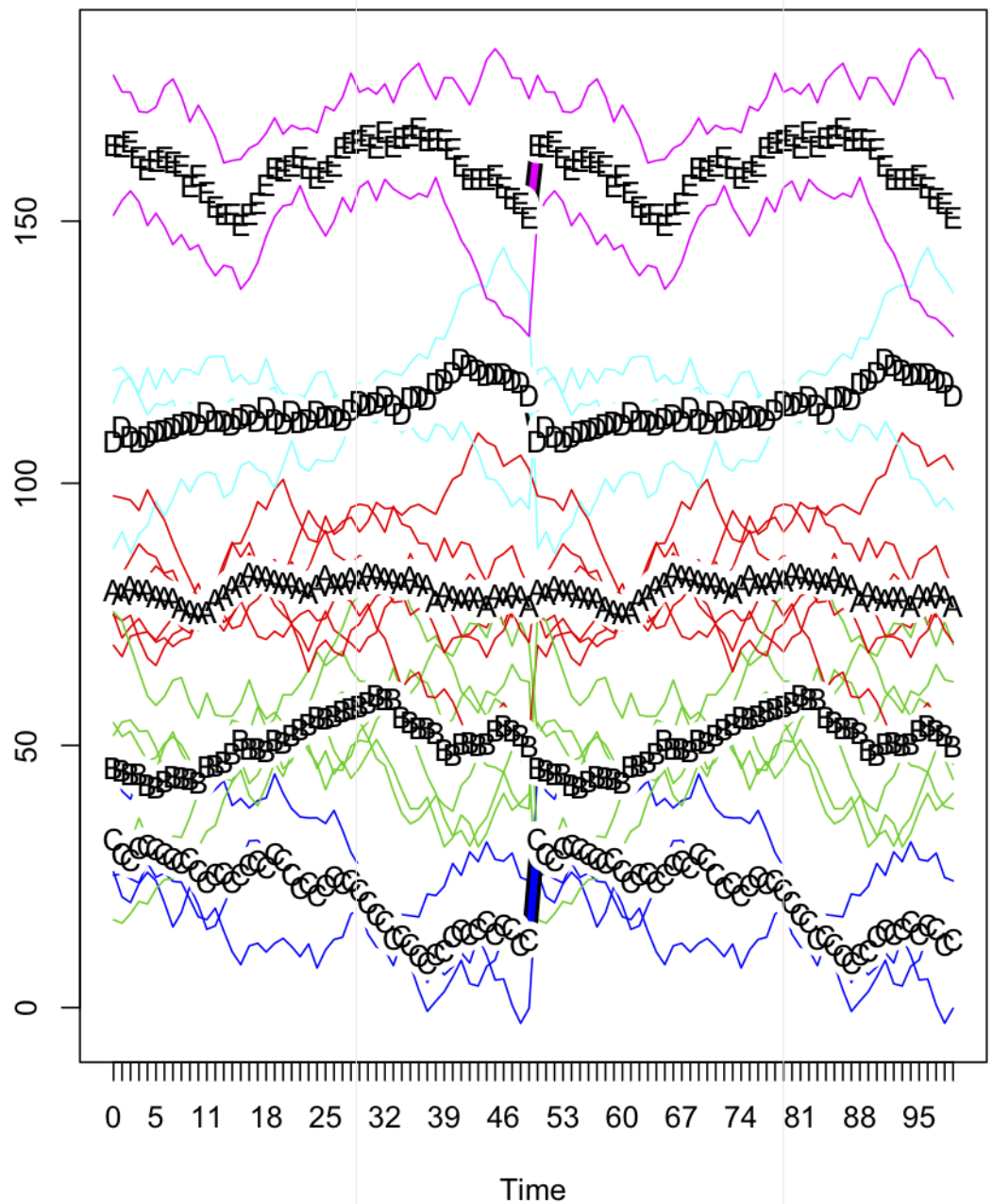
91 Is there a deal with label switching in mixture models?

Gilles Celeux also discusses switching, e.g.

G. Celeux, B.
the label switching problem in
Compstat 98,
(1998).

As a complement to @darrenjw's fine answer, here are two online papers that reviewed alternative strategies:

1. Jasra et al., [Markov Chain Monte Carlo Methods and the Label Switching Problem in Bayesian Mixture Modelling](#)



2. Sperrin et al., [Probabilistic relabelling strategies for the label switching problem in Bayesian mixture models](#)

92 Do working statisticians care about the difference between frequentist and bayesian inference?

I think bayesian statistics come into play in two different contexts.

On the one hand, some researcher/statistician are definitely convinced of the “bayesian spirit” and, acknowledging the limit of the classical frequentist hypothesis framework, have decided to concentrate on bayesian thinking. Studies in experimental psychology highlighting small effect sizes or borderline statistical significance are now increasingly relying on the bayesian framework. In this respect, I like to cite some of the extensive work of Bruno Lecoutre (1-4) who contributed to developing the use of fiducial risk and bayesian (M)ANOVA. I think the fact we can readily interpret a confidence interval in terms of probabilities applied on the parameter of interest (i.e. depending on the prior distribution) is a radical turn in statistical thinking. I can also imagine that everybody is actually aware of the ever growing work of Andrew Gelman in this domain, as pointed by @Skrikant, or of the incentive given by the [International Society for Bayesian Analysis](#) to use bayesian models. Frank Harrell also provides interesting outlines of [Bayesian Methods for Clinicians](#), as applied to [RCTs](#).

On the other hand, the bayesian approach has proved successful in diagnostic medicine (5), and is often used as an ultimate alternative where traditional statistics would fail, if applicable at all. I am thinking of a psychometrical paper (6) where authors were interested in assessing the agreement between radiologists about the severity of hip fractures from a very limited data set (12 doctors x 15 radiography) and use an Item Response model for polytomous items.

Finally, a recent 45-pages paper published in *Statistics in Medicine* provides an interesting overview of the “penetrance” of bayesian modeling in biostatistics:

Ashby, D (2006). [Bayesian statistics in medicine: a 25 year review](#). *Statistics in Medicine*, 25(21), 3589-631.

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ference in ANOVA: From significance tests to Bayesian procedures. *British Journal of Mathematical and Statistical Psychology*, 36, 252-268.

2. Lecoutre B., Lecoutre M.-P., Poitevineau J. (2001). Uses, abuses and misuses of significance tests in the scientific community: Won't the Bayesian choice be unavoidable? *International Statistical Review*, 69, 399-418.
3. Lecoutre B. (2006). Isn't everyone a Bayesian?. *Indian Bayesian Society News Letter*, III, 3-9.
4. Lecoutre B. (2006). And if you were a Bayesian without knowing it? In A. Mohammad-Djafari (Ed.): *26th Workshop on Bayesian Inference and Maximum Entropy Methods in Science and Engineering*. Melville : AIP Conference Proceedings Vol. 872, 15-22.
5. Broemeling, L.D. (2007). [Bayesian Biostatistics and Diagnostic Medicine](#). Chapman and Hall/CRC.
6. Baldwin, P., Bernstein, J., and Wainer, H. (2009). Hip psychometrics. *Statistics in Medicine*, 28(17), 2277-92.

93 What are the differences between Factor Analysis and Principal Component Analysis

You are right about your first point, although in FA you generally work with both (uniqueness and communality). The choice between PCA and FA is a long-standing debate among psychometricians. I don't quite follow your points, though. Rotation of principal axes can be applied whatever the method is used to constructed latent factors. In fact, most of the times this is the VARIMAX rotation (orthogonal rotation, considering uncorrelated factors) that is used, for practical reasons (easiest interpretation, easiest scoring rules or interpretation of factor scores, etc.), although oblique rotation (e.g. PROMAX) might probably better reflect the reality (latent constructs are often correlated one each other), at least in the tradition of FA where you assume that a latent construct is really at the heart of the observed inter-correlations between your variables. The point is that PCA followed by VARIMAX rotation somewhat distorts the interpretation of the linear combinations of the original variables in the “data analysis” tradition (see the work of Michel Tenenhaus). From a psychometrical perspective,

FA models are to be preferred since they explicitly account for measurement errors, while PCA doesn't care about that. Briefly stated, using PCA you are expressing each component (factor) as a linear combination of the variables, whereas in FA these are the variables that are expressed as linear combinations of the factors (including communalities and uniqueness components, as you said).

I recommend you to read first the following discussions about this topic:

- [What are the differences between Factor Analysis and Principal Component Analysis](#)
- [On the use of oblique rotation after PCA](#) – see reference therein

94 Excel as a statistics workbench

Incidentally, a question around the use of Google spreadsheets raised contrasting (hence, interesting) opinions about that, [Do some of you use Google Docs spreadsheet to conduct and share your statistical work with others?](#)

I have in mind an older paper which didn't seem so pessimist, but it is only marginally cited in the paper you mentioned: Keeling and Pavur, [A comparative study of the reliability of nine statistical software packages](#) (CSDA 2007 51: 3811). But now, I found yours on my hard drive. There was also a special issue in 2008, see [Special section on Microsoft Excel 2007](#), and more recently in the Journal of Statistical Software: [On the Numerical Accuracy of Spreadsheets](#).

I think it is a long-standing debate, and you will find varying papers/opinions about Excel reliability for statistical computing. I think there are different levels of discussion (what kind of analysis do you plan to do, do you rely on the internal solver, are there non-linear terms that enter a given model, etc.), and sources of numerical inaccuracy might arise as the result of *proper computing errors* or *design choices* issues; this is well summarized in

M. Altman, J. Gill & M.P. McDonald, *Numerical Issues in Statistical Computing for the Social Scientist*, Wiley, 2004.

Now, for exploratory data analysis, there are various alternatives that provide enhanced visualization capabilities, multivariate and dynamic graphics, e.g.

[GGobi](#) – but see related threads on this wiki.

But, clearly the first point you made addresses another issue (IMO), namely that of using a spreadsheet to deal with large data set: it is simply not possible to import a large csv file into Excel (I'm thinking of genomic data, but it applies to other kind of high-dimensional data). It has not been built for that purpose.

95 From a statistical perspective, can one infer causality using propensity scores with an observational study?

At the beginning of an article aiming at promoting the use of PSs in epidemiology, Oakes and Church (1) cited Hernán and Robins's claims about confounding effect in epidemiology (2):

Can you guarantee that the results from your observational study are unaffected by unmeasured confounding? The only answer an epidemiologist can provide is 'no'.

This is not just to say that we cannot ensure that results from observational studies are unbiased or useless (because, as @propofol said, their results can be useful for designing RCTs), but also that PSs do certainly not offer a complete solution to this problem, or at least do not necessarily yield better results than other matching or multivariate methods (see e.g. (10)).

Propensity scores (PS) are, by construction, *probabilistic* not *causal* indicators. The choice of the covariates that enter the propensity score function is a key element for ensuring its reliability, and their weakness, as has been said, mainly stands from not controlling for unobserved confounders (which is quite likely in retrospective or [case-control](#) studies). Others factors have to be considered: (a) model misspecification will impact direct effect estimates (not really more than in the OLS case, though), (b) there may be missing data at the level of the covariates, (c) PSs do not overcome synergistic effects which are known to affect causal interpretation (8,9).

As for references, I found Roger Newson's slides – [Causality, confounders, and propensity scores](#) – relatively well-balanced about the pros and cons of using propensity scores, with illustrations from real studies. There were also several good papers dis-

cussing the use of propensity scores in observational studies or environmental epidemiology two years ago in *Statistics in Medicine*, and I enclose a couple of them at the end (3-6). But I like Pearl's review (7) because it offers a larger perspective on causality issues (PSs are discussed p. 117 and 130). Obviously, you will find many more illustrations by looking at applied research. I would like to add two recent articles from William R Shadish that came across Andrew Gelman's website (11,12). The use of propensity scores is discussed, but the two papers more largely focus on causal inference in observational studies (and how it compare to randomized settings).

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96 Comparing mixed effect models with the same number of degrees of freedom

Still, you can compute confidence intervals for your fixed effects, and report AIC or BIC (see e.g. **Cnann et al.**, *Stat Med* 1997 16: 2349).

Now, you may be interested in taking a look at **Assessing model mimicry using the parametric bootstrap**, from Wagenmakers et al. which seems to more closely resemble your initial question about assessing the quality of two competing models.

Otherwise, the two papers about measures of explained variance in LMM that come to my mind are:

- Lloyd J. Edwards, Keith E. Muller, Russell D. Wolfinger, Bahjat F. Qaqish and Oliver Schabenberger (2008). **An R2 statistic for fixed effects in the linear mixed model**, *Statistics in Medicine*, 27(29), 6137-6157.
- Ronghui Xu (2003). Measuring explained variation in linear mixed effects models, *Statistics in Medicine*, 22(22), 3527-3541.

But maybe there are better options.

97 Calculating percentile of normal distribution

In Python, you can use the `stats` module from the `scipy` package (look for `cdf()`, as in the following [example](#)).

(It seems the `transcendental` package also includes usual cumulative distributions).

98 What are good basic statistics to use for Ordinal data?

For basic summaries, I agree that reporting frequency tables and some indication about central tendency is fine. For inference, a recent article published in PARE discussed t- vs. MWW-test, [Five-Point Likert Items: t test versus Mann-Whitney-Wilc](#)

For more elaborated treatment, I would recommend reading Agresti's review on ordered categorical variables:

Liu, Y and Agresti, A (2005). [The analysis of ordered categorical data: An overview and a survey of recent developments](#). *Sociedad de Estadística e Investigación Operativa Test*, 14(1), 1-73.

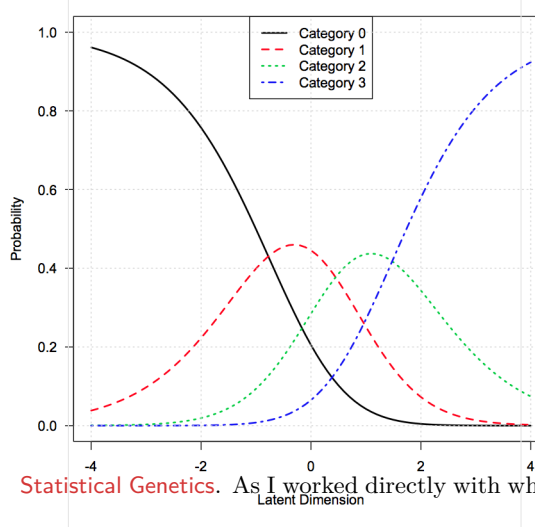
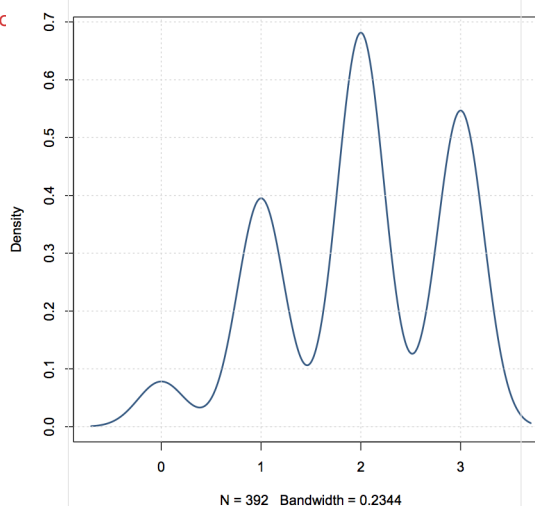
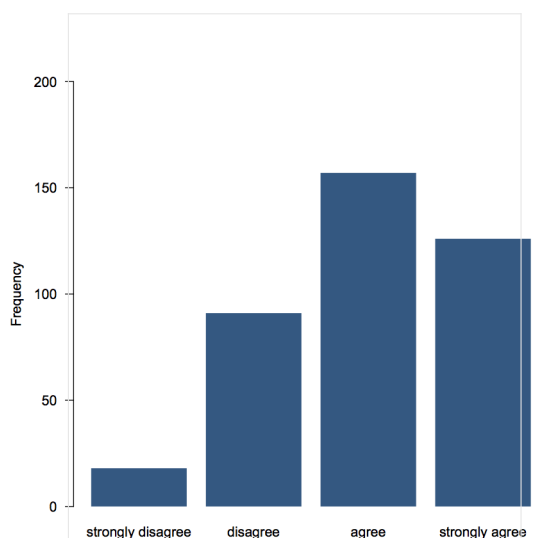
It largely extends beyond usual statistics, like threshold-based model (e.g. proportional odds-ratio), and is worth reading in place of Agresti's [CDA](#) book.

Below I show a picture of three different ways of treating a Likert item; from top to bottom, the “frequency” (nominal) view, the “numerical” view, and the “probabilistic” view (a [Partial Credit Model](#)):

The data comes from the `Science` data in the `ltm` package, where the item concerned *technology* (“New technology does not depend on basic scientific research”, with response “strongly disagree” to “strongly agree”, on a four-point scale)

99 How to Calculate the Pairwise LD for the given data?

There are various R/Bioconductor packages that allow you to compute pairwise correlation for SNPs in linkage disequilibrium, see the CRAN Task View



Statistical Genetics. As I worked directly with whole

genome scan, I've been mainly using `snpMatrix`, but `LDheatmap` or `mapLD` are fine. However, usually they expect genotype data (AA, AB, or BB), so I guess you will have to first convert your binary-encoded haplotype... About the filter on location, I also guess you just have to consider the pairwise R^2 or D' for proximal SNPs (usually, we draw a so-called heatmap of pairwise LD, which is roughly speaking the lower-diag elements of the correlation matrix, so you just have to consider the very first off diagonal elements).

Update

Now that I've read some papers, I'm not sure you will achieve your goals with the aforementioned method. To my knowledge, few packages allow to cope with multiallelic loci or haplotype blocks, one example being the `gap` package from JH Zhao (see also a review in the *Journal of Statistical Software*). The `LDkl()` function for example computes D' and ρ from a vector of haplotype frequencies, which can easily be plotted using `image()` or `levelplot()` from the `lattice` package.

100 Alternatives to classification trees with better predictive (e.g: CV) performance?

I think it would be worth giving a try to Random Forests (`randomForest`); some references were provided in response to related questions: *Feature selection for "final" model when performing cross-validation in machine learning*; *Can CART models be made robust?*. Boosting/bagging render them more stable than a single CART which is known to be very sensitive to small perturbations. Some authors argued that it performed as well as penalized SVM or *Gradient Boosting Machines* (see, e.g. Cutler et al., 2009). I think they certainly outperform NNs.

Boulesteix and Strobl provides a nice overview of several classifiers in *Optimal classifier selection and negative bias in error rate estimation: an empirical study on high-dimensional prediction* (BMC MRM 2009 9: 85). I've heard of another good study at the *IV EAM meeting*, which should be under review in *Statistics in Medicine*,

João Maroco, Dina Silva, Manuela Guerreiro, Alexandre de Mendonça. Do Random Forests Outperform Neural Networks,

Support Vector Machines and Discriminant Analysis classifiers? A case study in the evolution to dementia in elderly patients with cognitive complaints

I also like the `caret` package: it is well documented and allows to compare predictive accuracy of different classifiers on the same data set. It takes care of managing training /test samples, computing accuracy, etc in few user-friendly functions.

The `glmnet` package, from Friedman and coll., implements penalized GLM (see the review in the *Journal of Statistical Software*), so you remain in a well-known modeling framework.

Otherwise, you can also look for *association rules* based classifiers (see the CRAN Task View on *Machine Learning* or the *Top 10 algorithms in data mining* for a gentle introduction to some of them).

I'd like to mention another interesting approach that I plan to re-implement in R (actually, it's Matlab code) which is *Discriminant Correspondence Analysis* from Hervé Abdi. Although initially developed to cope with small-sample studies with a lot of explanatory variables (eventually grouped into coherent blocks), it seems to efficiently combine classical DA with data reduction techniques.

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101 Best practice when analysing pre-post treatment-control designs

There is a huge literature around this topic (change/gain scores), and I think the best references come from the biomedical domain, e.g.

Senn, S (2007). *Statistical issues in drug development*. Wiley (chap. 7 pp. 96-112)

In biomedical research, interesting work has also

been done in the study of **cross-over trials** (esp. in relation to *carry-over* effects, although I don't know how applicable it is to your study).

From Gain Score t to ANCOVA F (and vice versa), from Knapp & Schaffer, provides an interesting review of ANCOVA vs. t approach (the so-called Lord's Paradox). The simple analysis of change scores is not the recommended way for pre/post design according to Senn in his article **Change from baseline and analysis of covariance revisited** (Stat. Med. 2006 25(24)). Moreover, using a mixed-effects model (e.g. to account for the correlation between the two time points) is not better because you really need to use the "pre" measurement as a covariate to increase precision (through adjustment). Very briefly:

- The use of change scores (post – pre, or outcome – baseline) does not solve the problem of imbalance; the correlation between pre and post measurement is < 1 , and the correlation between pre and (post – pre) is generally negative – it follows that if the treatment (your group allocation) as measured by raw scores happens to be an unfair disadvantage compared to control, it will have an unfair advantage with change scores.
- The variance of the estimator used in ANCOVA is generally lower than that for raw or change scores (unless correlation between pre and post equals 1).
- If the pre/post relationships differ between the two groups (slope), it is not as much of a problem than for any other methods (the change scores approach also assumes that the relationship is identical between the two groups – the parallel slope hypothesis).
- Under the null hypothesis of equality of treatment (on the outcome), no interaction treatment x baseline is expected; it is dangerous to fit such a model, but in this case one must use centered baselines (otherwise, the treatment effect is estimated at the covariate origin).

I also like **Ten Difference Score Myths** from Edwards, although it focuses on difference scores in a different context; but here is an **annotated bibliography** on the analysis of pre-post change (unfortunately, it doesn't cover very recent work). Van Breukelen also compared ANOVA vs. ANCOVA in randomized and non-randomized setting, and his conclusions support the idea that ANCOVA is to be preferred, at least in randomized studies (which prevent from re-

gression to the mean effect).

102 Classification after factor analysis

Caution: I'm assuming that when you said "classification", you are rather referring to cluster analysis (as understood in French), that is an unsupervised method for allocating individuals in homogeneous groups without any prior information/label. It's not obvious to me how class membership might come into play in your question.

I'll take a different perspective from the other answers and suggest you to try to do a data reduction (through PCA) of your p variables followed by a mix of Ward's hierarchical and k-means clustering (this is called mixed clustering in the French literature, the basic idea is that HC is combined to a weighted k-means to consolidate the partition) on the first two or three factorial axes. This was proposed by Ludovic Lebart et coll. and is actually implemented in the **FactoClass** package.

The advantages are as follows:

- If any part of your survey is not clearly unidimensional, you will be able to gauge item contribution to the second axis, and this may help to flag those items for further inspection;
- Clustering is done on the PCA scores (or you can work with a multiple correspondence analysis, though in the case of binary items it amounts to yield the same results than a scaled PCA), and thanks to the mixed clustering the resulting partition is more stable and allow to spot potential extreme respondents; you can also introduce supplementary variable (like gender, SES or age), which is useful to inspect between-group homogeneity.

In this case, no rotation is supposed to be applied to the principal axes. Considering a subspace with $q < p$ allows to remove random fluctuations which often make the variance in the $p - q$ remaining axes. This can be viewed as some kind of "smoothing" on the data. Instead of PCA, as I said, you can use Multiple Correspondence Analysis (MCA), which is basically a non-linear PCA where numerical scores are assigned to respondents and modalities of dummy-coded variables. I have had some success using this method in characterizing clinical subgroups assessed on a wide range testing battery

for neuropsychology. It usually yields results that are interpretable (wrt. interpretation) (aka latent trait literature). The Factorial ANOVA is for the factorial clusters in the factor space.

Now, the problem is that the approach is that the dimensional representation (PCA or MCA) is used for identifying clusters. As discussed in Hwang (2014), any implementation of this approach is basically, the idea is to find a single step, criteria simultaneously for a criterion and the

References

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103 If an ANOVA effect is significant, the lack of a significant effect is the lack of a significant effect.

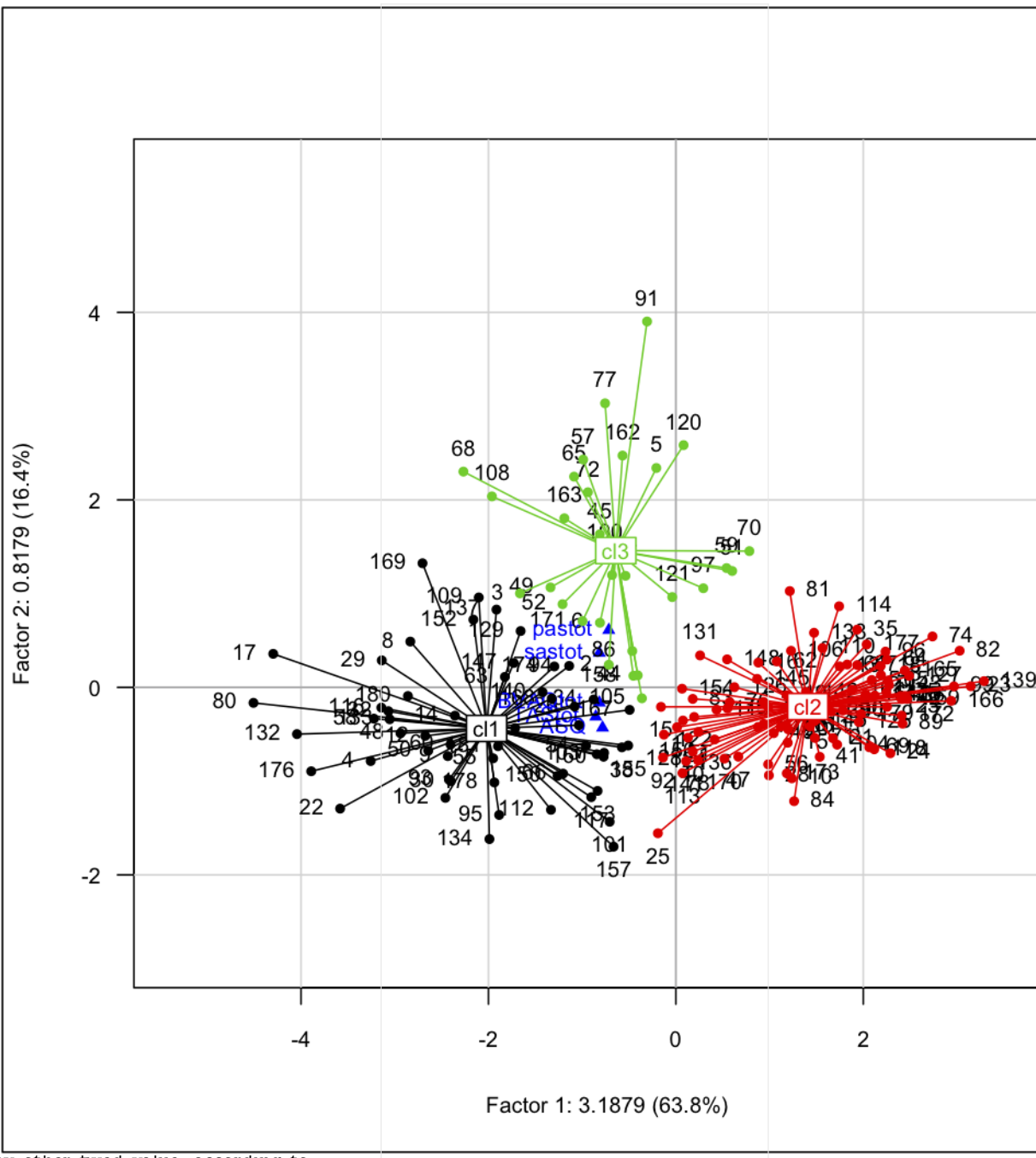
Well, it depends on the hypothesis or not. If you are encouraged to reject the null, you can simply reject the null (A:B terms) to get the effect size.

Now, the part of the effect size doesn't sound like it can prove that an observed effect is different from 0 (or any other fixed value, according to your alternative hypothesis), you cannot "accept" the null. If your test is non-significant, it simply means that you cannot reject H_0 . Non-significant results can also reflect lack of power (Type II error).

ent from 0 (or any other fixed value, according to your alternative hypothesis), you cannot "accept" the null. If your test is non-significant, it simply means that you cannot reject H_0 . Non-significant results can also reflect lack of power (Type II error).

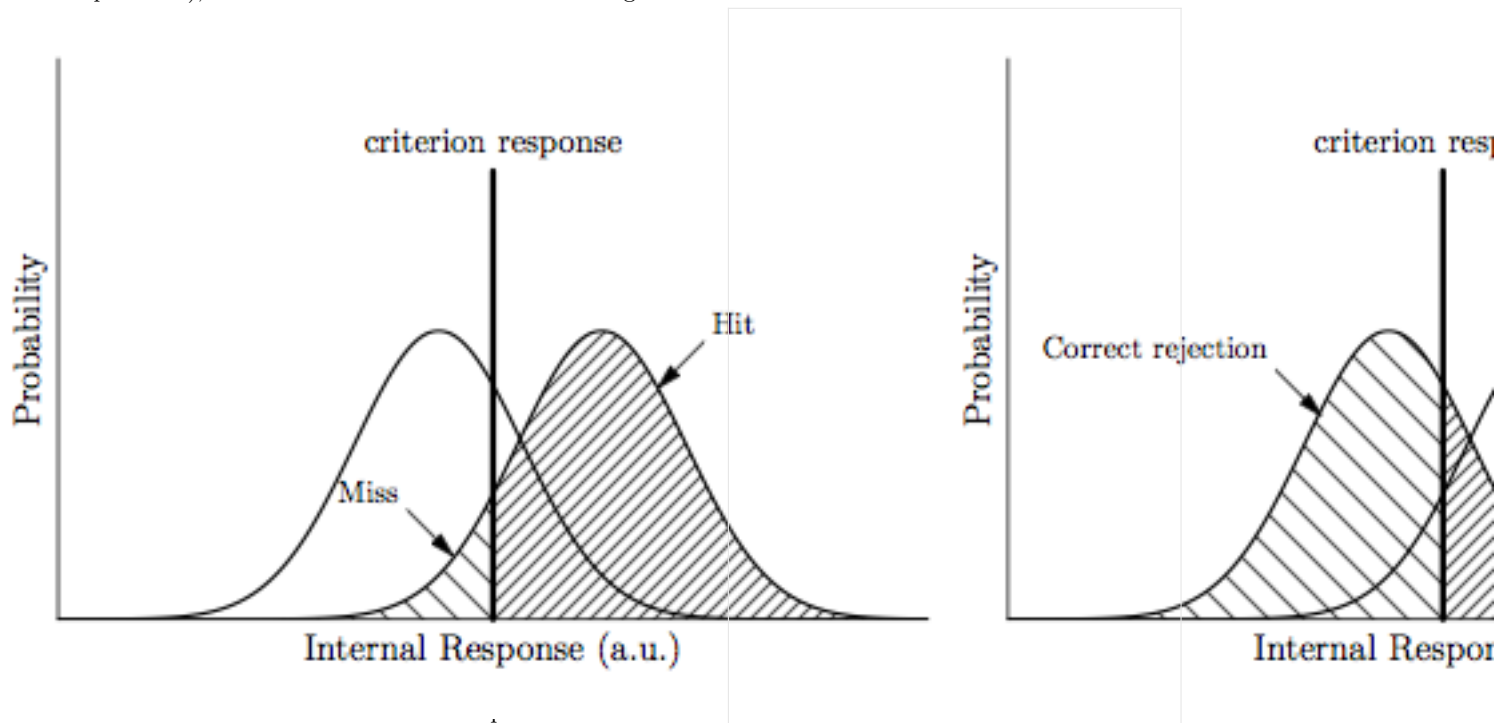
Also, rather than simply reporting crude p-values, it would be better (and it actually follows the APA recommendations) to also report some kind of effect size or difference of means, together with your inferential results.

Here is an example for reporting results from a factorial ANOVA (it has to be rework to fit your specific experimental design since your factors have a lot of levels):



A two-way analysis of variance yielded a main effect for A factor, $F(\nu_1, \nu_2) = 0.00$, $p < .05$, such that the average “value” was significantly higher in the a_1 condition (Mean=0.00, SD=0.00) compared to a_2 (Mean=0.00, SD=0.00) and a_3 (Mean=0.00, SD=0.00, Tukey HSD, all $p < 0.05$). The main effect of B was non-significant ($F(\nu_1, \nu_2) = 0.00$, $p = 0.00$), and no interaction effect was found sig-

Let an individual be facing a two-alternative choice experiment. Depending on the location of his internal criterion, his response may lead to Hit or False Alarm (response > criterion), or alternatively Correct Rejection or Miss (response < criterion). The corresponding probabilistic response curve resemble your situation.



zero effort to obtain a match. Here, I'm roughly quoting *Biometrics*, from Boulgouris et al. (chap. 26).

So, you may choose your cutoff by using standard ROC tools (search for “ROC analysis” on [Rseek](#)) to find the best compromise between FAR and TAR (this is not necessarily that cutoff that maximizes the AUC, it depends on your objectives).

Now, as has been highlighted in other responses, this compromise between FAR and TAR led to similar interpretation in psychophysics, classification, or biomedical science. It's just a matter of terminology, and we often speak of Hit rate vs. False Alarm rate; sensibility vs. specificity.

Note

Here are some pictures to complement other responses, which I hope will help you to draw the parallel with decision theory and statistical testing.

Most classical textbooks on Statistics provide a Table similar to the one below, where we describe the probabilities of incorrectly rejecting a null hypothesis (α) vs. falsely “accepting” the null (β) where in fact the alternative is true.

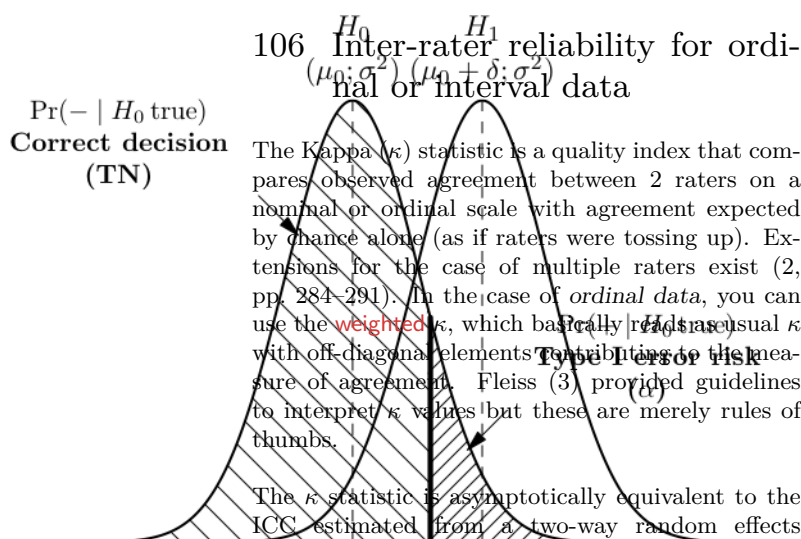
This leads to quite the same picture as with the psy-

Test

		Reality	
	H_0	H_1	
H_0	$1 - \alpha$ (TN)	β (FN)	
H_1	α (FP)	$1 - \beta$ (TP)	

In R, there is a function `pairwise.prop.test()` which allows any correction for multiple comparisons (single-step or step-down FWER methods or FDR-based), but it is quit what you already suggested (although Bonferroni is by far too conservative, but still very used in practice). A resampling approach, using permutation, might be interesting too. The `coin` R package provides a well-established testing framework in this respect, see §5 of [Implementing a Class of Permutation Tests: The coin Package](#), but I never had to deal with permutation tests on categorical data in a post-hoc way.

About the analysis of subdivided contingency tables, I generally consider specific associations as a guide to develop additional hypotheses (as for any unplanned comparisons), but this is another question. I generally just use visualization tools, like `mosaicplot` from [Michael Friendly](#), Pearson's residuals, and if I seek to explain specific patterns of association I use log-linear models.



chophysical threshold model:

105 Has anyone used the Marascuillo procedure for comparing multiple proportions?

Just a partial answer because I've never heard of this method. From what I read in the link you provided, it seems to be a single-step procedure (much like Bonferroni, except we rework the test statistics instead of the p-value) which is likely to be too conservative.

It should be noted that some psychometricians don't very much like κ because it is affected by the prevalence of the object of measurement much like predictive values are affected by the prevalence of the disease under consideration, and this can lead to paradoxical results.

Inter-rater reliability for k raters can be estimated

with Kendall's coefficient of concordance, W . When the number of items or units that are rated $n > 7$, $k(n-1)W \sim \chi^2(n-1)$. (2, pp. 269–270). This asymptotic approximation is valid for moderate value of n and k (6), but with less than 20 items F or permutation tests are more suitable (7). There is a close relationship between Spearman's ρ and Kendall's W statistic: W can be directly calculated from the mean of the pairwise Spearman correlations (for untied observations only).

Polychoric (ordinal data) correlation may also be used as a measure of inter-rater agreement. Indeed, they allow to

- estimate what would be the correlation if ratings were made on a continuous scale,
- test marginal homogeneity between raters.

In fact, it can be shown that it is a special case of latent trait modeling, which allows to relax distributional assumptions (4).

About *continuous* (or so assumed) measurements, the ICC which quantifies the proportion of variance attributable to the between-subject variation is fine. Again, bootstrapped CIs are recommended. As @ars said, there are basically two versions – agreement and consistency – that are applicable in the case of agreement studies (5), and that mainly differ on the way sum of squares are computed; the “consistency” ICC is generally estimated without considering the Item×Rater interaction. The ANOVA framework is useful with specific block design where one wants to minimize the number of ratings (**BIBD**) – in fact, this was one of the original motivation of Fleiss's work. It is also the best way to go for *multiple raters*. The natural extension of this approach is called the **Generalizability Theory**. A brief overview is given in **Rater Models: An Introduction**, otherwise the standard reference is Brennan's book, reviewed in **Psychometrika** 2006 71(3).

As for general references, I recommend chapter 3 of *Statistics in Psychiatry*, from Graham Dunn (Hodder Arnold, 2000). For a more complete treatment of reliability studies, the best reference to date is

Dunn, G (2004). *Design and Analysis of Reliability Studies*. Arnold. See the review in the **International Journal of Epidemiology**.

A good online introduction is available on John Uebersax's website, **Intraclass Correlation and Related Methods**; it includes a discussion of the pros and

cons of the ICC approach, especially with respect to ordinal scales.

Relevant R packages for two-way assessment (ordinal or continuous measurements) are found in the **Psychometrics** Task View; I generally use either the **psy**, **psych**, or **irr** packages. There's also the **concord** package but I never used it. For dealing with more than two raters, the **lme4** package is the way to go for it allows to easily incorporate random effects, but most of the reliability designs can be analysed using the **aoa** because we only need to estimate variance components.

References

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107 Logistic Regression: Which pseudo R-squared measure is the one to report (Cox & Snell or Nagelkerke)?

More details can be found in Harrell's book, *Regression Modeling Strategies* (pp. 203-205, 230-244, 247-249). For a more recent discussion, see also

Steyerberg, EW, Vickers, AJ, Cook, NR, Gerds, T, Gonen, M, Obuchowski, N, Pencina, MJ, and Kattan, MW (2010). *Assessing the Performance of Prediction Models, A Framework for Traditional and Novel Measures*. *Epidemiology*, 21(1), 128-138.

Both indices are measures of strength of association (i.e. whether any predictor is associated with the outcome, as for an LR test), and can be used to quantify predictive ability or model performance. A single predictor may have a significant effect on the outcome but it might not necessarily be so useful for *predicting individual response*, hence the need to assess model performance as a whole (wrt. the null model). The Nagelkerke R^2 is useful because it has a maximum value of 1.0, as Srikant said. This is just a normalized version of the R^2 computed from the likelihood ratio, $R^2_{LR} = 1 - \exp(-LR/n)$, which has connection with the Wald statistic for overall association, as originally proposed by Cox and Snell. Other indices of predictive ability are Brier score, the C index (concordance probability or ROC area), or Somers' D, the latter two providing a better measure of predictive discrimination.

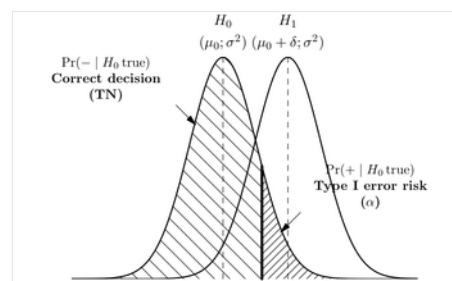
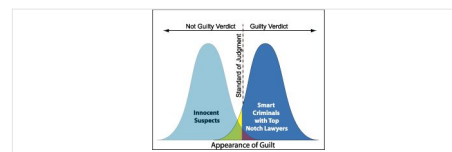
The only assumptions made in logistic regression are that of *linearity* and *additivity* (+ independence). Although many global goodness-of-fit tests (like the Hosmer & Lemeshow χ^2 test, but see my [comment](#) to @onestop) have been proposed, they generally lack power. For assessing model fit, it is better to rely on visual criteria (stratified estimates, nonparametric smoothing) that help to spot local or global departure between predicted and observed outcomes (e.g. non-linearity or interaction), and this is largely detailed in Harrell's [RMS handout](#). On a related subject (calibration tests), Steyerberg (*Clinical Prediction Models*, 2009) points to the same approach for assessing the agreement between observed outcomes and predicted probabilities:

Calibration is related to goodness-of-fit, which relates to the ability of a model to fit a given set of data. Typically, there is no single goodness-of-fit test that has good power against all kinds of lack of fit of a prediction model. Examples of lack of fit are missed non-linearities, interactions, or an inappropriate link function between the linear predictor and the outcome. Goodness-of-fit can be tested with a χ^2 statistic. (p. 274)

He also suggests to rely on the absolute difference between smoothed observed outcomes and predicted probabilities either visually, or with the so-

108 Is there a way to remember the definitions of Type I and Type II Errors?

I'll try not to be redundant with other responses (although it seems a little bit what J. M. already suggested), but I generally like showing the following two pictures:



109 Statistics and data mining software tools for dealing with large datasets

I'll second @suncoolsu comment: The dimensionality of your data set is not the only criterion that should orient you toward a specific software. For instance, if you're just planning to do unsupervised clustering or use PCA, there are several dedicated tools that cope with large data sets, as commonly encountered in genomic studies.

Now, R (64 bits) handles large data pretty well,

and you still have the option to use disk storage instead of RAM access, but see CRAN Task View [High-Performance and Parallel Computing with R](#). Standard GLM will easily accomodate 20,000 obs. (but see also [speedglm](#)) within reasonable time, as shown below:

```
> require(MASS)
> n <- 20000
> X <- mvrnorm(n, mu=c(0,0), Sigma=matrix(c(1,.8,.8,1), 2,
2))
> df <- cbind.data.frame(X, grp=gl(4, n/4), y=sample(c(0,1),
n, rep=TRUE))
> system.time(glm(y ~ ., data=df))
      user  system elapsed
   0.361   0.018   0.379
```

To give a more concrete illustration, I used R to process and analyse large genetic data (800 individuals x 800k [SNPs](#), where the main statistical model was a stratified GLM with several covariates (2 min); that was made possible thanks to efficient R and C codes available in the [snpMatrix](#) package (in comparison, the same kind of model took about 8 min using a dedicated C++ software ([plink](#))). I also worked on a clinical study (12k patients x 50 variables of interest) and R fits my needs too. Finally, as far as I know, the [lme4](#) package is the only software that allow to fit mixed-effects model with unbalanced and large data sets (as is the case in large-scale educational assessment).

Stata/SE is another software that can handle [large data set](#). SAS and SPSS are file based software, so they will handle large volumes of data. A comparative review of software for datamining is available in [Data Mining Tools: Which One is Best for CRM](#). For visualization, there are also plenty of options; maybe a good start is [Graphics of large datasets: visualizing a million](#) (reviewed in the JSS by P Murrell), and all related threads on this site.

110 Cox regression and time scale

Usually, age at baseline is used as a covariate (because it is often associated to disease/death), but it can be used as your time scale as well (I think it is used in some longitudinal studies, because you need to have enough people at risk along the time scale, but I can't remember actually – just found these slides about [Analysing cohort studies assuming a continuous time scale](#) which talk about cohort studies). In the interpretation, you should replace event time by age, and you might include age at diagnosis

as a covariate. This would make sense when you study age-specific mortality of a particular disease (as illustrated in these [slides](#)).

Maybe this article is interesting since it contrasts the two approaches, time-on-study vs. chronological age: [Time Scales in Cox Model: Effect of Variability Among Entry Ages on Coefficient Estimates](#). Here is another paper:

Cheung, YB, Gao, F, and Khoo, KS (2003). [Age at diagnosis and the choice of survival analysis methods in cancer epidemiology](#). *Journal of Clinical Epidemiology*, 56(1), 38-43.

But there are certainly better papers.

111 Relation between logistic regression coefficient and odds ratio in JMP

Ok, I drop a quick response. Your idea is correct in that the regression coefficient is the log of the OR. More precisely, if b is your regression coefficient, $\exp(b)$ is the odds ratio corresponding to a *one unit change* in your variable. So, to get back to the adjusted odds, you need to know what are the internal coding convention for your factor levels. Usually, for a binary variable it is 0/1 or 1/2. But if it happens that your levels are represented as -1/+1 (which I suspect here), then you have to multiply the regression coefficient by 2 when exponentiating.

The same would apply if you were working with a continuous variable, like age, and want to express the odds for 5 years ($\exp(5b)$) instead of 1 year ($\exp(b)$).

Update: I just found this about [JMP coding for nominal variables](#) (version < 7).

112 Repeatability and measurement error from and between observers

What you describe is a reliability study where each subject is going to be assessed by the same three raters on two occasions. Analysis can be done separately on the two outcomes (length and weight, though I assume they will be highly correlated and

you're not interested in how this correlation is reflected in raters' assessments). Estimating measurement reliability can be done in two ways:

- The original approach (as described in Fleiss, 1987) relies on the analysis of variance components through an ANOVA table, where we assume no subject by rater interaction (the corresponding SS is constrained to 0) – of course, you won't look at p -values, but at the MSs corresponding to relevant effects;
- A mixed-effects model allows to derive variance estimates, considering time as a fixed effect and subject and/or rater as random-effect(s) (the latter distinction depends on whether you consider that your three observers were taken or sampled from a pool of potential raters or not – if the rater effect is small, the two analyses will yield quite the same estimate for outcome reliability).

In both cases, you will be able to derive a single intraclass correlation coefficient, which is a measure of reliability of the assessments (under the Generalizability Theory, we would call them generalizability coefficients), which would answer your second question. The first question deals with a potential effect of time (considered as a fixed effect), which I discussed here, [Reliability in Elicitation Exercise](#). More details can be found in Dunn (1989) or Brennan (2001).

I have an [R example script](#) on Github which illustrates both approaches. I think it would not be too difficult to incorporate rater effects in the model.

References

1. Fleiss, J.L. (1987). *The design and analysis of clinical experiments*. New York: Wiley.
2. Dunn, G. (1989). *Design and analysis of reliability studies*. Oxford
3. Brennan, R.L. (2001). *Generalizability Theory*. Springer

113 Reliability in Elicitation Exercise

Maybe I misunderstood the question, but what you are describing sounds like a test-retest reliability study on your Q scores. You have a series of experts

each going to assess a number of items or questions, at two occasions (presumably fixed in time). So, basically you can assess the temporal stability of the judgments by computing an *intraclass correlation coefficient* (ICC), which will give you an idea of the variance attributable to subjects in the variability of observed scores (or, in other words of the closeness of the observations on the same subject relative to the closeness of observations on different subjects).

The ICC may easily be obtained from a mixed-effect model describing the measurement y_{ij} of subject i on occasion j as

$$y_{ij} = \mu + u_i + \varepsilon_{ij}, \quad \varepsilon \sim \mathcal{N}(0, \sigma^2)$$

where u_i is the difference between the overall mean and subject i 's mean measurement, and ε_{ij} is the measurement error for subject i on occasion j . Here, this is a random-effect model. Unlike a standard ANOVA with subjects as factor, we consider the u_i as random (i.i.d.) effects, $u_i \sim \mathcal{N}(0, \tau^2)$, independent of the error terms. Each measurement differ from the overall mean μ by the sum of the two error terms, among which the u_i is shared between occasion on the same subjects. The total variance is then $\tau^2 + \sigma^2$ and the proportion of the total variance that is accounted for by the subjects is

$$\rho = \frac{\tau^2}{\tau^2 + \sigma^2}$$

which is the ICC, or the reliability index from a psychometrical point of view. Note that this reliability is sample-dependent (as it depends on the between-subject variance). Instead of the mixed-effects model, we could derive the same results from a two-way ANOVA (subjects + time, as factors) and the corresponding Mean Squares. You will find additional references in those related questions: [Repeatability and measurement error from and between observers](#), and [Inter-rater reliability for ordinal or interval data](#).

In R, you can use the `icc()` function from the `psy` package; the random intercept model described above corresponds to the “agreement” ICC, while incorporating the time effect as a fixed factor would yield the “consistency” ICC. You can also use the `lmer()` function from the `lme4` package, or the `lme()` function from the `nlme` package. The latter has the advantage that you can easily obtain 95% CIs for the variance components (using the `intervals()` function). Dave Garson provided a nice overview (with SPSS illustrations) in [Reliability Analysis](#), and [Estimating Multilevel Models](#)

using SPSS, Stata, SAS, and R constitutes a useful tutorial, with applications in educational assessment. But the definitive reference is Shrout and Fleiss (1979), *Intraclass Correlations: Uses in Assessing Rater Reliability*, *Psychological Bulletin*, 86(2), 420-428.

I have also added an [example R script](#) on Github, that includes the ANOVA and mixed-effect approaches.

Also, should you add a constant value to all of the values taken at the second occasion, the Pearson correlation would remain identical (because it is based on deviations of the 1st and 2nd measurements from their *respective means*), whereas the reliability as computed through the random intercept model (or the agreement ICC) would decrease.

BTW, Cronbach's alpha is not very helpful in this case because it is merely a measure of the internal consistency (yet, another form of "reliability") of an unidimensional scale; it would have no meaning should it be computed on items underlying different constructs. Even if your questions survey a single domain, it's hard to imagine mixing the two series of measurements, and Cronbach's alpha should be computed on each set separately. Its associated 95% confidence interval (computed by bootstrap) should give an indication about the stability of the internal structure between the two test occasions.

As an example of applied work with ICC, I would suggest

Johnson, SR, Tomlinson, GA, Hawker, GA, Granton, JT, Grosbein, HA, and Feldman, BM (2010). [A valid and reliable belief elicitation method for Bayesian priors](#). *Journal of Clinical Epidemiology*, 63(4), 370-383.

114 Patient distance metrics

You asked a difficult question, but I'm a little bit surprised that the various clues that were suggested to you received so little attention. I upvoted all of them because I think they basically are useful responses, though in their actual form they call for further bibliographic work.

Disclaimer: I never had to deal with such a problem, but I regularly have to expose statistical results that may differ from physician's *a priori* beliefs and I learn a lot from unraveling their lines of reasoning.

Also, I have some background in teaching human decision/knowledge from the perspective of Artificial Intelligence and Cognitive Science, and I think what you asked is not so far from how experts actually decide that two objects are similar or not, based on their attributes and a common understanding of their relationships.

From your question, I noticed two interesting assertions. The first one related to how an expert assess the similarity or difference between two set of measurements:

I don't particularly care if there is some relation between attribute X and Y. What I care about is if a doctor thinks there is a relation between X and Y.

The second one,

How can I predict what they think the similarity is? Do they look at certain attributes?

looks like it is somewhat subsumed by the former, but it seems more closely related to what are the most salient attributes that allow to draw a clear separation between the objects of interest.

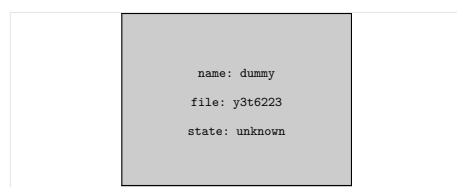
To the first question, I would answer: Well, if there is no characteristic or objective relationship between any two subjects, what would be the rationale for making up an hypothetical one? Rather, I think the question should be: If I only have limited resources (knowledge, time, data) to take a decision, how do I optimize my choice? To the second question, my answer is: Although it seems to partly contradicts your former assertion (if there is no relationship at all, it implies that the available attributes are not discriminative or useless), I think that most of the time this is a combination of attributes that makes sense, and not only how a given individual scores on a single attribute.

Let me dwell on these two points. Human beings have a limited or [bounded rationality](#), and can take a decision (often the right one) without examining all possible solutions. There is also a close connection with [abductive reasoning](#). It is well known that there is some variability between individual judgments, and even between judgments from the same expert at two occasions. This is what we are interested in in reliability studies. But you want to know how these experts elaborate their judgments. There is a huge amount of papers about that in cognitive psychology, especially on the fact that *relative judgments* are easier and more reliable

than *absolute* ones. Doctors' decisions are interesting in this respect because they are able to take a "good" decision with a limited amount of information, but at the same time they benefit from an ever growing internal knowledge base from which they can draw expected relationships (extrapolation). In other words, they have a built-in inference (assumed to be hypothetico-deductive) machinery and accumulate positive evidence or counterfactuals from their experience or practice. Reproducing this inferential ability and the use of declarative knowledge was the aim of several expert or **production rule** systems in the 70s, the most famous one being **MYCIN**, and more generally of Artificial Intelligence earlier in 1946 (Can we reproduce on an artificial system the intelligent behavior observed in man?). Automatic treatment of speech, problem solving, visual shape recognition are still active projects nowadays and they all have to do with identifying salient features and their relationships to make an appropriate decision (i.e., how far should two patterns be to be judged as the emanation of two distinct generating processes?).

In sum, our doctors are able to draw an optimal inference from a limited amount of data, compensating from noise that arises simply as a byproduct of individual variability (at the level of the patients). Thus, there is a clear connection with statistics and probability theory, and the question is what conscious or subconscious methodology help doctors forming their judgments. **Semantic networks** (SN), **belief networks**, and **decision trees** are all relevant to the question you asked. The paper you cited is about using an **ontology** as a basis of formal judgments, but it is no more than an extension of SNs, and many projects were initiated in this direction (I can think of the **Gene Ontology** for genomic studies, but many others exist in different domains).

Now, look at the following hierarchical classification of diagnostic categories (it is roughly taken from Dunn 1989, p. 25):



And now take a look at the **ICD classification**; I think it is not too far from this schematic classification. Mental disorders are organized into distinct categories, some of them being closer one to each

other. What render them similar is the closeness of their expression (phenotype) in any patient, and the fact that they share some similarities in their somatic/psychological etiology. Assessing whether two doctors would make the same diagnostic is a typical example of an *inter-rater agreement* study, where two psychiatrists are asked to place each of several patients in mutually exclusive categories. The hierarchical structure should be reflected in the disagreement between each doctor, that is they may not agree on the finer distinction between diagnostic classes (the leafs) but if they were to disagree between insomnia and schizophrenia, well it would be little bit disconcerting. . . How these two doctors decide on which class a given patient belongs to is no more than a clustering problem: How likely are two individuals, given a set of observed values on different attributes, to be similar enough so that I decide they share the same class membership?

Now, some attributes are more influential than others, and this is exactly what is reflected in the weight attributed to a given attribute in Latent Class Analysis (which can be thought of as a **probabilistic extension** of clustering methods like k-means), or the **variable importance** in Random Forests. We need to put things into boxes, because at first sight it's simpler. The problem is that often things overlap to some extent, so we need to consider different levels of categorization. In fact, *cluster analysis* is at the heart of the actual DSM categories, and many papers actually turn around assigning one patient to a specific syndromic category, based on the profile of his response to a battery of neuropsychological assessments. This merely looks like a *subtyping* approach; each time, we seek to refine a preliminary well-established diagnostic category, by adding exception rules or an additional relevant symptom or impairment.

A related topic is *decision trees* which are by far the most well understood statistical techniques by physicians. Most of the time, they described a nested series of boolean assertions (Do you have a sore throat? If yes, do you have a temperature? etc.; but look at an example of public **influenza diagnostic tree**) according to which we can form a decision regarding patients proximity (i.e. how similar patients are wrt. attributes considered for building the tree – the closer they are the more likely they are to end up in the same leaf). **Association rules** and the **C4.5 algorithm** rely quite on the same idea. On a related topic, there's the **patient rule-induction method** (PRIM). Now clearly, we must make a distinction between all those methods, that make an efficient use of a large body of data and incorpo-

rate bagging or boosting to compensate for model fragility or overfitting issues, and doctors who cannot process huge amount of data in an automatic and algorithmic manner. But, for small to moderate amount of descriptors, I think they perform quite good after all.

The yes-or-no approach is not the panacea, though. In behavioral genetics and psychiatry, it is commonly argued that the classification approach is probably not the best way to go, and that common diseases (learning disorders, depression, personality disorders, etc.) reflect a continuum rather than classes of opposite valence. Nobody's perfect!

In conclusion, I think doctors actually hold a kind of internalized inference engine that allows them to assign patients into distinctive classes that are characterized by a weighted combination of available evidences; in other words, they are able to organize their knowledge in an efficient manner, and these internal representations and the relationships they share may be augmented throughout experience. **Case-based reasoning** probably comes into play at some point too. All of this may be subjected to (a) revision with newly available data (we are not simply acting as definitive binary classifiers, and are able to incorporate new data in our decision making), and (b) subjective biases arising from past experience or wrong self-made association rules. However, they are prone to errors, as every decision systems...

All statistical techniques reflecting these steps – decisions trees, bagging/boosting, cluster analysis, latent cluster analysis – seems relevant to your questions, although they may be hard to instantiate in a single decision rule.

Here are a couple of references that might be helpful, as a first start to how doctors make their decisions:

- **A clinical decision support system for clinicians for determining appropriate radiologic imaging examination**
- Grzymala-Busse, JW. **Selected Algorithms of Machine Learning from Examples**. *Fundamenta Informaticae* 18 (1993), 193–207
- Santiago Medina, L, Kuntz, KM, and Pomeroy, S. **Children With Headache Suspected of Having a Brain Tumor: A Cost-Effectiveness Analysis of Diagnostic Strategies**. *Pediatrics* 108 (2001),

255-263

- **Building Better Algorithms for the Diagnosis of Nontraumatic Headache**
- Jenkins, J, Shields, M, Patterson, C, and Kee, F. **Decision making in asthma exacerbation: a clinical judgement analysis**. *Arch Dis Child* 92 (2007), 672–677
- Croskerry, P. **Achieving quality in clinical decision making: cognitive strategies and detection of bias**. *Acad Emerg Med* 9(11) (2002), 1184-204.
- Cahan, A, Gilon, D, Manor, O, and Paltiel. **Probabilistic reasoning and clinical decision-making: do doctors overestimate diagnostic probabilities?** *QJM* 96(10) (2003), 763-769
- Wegwarth, O, Gaissmaier, W, and Gigerenzer, G. **Smart strategies for doctors and doctors-in-training: heuristics in medicine**. *Medical Education* 43 (2009), 721–728

115 Incorporating boolean data into analysis

Ingo Ruczinski has contributed to promote the use of **Logic regression** for data set consisting of binary variables, with an emphasis on higher-order interaction terms. The main advantage compared to usual or penalized GLMs is that it is more parcimonious in terms of degrees of freedom. The outcome may be continuous or categorical, and continuous covariates can be added to the model (or the outcome can first be residualized on them if these are the binary predictors that are of interest).

The original paper

Ruczinski I, Kooperberg C, LeBlanc ML (2003). **Logic Regression**. *Journal of Computational and Graphical Statistics*, 12(3), 475-511.

includes several applications in biomedical studies, and a comparison of LR with **CART** and **MARS**. Although it has mainly been applied in large-scale genetic studies (e.g. genome-wide association studies), it should work with any binary variables whose combinations of interest can be expressed with a set of logical operators.

The **LogicReg** R package implements this technique; see also the related packages on CRAN and Bioconductor, esp. **LogicForest** which shares some ideas with Random Forests.

116 Where to cut a dendrogram?

There is no definitive answer since cluster analysis is essentially an exploratory approach; the interpretation of the resulting hierarchical structure is context-dependent and often several solutions are equally good from a theoretical point of view.

Several clues were given in a related question, **What stop-criteria for agglomerative hierarchical clustering are used in practice?** I generally use visual criteria, e.g. silhouette plots, and some kind of numerical criteria, like Dunn's validity index, Hubert's gamma, G2/G3 coefficient, or the corrected Rand index. Basically, we want to know how well the original distance matrix is approximated in the cluster space, so a measure of the **cophenetic correlation** is also useful. I also use k-means, with several starting values, and the **gap statistic** to determine the number of clusters that minimize the within-SS. The concordance with Ward hierarchical clustering gives an idea of the stability of the cluster solution (You can use `matchClasses()` in the **e1071** package for that).

You will find useful resources in the CRAN Task View **Cluster**, including **pvclust**, **fpc**, **clv**, among others. Also worth to give a try is the **clValid** package (described in the *Journal of Statistical Software*).

Now, if your clusters change over time, this is a bit more tricky; why choosing the first cluster-solution rather than another? Do you expect that some individuals move from one cluster to another as a result of an underlying process evolving with time?

There are some measure that try to match clusters that have a maximum absolute or relative overlap, as was suggested to you in your preceding question. Look at **Comparing Clusterings - An Overview** from Wagner and Wagner.

117 Choosing clustering method

There is no definitive answer to your question, as even within the same method the choice of the distance to represent individuals (dis)similarity may yield different result, e.g. when using euclidean vs. squared euclidean in hierarchical clustering. As an other example, for binary data, you can choose

the Jaccard index as a measure of similarity and proceed with classical hierarchical clustering; but there are alternative approaches, like the **Mona (Monothetic Analysis)** algorithm which only considers one variable at a time, while other hierarchical approaches (e.g. classical HC, Agnes, Diana) use all variables at each step. The k-means approach has been extended in various way, including partitioning around medoids (PAM) or representative objects rather than centroids (Kaufman and Rousseeuw, 1990), or fuzzy clustering (Chung and Lee, 1992). For instance, the main difference between the k-means and PAM is that PAM minimizes a sum of dissimilarities rather than a sum of squared euclidean distances; fuzzy clustering allows to consider "partial membership" (we associate to each observation a weight reflecting class membership). And for methods relying on a probabilistic framework, or so-called model-based clustering (or **latent profile analysis** for the psychometricians), there is a great package: **Mclust**. So definitively, you need to consider how to define the resemblance of individuals as well as the method for linking individuals together (recursive or iterative clustering, strict or fuzzy class membership, unsupervised or semi-supervised approach, etc.).

Usually, to assess cluster stability, it is interesting to compare several algorithm which basically "share" some similarity (e.g. k-means and hierarchical clustering, because euclidean distance work for both). For assessing the concordance between two cluster solutions, some pointers were suggested in response to this question, **Where to cut a dendrogram?** (see also the cross-references for other link on this website). If you are using R, you will see that several packages are already available in Task View on Cluster Analysis, and several packages include vignettes that explain specific methods or provide case studies.

Cluster Analysis: Basic Concepts and Algorithms provides a good overview of several techniques used in Cluster Analysis. As for a good recent book with R illustrations, I would recommend chapter 12 of Izenman, *Modern Multivariate Statistical Techniques* (Springer, 2008). A couple of other standard references is given below:

- Cormack, R., 1971. A review of classification. *Journal of the Royal Statistical Society, A* 134, 321–367.
- Everitt, B., 1974. *Cluster analysis*. London: Heinemann Educ. Books.

- Gordon, A., 1987. A review of hierarchical classification. *Journal of the Royal Statistical Society, A* 150, 119–137.
- Gordon, A., 1999. *Classification*, 2nd Edition. Chapman and Hall.
- Kaufman, L., Rousseeuw, P., 1990. *Finding Groups in Data: An Introduction to Cluster Analysis*. New York, Wiley.

118 What is a meaning of “p-value F” from Friedman test?

I generally used `friedman.test()` which doesn't return any F statistic. If you consider that you have b blocks, for which you assigned ranks to observations belonging to each of them, and that you sum these ranks for each of your a groups (let denote them sum R_i), then the Friedman statistic is defined as

$$F_r = \frac{12}{ba(a+1)} \sum_{i=1}^a R_i^2 - 3b(a+1)$$

and follows a $\chi^2(a-1)$, for a and b sufficiently large. Quoting Zar (*Biostatistical Analysis*, 4th ed., pp. 263-264), this approximation is conservative (hence, test has low power) and we can use an F-test, with

$$F_{\text{obs}} = \frac{(b-1)F_r}{b(a-1) - F_r}$$

which is to be compared to an F distribution with $a-1$ and $(a-1)(b-1)$ degrees of freedom.

119 The best measure of reliability for interval data between 0 and 1

Referring to your comments to @Henrik, I'm inclined to think that you rather have continuous measurements on a set of objects (here, your similarity measure) for 6 raters. You can compute an **intraclass correlation** coefficient, as described here **Reliability in Elicitation Exercise**. It will provide you with a measure of agreement (or concordance) between all 6 judges wrt. assessments they made, or more precisely the part of variance that is explained by between-rater variance. There's a working R script in appendix.

Note that this assumes that your measures are considered as real valued measurement (I refer to @onestop's comment), not really proportions of similarity or whatever between your paired sounds. I don't know of a specific version of the ICC for % or values bounded on an interval, only for binary or ranked data.

Update:

Following your comments about parameters of interest and language issue:

- There are many other online resources on the ICC; I think **David Howell** provides a gentle and well illustrated introduction to it. Its discussion generalize to k-sample (judges/raters) without any difficulty I think, or see this chapter from Sea and Fortna on **Psychometric Methods**. What you have to think to is mainly whether you want to consider your raters as an unique set of observers, not necessarily representative of all the raters that would have assess your object of measurement (this is called a fixed effect), or as a random sample of raters sampled from a larger (hypothetical) population of potential raters: in the former case, this corresponds to a one-way anova or a consistency ICC, in the latter case we talk about an agreement ICC.
- A colleague of mine successfully used **Kevin Brownhill's script** (from Matlab Central file exchange). The ICC you are interested in is then `cse=3` (if you consider that your raters are not representative of a more general population of raters).

120 How do you draw structural equation/MPLUS models?

I use the **psych** R package for CFA and John Fox's **sem** package with simple SEM. Note that the graphical backend is graphviz. I don't remember if the **lavaan** package provides similar or better facilities.

Otherwise, the **Mx software** for genetic modeling features a graphical interface in its Windows flavour, and you can export the model with path coefficients.

121 What graphical techniques are used in Structural Equation Modeling?

I worked with Laura Trinchera who contributed a nice R package for PLS-path modeling, `plspm`. It includes several graphical output for various kind of 2- and k-block data structures.

I just discovered the `plotSEMM` R package. It's more related to your second point, though, and is restricted to graphing bivariate relationships.

As for recent references on diagnostic plot for SEMs, here are two papers that may be interesting (for the second one, I just browsed the abstract recently but cannot find an ungated version):

1. Sanchez BN, Houseman EA, and Ryan LM. **Residual-Based Diagnostics for Structural Equation Models**. *Biometrics* (2009) 65, 104–115
2. Yuan KH and Hayashi K. **Fitting data to model: Structural equation modeling diagnosis using two scatter plots**, *Psychological Methods* (2010)

122 Data transformation for Principal Components Analysis from different likert scales

As suggested by @whuber, you can “abstract” the scale effect by working with a standardized version of your data. If you're willing to accept that an interval scale is the support of each of your item (i.e. the distance between every two response categories would have the same meaning for every respondents), then linear correlations are fine. But you can also compute **polychoric correlation** to better account for the discretization of a latent variable (see the R package `polycor`). Of note, it's a largely more computer-intensive job, but it works quite well in R.

Another possibility is to combine optimal scaling within your PCA, as implemented in the `homals` package. The idea is to find a suitable non-linear transformation of each scale, and this is very nicely described by Jan de Leeuw in the accompanying vignette or the JSS article, **Gifi Methods for Optimal Scaling in R: The Package homals**. There are several examples included.

For a more thorough understanding of this approach with any factorial method, see the work of **Yoshio Takane** in the 80s.

Similar points were raised by @Jeromy and @mbq on related questions, **Does it ever make sense to treat categorical data as continuous?**, **How can I use optimal scaling to scale an ordinal categorical variable?**

123 How does one calculate Cohen's d and confidence intervals after logit in Stata?

Cohen's d is not directly available in Stata, and you have to resort on external macros, e.g. `sizefx` (`ssc install sizefx`). It works fine if you have to series of values, but I found it less handy when you work with a full data set because there's no possibility to pass options to this command (e.g. `by()`).

Anyway, you can still use the original formula (with pooled SDs),

$$\delta_c = \frac{\bar{x}_1 - \bar{x}_2}{s_p}$$

where $s_p = \sqrt{\frac{(n_1-1)s_1^2 + (n_2-1)s_2^2}{(n_1+n_2-2)}}$.

Here is an example by hand:

```
. webuse lbw
. logit low age smoke
. graph box age, by(low)
. tabstat age, by(low) statistics(mean sd N)
```

```
Summary for variables: age
by categories of: low (birthweight<2500g)
```

low	mean	sd	N
0	23.66154	5.584522	130
1	22.30508	4.511496	59
Total	23.2381	5.298678	189

```
. display "Cohen's d: = " (23.66154-22.30508) / sqrt((129*(5.584522)^2+58*(4.511496)^2)/188)
```

```
Cohen's d: = .25714326
```

This is in agreement with what R would give:

```
library(MBESS)
res <- smd(Mean.1=23.66154, Mean.2=22.30508,
           s.1=5.584522, s.2=4.511496, n.1=130, n.2=59)
ci.smd(smd=res, n.1=130, n.2=59, conf.level=0.95)
```

that is an effect size of 0.257 with 95% CI [-0.052;0.566].

In contrast, `sizefx` gives results that differ a little (I have use `separate age`, `by(low)` and collapse the results in a new data window, here two columns labeled `age0` and `age1`), the ES version calculated above corresponding to what is referred to as Hedge's *g* below (unless I miss something in the `code` I read):

```
. sizefx age0 age1

Cohen's d and Hedges' g for: age0 vs. age1
Cohen's d statistic (pooled variance) = .26721576
Hedges' g statistic = .26494154

Effect size correlation (r) for: age0 vs. age1
ES correlation r = .13243109
```

124 Inter-rater reliability between similarity matrices

My first idea would be to try some kind of cluster analysis (e.g. [hierarchical clustering](#)) on each similarity matrix, and compare the classification trees across raters. We can derive a similarity index from all dendrograms, as discussed here, [A measure to describe the distribution of a dendrogram](#), or in this review, [Comparing Clusterings - An Overview](#) from Wagner and Wagner.

You benefit from working with already existing distance matrices, thus such methods will really reflect the nature of your data, and you can still derive a single numerical value to quantify the closeness of method-specific assessments. The following article may be interesting, if you need to refer to existing work:

Hamer, RM and Cunningham, JW. [Cluster Analyzing Profile Data Confounded with Interrater Differences: A Comparison of Profile Association Measures](#). *Applied Psychological Measurement* (1981) 5(1): 63-72.

Another approach would be to apply some kind of [Principal Component Analysis](#) on each similarity matrix, and keep only the first principal component

(the linear combination of all 100 items that account for the maximum of variance). More precisely, as you work with (dis)similarity indices or a particular distance/proximity metric, it is sometimes referred to as Principal Coordinate Analysis or [Multidimensional Scaling](#) (MDS), although PCA and MDS would yield similar results when dissimilarities are defined as euclidean distances. There is a working example in Izenman's book ([Modern Multivariate Statistical Techniques](#), chapter 13, "perceptions of color in human vision", pp. 468-470) and a discussion on so-called *all-pairs design* pp. 471-472. You can then compare the 6 linear combinations (i.e., the weights associated to each sound by rater-specific MDS) to assess their consistency across raters. There, an ICC (as described in my [previous answer](#)) could make sense, but I don't know of any application of it in this particular case.

125 How to create a barplot diagram where bars are side-by-side in R

I shall assume that you are able to import your data in R with `read.table()` or the short-hand `read.csv()` functions. Then you can apply any summary functions you want, for instance `table` or `mean`, as below:

```
x <- replicate(4, rnorm(100))
apply(x, 2, mean)
```

or

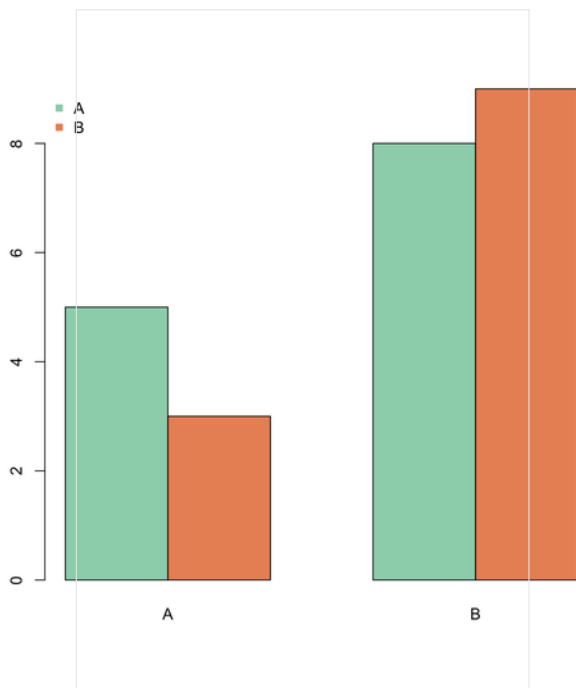
```
x <- replicate(2, sample(letters[1:2], 100, rep=T))
apply(x, 2, table)
```

The idea is to end up with a matrix or table for the summary values you want to display.

For the graphical output, look at the `barplot()` function with the option `beside=TRUE`, e.g.

```
barplot(matrix(c(5,3,8,9),nr=2), beside=T,
          col=c("aquamarine3","coral"),
          names.arg=LETTERS[1:2])
legend("topleft", c("A","B"), pch=15,
       col=c("aquamarine3","coral"),
       bty="n")
```

The `space` argument can be used to add an extra space between juxtaposed bars.



126 Comparing test-retest reliabilities

Both situations are specific cases of test-retest, except that the recall period is null in the first case you described. I would also expect a larger agreement in the former case, but that may be confounded with a learning or memory effect. A chance-corrected measure of agreement, like **Cohen's kappa**, can be used with binary variables, and bootstrapped confidence intervals might be compared in the two situations (this is better than using κ sampling variance directly). This should give an indication of the reliability of your measures, or in this case diagnostic agreement, at the two occasions. A **McNemar test** which tests for marginal homogeneity in matched pairs can also be used.

An approach based on the **intraclass correlation** is still valid and, provided your prevalence is not extreme, should be closed to

- a simple Pearson correlation (which, for binary data, is also called a **phi coefficient**) or the tetrachoric version suggested by @Skrikant,
- the aforementioned kappa (for a large sample, and assuming that the marginal distributions for case at the two occasions are the same, $\kappa \approx \text{ICC}$ from a one-way ANOVA).

About your bonus question, you generally need 3 time points to separate the lack of (temporal) stability – which can occur if the latent class or trait you are measuring evolve over time – from the lack of reliability (see for an illustration the model proposed by **Wiley and Wiley**, 1970, *American Sociological Review* 35).

127 Reorder categorical data in ggplot2

Would that help?

```
x <- gl(2, 20, 40, labels=c("K0", "WT"))
y <- rnorm(40)
qplot(x, y)
qplot(relevel(x, "WT"), y)
```

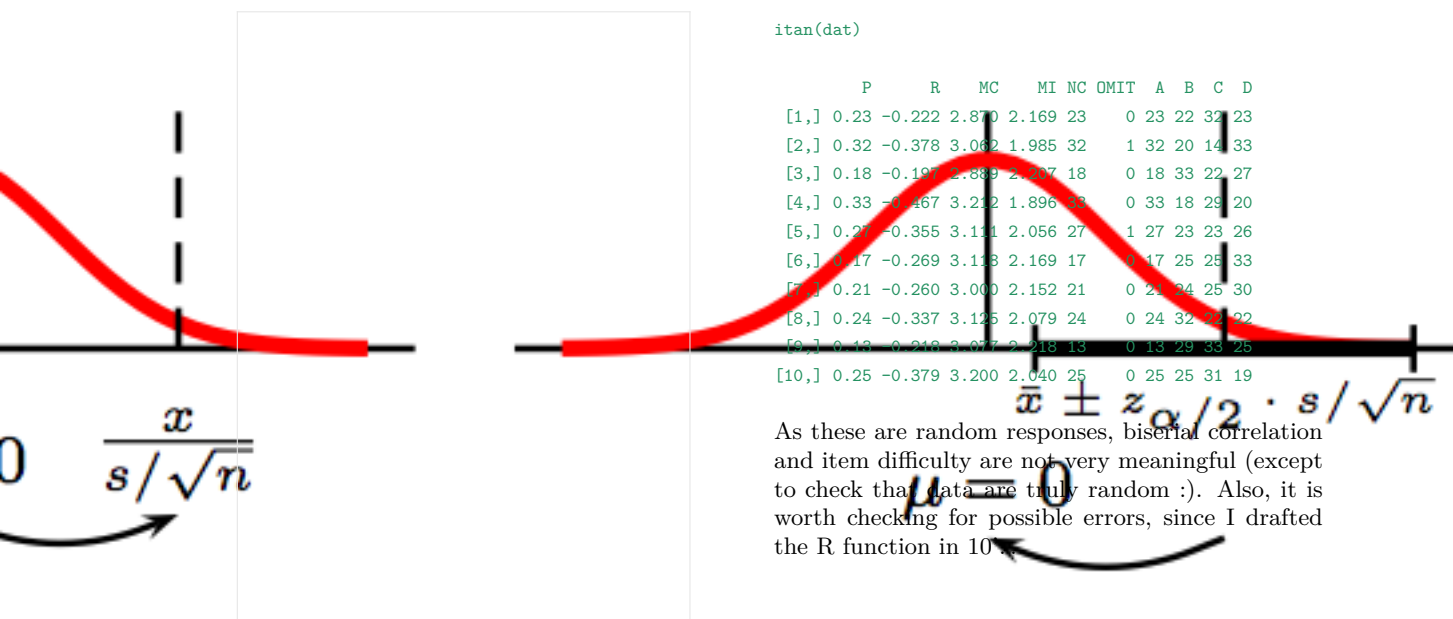
128 When are confidence intervals useful?

I like to think of CIs as some way to escape the Hypothesis Testing (HT) framework, at least the binary decision framework following **Neyman's** approach, and keep in line with theory of measurement in some way. More precisely, I view them as more close to the reliability of an estimation (a difference of means, for instance), and conversely HT are more close to hypothetico-deductive reasoning, with its pitfalls (we cannot accept the null, the alternative is often stochastic, etc.). Still, with both interval estimation and HT we have to rely on distribution assumptions most of the time (e.g. a sampling distribution under H_0), which allows to make inference from our sample to the general population or a representative one (at least in the frequentist approach).

In many context, CIs are complementary to usual HT, and I view them as in the following picture (it is under H_0):

that is, under the HT framework (left), you look at how far your statistic is from the null, while with CIs (right) you are looking at the null effect “from your statistic”, in a certain sense.

Also, note that for certain kind of statistic, like odds-ratio, HT are often meaningless and it is better to look at its associated CI which is asymmetrical and provide more relevant information as to the direction and precision of the association, if any.



129 Item Analysis for an R newbie

I can suggest you at least two packages that allow to perform these tasks: `psych` (`score.items`) and `ltm` (`descript`). The `CTT` package seems also to process MCQ but I have no experience with it. More information can be found on W Revelle's website, [The Personality Project](#), esp. the page dedicated to [psychometrics with R](#) which provides step-by-step instructions for importing, analyzing and report data. Also, the CRAN Task View on [Psychometrics](#) includes many additional resources.

As described in your link, MC stands for "Mean total raw score of the persons who answered the item with the correct response", and MI for "Mean total score of the persons who did not answer the item with the correct response". Point-biserial correlation ($R(IT)$) is also available in the `ltm` package (`biserial.cor`). This is basically an indicator of the discrimination power of the item (since it is the correlation of item and total score), and is related to the discrimination parameter of a 2-PL IRT model or factor loading in Factor Analysis.

If you really want to reproduce the table you show, I guess you will have to wrap some of this code with custom code, at least to output the same kind of table. I've made a [quick and dirty example](#) which reproduce your table:

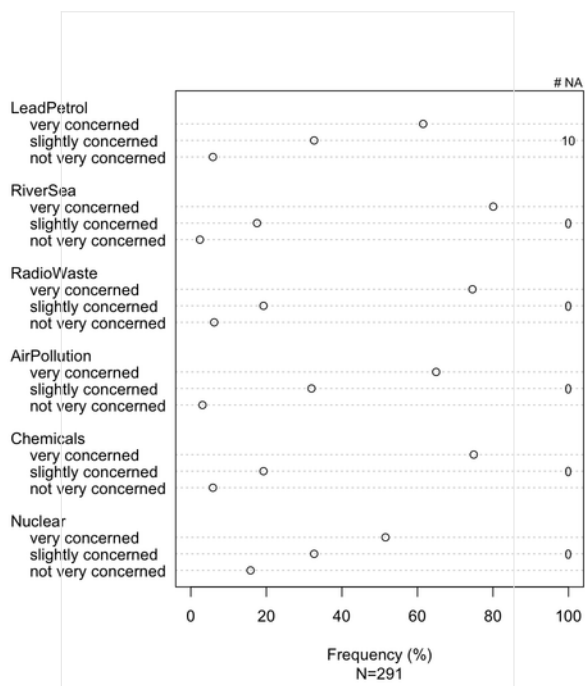
```
dat <- replicate(10, sample(LETTERS[1:4], 100, rep=TRUE))
dat[3,2] <- dat[67,5] <- NA
```

130 Visualizing Likert Item Response Data

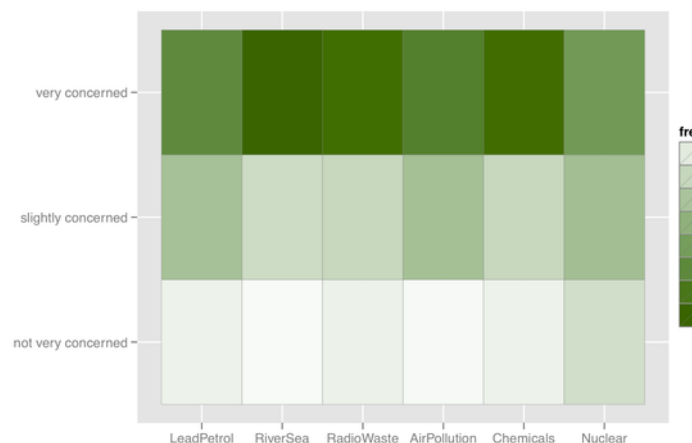
Stacked barcharts are generally well understood by non-statisticians, provided they are gently introduced. It is useful to scale them on a common metric (e.g. 0-100%), with a gradual color for each category if these are ordinal item (e.g. Likert). I prefer `dotchart` (Cleveland dot plot), when there are not too many items and no more than 3-5 responses categories. But it is really a matter of visual clarity. I generally provide % as it is a standardized measure, and only report both % and counts with non-stacked barchart. Here is an example of what I mean:

```
data(Environment, package="ltm")
Environment[sample(1:nrow(Environment), 10),1] <- NA
na.count <- apply(Environment, 2, function(x) sum(is.na(x)))
tab <- apply(Environment, 2, table)/
  apply(apply(Environment, 2, table), 2, sum)*100
dotchart(tab, xlim=c(0,100), xlab="Frequency (%)",
  sub=paste("N", nrow(Environment), sep=""))
text(100, c(2,7,12,17,22,27), rev(na.count), cex=.8)
mtext("# NA", side=3, line=0, at=100, cex=.8)
```

Better rendering could be achieved with `lattice` or `ggplot2`. All items have the same response categories in this particular example, but in more general case we might expect different ones, so that showing all of them would not seem redundant as is the case here. It would be possible, however, to give the same color to each response category so as to facilitate reading.



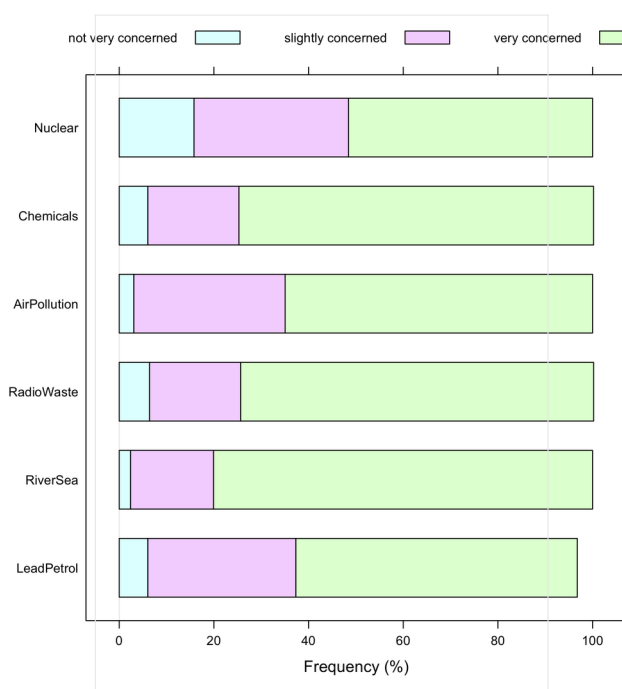
useful if there are many items with similar response



category.

Missing responses (esp. when non negligible or localized on specific item/question) should be reported, ideally for each item. Generally, % of responses for each category are computed without NA. This is what is usually done in survey or psychometrics (we speak of “expressed or observed responses”).

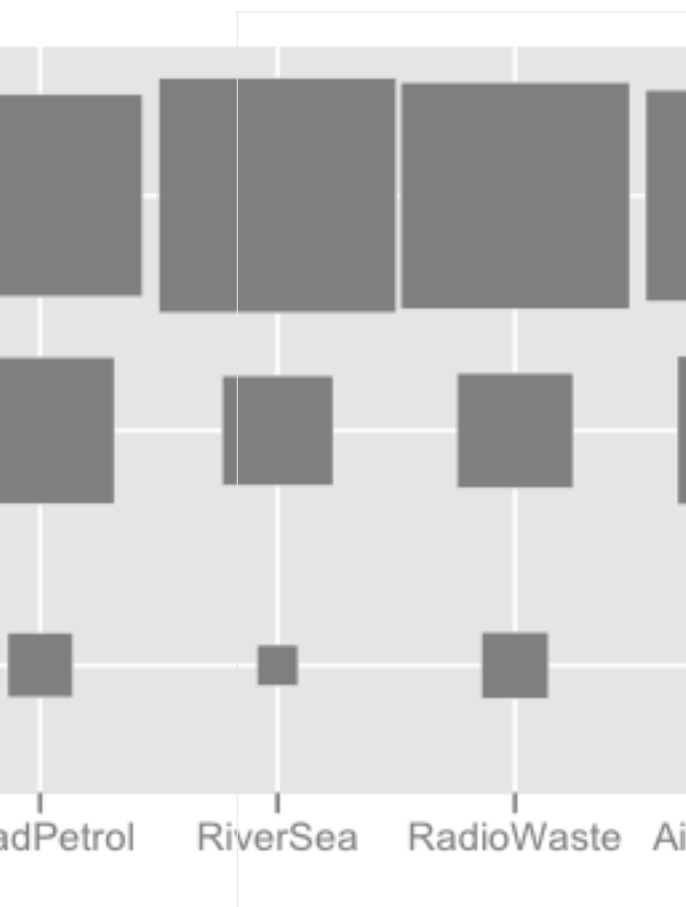
But I would say stacked barcharts are better when all items have the same response category, as they help to appreciate the frequency of one response modality across items:



P.S. I can think of more fancy things like the picture shown below (the first one was made by hand, the second is from `ggplot2`, `ggfluctuation(as.table(tab))`), but I don't think it convey as accurate information as dotplot or barchart since surface variations are



I can also think of some kind of heatmap, which is difficult to appreciate.



esting in this respect.

Of note, Kendall's tau (the a version) has connection to Somers' D (and Harrell's C) used for predictive modelling (see e.g., [Interpretation of Somers' D under four simple models](#) by RB Newson and reference 6 therein, and articles by Newson published in the Stata Journal 2006). An overview of rank-sum tests is provided in [Efficient Calculation of Jackknife Confidence Intervals for Rank Statistics](#), that was published in the JSS (2006).

132 What is your favorite, easy to use statistical analysis website or software package?

Your objectives seem rather vague, but I think the open-source [R statistical package](#) should fit your needs, and beyond. Although primarily a command line driven software, you will find several useful GUIs, e.g. [Rcommander](#) or [deducer](#) to help you start with.

The [CRAN](#) website contains everything you need to start with R, including a lot of [official](#) and [contributed](#) documentation.

R is made of several additional packages (a kind of extensions to the core statistical functions), and you will find interesting pointers on these related questions: [What R packages do you find most useful in your daily work?](#), [I just installed the latest version of R. What packages should I obtain?](#).

131 Kendall Tau or Spearman's rho?

I found that Spearman correlation is mostly used in place of usual linear correlation when working with integer valued scores on a measurement scale, when it has a moderate number of possible scores or when we don't want to make rely on assumptions about the bivariate relationships. As compared to Pearson coefficient, the interpretation of Kendall's tau seems to me less direct than that of Spearman's rho, in the sense that it quantifies the difference between the % of concordant and discordant pairs among all possible pairwise events. In my understanding, Kendall's tau more closely resembles [Goodman-Kruskal Gamma](#).

I just browsed an article from Larry Winner in the J. Statistics Educ. (2006) which discusses the use of both measures, [NASCAR Winston Cup Race Results for 1975-2003](#).

I also found [@onestop](#) answer about [Pearson's or Spearman's correlation with non-normal data](#) inter-

133 AIC and SC value

It is quite difficult to answer your question in a precise manner, but it seems to me you are comparing two criteria (information criteria and p-value) that don't give the same information. For all information criteria (AIC, or Schwarz criterion), the smaller they are the better the fit of your model is (from a statistical perspective) as they reflect a trade-off between the lack of fit and the number of parameters in the model; for example, the Akaike criterion reads $-2 \log(\ell) + 2k$, where k is the number of parameters. However, unlike AIC, SC is consistent: the probability of choosing incorrectly a bigger model converges to 0 as the sample size increases. They are used for comparing models, but you can well observe a model with significant predictors that provide poor fit (large residual deviance). If you can

achieve a different model with a lower AIC, this is suggestive of a poor model. And, if your sample size is large, p -values can still be low which doesn't give much information about model fit. At least, look if the AIC shows a significant decrease when comparing the model with an intercept only and the model with covariates. However, if your interest lies in finding the best subset of predictors, you definitely have to look at methods for variable selection.

I would suggest to look at *penalized regression*, which allows to perform variable selection to avoid overfitting issues. This is discussed in Frank Harrell's Regression Modeling Strategies (p. 207 ff.), or Moons et al., [Penalized maximum likelihood estimation to directly adjust diagnostic and prognostic prediction models for overoptimism: a clinical example](#), J Clin Epid (2004) 57(12).

See also the [Design](#) (`lrm`) and [stepPlr](#) (`step.plr`) R packages, or the [penalized](#) package. You may browse related questions on *variable selection* on this SE.

134 Sequential hypothesis testing in basic science

I don't know much of sequential tests and their application outside of interim analysis (Jennison and Turnbull, 2000) and computerized adaptive testing (van der Linden and Glas, 2010). One exception is in some fMRI studies that are associated to large costs and difficulty to enroll subjects. Basically, in this case sequential testing primarily aims at stopping the experiment earlier. So, I am not surprised that these very tailored approaches are not taught in usual statistical classes.

Sequential tests are not without their pitfalls, though (type I and II error have to be specified in advance, choice of the stopping rule and multiple look at results should be justified, p -values are not uniformly distributed under the null as in a fixed sample design, etc.). In most design, we work with a pre-specified experimental setting or a preliminary power study was carried out, to optimize some kind of cost-effectiveness criterion, in which case standard testing procedures apply.

I found, however, the following paper from Maik Dierkes about fixed vs. open sample design very interesting: [A claim for sequential designs of experiments](#).

135 Can cross validation be used for causal inference?

It seems to me that your question more generally addresses different flavour of validation for a predictive model: Cross-validation has somewhat more to do with *internal validity*, or at least the initial modelling stage, whereas drawing causal links on a wider population is more related to *external validity*. By that (and as an update following @Brett's nice remark), I mean that we usually build a model on a working sample, assuming an hypothetical conceptual model (i.e. we specify the relationships between predictors and the outcome(s) of interest), and we try to obtain reliable estimates with a minimal classification error rate or a minimal prediction error. Hopefully, the better the model performs, the better it will allow us to predict outcome(s) on unseen data; still, CV doesn't tell anything about the "validity" or adequacy of the hypothesized causal links. We could certainly achieve decent results with a model where some moderation and/or mediation effects are neglected or simply not known in advance.

My point is that whatever the method you use to validate your model (and holdout method is certainly not the best one, but still it is widely used in epidemiological study to alleviate the problems arising from stepwise model building), you work with the same sample (which we assume is representative of a larger population). On the contrary, generalizing the results and the causal links inferred this way to new samples or a plausibly related population is usually done by *replication studies*. This ensures that we can safely test the predictive ability of our model in a "superpopulation" which features a larger range of individual variations and may exhibit other potential factors of interest.

Your model might provide valid predictions for your working sample, and it includes all potential confounders you may have think of; however, it is possible that it will not perform as well with new data, just because other factors appear in the intervening causal path that were not identified when building the initial model. This may happen if some of the predictors and the causal links inferred therefrom depend on the particular trial centre where patients were recruited, for example.

In genetic epidemiology, many [genome-wide association studies](#) fail to replicate just because we are trying to model complex diseases with an oversimplified view on causal relationships between DNA markers and

the observed phenotype, while it is very likely that gene-gene (epistasis), gene-diseases (pleiotropy), gene-environment, and population substructure all come into play, but see for example [Validating, augmenting and refining genome-wide association signals](#) (Ioannidis et al., Nature Reviews Genetics, 2009 10). So, we can build-up a performant model to account for the observed cross-variations between a set of genetic markers (with very low and sparse effect size) and a multivariate pattern of observed phenotypes (e.g., volume of white/gray matter or localized activities in the brain as observed through fMRI, responses to neuropsychological assessment or personality inventory), still it won't perform as expected on an independent sample.

As for a general reference on this topic, can recommend chapter 17 and Part III of [Clinical Prediction Models](#), from EW Steyerberg (Springer, 2009). I also like the following article from Ioannidis:

Ioannidis, JPA, [Why Most Published Research Findings Are False?](#) PLoS Med. 2005 2(8): e124

136 Intra-class correlation

These are distinct ways of accounting for raters or items variance in overall variance, following [Shrout and Fleiss \(1979\)](#) (cases 1 to 3 in Table 1):

- *One-way random effects model*: raters are considered as sampled from a larger pool of potential raters, hence they are treated as random effects; the ICC is then interpreted as the % of total variance accounted for by subjects/items variance. This is called the consistency ICC.
- *Two-way random effects model*: both factors – raters and items/subjects – are viewed as random effects, and we have two variance components (or mean squares) in addition to the residual variance; we further assume that raters assess all items/subjects; the ICC gives in this case the % of variance attributable to raters + items/subjects.
- *Two-way mixed model*: contrary to the one-way approach, here raters are considered as fixed effects (no generalization beyond the sample at hand) but items/subjects are treated as random effects; the unit of analysis may be the individual or the average ratings.

I would say raters have to be entered as columns,

although I'm not a specialist of SPSS. Dave Garson's [dedicated website](#) is worth looking at for those working with SPSS. There is also a complete on-line tutorial on [reliability analysis](#) (Robert A. Yaffee).

For theoretical consideration about the mixed-effect approach, please consider reading my answer to this related question: [Reliability in Elicitation Exercise](#).

137 Chi-square test for equality of distributions: how many zeroes does it tolerate?

The usual guidelines are that the expected counts should be greater than 5, but it can be somewhat relaxed as discussed in the following article:

Campbell , I, [Chi-squared and Fisher-Irwin tests of two-by-two tables with small sample recommendations](#), Statistics in Medicine (2007) 26(19): 3661–3675.

See also Ian Campbell's [homepage](#).

Note that in R, there's always the possibility to compute *p*-value by a Monte Carlo approach (`chisq.test(..., sim=TRUE)`), instead of relying on the asymptotic distribution.

In your case, it appears that about 80% of the expected counts are below 5, and 40% are below 1. Would it make sense to aggregate some of the observed phenotypes?

138 Testing nonlinearity in logistic regression

I would suggest to use restricted cubic splines (`rCs` in R, see the [Hmisc](#) and [Design](#) packages for examples of use), instead of adding power of *X* in your model. This approach is the one that is recommended by Frank Harrell, for instance, and you will find a nice illustration in his handouts (§2.5 and chap. 9) on *Regression Modeling Strategies* (see the [companion website](#)).

You can compare the results with your Box-Tidwell test by using the `boxTidwell()` in the [car](#) package.

Transforming continuous predictors into categorical ones is generally not a good idea, see e.g. [Problems Caused by Categorizing Continuous Variables](#).

139 How to perform t-test with huge samples?

The t distribution tend to the z (gaussian) distribution when n is large (in fact, when $n > 30$, they are almost identical, see the picture provided by @on-estop). In your case, I would say that n is VERY large, so that you can just use a z -test. As a consequence of the sample size, any VERY small differences will be declared significant. So, it is worth asking yourself if these tests (with the full data set) are really interesting.

Just to be sure, as your data set includes 25 variables, you are making 25 tests? If this is the case, you probably need to correct for multiple comparisons so as not to inflate the type I error rate (see related thread on this site).

BTW, the R software would gives you the p-values you are looking for, no need to rely on Tables:

```
> x1 <- rnorm(n=38704)
> x2 <- rnorm(n=1313662, mean=.1)
> t.test(x1, x2, var.equal=TRUE)
```

Two Sample t-test

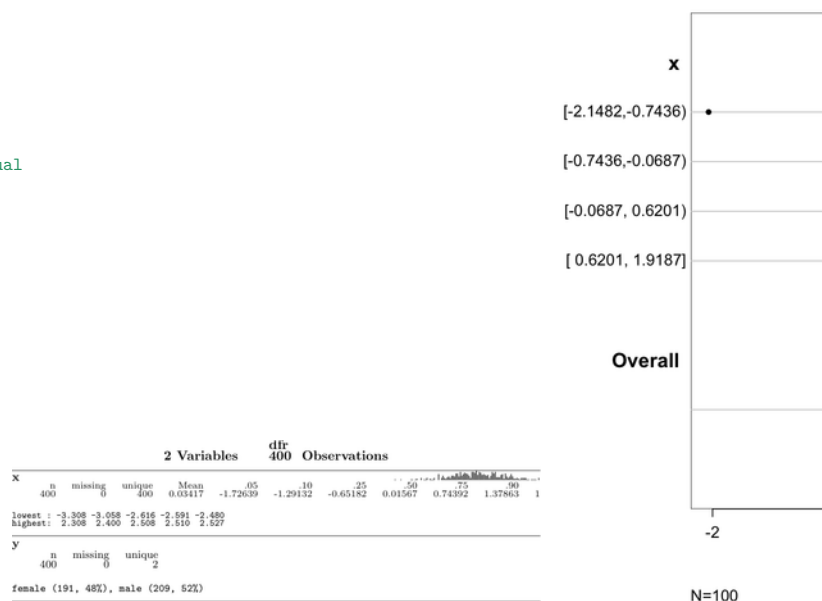
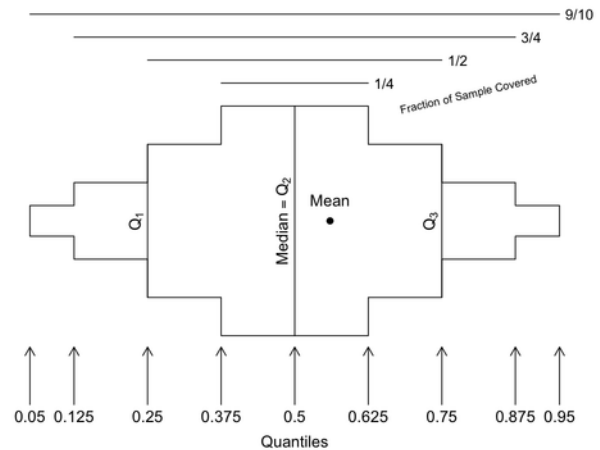
```
data: x1 and x2
t = -17.9156, df = 1352364, p-value < 2.2e-16
alternative hypothesis: true difference in means is not equal
to 0
95 percent confidence interval:
 -0.1024183 -0.0822190
sample estimates:
 mean of x mean of y
0.007137404 0.099456039
```

140 Graphical data overview (summary) function in R

Frank Harrell's **Hmisc** package has some basic graphics with options for annotation. But I think that was you're looking for is in the **Design** package: check the `summary.formula()` and related `plot` wrap functions. I also like the `describe()` function.

For additional information, have a look at the [The Hmisc Library](#) or [An Introduction to S-Plus and the Hmisc and Design Libraries](#).

Here are some pictures taken from the on-line help (`bpplt`, `describe`, and `plot(summary(...))`):



Many other examples can be browsed on-line on the [R Graphical Manual](#), see [Hmisc](#) and [Design](#).

141 Interpreting PCA scores

Basically, the factor scores are computed as the

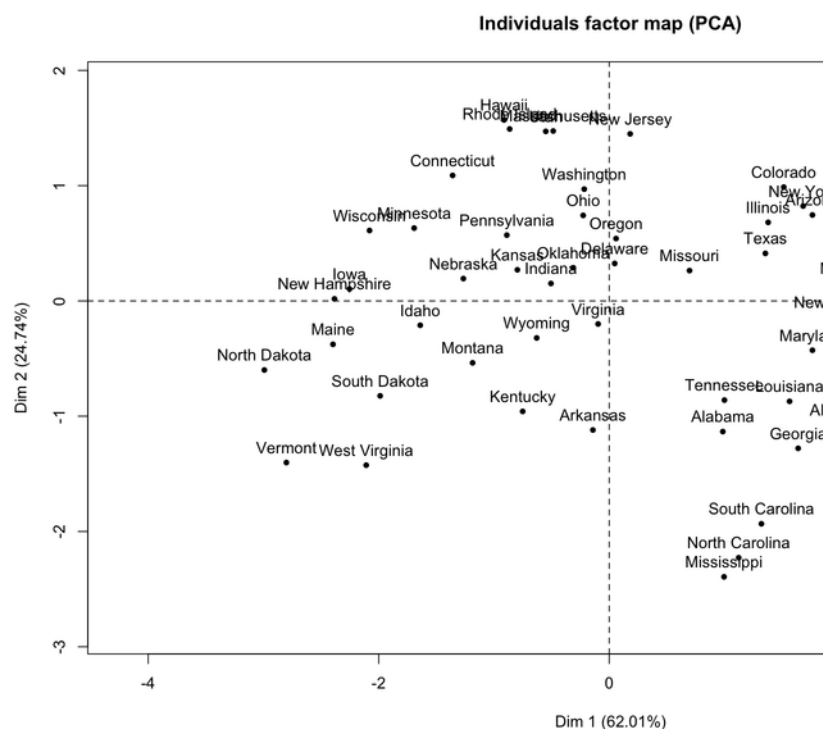
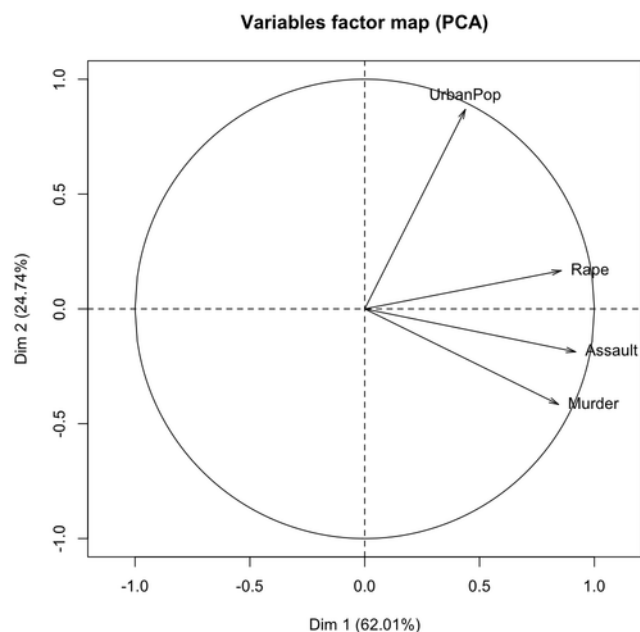
raw responses weighted by the factor loadings. So, you need to look at the factor loadings of your first dimension to see how each variable relate to the principal component. Observing high positive (resp. negative) loadings associated to specific variables means that these variables contribute positively (resp. negatively) to this component; hence, people scoring high on these variables will tend to have higher (resp. lower) factor scores on this particular dimension.

Drawing the correlation circle is useful to have a general idea of the variables that contribute “positively” vs. “negatively” (if any) to the first principal axis, but if you are using R you may have a look at the **FactoMineR** package and the `dimdesc()` function.

Here is an example with the **USArrests** data:

```
> data(USArrests)
> library(FactoMineR)
> res <- PCA(USArrests)
> dimdesc(res, axes=1) # show correlation of variables with
1st axis
$Dim.1
$Dim.1$quanti
      correlation p.value
Assault      0.918 5.76e-21
Rape         0.856 2.40e-15
Murder       0.844 1.39e-14
UrbanPop     0.438 1.46e-03
> res$var$coord # show loadings associated to each axis
      Dim.1 Dim.2 Dim.3 Dim.4
Murder  0.844 -0.416  0.204  0.2704
Assault  0.918 -0.187  0.160 -0.3096
UrbanPop 0.438  0.868  0.226  0.0558
Rape     0.856  0.166 -0.488  0.0371
```

As can be seen from the latest result, the first dimension mainly reflects violent acts (of any kind). If we look at the individual map, it is clear that states located on the right are those where such acts are most frequent.



You may also be interested in this related question:
[What are principal component scores?](#)

142 Cross tabulation of two categorical variables: recommended techniques

Arguably, the question is not very precise. Rather than enumerating all measures of association for 2×2 tables, I shall concentrate on the way such measures may be constructed and how to select the one that is most appropriate with respect to hypothesis or constraints relevant to a cross-classification.

The very first questions to ask are: what does the table reflect (concordance, agreement, association between two attributes, etc.), do you seek an overall measure of association or do you think one of the two variables plays a specific role (which would justify the search for an “oriented” association), do you consider either or both of the margins fixed (row and/or columns totals)? All of this impact on the method to choose and the way to interpret the results.

The 2×2 case

Two-by-two tables are often treated separately from $I \times J$ tables because we often consider that variables play a symmetric role in this particular case. Obviously, this is not always the case: cross-classification of exposure and disease, as commonly found in epidemiological studies, is an example where both variables play a distinct role, which may lend to more than a simple interpretation in terms of association. Another one is 2×2 tables constructed for studying the screening properties of a given diagnostic instrument. Although the odds-ratio (compared to, e.g. the relative risk) keeps its nice properties, we may be interested in predictive/negative positive values or specificity/sensibility, which means working with other quantities of interest. Hence, the need to specify whether the problem at hand implies two variables that are purely acting in a symmetrical way, or not, because it influences the way we interpret the results or derive a useful measure of association, agreement, or discrimination.

For the sake of clarity, I will consider that data (counts) are arranged in the following way:

p_{11}	p_{12}	$p_{1\cdot}$
p_{21}	p_{22}	$p_{2\cdot}$
$p_{\cdot 1}$	$p_{\cdot 2}$	

Basically, measures of association for 2×2 tables can be grouped in two classes: those relying on (a) (a function of) the cross-product ratio and those based on (b) the product-moment (Pearson) correlation,

or a function thereof.

The cross-product ratio, mostly known as the odds-ratio, is simply $\alpha = p_{11}p_{22}/p_{12}p_{21}$. It is invariant under rows *and* columns interchange, and transformations of margins that preserves $\sum_{i,j} p_{ij} = 1$. In epidemiology, we usually think of it as a measure of association where rows (or columns) are fixed: p_{11}/p_{12} is then the odds of being in the first column (e.g., diseased) conditional on being in the first row (e.g., exposed), and likewise p_{21}/p_{22} is the odds for the second row, or in other words

$$\alpha = \frac{p_{11}/p_{12}}{p_{21}/p_{22}}.$$

Yule’s $Q = (\alpha - 1)/(\alpha + 1)$ fall into the former case, (a). Yule also proposed a measure of “colligation”, Y , as $(\sqrt{\alpha} - 1)/(\sqrt{\alpha} + 1)$. Yule’s Q can be interpreted as the difference between conditional probabilities of like and unlike “orders” for two individuals chosen at random; it is identical to Goodman and Kruskal’s γ measure of association for $I \times J$ tables.

For (b), we can derive a correlation coefficient for a 2×2 table by thinking of the table as a combination of each of two variables scores (taking value 0 and 1, for the first and second row/column, resp.). Then, the coefficient ρ is defined as the covariance divided by the square root of the product of the variances:

$$\rho = \frac{p_{22} - p_{2\cdot}p_{\cdot 2}}{\sqrt{p_{1\cdot}p_{2\cdot}p_{\cdot 1}p_{\cdot 2}}},$$

which is equivalent to putting $p_{11}p_{22} - p_{21}p_{12}$ in the numerator. Plugging in the observed counts, Pearson’s r is the MLE of ρ under a multinomial sampling model. It is invariant under rows *and* columns interchange, and positive linear transformation.

It can be shown (Yule, 1912) that ρ is identical to Yule’s Y if we standardize our table such that row and column margins sum to 1/2, i.e. $p_{11}^* = p_{22}^* = 0.5(\sqrt{\alpha}/(\sqrt{\alpha} + 1))$ and $p_{12}^* = p_{21}^* = 0.5(1/(\sqrt{\alpha} + 1))$. By doing this, we remove the information coming from the margins, such that $Y = 2(p_{11}^* - p_{12}^*)$.

Correlation-based measures are connected to the usual Pearson’s chi-square statistic, since

$$\Phi^2 = \sum_{i=1}^2 \sum_{j=1}^2 \frac{(p_{ij} - p_{i\cdot}p_{\cdot j})^2}{p_{i\cdot}p_{\cdot j}},$$

that is,

$$\Phi^2 = \frac{(p_{11}p_{22} - p_{21}p_{12})^2}{p_{1.}p_{2.}p_{.1}p_{.2}} = \rho^2.$$

In a 2×2 table, we thus have $r^2 = \chi^2/N$. Pearson also proposed to use $\sqrt{\rho^2/(1 + \rho^2)}$ as a measure of association, and he coined it the *coefficient of mean square contingency*.

As to how to choose the correct measure (a vs. b), it clearly depends on whether we want to be sensitive to marginal totals (in this case, ρ cannot take its full range of possible values in $[-1; 1]$), and whether we consider that we observe a full association even if one of the four cells is zero (in this case, ρ cannot take the value $+1$ or -1 if only one of the cells is zero, which is not the case of Yule's Q). Of note, correlation-based measures are better if they are used in a correlation matrix (e.g., for factor analysis), because we cannot guarantee that a matrix composed of Yule's Q coefficient will be positive definite.

The $I \times J$ case

Like for the 2×2 case, we can derive measures of association based on different quantities. Measures based on chi-square include

- Pearson's P coefficient based on Φ^2 (see above), $\sqrt{\Phi^2/(\Phi^2 + 1)}$ (to overcome the fact that Φ^2 no longer lies in $[0; 1]$ when I or $J > 2$);
- Tschuprow's $T = \left(\Phi^2/\sqrt{(I-1)(J-1)}\right)^{1/2}$, which behaves better than P in square tables (in that it can reach a maximum value of 1, for full or complete association);
- Cramer's V is another derivation, and $V = \left(\Phi^2/\min(I-1, J-1)\right)^{1/2}$ (we have $V \geq T$ for all $I, J > 2$).

These measures are all measures of association where none of the variables plays a specific role. In case a χ^2 test is significant, it is more interesting to look at how the expected counts depart from the observed counts (i.e. look at the Pearson residuals) in all (i, j) cells, or to use something like a **mosaic plot**.

Goodman and Kruskal (1954) also proposed a predictive measure of association between rows and columns, or more specifically a measure of proportional reduction in error in predicting one column category when the row category is known as op-

posed to the case when the latter one is unknown. This is called $\lambda_{C|R}$ and its MLE is

$$\hat{\lambda}_{C|R} = \frac{\sum_{i=1}^I x_{im} - x_{.m}}{N - x_{.m}}$$

where x_{im} and $x_{.m}$ stand for the maximum for the i th row and the column totals. This measure is interesting because it has a nicer interpretation than χ^2 -based measure, but it also has some drawbacks: when there is statistical independence, $\lambda_{C|R}$ is not necessarily zero, for instance.

A measure of the proportion of explained variance (derived from Gini's total variation) may be derived from the total sum of squares (SS) in an $I \times J$ table

$$\text{TSS} = \frac{N}{2} - \frac{1}{2N} \sum_{i=1}^I x_i^2,$$

which can be partitioned as a within- and between-group SS. Of interest here is the variance explained by considering the different categories (BSS) divided by the total variance, TSS. Like in the ANOVA framework, we have $\text{BSS} = \text{TSS} - \text{WSS}$, where

$$\text{WSS} = \frac{N}{2} - \frac{1}{2} \sum_{j=1}^J \frac{1}{x_{.j}} \sum_{i=1}^I x_{ij}^2,$$

so that we can derive BSS/TSS as

$$\hat{\tau}_{R|C} = \frac{\sum_j \frac{1}{x_{.j}} \sum_i x_{ij}^2 - \frac{1}{N} \sum_i x_i^2}{N - \frac{1}{N} \sum_i x_i^2}.$$

This measure can be interpreted as “the relative decrease in the proportion of incorrect predictions when we go from predicting the row category based only on the row marginal probabilities to predicting the row category based on the conditional proportions $p_{ij}/p_{.j}$ ” (Bishop et al., 2007, p. 391).

Finally, measures based on the cross-product ratios are also available, as well as measures of agreement for ordinal variables, but I realize now that I need to stop (and thank the reader who reached the end of this overview).

A thorough overview of measures of association may be found in Bishop et al. (2007), from which I grabbed most of the above discussion, and of course Agresti (2002), about which Laura Thompson made a complete R adaptation in his textbook **R (and S-PLUS) Manual to Accompany Agresti's Categorical Data Analysis**.

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143 Does it ever make sense to treat categorical data as continuous?

I will assume that a “categorical” variable actually stands for an ordinal variable, otherwise it doesn’t make much sense to treat it as a continuous one, unless it’s a binary variable (coded 0/1) as pointed by @Rob. Then, I would say that the problem is not that much the way we treat the variable, although many models for categorical data analysis have been developed so far—see e.g., [The analysis of ordered categorical data: An overview and a survey of recent developments](#) from Liu and Agresti—, than the underlying measurement scale we assume. My response will focus on this second point, although I will first briefly discuss the assignment of numerical scores to variable categories or levels.

By using a simple numerical recoding of an ordinal variable, you are assuming that the variable has interval properties (in the sense of the classification given by Stevens, 1946). From a measurement theory perspective (in psychology), this may often be a too strong assumption, but for basic study (i.e. where a single item is used to express one’s opinion about a daily activity with clear wording) any monotone scores should give comparable results. Cochran (1954) already pointed that

any set of scores gives a *valid* test, provided that they are constructed without consulting the results of the experiment. If the set of scores is poor, in that it badly distorts a numerical scale that really does underlie the ordered classification, the test will not be sen-

sitive. The scores should therefore embody the best insight available about the way in which the classification was constructed and used.

(Many thanks to @whuber for reminding me about this throughout one of his comments, which lead me to re-read Agresti’s book from which this citation comes from)

Actually, several tests treat implicitly such variables as interval scales; for example, the M^2 statistic for testing a linear trend (as an alternative to simple independence) is based on a correlational approach ($M^2 = (n - 1)r^2$, Agresti, 2002, p. 87).

Well, you can also decide to recode your variable on an irregular range, or aggregate some of its levels, but in this case strong imbalance between recoded categories may distort statistical tests, e.g. the aforementioned trend test. A nice alternative for assigning distance between categories was already proposed by @Jeromy, namely optimal scaling.

Now, let’s discuss the second point I made, that of the underlying measurement model. I’m always hesitating about adding the “psychometrics” tag when I see this kind of question, because the construction and analysis of measurement scales come under Psychometric Theory (Nunnally and Bernstein, 1994, for a neat overview). I will not dwell on all the models that are actually headed under the [Item Response Theory](#), and I kindly refer the interested reader to I. Partchev’s tutorial, [A visual guide to item response theory](#), for a gentle introduction to IRT, and to references (5-8) listed at the end for possible IRT taxonomies. Very briefly, the idea is that rather than assigning arbitrary distances between variable categories, you assume a latent scale and estimate their location on that continuum, together with individuals ability or liability. A simple example is worth several mathematical notation, so let’s consider the following item (coming from the [EORTC QLQ-C30](#) health-related quality of life questionnaire):

Did you worry?

which is coded on a four-point scale, ranging from “Not at all” to “Very much”. Raw scores are computed by assigning a score of 1 to 4. Scores on items belonging to the same scale can then be added together to yield a so-called scale score, which denotes one’s *rank* on the underlying construct (here, a mental health component). Such summated scale scores are very practical because of scoring easiness

(for the practitioner or nurse), but they are nothing more than a discrete (ordered) scale.

We can also consider that the probability of endorsing a given response category obeys some kind of a logistic model, as described in I. Partchev's tutorial, referred above. Basically, the idea is that of a kind of threshold model (which lead to equivalent formulation in terms of the proportional or cumulative odds models) and we model the odds of being in one response category rather the preceding one or the odds of scoring above a certain category, conditional on subjects' location on the latent trait. In addition, we may impose that response categories are equally spaced on the latent scale (this is the Rating Scale model)—which is the way we do by assigning regularly spaced numerical scores— or not (this is the Partial Credit model).

Clearly, we are not adding very much to Classical Test Theory, where ordinal variable are treated as numerical ones. However, we introduce a probabilistic model, where we assume a continuous scale (with interval properties) and where specific errors of measurement can be accounted for, and we can plug these factorial scores in any regression model.

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Modeling: The eRm Package for the Application of IRT Models in R. *Journal of Statistical Software*, **20**(9), 2007.

144 Using lmer for prediction

Expressing factors relationships using R formulas follows from Wilkinson's notation, where '*' denotes crossing and '/' nesting, but there are some particularities in the way formula for mixed-effects models, or more generally random effects, are handled. For example, two crossed random effects might be represented as `(1|x1)+(1|x2)`. I have interpreted your description as a case of nesting, much like classes are nested in schools (nested in states, etc.), so a basic formula with `lmer` would look like (unless otherwise stated, a `gaussian` family is used by default):

```
y ~ x + (1|A:B) + (1|A)
```

where A and B correspond to your inner and outer factors, respectively. A is nested in B, and both are treated as random factors. In the older `nlme` package, this would correspond to something like `lme(y ~ x, random=~ 1 | A/B)`. If A was to be considered as a fixed factor, the formula should read `y ~ x + A + (1|A:B)`.

But it is worth checking more precisely D. Bates' specifications for the `lme4` package, e.g. in his forthcoming textbook, *lme4: Mixed-effects Modeling with R*, or the many handouts available on the same webpage. In particular, there is an example for such nesting relations in *Fitting Linear Mixed-Effects Models, the lme4 Package in R*. John Maindonald's tutorial also provides a nice overview: *The Anatomy of a Mixed Model Analysis, with R's lme4 Package*. Finally, section 3 of the R vignette on *lme4 implementation* includes an example of the analysis of a nested structure.

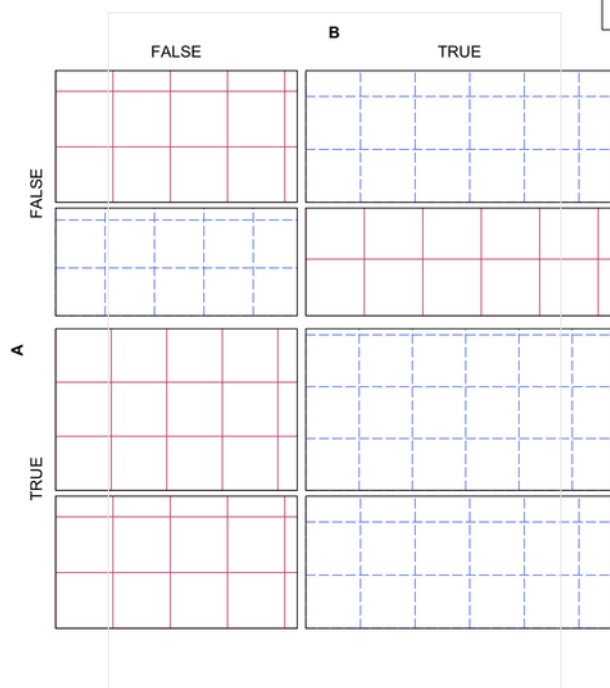
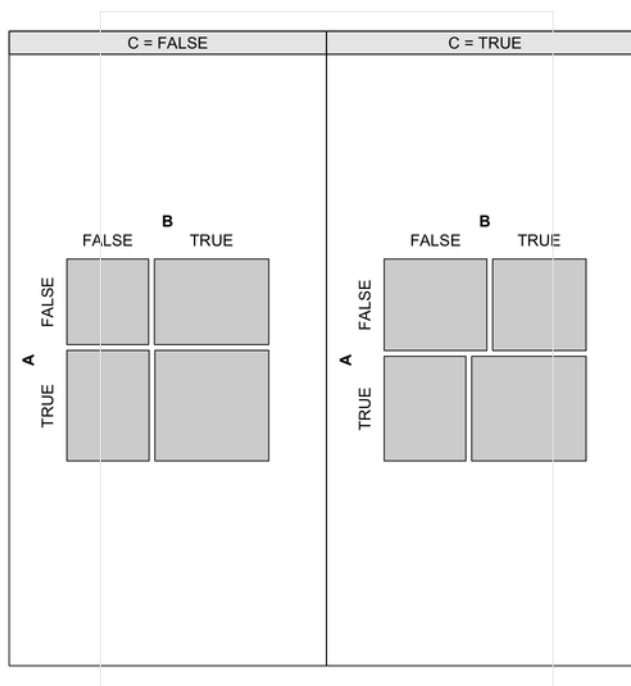
There is no `predict()` function in `lme4`, and you have to compute yourself predicted individual values using the estimated fixed (see `?fixeff`) and random (see `?ranef`) effects, but see also this thread on the *lack of predict function in lme4*. Still, you can generate a sample from the posterior distribution using the `mcmcscamp()` function. Sometimes, it might clash, though. See the *sig-me* mailing list for more updated information.

145 How to visualize 3D contingency matrix?

I would try some kind of 3D heatmap, `mosaic p` or a `sieve plot` (available in the `vcd` package). Is the base `mosaicplot()` function working with three way table? (at least `mosaic3d()` in the `vcdExtra` package should work, see e.g. <http://datavis.ca/R>)

Here's an example (including a conditional plot):

```
A <- sample(c(T,F), 100, replace=T)
B <- sample(c(T,F), 100, replace=T)
C <- sample(c(T,F), 100, replace=T)
tab <- table(A,B,C)
library(vcd)
sieve(tab, shade=TRUE)
cotabplot(tab)
library(vcdExtra)
mosaic3d(tab, type="expected", box=TRUE)
```



Actually, the rendering of `mosaic3d()` rely on the `rgl` package, so it is hard to give a pretty picture of the result.

146 Training a model

Although the `curse of dimensionality` and multi-

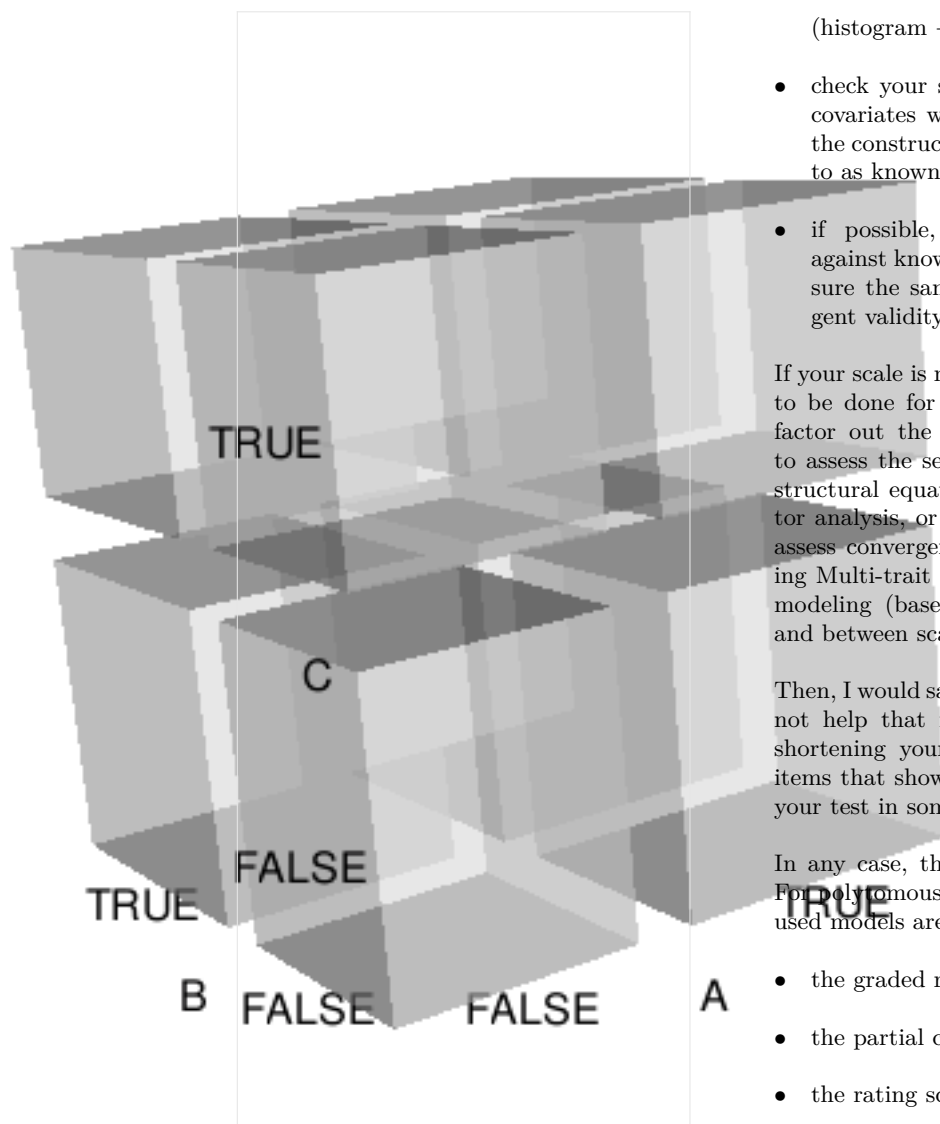
collinearity are distinct issues, cross-validation is used for building a predictive model: we usually estimate parameters of our model on training samples, and assess its generalizability on test samples. This yields a measure of model performance, which can be a % of prediction accuracy if we work with a classification model, or an RMSEA if it is a regression model. The idea is that the better the model performs, the better it will allow us to predict outcome(s) on unseen data. To overcome the problem of overfitting when building a predictive model, we may also introduce some kind of variable or **feature selection**.

Cross-validation may be done in various way (split or holdout method, k-fold, leave-one-out, etc.) but the general idea to keep in mind is that the final model is assessed on individuals who do not participate to its construction.

You may find additional information by looking at the “**cross-validation**” or “**feature selection**” tags.

147 Validating questionnaires

I will assume that your questionnaire is to be considered as one unidimensional scale (otherwise, Cronbach's alpha doesn't make very much sense). It is worth running an exploratory factor analysis to check for that. It will also allow you to see how



items relate to the scale (i.e., through their loadings).

Basic steps for validating your items and your scale should include:

- a complete report on the items' basic statistics (range, quartiles, central tendency, ceiling and floor effects if any);
- checking the internal consistency as you've done with your alpha (best, give 95% confidence intervals, because it is sample-dependent);
- describe your summary measure (e.g., total or mean score, aka scale score) with usual statistics

(histogram + density, quantiles etc.);

- check your summary responses against specific covariates which are supposed to be related to the construct you are assessing – this is referred to as known-group validity;
- if possible, check your summary responses against known instruments that purport to measure the same construct (**concurrent** or convergent validity).

If your scale is not unidimensional, these steps have to be done for each subscale, and you could also factor out the correlation matrix of your factors to assess the second-order factor structure (or use structural equation modeling, or confirmatory factor analysis, or whatever you want). You can also assess convergent and discriminant validity by using Multi-trait scaling or Multi-trait multi-method modeling (based on interitem correlations within and between scales), or, again, SEMs.

Then, I would say that Item Response Theory would not help that much unless you are interested in shortening your questionnaire, filtering out some items that show **differential item functioning**, or use your test in some kind of a **computer adaptive test**.

In any case, the **Rasch model** is for binary items. For polytomous ordered items, the most commonly used models are :

- the graded response model
- the partial credit model
- the rating scale model.

Only the latter two are from the Rasch family, and they basically use an adjacent odds formulation, with the idea that subject has to “pass” several thresholds to endorse a given response category. The difference between these two models is that the PCM does not impose that thresholds are equally spaced on the theta (*ability*, or subject location on the latent trait) scale. The graded response model relies on a cumulative odds formulation. Be aware that these models all suppose that the scale is unidimensional; i.e., there's only one latent trait. There are additional assumptions like, e.g., local independence (i.e., the correlations between responses are explained by variation on the ability scale).

Anyway, you will find a very complete documentation and useful clues to apply psychometric meth-

ods in R in volume 20 of the Journal of Statistical Software: [Special Volume: Psychometrics in R](#). Basically, the most interesting R packages that I use in my daily work are: [ltm](#), [eRm](#), [psych](#), [psy](#). Others are referenced on the CRAN task view [Psychometrics](#). Other resources of interest are:

- [Notes on the use of R for psychology experiments and questionnaires](#)
- [Using R for psychological research](#) (W. Revelle is actually writing a book on [psychometrics in R](#))
- the [PsychoR](#) project (it does not focus on IRT and scale development, though).

A good review on the use of FA vs. IRT in scale development can be found in Scale construction and evaluation in practice: [A review of factor analysis versus item response theory applications](#), by ten Holt et al (Psychological Test and Assessment Modeling (2010) 52(3): 272-297).

148 When to use regularization methods for regression?

Short answer: Whenever you are facing one of these situations: large number of variables or low ratio of no. observations to no. variables (including the $n \ll p$ case), high collinearity, seeking for a sparse solution (i.e., embed feature selection when estimating model parameters), or accounting for variables grouping in high-dimensional data set.

Ridge regression generally yields better predictions than OLS solution, through a better compromise between bias and variance. Its main drawback is that all predictors are kept in the model, so it is not very interesting if you seek a parcimonious model or want to apply some kind of feature selection.

To achieve sparsity, the lasso is more appropriate but it will not necessarily yield good results in presence of high collinearity (it has been observed that if predictors are highly correlated, the prediction performance of the lasso is dominated by ridge regression). The second problem with L1 penalty is that the lasso solution is not uniquely determined when the number of variables is greater than the number of subjects (this is not the case of ridge regression). The last drawback of lasso is that it tends to select only one variable among a group of predictors with high pairwise correlations. In this case, there are alternative solutions like the [group](#)

(i.e., achieve shrinkage on block of covariates, that is some blocks of regression coefficients are exactly zero) or [fused](#) lasso. The [Graphical Lasso](#) also offers promising features for GGMs (see the R [glasso](#) package).

But, definitely, the *elasticnet* criteria, which is a combination of L1 and L2 penalties achieve both shrinkage and automatic variable selection, and it allows to keep $m > p$ variables in the case where $n \ll p$. Following Zou and Hastie (2005), it is defined as the argument that minimizes (over β)

$$L(\lambda_1, \lambda_2, \beta) = \|Y - X\beta\|^2 + \lambda_2 \|\beta\|^2 + \lambda_1 \|\beta\|_1$$

where $\|\beta\|^2 = \sum_{j=1}^p \beta_j^2$ and $\|\beta\|_1 = \sum_{j=1}^p |\beta_j|$.

The lasso can be computed with an algorithm based on coordinate descent as described in the recent paper by Friedman and coll., [Regularization Paths for Generalized Linear Models via Coordinate Descent](#) (JSS, 2010) or the LARS algorithm. In R, the [penalized](#), [lars](#) or [biglars](#), and [glmnet](#) packages are useful packages; in Python, there's the [scikit.learn](#) toolkit, with extensive [documentation](#) on the algorithms used to apply all three kind of regularization schemes.

As for general references, the [Lasso page](#) contains most of what is needed to get started with lasso regression and technical details about L1-penalty, and this related question features essential references, [When should I use lasso vs ridge?](#)

149 Principal Component Analysis among matrices

I don't know if this is exactly what you are looking for (esp. I don't know how large is n and what you intend to do with these results), however I have successfully used [coinertia analysis](#) when I was working with two data sets (same observations in rows), and for more than two data sets there are K-table methods, as implemented in the [ade4](#) R package. [An introduction to K-table analyses](#) outlines the main principles. When the objective is to link two or more Tables, [Generalized Canonical Correlation Analysis](#) is also an option.

It seems to me that you can choose non-euclidean metric, provided it has some meaning for the data at hand and the interpretation of the factorial space. You can see an example with the use of [kdist\(\)](#) in [ade4](#) for applying an PCA on different distance ma-

trices. Jolliffe’s book on *Principal component analysis* should provide additional hints about this (but I didn’t check). There’s also all the work made in the spirit of Gifi on non-linear methods (in R, a lot of packages have been developed by Jan de Leeuw, see the [PsychoR](#) project).

150 What is the difference between effectiveness and efficacy in determining the benefit of therapy ‘A’ on condition ‘B’?

I’m not a specialist of this domain in epidemiological studies, but it seems to me that *efficacy* has to do with the observed effect in a controlled setting, like a randomized clinical trial, whereas *effectiveness* has more to do with a larger range of outcomes or environmental factors (potentially unobserved or non manipulated in the RCT), hence it has a wider scope. At least, I’ve often heard of *cost-effectiveness* studies in pharmacoeconomics, and treatment efficacy (e.g., when comparing two treatment arms).

Quoting this article [Efficacy, effectiveness, efficiency](#),

- efficacy is “the extent to which a drug has the ability to bring about its intended effect under ideal circumstances, such as in a randomised clinical trial”
- effectiveness is “the extent to which a drug achieves its intended effect in the usual clinical setting”

As for other references, I would suggest starting with [Pitfalls of Multisite Randomized Clinical Trials of Efficacy and Effectiveness](#) from Helena C Kraemer (*Schizophrenia Bulletin* 26(3), 2000), and references therein. For example, it is read that “efficacy and effectiveness are opposite extremes on a complex multidimensional continuum of decision making in research design”.

Note

Coming back from the [ISPOR](#) 13th European conference, I’ve heard that the European Federation of Pharmaceutical Industries and Associations ([EFPIA](#)) considers there’s now agreement on the following definitions:

- *relative efficacy* can be defined as the extent to which an intervention does more good than harm, under ideal circumstances, compared to

one or more alternative interventions;

- *relative effectiveness* can be defined as the extent to which an intervention does more good than harm compared to one or more alternatives for achieving the desired results when provided under the usual circumstances of health care practice.

151 What is a good use of the ‘comment’ function in R?

To second @ucfagls, Frank Harrell has developed efficient ways to handle annotated data.frame in R in his [Hmisc](#) package. For example, the [label\(\)](#) and [units\(\)](#) functions allow to add dedicated attributes to R objects. I find them very handy when producing summary of data.frame (e.g., with [describe\(\)](#)).

Another useful way of using such an extra attribute is to apply a timestamp on a data set. I also add an attribute for things like random seed, fold number (when I use k-fold or LOO cross-validation).

152 Can one validly reduce the numbers of items in a published Likert-scale?

Although there is still some information lacking (No. individuals and items per subscale), here are some general hints about scale reduction. Also, since you are working at the questionnaire level, I don’t see why its length matters so much (after all, you will just give summary statistics, like total or mean scores).

I shall assume that (a) you have a set of K items measuring some construct related to morale, (b) your “unidimensional” scale is a second-order factor that might be subdivided into different facets, (c) you would like to reduce your scale to $k < K$ items so as to summarize with sufficient accuracy subjects’ totalled scale scores while preserving the content validity of the scale.

About content/construct validity of this validated scale: The number of items has certainly been chosen so as to best reflect the construct of interest. By shortening the questionnaire, you are actually reducing construct coverage. It would be good to check that the factor structure remains the same when considering only half of the items (which

could also impact the way you select them, after all). This can be done using traditional FA techniques. You hold the responsibility of interpreting the scale in a spirit similar to that of the authors.

About scores reliability: Although it is a sample-dependent measure, scores reliability decreases when decreasing the number of items (cf. [Spearman-Brown formula](#)); another way to see that is that the standard error of measurement (SEM) will increase, but see [An NCME Instructional Module on Standard Error of Measurement](#), by Leo M Harvill. Needless to say, it applies to every indicator that depends on the number of items (e.g., Cronbach's alpha which can be used to estimate one form of reliability, namely the internal consistency). Hopefully, this will not impact any between-group comparisons based on raw scores.

So, my recommendations (the easiest way) would be:

1. Select your items so as to maximise construct coverage; check the dimensionality with FA and coverage with univariate responses distributions;
2. Compare average interitem correlations to previously reported ones;
3. Compute internal consistency for the full scale and your composites; check that they are in agreement with published statistics on the original scale (no need to test anything, these are sample-dependent measures);
4. Test the linear (or polychoric, or rank) correlations between original and reduced (sub)scores, to ensure that they are comparable (i.e., that individuals locations on the latent trait do not vary to a great extent, as objectivated through the raw scores);
5. If you have an external subject-specific variable (e.g., gender, age, or best a measure related to morale), compare [known-group validity](#) between the two forms.

The hard way would be to rely on [Item Response Theory](#) to select those items that carry the maximum of information on the latent trait – scale reduction is actually one of its best application. Models for polytomous items were partly described in this thread, [Validating questionnaires](#).

Update after your 2nd update

1. Forget about any IRT models for polytomous items with so few subjects.
2. Factor Analysis will also suffer from such a low sample size; you will get unreliable factor loadings estimates.
3. 30 items divided by 2 = 15 items (it's easy to get an idea of the increase in the corresponding SEM for the total score), but it will definitively get worse if you consider subscales (this was actually my 2nd question—No. items per subscale, if any)

153 What is the difference between statistics and biostatistics?

When I look at the Wikipedia entry for [biostatistics](#), the relation to *biometrics* doesn't seem so obvious to me since, historically, biometrics was more concerned with characterizing individuals by some phenotypes of interest, with large applications in population genetics (as exemplified by the work of Fisher), whereas part of this discipline now focus on biometric systems (whose objectives are the “recognition or identification of individuals based on some physical or behavioral characteristics that are intrinsically unique for each individual”, according to Boulgouris et al., *Biometrics*, 2010). Anyway, there still are reviews like [Biometrika](#) and [Biometrics](#); although I read the latter on an irregular basis, most articles focus on “biostatistical” theoretical or applied work. The same applies for [Biostatistics](#). By “biostatistical” applications, I mean that it has to do with applications or models related to the biomedical domain, in a wide sense (biology, health science, genetics, etc.).

According to the *Encyclopedia of Biostatistics* (2005, 2nd ed.),

(...) As is clear from the above examples, biostatistics is problem oriented. It is specifically directed to questions that arise in biomedical science. The methods of biostatistics are the methods of statistics – concepts directed at variation in observations and methods for extracting information from observations in the face of variation from various sources, but notably from variation in the responses of living organisms and particularly human beings under study. Biostatistical activity spans a broad range of sci-

entific inquiry, from the basic structure and functions of human beings, through the interactions of human beings with their environment, including problems of environmental toxicities and sanitation, health enhancement and education, disease prevention and therapy, the organization of health care systems and health care financing.

In sum, I think that Biostatistics is part of a super-family—Statistics—, and share most of its methods, but has a more focused area of interest (hence, an historical background, specific designs, and a general theoretical framework) and dedicated modeling strategies.

154 Make R report error on using non-existent column name in a data frame

Maybe, you can enclose your code into try-catch blocks, see `?try` and the associated examples. It is easy to test for the class of the results (“try-error”) in turn, e.g.

```
> res <- try(log("A"), silent=TRUE)
> class(res)
[1] "try-error"
```

You can also test directly for the correct spelling, by first listing the variables of interest—in your case, `MYVAR` and `WEIGHT`—and test that they are part of the data.frame `df`, e.g.

```
df <- data.frame(x=rnorm(100), g1=g1(2, 50), g2=g1(5,20))
sel.vars <- c("x","g2")
ifelse(all(sel.vars %in% colnames(df)), <compute things here>,
"fail")
```

155 Estimating the probability of a person getting a question right

If I understand your question correctly, you have a set of items (pass-fail) and you want to assess the probability of endorsing the k th item given its preceding responses? If that’s the case, what is usually done in psychometrics for educational assessment is to rely on **Item Response Model**, like the **Rasch Model**. In short, you model the probability of endorsing an item as a function of item difficulty and person ability (the more proficient an individual is, the more likely his response to an easy item will be

correct). This assumes that the content you are assessing is unidimensional, and that the items can be ordered by difficulty on that scale. A **Guttman model** is rarely applicable, so we may allow for some “imperfect” response patterns (e.g., 111011101110000, the 4th and 8th items were failed although the examinee reached the 11th item before giving up), but the sum score is a sufficient statistic for the Rasch Model. Under this approach, you need to have responses from other individuals on the same set of items. To get an idea, look at the **LSAT** data set and the way it is analysed in the **ltm** R package.

I have described a psychometrical model, not a purely probabilistic framework for estimating the probability of failing after the k th item, with all other items right (this would follow a geometric law).

156 Latest article or new development in cross validation?

Your question is not really precise, but I think the **caret** package and its associated vignettes may be a good start. Quoting the website, it is

a set of functions that attempt to streamline the process for creating predictive models.

In fact, it depends on a lot of other R packages dedicated to ML (see the list of suggested packages on **CRAN**), but it definitively simplifies the management of cross-validation scheme (k-fold, leave-one-out).

The Elements of Statistical Learning (Hastie et al.) is available on-line in its second edition, and all illustrations are done in R. Cross-validation is described at length in Chapter 7.

157 Tips and tricks to get started with statistical modeling?

As for (on-line) references, I would recommend looking at Andrew Moore’s tutorial slides on **Statistical Data Mining**.

There are many textbooks on data mining and machine learning; maybe a good starting point is **Principles of Data Mining**, by Hand et al., and **Introduction to Machine Learning**, by Alpaydin.

158 Who to follow on github to learn about best practice in data analysis?

I also follow [John Myles White's](#) GitHub [repository](#). There are several data-oriented projects, but also interesting stuff for R developers:

- [ProjectTemplate](#), a template system for building R project;
- [log4r](#), a logging system.

159 R and as.numeric()

Would that help?

```
> a <- as.data.frame(matrix(scan("1.txt", what="character",
                                na.strings=c("NA",paste("V",1:6,sep="")),
                                nc=13, byrow=T))
> class(a[,1])
[1] "factor"
> for (i in 1:ncol(a)) a[,i] <- as.numeric(as.character(a[,i]))
> class(a[,1])
[1] "numeric"
> summary(a) # should work here
```

The way you import data doesn't matter so much; I think the critical part is to convert value as character then as numeric (this allows to convert levels of a factor to their numerical counterparts).

160 Small sample linear regression: Where to start

I find @ucfagls's idea most appropriate here, since you have very few observations and a lot of variables. Ridge regression should do its job for prediction purpose.

Another way to analyse the data would be to rely on [PLS regression](#) (in this case, PLS1), which bears some idea with regression on PCA scores but seems more interesting in your case. As multicollinearity might be an issue there, you can look at *sparse* solution (see e.g., the [spl](#)s or the [mixOmics](#) R packages).

161 What tool do you use to communicate your survey questionnaires?

A combination of Perl + CGI is generally interesting for small surveys/questionnaires (because I hate PHP + MySQL). A gentle introduction can be found in [How to Conduct Behavioral Research over the Internet: A Beginner's Guide to Html and Cgi/Perl](#).

Now, I think that Ruby and Rails should provide very handy tools for that particular purpose. I can think of [surveyor](#), for example. I'm quite sure there are similar tools in Python.

As for an all-in-one system (no need to program anything, multiple and linked form available, automatic mailing, etc.), there's [Lime Survey](#).

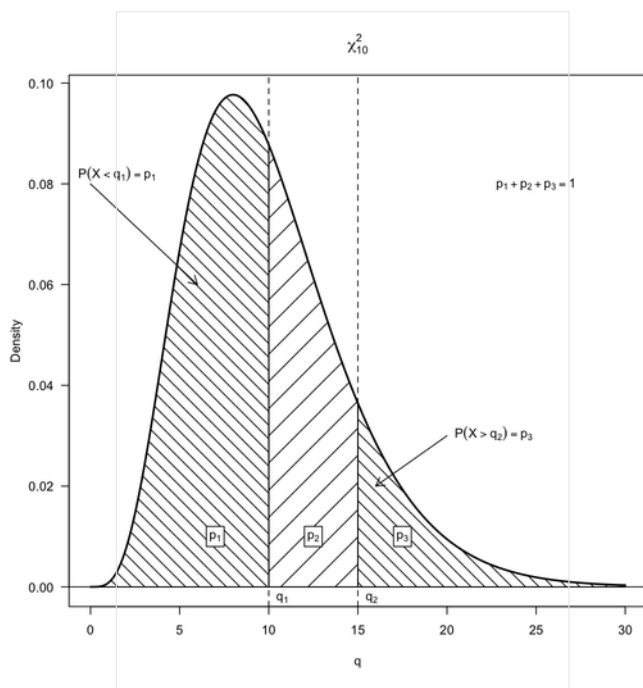
For off-line questionnaires, I would prefer *L^AT_EX* or Docbook.

162 Calculating quantiles for chi squared distribution

By hand, you need to refer to a tabulated distribution of the χ^2 , which should be found easily on the web (e.g., [this one](#)). Let x denotes the quantile of interest, and v the degrees of freedom of the chi-square distribution. You just have to know that the total area under the curve (i.e., the density) equals 1 (this will help you to work through the third case), and that such Tables generally gives $P(X < x) = p$, for a certain v . Knowing x (resp. p), you can find the approximated value of p (resp. x). In the aforementioned Table, the first cell reads: $P(X < 1.32) = 0.25$, for a 1-df chi-square.

If you have R, the [qchisq\(\)](#) function gives you the requested quantiles (look at the on-line help to be sure of what is returned, esp. the [lower.tail](#) argument). For the preceding example, we would use [qchisq\(0.25, 1, lower.tail=FALSE\)](#).

It is always a good idea to draw the corresponding density curve, as illustrated below. Note that p_3 is also $1 - P(X < q_2)$.



163 Typographic conventions for width of figures in LaTeX data analysis reports

I'll second @onestop comment about the fact that this question seems marginally related to statistical analysis or reporting.

That being said, I can't refrain from thinking of Ed. Tufte's work on the display of quantitative information, especially the design of his books which mixes different graphics layouts: some figures or tables are put in the margin, other in the body with caption in the margin, and large figures may extend beyond the body (full page width). The [tufte-latex](#) project offers $L^A_T_E_X$ classes for articles/handouts and books in the spirit of Ed. Tufte's design. Some examples are included on the project page; I particularly like the [example handout](#). On a related point, I also like the [tutorial](#) from the [vegan](#) R package.

My personal approach is to use 80% or 100% of text width (and keep it consistent across all the document), but I often play with the `width`, `height`, and `cex` arguments of `pdf()` when exporting figure so as to get the most clean and readable figure. It also happens to me to rely on a different layout—figure 60% and caption 40%, side by side, 100% of text width—for small illustrations or graphics.

164 Inserting small arrows (triangles) in bottom axis pointing up with R

I prefer to use dedicated symbol for that purpose. For example, use `points()` with `pch=17` (filled triangle) like in the example below:

```
dd <- dotplot(m~i|p,data=res.EQ.long,subset=i %in% eq5d.items.names[1:5],
  ylab="Score",ylim=c(0.5,3.5),
  scales=list(x=list(rot=45,at=1:5,
    labels=eq5d.items.names[1:5],cex=c(.8,.8))),
  main="EQ-5D",
  panel=function(x,y,subscripts,...){
    panel.dotplot(x,y,...)
    panel.lines(x=c(1,5),y=3,lwd=1.5, col="gray50")
    panel.segments(x0=c(1,3,5),y0=rep(2.95,3),
      x1=c(1,3,5),y1=rep(3.05,3),
      lwd=1.5, col="gray50")
    panel.text(x=c(1,3,5),y=3,c(0,50,100),cex=.6,pos=3)
    panel.text(x=3,y=3.2,"Health State",cex=.6,pos=3)
    panel.points(x=(4*ref[packet.number()])/100+1,y=2.9,
      pch=17)
    panel.points(x=1:5,y=res.EQ2[1:5,packet.number()],
      col="red",pch=19,cex=.6)
  },
  key=list(text=list(c("Mean","Median")),x=0.7,y=0.9,
    lines=list(col=c("blue","red"),type="p",
      pch=19,cex=.8)))

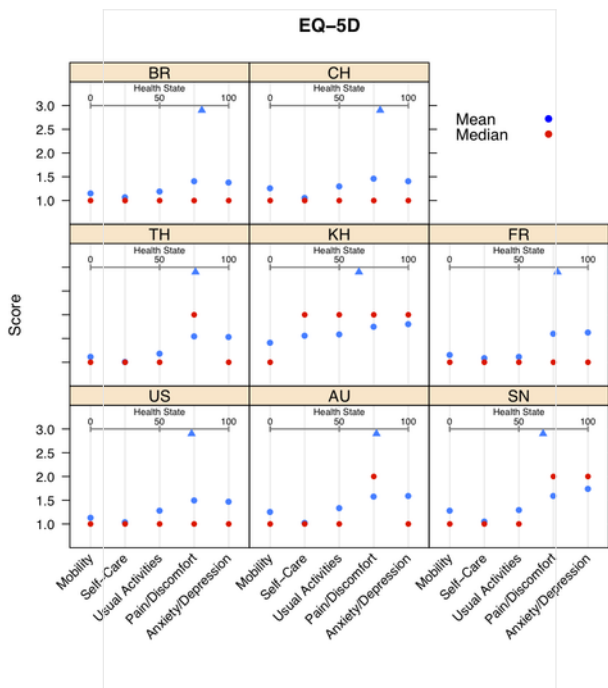
print(dd)
```

The complete list of symbols can be obtained with `show.pch()` from the [Hmisc](#) package.

165 What's the relation between hierarchical models, neural networks, graphical models, bayesian networks?

As [@carlosdc](#) said, a bayesian network is a type of Graphical Model (i.e., a directed acyclic graph (DAG) whose structure defines a set of conditional independence properties). [Hierarchical Bayes Models](#) can also be represented as DAGs; [Hierarchical Naive Bayes Classifiers for uncertain data](#), by Bellazzi et al., provides a good introduction to classification with such models. About hierarchical models, I think many articles can be retrieved by googling with appropriate keywords; for example, I found this one:

C. H. Jackson, N. G. Best and S. Richardson. [Bayesian graphical models for regression](#)



on multiple data sets with different variables.
Biostatistics (2008) 10(2): 335-351.

Michael I. Jordan has a nice tutorial on [Graphical Models](#), with various applications based on the factorial [Hidden Markov model](#) in bioinformatics or natural language processing. His book, [Learning in Graphical Models](#) (MIT Press, 1998), is also worth reading (there's an application of GMs to structural modeling with [BUGS](#) code, pp. 575-598)

166 What is the difference between the “coef” and “(exp)coef” output of coxph in R?

If you have a single explanatory variable, say treatment group, a Cox's regression model is fitted with `coxph()`; the coefficient (`coef`) reads as a regression coefficient (in the context of the Cox model, described hereafter) and its exponential gives you the hazard in the treatment group (compared to the control or placebo group). For example, if $\hat{\beta} = -1.80$, then the hazard is $\exp(-1.80) = 0.165$, that is 16.5%.

As you may know, the hazard function is modeled as

$$h(t) = h_0(t) \exp(\beta'x)$$

where $h_0(t)$ is the baseline hazard. The hazards depend multiplicatively on the covariates, and $\exp(\beta_1)$ is the ratio of the hazards between two individuals whose values of x_1 differ by one unit when all other covariates are held constant. The ratio of the hazards of any two individuals i and j is $\exp(\beta'(x_i - x_j))$, and is called the hazard ratio (or incidence rate ratio). This ratio is assumed to be constant over time, hence the name of *proportional hazard*.

To echo your preceding question about [survreg](#), here the form of $h_0(t)$ is left unspecified; more precisely, this is a semi-parametric model in that only the effects of covariates are parametrized, and not the hazard function. In other words, we don't make any distribution assumption about survival times.

The regression parameters are estimated by maximizing the partial log-likelihood defined by

$$\ell = \sum_f \log \left(\frac{\exp(\beta'x_f)}{\sum_{r(f)} \exp(\beta'x_r)} \right)$$

where the first summation is over all deaths or failures f , and the second summation is over all subjects $r(f)$ still alive (but at risk) at the time of failure – this is known as the *risk set*. In other words, ℓ can be interpreted as the log profile likelihood for β after eliminating $h_0(t)$ (or in other words, the LL where the $h_0(t)$ have been replaced by functions of β that maximize the likelihood with respect to $h_0(t)$ for a fixed vector β).

About censoring, it is not clear whether you refer to left censoring (as might be the case if we consider an origin for the time scale that is earlier than the time when observation began, also called *delayed entry*), or right-censoring. In any case, more details about the computation of the regression coefficients and how the [survival](#) package handles censoring can be found in Therneau and Grambsch, [Modeling Survival Data](#) (Springer, 2000). [Terry Therneau](#) is the author of the former S package. An [online tutorial](#) is available.

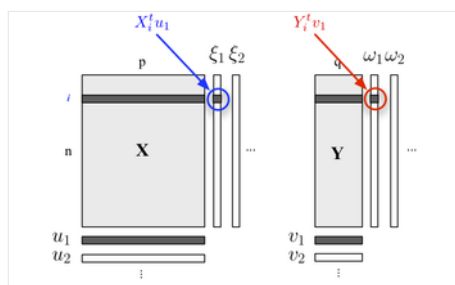
[Survival Analysis in R](#), by David Diez, provides a good introduction to Survival Analysis in R. A brief overview of χ^2 tests for regression parameters is given p. 10. Hopefully, this should help clarifying the on-line help quoted by [@onestop](#), “coefficients the coefficients of the linear predictor, which multiply the columns of the model matrix.” For an applied textbook, I recommend [Analyzing Medical Data Using S-PLUS](#), by Everitt and Rabe-Hesketh

(Springer, 2001, chap. 16 and 17), from which most of the above comes from. Another useful reference is John Fox's appendix on [Cox Proportional-Hazards Regression for Survival Data](#).

167 Regression with multiple dependent variables?

@Brett's response is fine.

If you are interested in describing your two-block structure, you could also use [PLS regression](#). Basically, it is a regression framework which relies on the idea of building successive (orthogonal) linear combinations of the variables belonging to each block such that their covariance is maximal. Here we consider that one block X contains explanatory variables, and the other block Y responses variables, as shown below:



We seek “latent variables” who account for a maximum of information (in a linear fashion) included in the X block while allowing to predict the Y block with minimal error. The u_j and v_j are the loadings (i.e., linear combinations) associated to each dimension. The optimization criteria reads

$$\max_{|u_h|=1, |v_h|=1} \text{cov}(X_{h-1}u_h, Yv_h) \quad (\equiv \max \text{cov}(\xi_h, \omega_h))$$

where X_{h-1} stands for the deflated (i.e., residualized) X block, after the h th regression.

The correlation between factorial scores on the first dimension (ξ_1 and ω_1) reflects the magnitude of the X - Y link.

168 Calculation of incidence rate for epidemiological study in hospital

It is commonly admitted that the denominator for IRs is the “population at risk” (i.e., all individuals

in which the studied event(s) may occur). Although your first formula is generally used, I found in *The new public health*, by Tulchinsky and Varavikova (Elsevier, 2009, 2nd. ed., [p. 84](#)) that a distinction is made between *ordinary incidence rate*, where the average size of the population in the fixed period of time is used in the denominator, and *person-time incidence rate*, with PT at risk in the denominator.

Obviously, when individuals not at risk of the disease are included in the denominator, the resultant measure of disease frequency will underestimate the true incidence of disease in the population under investigation, but see [Numerators, denominators and populations at risk](#).

169 How does one do a Type-III SS ANOVA in R with contrast codes?

Type III sum of squares for ANOVA are readily available through the [Anova\(\)](#) function from the [car](#) package.

Contrast coding can be done in several ways, using [C\(\)](#), the [contr.*](#) family (as indicated by @nico), or directly the [contrasts\(\)](#) function/argument. This is detailed in §6.2 (pp. 144-151) of *Modern Applied Statistics with S* (Springer, 2002, 4th ed.). Note that [aov\(\)](#) is just a wrapper function for the [lm\(\)](#) function. It is interesting when one wants to control the error term of the model (like in a within-subject design), but otherwise they both yield the same results (and whatever the way you fit your model, you still can output ANOVA or LM-like summaries with [summary.aov](#) or [summary.lm](#)).

I don't have SPSS to compare the two outputs, but something like

```
> library(car)
> sample.data <- data.frame(IV=factor(rep(1:4,each=20)),
                             DV=rep(c(-3,-3,1,3),each=20)+rnorm(80))
> Anova(lm1 <- lm(DV ~ IV, data=sample.data,
                  contrasts=list(IV=contr.poly)), type="III")
Anova Table (Type III tests)
```

```
Response: DV
              Sum Sq Df F value    Pr(>F)
(Intercept)  18.08   1  21.815 1.27e-05 ***
IV           567.05   3 228.046 < 2.2e-16 ***
Residuals    62.99  76
---
```

```
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

is worth to try in first instance.

About factor coding in R vs. SAS: R considers the baseline or reference level as the first level in lexicographic order, whereas SAS considers the last one. So, to get comparable results, either you have to use `contr.SAS()` or to `relevel()` your R factor.

170 In R, does “glmnet” fit an intercept?

Yes, an intercept is included in a `glmnet` model, but it is not regularized (cf. [Regularization Paths for Generalized Linear Models via Coordinate Descent](#), p. 13). More details about the implementation could certainly be obtained by carefully looking at the code (for a gaussian family, it is the `elnet()` function that is called by `glmnet()`), but it is in Fortran.

You could try the `penalized` package, which allows to remove the intercept by passing `unpenalized = ~0` to `penalized()`.

```
> x <- matrix(rnorm(100*20),100,20)
> y <- rnorm(100)
> fit1 <- penalized(y, penalized=x, unpenalized=~0,
+                 standardize=TRUE)
> fit2 <- lm(y ~ 0+x)
> plot((coef(fit1) + coef(fit2))/2, coef(fit2)-coef(fit1))
```

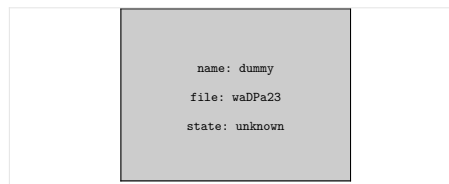
To get Lasso regularization, you might try something like

```
> fit1b <- penalized(y, penalized=x, unpenalized=~0,
+                 standardize=TRUE, lambda1=1, steps=20)
> show(fit1b)
> plotpath(fit1b)
```

As can be seen in the next figure, there is little differences between the regression parameters computed with both methods (left), and you can plot the Lasso path solution very easily (right).

171 Yates continuity correction for 2 x 2 contingency tables

Yates' correction results in tests that are more conservative as with Fisher's “exact” tests.



Here is an online tutorial on the use of [Yates's continuity correction](#), by Stefanescu et al, which clearly points to various flaws of systematic correction for continuity (pp. 4-6). Quoting Agresti (CDA 2002), “Yates (1934) mentioned that Fisher suggested the hypergeometric to him for an exact test”, which led to the continuity-corrected version of the χ^2 . Agresti also indicated that Fisher's test is a good alternative now that computers can do it even for large samples (p. 103). Now, the point is that choosing a test really depends on the question that is asked and the assumptions that are made by each of them (e.g., in the case of the Fisher's test we assume that margins are fixed).

In your case, Fisher test and corrected χ^2 agree and yield p -value above 5%. In the case of the ordinary χ^2 , if p -values are computed using a Monte Carlo approach (see `simulate.p.value`), then it fails to reach significance too.

Other useful references dealing with small sample size issues and the overuse of Fisher's test, include:

- I. Campbell, [Chi-squared and Fisher-Irwin tests of two-by-two tables with small sample recommendations](#), *Statistics in Medicine* 26(19): 3661–3675, 2007.
- Mark G. Haviland, [Yates's correction for continuity and the analysis of \$2 \times 2\$ contingency tables](#), *Statistics in Medicine* 9(4): 363–367, 1990.

172 Tutorials on object-oriented programming in R

In addition to [@suncoolsu](#) excellent response, there is [A \(Not So\) Short Introduction to S4](#), by Christophe Genolini. It is available on CRAN website.

173 Java implementations of the lasso

About clean implementation in Python, there is the `scikit.learn` toolkit. The [L1/L2 regularization scheme](#) (incl. `elasticnet`) works great with GLM (LARS

and coordinate descent algorithms available). Don't know about Java implementation.

174 Good econometrics textbooks?

Definitively [Econometric Analysis](#), by Greene. I'm not an econometrician, but I found this book very useful and well written.

175 Establishing that the population sampled of two separate surveys is the same

I think that for subject-specific characteristics, like demographic data, you can proceed the usual way (t-test, etc.). This will help showing that your samples don't differ according to these variables. About self-reported attitude data, if you have very few items, skip to step 2, otherwise step 1 might be appropriate.

1. Assessing measurement equivalence

Rather than saying that the two populations (or actually, samples) are "the same", I would say you have to show that your two questionnaires are assessing the same construct(s). This is what is done in cross-cultural surveys or international clinical trials where health-related quality of life is used as a secondary endpoint, for example. In each case, we have a set of items that purports to assess different dimensions, and we want to demonstrate whether we are measuring individuals in the same way irrespective of their country. When dealing with uni- or multidimensional scales, it is known as *measurement invariance* in psychometrics, that is you want to show that the factorial structure is comparable between the two groups. But, the same remark would apply as well if we were considering longitudinal data (I interpret your question as involving different samples at each time point). A *multi-group confirmatory factor analysis* is appropriate in this case. Standard references include:

- Meredith, W (1993). Measurement invariance, factor analysis, and factorial invariance. *Psychometrika*, 58, 525-543.
- Vandenberg, RJ and Lance, CE (2000). A review and synthesis of the measurement invariance literature: Suggestions, practices, and recommendations for organizational research. *Organizational Research Methods*, 3, 4-70.

In R, the [lavaan](#) package provides facilities for that kind of analysis, but see the documentation: [lavaan: an R package for structural equation modeling and more](#) (§6.2). Otherwise, you have to resort on [Mplus](#) or a good software for SEMs. [Studying Measurement Invariance Using Confirmatory Factor Analysis](#) provides illustration with [LISREL](#) syntax.

You may want to consider data from 6 to 12 months (to collect 1 or 2 waves for survey A). After that, I think you can just pool your data.

2. Assessing group comparability

Now, if you cannot define a clear construct common to those two questionnaires, or if you have so few items that it would make no sense to consider a scale, then you can rely on basic group statistics for each item (using e.g., t-test, trend test for ordinal data, tests for nominal data, etc.). In this case, you are essentially studying between-group differences. This basically tells you whether (aggregated) scores differ, but not whether items are perceived as having the same meaning (or underlying the same construct) across the two groups.

176 R resources in non-English languages

All RSS feeds I follow are in English actually, so I'll just point to tutorials available in French, or made by French researchers.

Apart from the [Contributed Documentation](#) on CRAN, I often browse the R website hosted at the [bioinformatics lab](#) in Lyon (France); it is mostly in French, but it also includes english material. I also like [Philippe Besse](#) resources (SAS + R).

177 Internal reliability for an ordinal scale

From a practical perspective, I don't see any obvious reason to not use Cronbach's alpha with ordinal items (e.g., Likert-type items), as is commonly done in most of the studies. It is a lower bound for reliability, and is essentially used as an indicator of internal consistency of a test or questionnaire. The usual assumptions pertaining to a correct interpretation of its value are as follows: (i) no residual correlations, (ii) items have identical loadings, and (iii) the scale is unidimensional. In fact, the

sole case where alpha will be essentially the same as reliability is the case of uniformly high factor loadings, no error covariances, and unidimensional instrument (1).

However, we can speak of an *ordinal reliability alpha*. For instance, Zumbo et coll. (2) use a polychoric correlation matrix input to calculate alpha parallel to Cronbach. Their simulation studies lead them to conclude that ordinal reliability alpha provides “consistently suitable estimates of the theoretical reliability, regardless of the magnitude of the theoretical reliability, the number of scale points, and the skewness of the scale point distributions. In contrast, coefficient alpha is in general a negatively biased estimate of reliability” for ordinal data (p. 21). Ordinal reliability alpha will normally be higher than the corresponding Cronbach’s alpha.

Otherwise, the usual Cronbach’s α is influenced by the number of items in the test and interitem correlations (for a fixed sample size $N = 300$, even with modest–albeit perfect–correlation between items, e.g. $\rho = 0.35$, Cronbach’s α would still be at 0.943 with 30 items, and 0.910 with 20 items). There’re subtle issues with Cronbach’s α and departure from the unidimensionality assumption (systematic errors can greatly inflate the estimate of alpha, especially with large sample sizes) or the presence of inconsistent responses (random responses may inflate Cronbach’s alpha when their mean differ from that of the true responses). If the variables being tested are all dichotomous, Cronbach’s alpha is the same as Kuder-Richardson coefficient (3).

Of note, there are alternative ways to estimate the reliability of test scores, see e.g., Zinbarg et al. (4).

A good review is

Bruce Thompson. *Score Reliability. Contemporary Thinking on Reliability issues*. Sage Publications, 2003.

References

1. T Raykov. *Scale reliability, Cronbach's coefficient alpha, and violations of essential tau-equivalence for fixed congeneric components*. *Multivariate Behavioral Research*, **32**: 329-254, 1997.
2. B D Zumbo, A M Gadermann, and C Zeisser. *Ordinal versions of coefficients alpha and theta for likert rating scales*. *Journal of Modern Applied Statistical Methods*, **6**: 21-29, 2007.

3. G F Kuder and M W Richardson. *The theory of the estimation of test reliability*. *Psychometrika*, **2**: 151-160, 1937.
4. R E Zinbarg, W Revelle, I Yovel, and W Li. *Cronbach's α , Revelle's β , and McDonald's ω_h : Their relations with each other and two alternative conceptualizations of reliability*. *Psychometrika*, **70**(1): 123-133, 2005.

178 How to do weighted pair hierarchical clustering in R?

About your first question, it seems that the `mcquitty` option corresponds to WPGMA clustering, while `average` is for UPGMA. It is just by looking at the *source code*, so it is worth to double check it. But it also referred as is in the `upgma()` function from the `phangorn` package.

About your second question, I think you just have to subset your genes by the group labels found after `cutree`, and then plot expression profiles as usual.

179 What is the difference between fixed effect, random effect and mixed effect models?

Not really a formal definition, but I like the following slides: *Mixed models and why sociolinguists should use them*, from Daniel Ezra Johnson. A brief recap’ is offered on slide 4. Although it mostly focused on psycholinguistic studies, it is very useful as a first step.

180 Automatically produce summary by factor variable in R

Check out the `by()` or `tapply()` functions. Basically,

```
tapply(y, g, mean)
```

will give you the mean of `y` by levels of `g`. If you want to get a data.frame from the resulting aggregated measures, use `aggregate()`.

A more elaborated solution is available through the `summary.formula()` function in the `Hmisc` package.

181 Logistic Regression: Classification Tables a la SPSS in R

Thomas D. Fletcher has a function called `ClassLog()` (for “classification analysis for a logistic regression model”) in his `QuantPsyc` package. However, I like @caracal’s response because it is self-made and easily customizable.

182 Longitudinal Data analysis by multilevel modeling in R

The `intervals()` function should provide you with $100(1-\alpha)$ confidence intervals for the random effects in your model, see `help(intervals.lme)` for more information. You can also test if any of the variance components can be dropped from the model by using `anova()` (which amounts to do an LRT between two nested models).

183 Should I use factor analysis on my data?

As an alternative to CA suggested by @Brandon, you could also try `Multiple Correspondence Analysis` which has the advantage of considering all types of games at the same time (unlike CA), which are probably scored as binary variable (in this case, the MCA solution will be close to the PCA one-factor scores and eigenvalues are linearly related). Basically this will give you an idea of how games group together, if any. At the same time, you can use your “hardcore gamer” status (yes/no) as an illustrative variable (i.e., this variable will not participate to the construction of the factorial axes), which will help you identifying how it related to these clusters of variables.

The `FactoMineR` R package offers all of what is needed for such kind of analysis, see `MCA()`.

As you didn’t say anything about your sample size, it’s hard to suggest confirmatory or model-based approaches, like logistic regression or latent class regression. But you can look at `Random Forests` and try to identify the “best” variables that allow to predict your outcome with a minimal classification error rate (see the `randomForest` package).

184 Creating plots for String type columns in R

It seems the `barplot()` will be your friend in that case, e.g.

```
x <- sample(c("Win", "Linux", "Mac"), 100, replace=TRUE)
barplot(table(x))
```

This will work for variables of type `character` or `factor`. Another option is to use Cleveland’s `dotplot`, see `dotchart()` (or `dotplot()` in the `lattice` package).

Update

You could replace `table(x)` by `table(x)/sum(table(x))*100` to express data as % rather than counts. I know there are more elegant solutions in additional packages, but I can’t remember their names actually. The `table()` function will also work for two-way classification, and marginal totals can easily be computed in a similar way; e.g. `apply(table(x, y), 1, sum)` gives rows marginal frequencies.

185 How do you test an implementation of k-means?

The k-means includes a stochastic component, so it is very unlikely you will get the same result unless you have exactly the same implementation and use the same starting configuration. However, you could see if your results are in agreement with well-known implementations (don’t know about Matlab, but implementation of k-means algorithm in R is well explained, see `Hartigan & Wong, 1979`).

As for comparing two series of results, there still is an issue with label switching if it is to be run multiple times. Again, in the `e1071` R package, there is a very handy function (`matchClasses()`) that might be used to find the ‘best’ mapping between two categories in a two-way classification table. Basically, the idea is to rearrange the rows so as to maximise their agreement with columns, or use a greedy approach and permute rows and columns until the sum of on the diagonal (raw agreement) is maximal. Coefficient of agreement like the `Kappa` statistic are also provided.

Finally, about how to benchmark your implementation, there are a lot of freely available data, or you can simulate a dedicated data set (e.g., through a

finite mixture model, see the [MixSim](#) package).

186 R getting share of users with multiple of an element

You could try something like

```
> df <- data.frame(User=sample(LETTERS[1:10], 100, rep=T),
  OS=sample(c("Win", "Lin", "Mac"), 100, rep=T))
> (res <- with(df, tapply(OS, User, function(x) length(unique(x)))))
A B C D E F G H I J
2 3 3 3 3 3 3 3 3 3
> barplot(table(res)) # for counts
> barplot(table(ifelse(res==1, "1", "2+")))
```

Replace `table()` by `prop.table()` if you want proportions instead of counts, as suggested by [@Chase](#) in a comment to your preceding question.

187 Coordinate descent for the lasso or elastic net

I earlier suggested the recent paper by Friedman and coll., [Regularization Paths for Generalized Linear Models via Coordinate Descent](#), published in the Journal of Statistical Software (2010). Here are some other references that might be useful:

- [Pathwise coordinate optimization](#), by Friedman and coll.
- [Fast Regularization Paths via Coordinate Descent](#), by Hastie (UseR! 2009)
- [Coordinate descent algorithms for lasso penalized regression](#), by Wu and Lange (Ann. Appl. Stat. 2(1): 224-244, 2008; also on available on [arXiv.org](#))
- [Coordinate Descent for Sparse Solutions of Underdetermined Linear Systems of Equations](#), by Yagle (a bit too complex for me)

188 Identifying Interaction Effects

Cox and Wermuth (1996) or Cox (1984) discussed some methods for detecting interactions. The problem is usually how general the interaction terms should be. Basically, we (a) fit (and test) all second-order interaction terms, one at a time, and (b) plot

their corresponding p-values (i.e., the No. terms as a function of $1-p$). The idea is then to look if a certain number of interaction terms should be retained: Under the assumption that all interaction terms are null the distribution of the p-values should be uniform (or equivalently, the points on the scatterplot should be roughly distributed along a line passing through the origin).

Now, as [@Gavin](#) said, fitting many (if not all) interactions might lead to overfitting, but it is also useless in a certain sense (some high-order interaction terms often have no sense at all). However, this has to do with interpretation, not detection of interactions, and a good review was already provided by Cox in [Interpretation of interaction: A review](#) (*The Annals of Applied Statistics* 2007, 1(2), 371–385)—it includes references cited above. Other lines of research worth to look at are study of [epistatic effects](#) in genetic studies, in particular methods based on graphical models (e.g., [An efficient method for identifying statistical interactors in gene association networks](#)).

189 Is interaction possible between two continuous variables?

Yes, why not? The same consideration as for categorical variables would apply in this case: The effect of X_1 on the outcome Y is not the same depending on the value of X_2 . To help visualize it, you can think of the values taken by X_1 when X_2 takes high or low values. Contrary to categorical variables, here interaction is just represented by the product of X_1 and X_2 . Of note, it's better to center your two variables first (so that the coefficient for say X_1 reads as the effect of X_1 when X_2 is at its sample mean).

As kindly suggested by [@whuber](#), an easy way to see how X_1 varies with Y as a function of X_2 when an interaction term is included, is to write down the model $\mathbb{E}(Y|X) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \beta_3 X_1 X_2$.

Then, it can be seen that the effect of a one-unit increase in X_1 when X_2 is held constant may be expressed as:

$$\begin{aligned} \mathbb{E}(Y|X_1+1, X_2) - \mathbb{E}(Y|X_1, X_2) &= \beta_0 + \beta_1(X_1+1) + \beta_2 X_2 + \beta_3(X_1+1)X_2 \\ &\quad - (\beta_0 + \beta_1 X_1 + \beta_2 X_2 + \beta_3 X_1 X_2) \\ &= \beta_1 + \beta_3 X_2 \end{aligned}$$

Likewise, the effect when X_2 is increased by one

unit while holding
demonstrates why
effects of X_1 (β_1)
will even be mor
are highly correla
in mind the linear
in such a linear m

You can have a l
and interpreting i
Stephen G. West
Publications, 199
ent kind of interac
(This is probably
able through Goo

Here is a toy exam

```
library(mvtnorm)
set.seed(101)
n <- 300
S <- matrix(c(1,.2,
              .2,1,
              .4,0,
              0,0,1),
            nr=4, nc=4,
            dimnames=list(c("x1", "x2", "y", "x1x2"),
                          c("x1", "x2", "y", "x1x2")))
X <- as.data.frame(
  colnames(X) <- c("x1", "x2", "y", "x1x2")
  summary(lm(y~x1+x2+
    pairs(X)
```

where the output

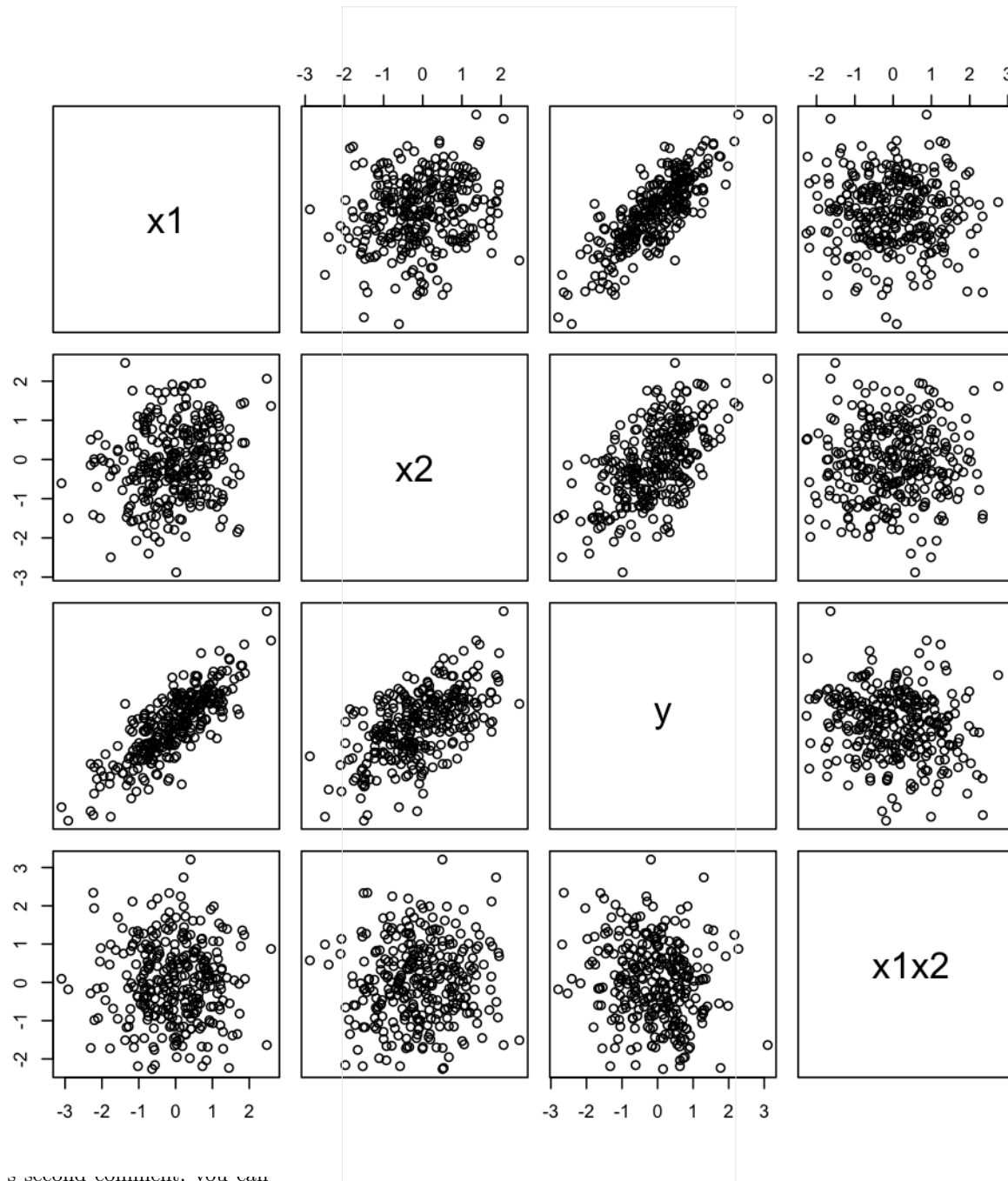
```
Coefficients:
            Estim
(Intercept) -0.0105
x1           0.7149
x2           0.4370
x1x2        -0.1762
---
Signif. codes:  0 '1
' 1

Residual standard e
Multiple R-squared:
F-statistic: 743.2
```

And here is how t

To illustrate @whuber's second comment, you can
always look at the variations of Y as a function of
 X_2 at different values of X_1 (e.g., terciles or deciles);
trellis displays are useful in this case. With the data
above, we would proceed as follows:

```
library(Hmisc)
X$x1b <- cut2(X$x1, g=5) # consider 5 quantiles (60 obs. per
group)
coplot(y~x2|x1b, data=X, panel = panel.smooth)
```



190 Stacked bar plot

I doubt you fill find a suitable range of distinct
colours with so much categories. Anyway, here are
some ideas:

Given : x1b

any one of the diverging or sequential palettes. However, the number of colour is limited, so you will need to recycle them (e.g., every 6 items)

Here is an illustration:
[0.764, 2.590]

[0.228, 0.764]

[-0.203, 0.228]

[-0.824,-0.203]

[-3.095,-0.824]

```
library(RColorBrewer)
x <- sample(LETTERS[1:20], 100, replace=TRUE)
tab <- as.matrix(table(x))
my.col <- brewer.pal(6, "BrBG") # or brewer.pal(6, "Blues")
barplot(tab, col=my.col)
```

There is also the [colorspace](#) package, which has a nice accompanying vignette about the design of good color schemes. Check also Ross Ihaka's course on [Topic in Computational Data Analysis and Graphics](#).

Now, a better way to display such data is probably to use a so-called Cleveland dot plot, i.e.

```
dotchart(tab)
```

191 Computing the decision boundary of a linear SVM model

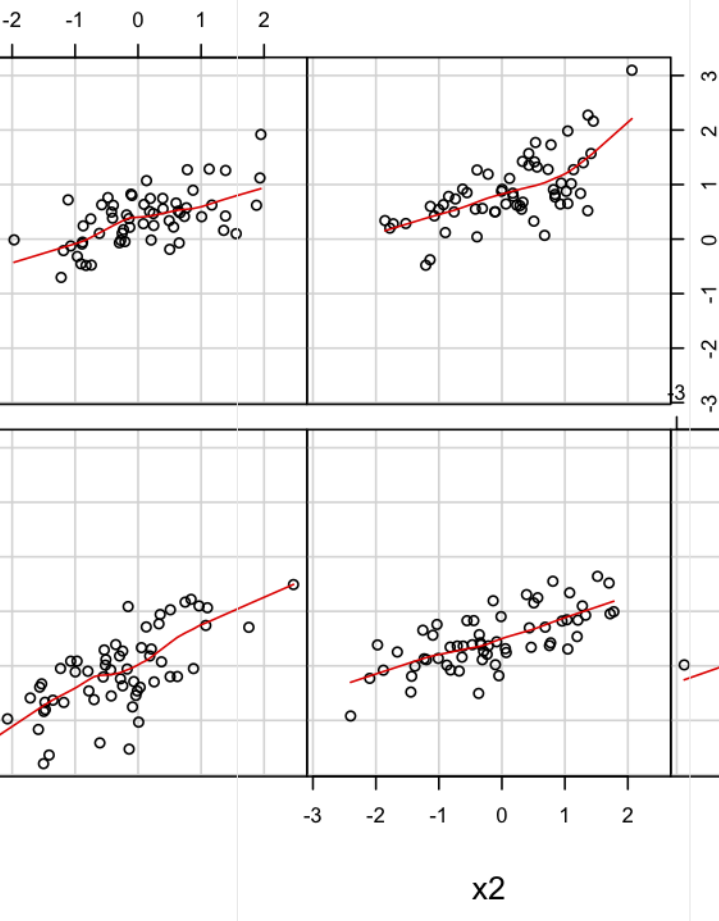
The [Elements of Statistical Learning](#), from Hastie et al., has a complete chapter on support vector classifiers and SVMs (in your case, start page 418 on the 2nd edition). Another good tutorial is [Support Vector Machines in R](#), by David Meyer.

Unless I misunderstood your question, the decision boundary (or hyperplane) is defined by $x^T \beta + \beta_0 = 0$ (with $\|\beta\| = 1$, and β_0 the intercept term), or as @chony said a linear combination of the support vectors. The margin is then $2/\|\beta\|$, following Hastie et al. notations.

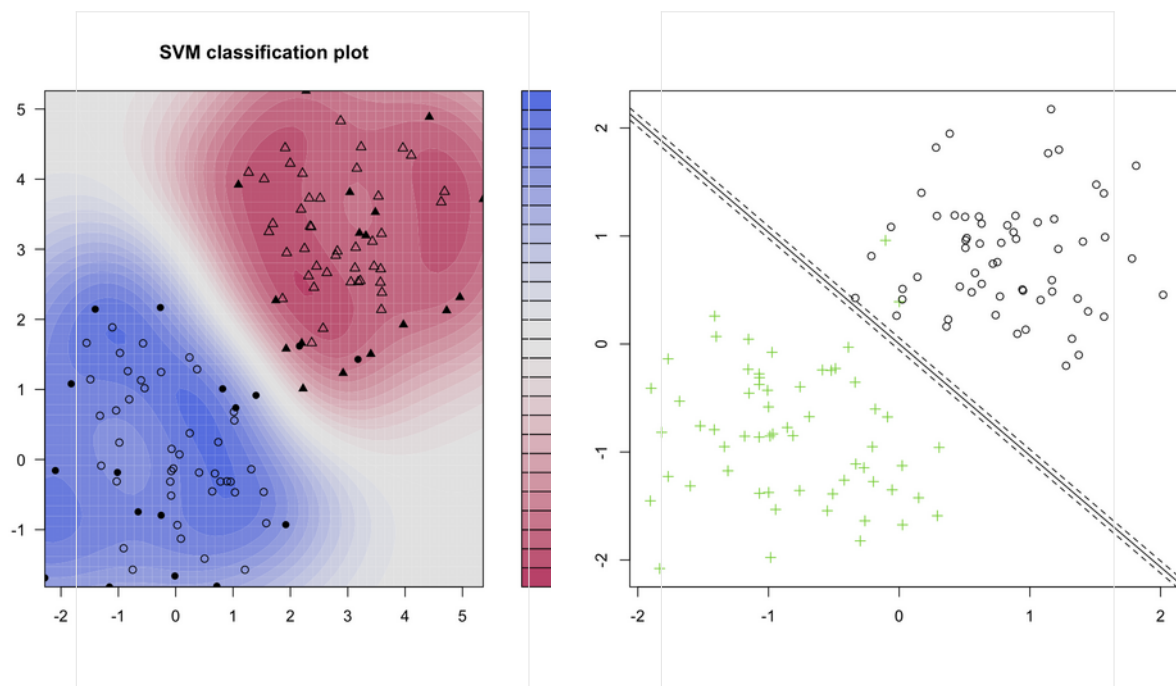
From the on-line help of `ksvm()` in the [kernlab](#) R package, but see also [kernlab – An S4 Package for Kernel Methods in R](#), here is a toy example:

```
set.seed(101)
x <- rbind(matrix(rnorm(120),,2),matrix(rnorm(120,mean=3),,2))
y <- matrix(c(rep(1,60),rep(-1,60)))
svp <- ksvm(x,y,type="C-svc")
plot(svp,data=x)
```

Note that for the sake of clarity, we don't consider train and test samples. Results are shown below, where color shading helps visualizing the fitted decision values; values around 0 are on the decision



1. For stacked barchart, you need `barplot()` with `beside=FALSE` (which is the default) – this is in base R (@Chase's solution with `ggplot2` is good too)
2. For generating a color ramp, you can use the [RColorBrewer](#) package; the example shown by @fRed can be reproduced with `brewer.pal` and



boundary.

Calling `attributes(svp)` gives you attributes that you can access, e.g.

```
alpha(svp) # support vectors whose indices may be
            # found with alphaindex(svp)
b(svp)     # (negative) intercept
```

So, to display the decision boundary, with its corresponding margin, let's try the following (in the rescaled space), which is largely inspired from a tutorial on SVM made some time ago by [Jean-Philippe Vert](#):

```
plot(scale(x), col=y+2, pch=y+2, xlab="", ylab="")
w <- colSums(coef(svp)[[1]] * x[unlist(alphaindex(svp)),])
b <- b(svp)
abline(b/w[1], -w[2]/w[1])
abline((b+1)/w[1], -w[2]/w[1], lty=2)
abline((b-1)/w[1], -w[2]/w[1], lty=2)
```

And here it is:

192 Boxplots as tables

I tend to think that boxplots will convey more effective information if there are numerous empirical distributions that you want to summarize into a single figure. If you only have two or three groups, ed-

itors may ask you to provide numerical summaries instead, either because it is more suitable for the journal policy, or because readers won't gain much insight into the data from a figure. If you provide the three quartiles, range, and optionally the mean \pm SD, then an advertised reader should have a clear idea of the shape of the distribution (symmetry, presence of outlying values, etc.).

I would suggest two critical reviews by Andrew Gelman (the first goes the other way around, but still it provides insightful ideas):

1. Gelman, A., Pasarica, C., and Dodhia, R. [Let's practice what we preach](#). The American Statistician (2002) 56(2): 121-130.
2. Gelman, A. [Why Tables are Really Much Better than Graphs](#). (also discussed on his blog)

193 Social network datasets

Just found this: [476 million Twitter tweets](#) (via [@yarapavan](#)).

194 Appropriate test for multivariate experiment result with unknown distributions

Well, following your update, it seems you are dealing with a factorial experiment (*factorial* means that every factors are crossed, or, in other words, each unit is subjected to every possible combination of your factors), with five replicates. Let assume that these are not the same statistical units whose temperature is repeatedly measured across each of the 12 combinations (for the sake of clarity).

An **ANalysis Of VAriance** (ANOVA) seems to be the most appropriate method to deal with this design. Basically, it will allow you to estimate the contribution of each source of variance (decay, particles, and velocity) wrt. the total variance in the observed temperature. What is not explained by these factors is called the residual variance (what you call the ‘random effect’). A full additive model (i.e., without modeling interaction between your factors) will read something like

$$y_{ijkl} = \mu + \alpha_i + \beta_j + \gamma_k + \varepsilon_{ijkl},$$

where y_{ijkl} is the temperature for unit l when considering levels $i = 1 \dots a$, $j = 1 \dots b$, and $k = 1 \dots c$, of factors α (decay), β (particles), and γ (velocity); the ε_{ijk} are the residuals assumed to follow a gaussian distribution of unknown variance, σ^2 . They can be viewed as random fluctuations around μ , the overall mean, and reflect the between-unit variations that are not accounted for by the other factors. The α_i , β_j , and γ_k can be viewed as factor-specific deviations from the overall mean μ .

The so-called *main effect* of decay, particles, and velocity will be estimated by forming a ratio between the variance that they account for (known as *mean squares*) and the residual variance (what is left after considering all variance explained by those factors), which is known to follow a Fisher-Snedecor (F) distribution, with $d-1$ and $N-abc$ degrees of freedom, where $d = a$, b , or c stands for the number of levels of α (decay), β (particles), and γ (velocity). A significant effect (following an **hypothesis test** of a null effect, i.e. $H_0 : \mu_i = \mu_j \forall i \neq j$ vs. H_1 : at least two of the μ_i ’s differ) would indicate that the factor under consideration has a significant effect on the outcome. This is readily obtained by any statistical software. For instance, in R you would use something like

```
summary(aov(temperature ~ decay + particles + velocity, data=df))
```

provided temperature and factor levels are organized in four columns, in a data.frame named `df`, as suggested below:

```
t1 0.1 10 30
t2 0.1 10 30
t3 0.1 10 30
t4 0.1 10 30
t5 0.1 10 30
t6 0.2 10 30
t7 0.2 10 30
...
t60 0.3 100 70
```

The effect of any of the three factors can also be summarized under an equation like the one you referred to by simply calling (again under R):

```
summary.lm(aov(temperature ~ decay + particles + velocity))
```

This follows from the fact that an ANOVA is nothing more than a **Linear Model** that you may have heard about (think of a regression model where the explanatory variables are all categorical).

Should you want to account for possible interactions between all three factors, you need to add three second-order and one three-order interaction terms. If any of these effects prove to be significant, this would mean that the effect of the corresponding factors cannot be considered in isolation one from the other (e.g., the effect of decay on temperature is not the same depending on the number of particles).

As for references, I would suggest starting with online tutorial or textbook like **Three-Way ANOVA**, by Barry Cohen, or **Practical Regression and Anova using R**, by John MainDonald (but see also other textbooks available on **CRAN documentation**). The definitive reference is Montgomery, **Design and Analysis of Experiments** (Wiley, 2005).

195 Correction due to rounding error

I understand the question as one where we know the theoretical distribution of students height with some precision (i.e., with one decimal place). In the present case, this is a gaussian distribution with parameters $\mathcal{N}(174.5; 6.9^2)$.

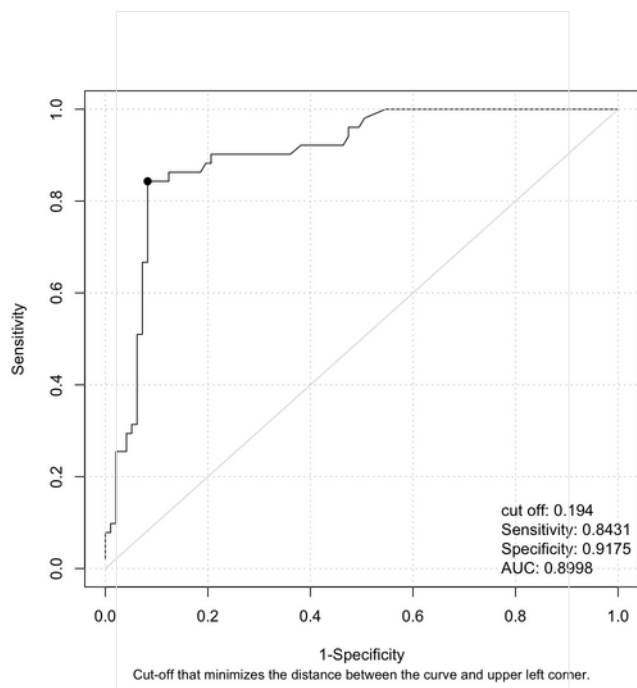
Now, we have empirical measurements of students height on small samples ($n = 25$), but results are rounded to the nearest integer due to possible mea-

surement error or imperfect measurement device.

So, my understanding is that the question is really to assess $\Pr(X > 176)$ or $\Pr(Z > \frac{176-174.5}{6.9})$ if you refer to the standardized $\mathcal{N}(0;1)$ distribution, and not $\Pr(X > 176.5)$ as you suggested.

196 Choosing the right threshold for a biometric trait authentication system

Generally, the cut-off value is chosen such as to maximize the compromise between sensitivity (Se) and specificity (Sp). You can generate a regular sequence of thresholds and plot the resulting ROC curve, as shown below, based on the [DiagnosisMed](#) R package.



Actually, the raw data looks like

	test.values	TP	FN	FP	TN	Sensitivity	Specificity
1	0.037	51	0	97	0	1	0.0000
2	0.038	51	0	96	1	1	0.0103
3	0.039	51	0	91	6	1	0.0619
4	0.040	51	0	84	13	1	0.1340
5	0.041	51	0	74	23	1	0.2371
6	0.042	51	0	67	30	1	0.3093

and the optimal threshold is found as

	test.values	TP	FN	FP	TN	Sensitivity	Specificity
47	0.194	43	8	8	89	0.8431	0.9175

To sum up, I would suggest to generate a regular sequence of possible thresholds and compute Se and Sp in each case; then, choose the one that maximize Se and (1-Sp) (or use other criteria if you want to minimize FP or FN rates).

197 Explanation of finite correction factor

The threshold is chosen such that it ensures convergence of the [hypergeometric distribution](#) ($\sqrt{\frac{N-n}{N-1}}$ is its SD), instead of a binomial distribution (for sampling with replacement), to a normal distribution (this is the Central Limit Theorem, see e.g., [The Normal Curve, the Central Limit Theorem, and Markov's and Chebychev's Inequalities for Random Variables](#)). In other words, when $n/N \leq 0.05$ (i.e., n is not 'too large' compared to N), the FPC can safely be ignored; it is easy to see how the correction factor evolves with varying n for a fixed N : with $N = 10,000$, we have $FPC = .9995$ when $n = 10$ while $FPC = .3162$ when $n = 9,000$. When $N \rightarrow \infty$, the FPC approaches 1 and we are close to the situation of sampling with replacement (i.e., like with an infinite population).

To understand this results, a good starting point is to read some online tutorials on sampling theory where sampling is done without replacement ([simple random sampling](#)). This online tutorial on [Nonparametric statistics](#) has an illustration on computing the expectation and variance for a total.

You will notice that some authors use N instead of $N - 1$ in the denominator of the FPC; in fact, it depends on whether you work with the sample or population statistic: for the variance, it will be N instead of $N - 1$ if you are interested in S^2 rather than σ^2 .

As for online references, I can suggest you

- [Estimation and statistical inference](#)
- [A new look at inference for the Hypergeometric Distribution](#)
- [Finite Population Sampling with Application to the](#)

Hypergeometric Distribution

- Simple random sampling

198 Nonparametric Bayesian analysis in R

Here are some online resources I found interesting without going into detail (and I'm not a specialist of this topic):

- [Hierarchical Dirichlet Processes](#), by Teh et al. (2005)
- [Dirichlet Processes A gentle tutorial](#), by El-Arini (2008)
- [Bayesian Nonparametrics](#), by Rosasco (2010)
- [Non-parametric Bayesian Methods](#), by Ghahmani (2005)

The definitive reference seems to be

N. Hjort, C. Holmes, P. Müller, and S. Walker, editors. *Bayesian Nonparametrics*. Number 28 in Cambridge Series in Statistical and Probabilistic Mathematics. Cambridge University Press, 2010.

About R, there seems to be some other package worth to explore if the [DPpackage](#) does not suit your needs, e.g. [dpmixsim](#), [BHC](#), or [mbsc](#) found [Rseek.org](#).

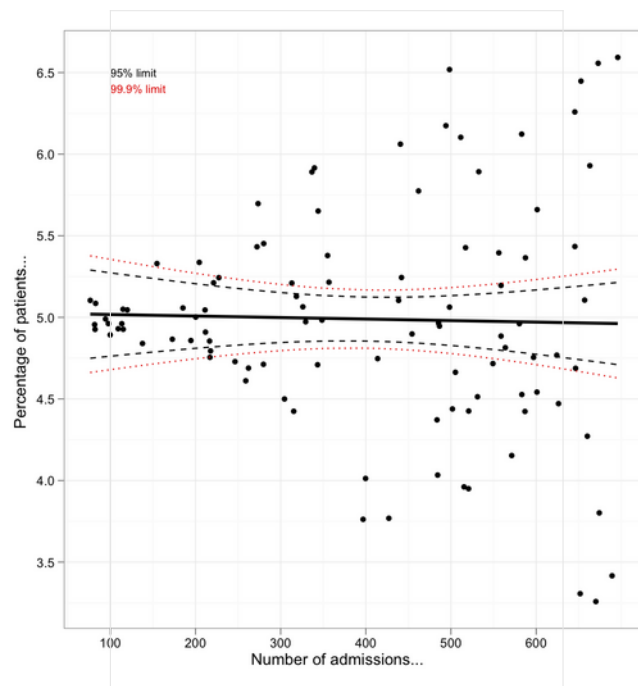
199 How to draw funnel plot using ggplot2 in R?

Although there's room for improvement, here is a small attempt with simulated (heteroscedastic) data:

```
library(ggplot2)
set.seed(101)
x <- runif(100, min=1, max=10)
y <- rnorm(length(x), mean=5, sd=0.1*x)
df <- data.frame(x=x*70, y=y)
m <- lm(y ~ x, data=df)
fit95 <- predict(m, interval="conf", level=.95)
fit99 <- predict(m, interval="conf", level=.999)
df <- cbind.data.frame(df,
  lwr95=fit95[, "lwr"], upr95=fit95[, "upr"],
```

```
lwr99=fit99[, "lwr"], upr99=fit99[, "upr"])
```

```
p <- ggplot(df, aes(x, y))
p + geom_point() +
  geom_smooth(method="lm", colour="black", lwd=1.1, se=FALSE) +
  geom_line(aes(y = upr95), color="black", linetype=2) +
  geom_line(aes(y = lwr95), color="black", linetype=2) +
  geom_line(aes(y = upr99), color="red", linetype=3) +
  geom_line(aes(y = lwr99), color="red", linetype=3) +
  annotate("text", 100, 6.5, label="95% limit", colour="black",
    size=3, hjust=0) +
  annotate("text", 100, 6.4, label="99.9% limit", colour="red",
    size=3, hjust=0) +
  labs(x="No. admissions...", y="Percentage of patients...")
+ theme_bw()
```



200 How do you calculate simple statistics for left censored data in R?

I will let other suggest better alternatives to [NADA](#), but it seems the package is still available on CRAN, in the [Archive](#) section. The last version is from May, 2009.

Installation went fine for me, using

```
R CMD install NADA_1.5-2.tar.gz
```

Under Windows, I guess you can just download the `tgz` and use built-in install facilities.

201 Testing paired frequencies for independence

Log-linear models might be another option to look at, if you want to study your two-way data structure.

If you assume that the two samples are matched (i.e., there is some kind of dependency between the two series of locations) and you take into consideration that data are actually counts that can be considered as scores or ordered responses (as suggested by @caracal), then you can also look at marginal models for matched-pairs, which usually involve the analysis of a square contingency table. It may not be necessarily the case that you end up with such a square Table, but we can also decide of an upper-bound for the number of, e.g. passive sentences. Anyway, models for matched pairs are well explained in Chapter 10 of Agresti, *Categorical Data Analysis*; relevant models for ordinal categories in square tables are testing for *quasi-symmetry* (the difference in the effect of a category from one case to the other follows a linear trend in the category scores), *conditional symmetry* ($\pi_{ab} < \pi_{ba}$ or $\pi_{ab} > \pi_{ba}$, $\forall a, b$), and *quasi-uniform association* (linear-by-linear association off the main diagonal, which in the case of equal-interval scores means an uniform local association). Ordinal quasi-symmetry (OQS) is a special case of linear logit model, and it can be compared to a simpler model where only *marginal homogeneity* holds with an LR test, because ordinal quasi-symmetry + marginal homogeneity = symmetry.

Following Agresti's notation (p. 429), we consider $u_1 \leq \dots \leq u_I$ ordered scores for variable X (in rows) and variable Y (in columns); a or b denotes any row or column. The OQS model reads as the following log-linear model:

$$\log \mu_{ab} = \lambda + \lambda_a + \lambda_b + \beta u_b + \lambda_{ab}$$

where $\lambda_{ab} = \lambda_{ba}$ for all $a < b$. Compared to the usual QS model for nominal data which is $\log \mu_{ab} = \lambda + \lambda_a^X + \lambda_b^Y + \lambda_{ab}$, where $\lambda_{ab} = 0$ would mean independence between the two variables, in the OQS model we impose $\lambda_b^Y - \lambda_b^X = \beta u_b$ (hence introducing the idea of a linear trend). The equivalent logit

representation is $\log(\pi_{ab}/\pi_{ba}) = \beta(u_b - u_a)$, for $a \leq b$.

If $\beta = 0$, then we have symmetry as a special case of this model. If $\beta \neq 0$, then we have stochastically ordered margins, that is $\beta > 0$ means that column mean is higher compared to row mean (and the greater $|\beta|$, the greater the differences between the two joint probabilities distributions π_{ab} and π_{ba} are, which will be reflected in the differences between row and column marginal distributions). A test of $\beta = 0$ corresponds to a test of marginal homogeneity. The interpretation of the estimated β is straightforward: the estimated probability that score on variable X is x units more positive than the score on Y is $\exp(\hat{\beta}x)$ times the reverse probability. In your particular case, it means that $\hat{\beta}$ might allow to quantify the influence that one particular speaker exerts on the other.

Of note, all R code was made available by Laura Thompson in her *S Manual to Accompany Agresti's Categorical Data Analysis*.

Hereafter, I provide some example R code so that you can play with it on your own data. So, let's try to generate some data first:

```
set.seed(56)
d <- as.data.frame(replicate(2, rpois(420, 1.5)))
colnames(d) <- paste("S", 1:2, sep="")
d.tab <- table(d$S1, d$S2, dnn=names(d)) # or xtabs(~S1+S2,
d)
library(vcdExtra)
structable(~S1+S2, data=d)
# library(ggplot2)
# ggfluctuation(d.tab, type="color") + labs(x="S1", y="S2")
+ theme_bw()
```

Visually, the cross-classification looks like this:

	S2	0	1	2	3	4	5	6
S1								
0		17	35	31	8	7	3	0
1		41	41	30	23	7	2	0
2		19	43	18	18	5	0	1
3		11	21	9	15	2	1	0
4		0	3	4	1	0	0	0
5		1	0	0	2	0	0	0
6		0	0	0	1	0	0	0

Now, we can fit the OQS model. Unlike Laura Thompson which used the base `glm()` function and a custom design matrix for symmetry, we can rely on the `gnm` package; we need, however, to add a vec-

tor for numerical scores to estimate β in the above model.

```
library(gnm)
d.long <- data.frame(counts=c(d.tab), S1=gl(7,1,7*7,labels=0:6),
                      S2=gl(7,7,7*7,labels=0:6))
d.long$scores <- rep(0:6, each=7)
summary(mod.oqs <- gnm(counts=scores+Symm(S1,S2), data=d.long,
                      family=poisson))
anova(mod.oqs)
```

Here, we have $\hat{\beta} = 0.123$, and thus the probability that Speaker B scores 4 when Speaker A scores 3 is $\exp(0.123) = 1.13$ times the probability that Speaker B have a score of 3 while Speaker A have a score of 4.

I recently came across the **catspec** R package which seems to offer similar facilities, but I didn't try it. There was a good tutorial at UseR! 2009 about all this stuff: [Introduction to Generalized Nonlinear Models in R](#), but see also the accompanying vignette, [Generalized nonlinear models in R: An overview of the gnm package](#).

If you want to grasp the idea with real data, there are a lot of examples with real data sets in the **vcdExtra** package from Michael Friendly. About the OQS model, Agresti used data on Premarital Sex and Extramarital sex (Table 10.5, p. 421). Results are discussed in §10.4.7 (p. 430), and $\hat{\beta}$ was estimated at -2.86. The code below allow (partly grabbed from Thompson's textbook) to reproduce these results. We would need to relevel factor levels so as to set the same baseline than Agresti.

```
table.10.5 <- data.frame(expand.grid(PreSex=factor(1:4),
                                     ExSex=factor(1:4)),
                        counts=c(144,33,84,126,2,4,14,29,0,2,8,25,0,0,1,3))
table.10.5$scores <- rep(1:4,each=4)
summary(mod.oqs <- gnm(counts=scores+Symm(PreSex,ExSex), data=table.10.5,
                      family=poisson)) # beta = -2.857
anova(mod.oqs) # G^2(5)=2.10
```

202 Two-sample test for multivariate normal distributions under the assumption that means are the same

The **Mauchly's test** allows to test if a given covariance matrix is proportional to a reference (identity or other) and is available through **mauchly.test()** under R. It is mostly used in repeated-measures de-

sign (to test (1) if the dependent variable VC matrices are equal or homogeneous, and (2) whether the correlations between the levels of the within-subjects variable are comparable-altogether, this is known as the *sphericity assumption*).

Box's M statistic is used (in MANOVA or LDA) to test for homogeneity of covariance matrices, but as it is very sensitive to normality it will often reject the null (**R code** not available in standard packages).

Covariance structure models as found in **Structural Equation Modeling** are also an option for more complex stuff (although in multigroup analysis testing for the equality of covariances makes little sense if the variances are not equal), but I have no references to offer actually.

I guess any textbook on multivariate data analysis would have additional details on these procedures. I also found this article for the case where normality assumption is not met:

Aslam, S and Rocke, DM. **A robust testing procedure for the equality of covariance matrices**, Computational Statistics & Data Analysis 49 (2005) 863-874

203 How to visualize what ANOVA does?

Personally, I like introducing linear regression and ANOVA by showing that it is all the same and that linear models amount to partition the total variance: We have some kind of variance in the outcome that can be explained by the factors of interest, plus the unexplained part (called the 'residual'). I generally use the following illustration (gray line for total variability, black lines for group- or individual value specific variability) :

I also like the **heplots** R package, from Michael Friendly and John Fox, but see also **Visual Hypothesis Tests in Multivariate Linear Models: The heplots Package for R**.

Standard ways to explain what ANOVA actually does, especially in the Linear Model framework, are really well explained in **Plane answers to complex questions**, by Christensen, but there are very few illustrations. Saville and Wood's **Statistical methods: The geometric approach** has some examples, but mainly on regression. In Montgomery's **Design and**

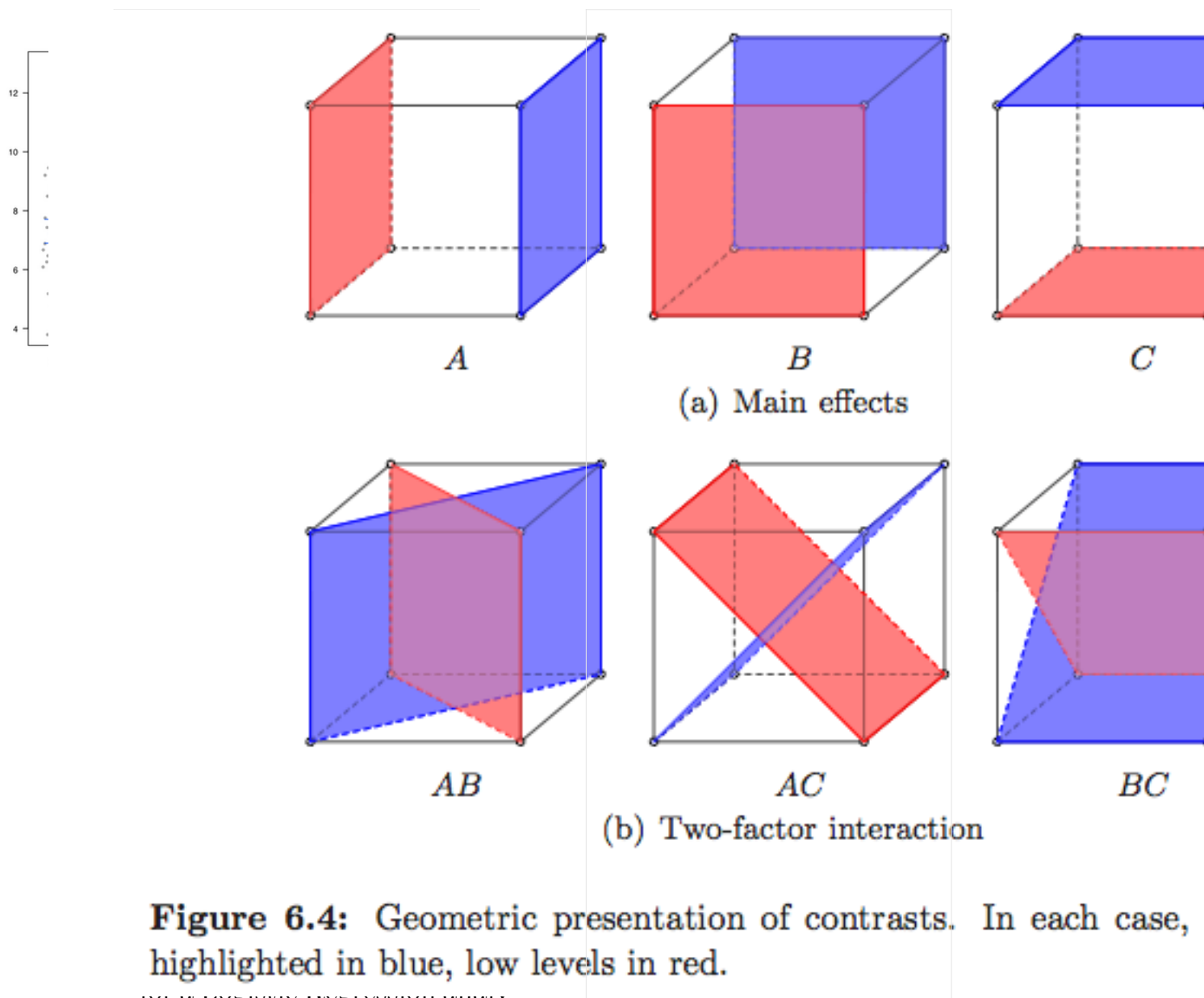


Figure 6.4: Geometric presentation of contrasts. In each case, highlighted in blue, low levels in red.

for a logistic regression analysis

Take a look at the `lrm()` function from the **Design** package. It features everything you need for fitting GLM. The Hosmer and Lemeshow test has limited power and depends on arbitrary discretization; it is discussed in Harrell, *Regression Modeling Strategies* (p. 231) and on the **R-help** mailing-list. There is also a comparison of GoF tests for logistic regression in *A comparison of goodness-of-fit tests for the logistic regression model*, Stat. Med. 1997 16(9):965.

Here is an example of use:

```
library(Design) # depends on Hmisc
x1 <- rnorm(500)
x2 <- rnorm(500)
L <- x1+abs(x2)
y <- ifelse(runif(500)<=plogis(L), 1, 0)
f <- lrm(y ~ x1+x2, x=TRUE, y=TRUE)
resid(f, 'gof')
```

which yields something like

Sum of squared errors	Expected value H0
100.33517914	100.37281429
Z	P
-0.09998187	0.92035872

likelihood method, with applications in veterinary epidemiology. You can also take a look at Venables and Ripley's MASS book, §8.4, pp. 220-221.

but see `help(residuals.lrm)` for additional help.

The following thread contains critical discussions that might also be helpful: Logistic Regression: Which pseudo R-squared measure is the one to report (Cox & Snell or Nagelkerke)?

205 Why is there a difference between manually calculating a logistic regression 95% confidence interval, and using the `confint()` function in R?

After having fetched the data from the [accompaning website](#), here is how I would do:

```
chdage <- read.table("chdage.dat", header=F, col.names=c("id", "age", "dage"))
chdage$aged <- ifelse(chdage$dage>=55, 1, 0)
mod.lr <- glm(chd ~ aged, data=chdage, family=binomial)
summary(mod.lr)
```

The 95% CIs based on profile likelihood is obtained with

```
require(MASS)
exp(confint(mod.lr))
```

This often is the default if package `MASS` is automatically loaded. In this case, I get

	2.5 %	97.5 %
(Intercept)	0.2566283	0.7013384
aged	3.0293727	24.7013080

Now, if I want to compare with 95% Wald CIs (based on asymptotic normality) like the one you computed by hand, I would use `confint.default()` instead; this yields:

	2.5 %	97.5 %
(Intercept)	0.2616579	0.7111663
aged	2.8795652	22.8614705

Wald CIs are good in most situation, although profile likelihood-based may be useful with complex sampling strategies. If you want to grasp the idea of how they work, here is a brief overview of the main principles: [Confidence intervals by the profile](#)

206 Which test to find out best concentration (the one having maximum effect)?

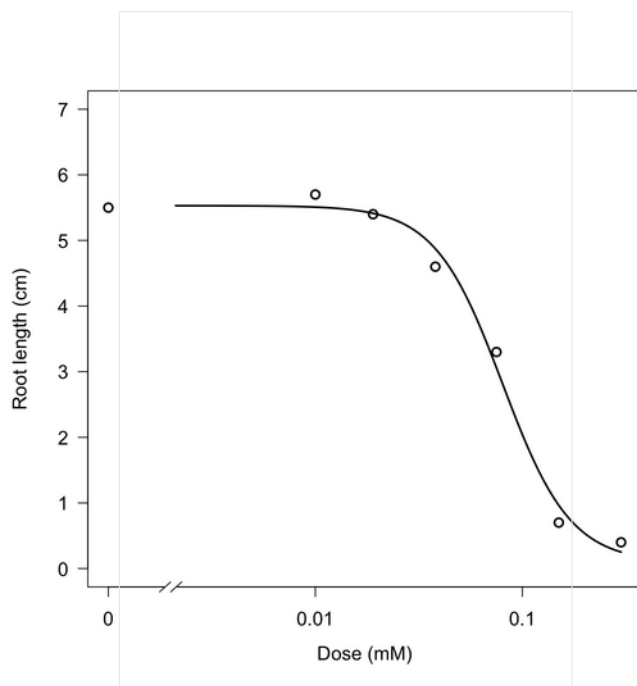
Are you not looking after method for studying dose-response relationship? From the description you gave, I guess your growth curves are highly non-linear. Moreover, the problem of using simple ANOVA to compare two growth curve (i.e., equivalent to a time course curve) is that it would be blind to the fact that the different doses are administered in a sequential order, though it should give a reasonable first idea of *some kind* of differences between the two, if any. However, multiple comparisons are likely not to answer the question you really want to ask since we often expect very few differences at baseline and larger ones at intermediate concentration levels, whereas what you generally want to assess is what's the lowest concentration level that yields a significant difference between the two products.

If you are using R, the `drc` package (see also [www.bioassay.dk](#)) allows to fit various models to DR data, and plot the resulting DR curve like the one shown below. It was generated from the on-line help with data `secalonic` whose description is:

Data stem from an experiment assessing the inhibitory effect of secalonic acids on plant growth.

Gong, X. and Zeng, R. and Luo, S. and Yong, C. and Zheng, Q. (2004) Two new secalonic acids from *Aspergillus Japonicus* and their allelopathic effects on higher plants, *Proceedings of International Symposium on Allelopathy Research and Application*, 27-29 April, Shanshui, Guangdong, China (Editors: R. Zeng and S. Luo), 209-217.

Ritz, C (2009) Towards a unified approach to dose-response modeling in ecotoxicology To appear in *Environ Toxicol Chem*.



207 How to choose nlme or lme4 R library for mixed effects models?

Both packages use `Lattice` as the backend, but `nlme` has some nice features like `groupedData()` and `lmList()` that are lacking in `lme4` (IMO). From a practical perspective, the two most important criteria seem, however, that

1. `lme4` extends `nlme` with other link functions: in `nlme`, you cannot fit outcomes whose distribution is not gaussian, `lme4` can be used to fit mixed-effects logistic regression, for example.
2. in `nlme`, it is possible to specify the variance-covariance matrix for the random effects (e.g. an `AR(1)`); it is not possible in `lme4`.

Now, `lme4` can easily handle very huge number of random effects (hence, No. individuals in a given study) thanks to its C part and the use of sparse matrices. The `nlme` package has somewhat been superseded by `lme4` so I won't expect people spending much time developing add-ons on top of `nlme`. Personally, when I have a continuous response in my model, I tend to use both packages, but I'm now versed to the `lme4` way for fitting GLMM.

Rather than buying a book, take a look first at

the Doug Bates' draft book on R-forge: [lme4: Mixed-effects Modeling with R](#).

208 What does the Psi term in factor analysis signify?

Following Bishop's notation, the FA model is written as (Eq. 12.64, p. 584):

$$p(x|z) = \mathcal{N}(x|\mathbf{W}z + \mu, \psi)$$

where ψ is a $D \times D$ diagonal matrix of so-called variable uniquenesses, that is the variance not accounted for by the latent factors, whereas \mathbf{W} reflects factor loadings λ_i , that is the correlation of variable i with factors represented in z (more exactly, the square of λ_i is the variance explained by the latent factor).

If you're not familiar with the FA literature, I would suggest "lighter" approach, e.g. William Revelle has good tutorials on his website [personality-project.org](#); especially, I would suggest Chapter 6 of his forthcoming book on Psychometric methods entitled [Constructs, Components, and Factor models](#). You will shortly understand the relations between PCA and FA. Specifically, with PCA, we are constructing linear combinations of observed variables (this yields a composite variable), whereas in FA we are expressing each variable as a weighted combination of hypothesized latent factors (where weights are called loadings) plus an error term (the ψ in the above formula). In sum, the FA model incorporates a model for noise—this is what is expressed in Equation 12.65; but see [What are the differences between Factor Analysis and Principal Component Analysis](#), for additional discussion.

209 How to compute the confidence intervals on regression coefficients in PLS?

Do you know this article: [PLS-regression: a basic tool of chemometrics](#)? Deriving SE and CI for the PLS parameters is described in §3.11.

I generally rely on Bootstrap for computing CIs, as suggested in e.g., Abdi, H. [Partial least squares regression and projection on latent structure regression \(PLS Regression\)](#). I seem to remember there are theoretical solutions discussed in Tenenhaus M. (1998) *La régression PLS: Théorie et pratique* (Technip),

but I cannot check for now as I don't have the book. For now, there are some useful R packages, like [plsRglm](#).

P.S. I just discovered [Nicole Krämer](#)'s article, in reference to the [plsdo](#) R package.

210 Logit with ordinal independent variables

To add to @dmk38's response, "any set of scores gives a *valid* test, provided they are constructed without consulting the results of the experiment. If the set of scores is poor, in that it badly distorts a numerical scale that really does underlie the ordered classification, the test will not be sensitive. The scores should therefore embody the best insight available about the way in which the classification was constructed and used." (Cochran, 1954, cited by Agresti, 2002, pp. 88-89). In other words, treating an ordered factor as a numerically scored variable is merely a modelling issue. Provided it makes sense, this will only impact the way you interpret the result, and there is no definitive rule of thumb on how to choose the best representation for an ordinal variable.

Consider the following example on maternal Alcohol consumption and presence or absence of congenital malformation (Agresti, [Categorical Data Analysis](#), Table 3.7 p.89):

```

      0    <1 1-2 3-5 6+
Absent 17066 14464 788 126 37
Present 48    38   5   1   1

```

In this particular case, we can model the outcome using logistic regression or simple association table. Let's do it in R:

```

tab3.7 <- matrix(c(17066,48,14464,38,788,5,126,1,37,1), nr=2,
  dimnames=list(c("Absent","Present"),
    c("0","<1","1-2","3-5","6+")))
library(vcd)
assocstats(tab3.7)

```

Usual χ^2 (12.08, $p=0.016751$) or LR (6.20, $p=0.184562$) statistic (with 4 df) do not account for the ordered levels in Alcohol consumption.

Treating both variables as ordinal with equally spaced scores (this has no impact for binary variables, like malformation, and we choose the baseline as 0=absent), we could test for a linear by linear as-

sociation. Let's first construct an exploded version of this contingency Table:

```

library(reshape)
tab3.7.df <- untable(data.frame(malform=gl(2,1,10,labels=0:1),
  alcohol=gl(5,2,10,labels=colnames(tab3.7))),
  c(tab3.7))
# xtabs(~malform+alcohol, tab3.7.df) # check

```

Then we can test for a linear association using

```

library(coin)
#lbl_test(as.table(tab3.7))
lbl_test(malform ~ alcohol, data=tab3.7.df)

```

which yields $\chi^2(1) = 1.83$ with $p = 0.1764$. Note that this statistic is simply the correlation between the two series of scores (that Agresti called $M^2 = (n-1)r^2$), which is readily computed as

```
cor(sapply(tab3.7.df, as.numeric))[1,2]^2*(32574-1)
```

As can be seen, there is not much evidence of a clear association between the two variables. As done by Agresti, if we choose to recode Alcohol levels as $\{0,0.5,1.5,4,7\}$, that is using mid-range values for an hypothesized continuous scale with the last score being somewhat purely arbitrary, then we would conclude to a larger effect of maternal Alcohol consumption on the development of congenital malformation:

```

lbl_test(malform ~ alcohol, data=tab3.7.df,
  scores=list(alcohol=c(0,0.5,1.5,4,7)))

```

yields a test statistic of 6.57 with an associated p -value of 0.01037.

There are alternative coding schemes, including *midranks* (in which case, we fall back to Spearman ρ instead of Pearson r) that is discussed by Agresti, but I hope you catch the general idea here: It is best to select scores that actually reflect a reasonable measures of the distance between adjacent categories of your ordinal variable, and equal spacing is often a good compromise (in the absence of theoretical justification).

Using the GLM approach, we would proceed as follows. But first check how Alcohol is encoded in R:

```
class(tab3.7.df$alcohol)
```

It is a simple unordered factor ("**factor**"), hence a nominal predictor. Now, here are three models were we consider Alcohol as a nominal, ordinal or

continuous predictor.

```
summary(mod1 <- glm(malform ~ alcohol, data=tab3.7.df,
                    family=binomial))
summary(mod2 <- glm(malform ~ ordered(alcohol), data=tab3.7,
                    family=binomial))
summary(mod3 <- glm(malform ~ as.numeric(alcohol), data=tab3,
                    family=binomial))
```

The last case implicitly assumes an equal-interval scale, and the $\hat{\beta}$ is interpreted as @dmk38 did: it reflects the effect of a one-unit increase in Alcohol on the outcome through the logit link, that is the increase in probability of observing a malformation (compared to no malformation, i.e. the odds-ratio) is $\exp(\hat{\theta}) = \exp(0.228) = 1.256$. The Wald test is not significant at the usual 5% level. In this case, the design matrix only includes 2 columns: the first is a constant column of 1's for the intercept, the second is the numerical value (1 to 5) for the predictor, as in a simple linear regression. In sum, this model tests for a linear effect of Alcohol on the outcome (on the logit scale).

However, in the two other cases (`mod1` and `mod2`), we get different output because the design matrix used to model the predictor differs, as can be checked by using:

```
model.matrix(mod1)
model.matrix(mod2)
```

We can see that the associated design matrix for `mod1` includes dummy variables for the $k-1$ levels of Alcohol (0 is always the baseline) after the intercept term in the first column, whereas in the case of `mod2` we have four columns of contrast-coded effects (after the column of 1's for the intercept). The coefficient for the category "3-5" is estimated at 1.03736 under `mod1`, but 0.01633 under `mod2`. Note that AIC and other likelihood-based measures remain, however, identical between these two models.

You can try assigning new scores to Alcohol and see how it will impact the predicted probability of a malformation.

211 Confirmatory and explanatory factor analysis

What is the rationale of applying an exploratory/unsupervised method (PCA or FA with VARIMAX rotation) after having tested a confirmatory model, especially

if this is done on the same sample?

In your CFA model, you impose constraints on your pattern matrix, e.g. some items are supposed to load on one factor but not on the others. A large modification index indicates that freeing a parameter or removing an equality constraint could result in better model fit. Item loadings are already available through your model fit.

On the contrary, in PCA or FA there is no such constraint, even following an orthogonal rotation (whose purpose is just to make factor more interpretable in that items would generally tend to load more heavily on a factor than on several ones). But, it is worth noting that these models are conceptually and mathematically different: the FA model is a measurement model, where we assume that there is some unique error attached to each item; this is not the case under the PCA framework. It is thus not surprising that you failed to replicate your factor structure, which may be an indication that there are possible item cross-loading, low item reliability, low stability in your factor structure, or the existence of a higher-order factor structure, that is enhanced by your low sample size.

In both case, but especially CFA, $N = 96$ is a very limited sample size. Although some authors have suggested a ratio individuals:items of 5 to 10, this is merely the number of dimensions that is important. In your case, the estimation of your parameters will be noisy, and in the case of PCA you may expect fluctuations in your estimated loadings (just try bootstrap to get an idea of 95% CIs).

212 Is there a quote like this from some statistician?

Maybe you are after this talk?

Tutorial: Methods for Reproducible Research, by Roger D. Peng (slide 3)

Also, papers on Reproducible research written by de Leeuw that I am aware of are **Reproducible Research: the Bottom Line**, and **Statistical Software – Overview**. But a quick check didn't reveal any citation like the one you show.

213 Get the number of parameters of a linear model

Try something like:

```
> x <- replicate(2, rnorm(100))
> y <- 1.2*x[,1]+rnorm(100)
> summary(lm.fit <- lm(y~x))
> length(lm.fit$coefficients)
[1] 3
> # or
> length(coef(lm.fit))
[1] 3
```

You can have a better idea of what an R object includes with

```
> str(lm.fit)
```

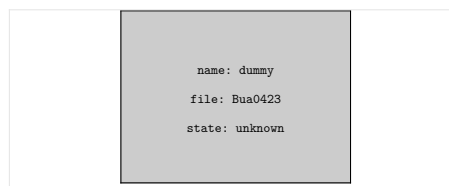
214 Probability and limits

The two key elements that you need to be familiar with before working through this exercise are:

- The **sampling distribution** of a mean: in your case, the average of your four measurements will follow a normal distribution, with a mean that equals that of the parent distribution but with a standard deviation of $3/\sqrt{4}$.
- How to translate probabilistic assertions in terms of the underlying **Probability Density Function**: given that there is an infinite number of possible PDFs for a distribution described by two parameters (its location and shape), it is often more convenient to work with the standardized normal distribution which is simply $\mathcal{N}(0; 1)$, because if $X \sim \mathcal{N}(\mu; \sigma)$, then we know that $Z = \frac{X-\mu}{\sigma} \sim \mathcal{N}(0; 1)$.

Then, you just have to figure out how to find z_1 and z_2 such that $\Pr(z_1 \leq Z \leq z_2) = 1 - 0.01$. It often helps to draw the graph of $\Pr(Z \leq z)$ which is bell-shaped, centered on its mean, and whose total area equals 1. For instance, for a given quantile z_1 , $\Pr(Z < z_1) = p_1$ where p_1 is the shaded area shown below (here, $z_1 = -1$, that is 1 standard deviation below the mean):

As the total area equals 1, the remaining (unshaded) area equals $1-p_1$. Likewise, you may readily express any bounded area as a sum or difference of such inequalities.



215 Automating model selection criteria production

Ok, here is another way (which is certainly not as elegant or concise than the other ones, but it has the merit of not requiring Emacs to check parenthesis matches :-)

Let's say we have a vector of predictors of interest like this

```
> Xs <- paste("X", 1:4, sep="")
```

Then, we can just use

```
> allXs <- lapply(seq(along=Xs),
                    function(x) combn(Xs, x, simplify=FALSE,
                                       FUN=paste, collapse="+"))
```

where

```
> unlist(allXs)
```

gives you all 15 combinations of the X's.

Another option is to just change the right-hand side of a formula, say

```
fm <- as.formula(paste("Y ~ ", paste(Xs, collapse="+")))
```

so as to reflect the different combinations that you enumerate in your **comb_list** object. This can be done using the **Formula** package:

```
> fm <- as.formula(paste("Y ~ ", paste(Xs, collapse="|")))
> fm2 <- Formula(fm)
> foo <- function(x) formula(fm2, rhs=x, collapse=TRUE)
> foo(1:2)
Y ~ X1 + X2
<environment: 0x102593040>
> foo(c(1,3,4))
Y ~ X1 + X3 + X4
<environment: 0x1021d02a0>
```

216 p-vector and K-vector

It's merely some generic notation for a vector of p attributes or variables observed on $i = 1, \dots, N$ individuals, so that you can define $X^T = (X_1, X_2, \dots, X_p)$ as a vector of inputs, in the feature (or input) space (and each individual will have one such vector of observed inputs).

The K notation seems to be reserved to the output space: in a classical linear regression model where $Y = X\beta$, Y is a scalar ($K = 1$), whereas in a multivariate setting (say, you record weight, height, and color) it could be a K -vector (i.e., 3-vector with my example).

217 R package for combining factor levels for datamining?

It seems it's just a matter of "releveling" the factor; no need to compute partial sums or make a copy of the original vector. E.g.,

```
set.seed(101)
a <- factor(LETTERS[sample(5, 150, replace=TRUE,
                           prob=c(.1, .15, rep(.75/3,3))]])
p <- 1/5
lf <- names(which(prop.table(table(a)) < p))
levels(a)[levels(a) %in% lf] <- "Other"
```

Here, the original factor levels are distributed as follows:

```
A B C D E
18 23 35 36 38
```

and then it becomes

```
Other C D E
41 35 36 38
```

It may be conveniently wrapped into a function. There is a `combine_factor()` function in the `reshape` package, so I guess it could be useful too.

Also, as you seem interested in data mining, you might have a look at the `caret` package. It has a lot of useful features for data preprocessing, including functions like `nearZeroVar()` that allows to flag predictors with very imbalanced distribution of observed values (See the vignette, [example data, pre-processing functions, visualizations and other functions](#), p. 5, for example of use).

218 Effect size of Spearman's rank test

I see no obvious reason not to do so. As far as I know, we usually make a distinction between two kind of effect size (ES) measures for qualifying the strength of an observed association: ES based on d (difference of means) and ES based on r (correlation). The latter includes Pearson's r , but also Spearman's ρ , Kendall's τ , or the multiple correlation coefficient.

As for their interpretation, I think it mainly depends on the field you are working in: A correlation of .20 would certainly not be interpreted in the same way in psychological vs. software engineering studies. Don't forget that Cohen's three-way classification—small, medium, large—was based on behavioral data, as discussed in Kraemer et al. (2003), p. 1526. In their Table 1, they made no distinction about the different types of ES measures belonging to the r family. There have by no way an absolute meaning and should be interpreted with reference to established results or literature review.

I would like to add some other references that provide useful reviews of common ES measures and their interpretation.

References

1. Helena C. Kraemer, George A. Morgan, Nancy L. Leech, Jeffrey A. Gliner, Jerry J. Vaske, and Robert J. Harmon (2003). [Measures of Clinical Significance](#). *J Am Acad Child Adolesc Psychiatry*, 42(12), 1524-1529.
2. Christopher J. Ferguson (2009). [An Effect Size Primer: A Guide for Clinicians and Researchers](#). *Professional Psychology: Research and Practice*, 40(5), 532-538.
3. Edward F. Fern and Kent B. Monroe (1996). [Effect-Size Estimates: Issues and Problems in Interpretation](#). *Journal of Consumer Research*, 23, 89-105.
4. Daniel J. Denis (2003). [Alternatives to Null Hypothesis Significance Testing](#). *Theory and Science*, 4(1).
5. Paul D. Ellis (2010). [The Essential Guide to Effect Sizes](#). Cambridge University Press. – just browsed the TOC

219 What logistic regression is best to use?

Logistic regression is to be used when the outcome of interest is a binary variable (e.g., success/failure), whereas multinomial logistic is reserved for the case of a multi-category response variable (e.g., blue/red/green). In both cases, the response variable to be predicted is a categorical variable. The predictors might be categorical or continuous variables.

From what you described, you are interested in predicting scores on a social attribution task based on observed scores on a reading scores. If both sets of scores are numerical, then it is simply a linear regression model. In R, it is something like

```
# fake data
x <- rnorm(100)
y <- 1.2*x + rnorm(100)
# model fit
summary(lm(y ~ x))
```

The output reads:

```
Call:
lm(formula = y ~ x)

Residuals:
    Min       1Q   Median       3Q      Max
-2.6635 -0.6227 -0.1589  0.6360  2.2494

Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.03769    0.10042   -0.375    0.708
x             1.31590    0.10796   12.188 <2e-16 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 1.003 on 98 degrees of freedom
Multiple R-squared:  0.6025, Adjusted R-squared:  0.5985
F-statistic: 148.6 on 1 and 98 DF, p-value: < 2.2e-16
```

Here an increase of one unit in x is associated to an increase of 1.32 units in y . This is basically what the regression coefficient tells you when you assume a model like $E(y|x) = \beta_0 + \beta_1 x + \varepsilon$, where ε are independent and identically distributed as a centered gaussian with variance σ^2 (unknown).

Now, you may still be interested in a binary outcome. For instance, I can define “Success” as $y > 0$. In this case, a logistic regression would look like

```
yb <- ifelse(y>0, 1, 0)
summary(glm(yb ~ x, family=binomial))
```

where the regression coefficient gives you the log of the odds of passing the exam (compared to failing to reach a score of 0):

```
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) -0.5047    0.3194  -1.580    0.114
x             3.3959    0.6871   4.943  7.7e-07 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Edit

Some pointers with multinomial logistic regression have already been given. What I wonder is the outcome you want to model: If you’re interested in whether the autistic group has lower scores on the attribution task (as would be expected from the literature), then in a logistic model both scores will be continuous predictors, and reading scores will be considered as a covariate; in other words, you model the odds of being in one of the diagnostic class as a function of attrition task, after adjusting for baseline differences in language proficiency. But it seems it would make sense only if the diagnostic categories are not *a priori* defined and it does not directly answer the question you are asking (whether language is a predictor for attrition that may act differentially according to the diagnostic group); otherwise, I would rather model the attrition scores as a function of a grouping variable (diagnostic category) + reading scores, which is basically an **ANCOVA** model.

Both models are available in R: `mlogit()` in the package **mlogit** for multinomial logistic regression, and `lm()` for ANCOVA.

220 Can principal component analysis be applied to categorical data?

Although a PCA applied on binary data would yield results comparable to those obtained from a **Multiple Correspondence Analysis** (factor scores and eigenvalues are linearly related), there are more appropriate techniques to deal with mixed data types, namely Multiple Factor Analysis for mixed data available in the **FactoMineR** R package (`AFDM()`). If your

variables can be considered as structured subsets of descriptive attributes, then **Multiple Factor Analysis** (**MFA()**) is also an option.

The challenge with categorical variables is to find a suitable way to represent distances between variable categories and individuals in the factorial space. To overcome this problem, you can look for a non-linear transformation of each variable—whether it be nominal, ordinal, polynomial, or numerical—with optimal scaling. This is well explained in **Gifi Methods for Optimal Scaling in R: The Package homals**, and an implementation is available in the corresponding R package **homals**.

221 R-square from rpart model

The advantage of R is that most of the time you can easily access the source code. So in your case, start with

```
> rsq.rpart
```

(without parenthesis) to see what the function actually does. The R^2 values are obtained as

```
tmp <- printcp(fit)
rsq.val <- 1-tmp[,c(3,4)]
```

where for each row (aka, No. splits) we have the “apparent” and “relative” (wrt. cross-validation) statistics.

222 Annotating graphs in R

A quick and dirty way to paste some text and numerical results along the labels of your legend is to simply rename the factor levels. For instance,

```
df <- data.frame(x=rnorm(100), y=rnorm(100), f=gl(2,50))
df$f2 <- df$f
levels(df$f2) <- paste(levels(df$f), tapply(df$y, df$f, mean),
  sep=": ")
p <- ggplot(data=df) + geom_point(aes(x=x, y=y, color=f2))
p + opts(legend.position = 'bottom', legend.title=NULL)
```

You can add whatever you want into the new labels, such as mean, min, max, etc. (e.g., create a custom function, inspired from **summary()** that returns the values you want, and append them to **c("In", "Out")**).

223 Good text for resampling?

As for a good reference, I would recommend Philip Good, *Resampling Methods: A Practical Guide to Data Analysis* (Birkhäuser Boston, 2005, 3rd ed.) for an applied companion textbook. And here is **An Annotated Bibliography for Bootstrap Resampling. Resampling methods: Concepts, Applications, and Justification** also provides a good start.

There are many R packages that facilitate the use of resampling techniques:

- **boot**, for bootstrapping – but see also P. Burns, *The Statistical Bootstrap and Other Resampling Methods*, for illustrations
- **coin**, for permutation tests (bit see the accompanying **vignette** which includes extensive help)

(There are many other packages...)

224 What are some alternatives to a boxplot?

A boxplot isn't that complicated. After all, you just need to compute the three **quartiles**, and the min and max which define the range; a subtlety arises when we want to draw the whiskers and various methods have been proposed. For instance, in a **Tukey boxplot** values outside 1.5 times the inter-quartile from the first or third quartile would be considered as outliers and displayed as simple points. See also **Methods for Presenting Statistical Information: The Box Plot for a good overview**, by Kristin Potter. The R software implements a slightly different rule but the source code is available if you want to study it (see the **boxplot()** and **boxplot.stats()** functions). However, it is not very useful when the interest is in identifying outliers from a very skewed distribution (but see, **An adjusted boxplot for skewed distributions**, by Hubert and Vandervieren, CSDA 2008 52(12)).

As far as online visualization is concerned, I would suggest taking a look at **Protovis** which is a plugin-free js toolbox for interactive web displays. The **examples** page has very illustrations of what can be achieved with it, in very few lines.

225 Should I re-shuffle my data?

As you already use a holdout sample, I would say you should keep it and build your new models on the same training sample so that all models will consider the same relationships between features. In addition, if you perform feature selection, the samples must be left out before any of these filtering stages; that is, feature selection must be included in the cross-validation loop.

Of note, there are more powerful methods than a 0.67/0.33 split for model selection, namely k-fold cross-validation or leave-one-out. See e.g. [The Elements of Statistical Learning](#) (§7.10, pp. 241-248), www.modelselection.org or [A survey of cross-validation procedures for model selection](#) by Arlot and Celisse (more advanced mathematical background required).

226 MCMC method — good sources

For online tutorials, there are

- [A tutorial in MCMC](#), by Sahut (2000)
- [Tutorial on Markov Chain Monte Carlo](#), by Hanson (2000)
- [Markov Chain Monte Carlo for Computer Vision](#), by Zhu et al. (2005)
- [Introduction to Markov Chain Monte Carlo simulations and their statistical analysis](#), by Berg (2004).

[Practical Markov Chain Monte Carlo](#), by Geyer (*Stat. Science*, 1992), is also a good starting point, and you can look at the [MCMCpack](#) or [mcmc](#) R packages for illustrations.

227 What are the practical & interpretation differences between alternatives and logistic regression?

Disclaimer: It is certainly far from being a full answer to the question!

I think there are at least two levels to consider before establishing a distinction between all such methods:

- **whether a single model is fitted or not:** This helps opposing methods like logistic regression vs. RF or [Gradient Boosting](#) (or more generally [Ensemble methods](#)), and also put emphasis on parameters estimation (with associated asymptotic or bootstrap confidence intervals) vs. classification or prediction accuracy computation;
- **whether all variables are considered or not:** This is the basis of feature selection, in the sense that penalization or regularization allows to cope with “irregular” data sets (e.g., large p and/or small n) and improve generalizability of the findings.

Here are few other points that I think are relevant to the question.

In case we consider several models—the same model is fitted on different subsets (individuals and/or variables) of the available data, or different competitive models are fitted on the same data set—, [cross-validation](#) can be used to avoid overfitting and perform model or feature selection, although CV is not limited to this particular cases (it can be used with [GAMs](#) or penalized GLMs, for instance). Also, there is the traditional interpretation issue: more complex models often implies more complex interpretation (more parameters, more stringent assumptions, etc.).

Gradient boosting and RFs overcome the limitations of a single decision tree, thanks to [Boosting](#) whose main idea is to combine the output of several weak learning algorithms in order to build a more accurate and stable decision rule, and [Bagging](#) where we “average” results over resampled data sets. Altogether, they are often viewed as some kind of black boxes in comparison to more “classical” models where clear specifications for the model are provided (I can think of three classes of models: [parameteric](#), [semi-parametric](#), [non-parametric](#)), but I think the discussion held under this other thread [The Two Cultures: statistics vs. machine learning?](#) provide interesting viewpoints.

Here are a couple of papers about feature selection and some ML techniques:

1. Saeys, Y, Inza, I, and Larrañaga, P. [A review of feature selection techniques in bioinformatics](#), *Bioinformatics* (2007) 23(19): 2507-2517.
2. Dougherty, ER, Hua J, and Sima, C. [Performance](#)

of [Feature Selection Methods](#), Current Genomics (2009) 10(6): 365–374.

3. Boulesteix, A-L and Strobl, C. [Optimal classifier selection and negative bias in error rate estimation: an empirical study on high-dimensional prediction](#), BMC Medical Research Methodology (2009) 9:85.
4. Caruana, R and Niculescu-Mizil, A. [An Empirical Comparison of Supervised Learning Algorithms](#). Proceedings of the 23rd International Conference on Machine Learning (2006).
5. Friedman, J, Hastie, T, and Tibshirani, R. [Additive logistic regression: A statistical view of boosting](#), Ann. Statist. (2000) 28(2):337-407. (With discussion)
6. Olden, JD, Lawler, JJ, and Poff, NL. [Machine learning methods without tears: a primer for ecologists](#), Q Rev Biol. (2008) 83(2):171-93.

And of course, [The Elements of Statistical Learning](#), by Hastie and coll., is full of illustrations and references. Also be sure to check the [Statistical Data Mining Tutorials](#), from Andrew Moore.

228 What kinds of things can I predict with a naive Bayesian classifier?

[The Elements of Statistical Learning](#), by Hastie et al. has a lot of illustrations of Machine Learning applications, and all [data sets](#) are available on the companion website, including data on spam as on the [Ruby Classifier](#) webpage.

As for a gentle introduction to Bayes classifier, I would suggest to look at the following tutorial from Andrew Moore: [A Short Intro to Naive Bayesian Classifiers](#) (many [other tutorials](#) are also available).

229 Regression with an unknown dependent variable - estimating “likelihood” to do something

I agree with @Aniko that you won’t be able to “predict” anything without an outcome. Now @Andy

suggestion makes sense, provided you find a users database sharing similar characteristics. As an example of related studies, I guess you might find interesting google hits on users’ characteristics in studies on Twitter, Facebook or other social networks or community driven sites. Here is a couple of references that I just found: [Comparing community structure to characteristics in online collegiate social networks](#) (Traud et al., [arXiv:0809.0690](#)); [Social Computing Privacy Concerns: Antecedents & Effects](#) (Nov and Wattal, CHI 2009).

Aside from users’ characteristics, willingness to participate or contribute to a social network is also a function of its characteristics (number of members, contact frequency, content quality, etc.).

Finally, you still can do some kind of exploratory analysis on your data set. Specifically, a cluster analysis would help to highlight groups of individuals sharing similar response profiles. Another approach would be to use [Multiple correspondence analysis](#) to uncover potentially interesting patterns in your data. Those features might be related to already published results in a further step. Obviously, you can’t regress an outcome that is still to be observed from this, but you will certainly get a better idea of how structured your data are.

230 How should I analyze repeated-measures individual differences experiments?

There were already some useful comments, that are probably waiting for some updates in the question, so I will just drop some general online references:

- [Practical Data Analysis for the Language Sciences with R](#), Baayen (2008)
- [Categorical data analysis: Away from ANOVAs \(transformation or not\) and towards logit mixed models](#), Jaeger (J. Mem. Language 2008 59(4))

Examples using R may be found on Doug Bates’ [lme4 - Mixed-effects models project](#).

231 An R function for performing searches

You should first replace your for loop with something like `apply(d7_dataset, 1, foo)`, where `foo()`

is either your function or something along those lines, e.g. `gregexpr()`. The result of `gregexpr()` is a list of numeric vectors with attributes similar to `regexpr()` but giving all matches in each element.

On a related point, there was another function that was proposed as a more user-friendly alternative to `gregexpr()`: `easyGregexpr`.

The following example gives you the number of matches for each row when considering a list of three motifs (in a 10x100 matrix):

```
dd <- replicate(100, replicate(10, paste(sample(letters[1:4], 2),
                                           collapse="")))
pat <- list("aa", "ab", "cd")
foo <- function(d, p) apply(d, 1, function(x) length(grep(p, x)))
lapply(pat, function(x) foo(dd, x))
```

232 R: update a graph dynamically

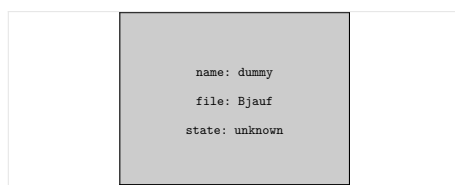
For offline visualization, you can generate PNG files and convert them to an animated GIF using `ImageMagick`. I used it for demonstration (this redraw all data, though):

```
source(url("http://aliquote.org/pub/spin_plot.R"))
dd <- replicate(3, rnorm(100))
spin.plot(dd)
```

This generates several PNG files, prefixed with `fig`. Then, on an `un*x` shell,

```
convert -delay 20 -loop 0 fig*.png sequence.gif
```

gives this animation (which is inspired from *Modern Applied Biostatistical Methods using S-Plus*, S. Selvin, 1998):



Another option which looks much more promising is to rely on the `animation` package. There is an example with a `Moving Window Auto-Regression` that should let you get started with.

233 How do I interpret $\text{Exp}(B)$ in Cox Regression?

Generally speaking, $\exp(\hat{\beta}_1)$ is the ratio of the hazards between two individuals whose values of x_1 differ by one unit when all other covariates are held constant. The parallel with other linear models is that in Cox regression the hazard function is modeled as $h(t) = h_0(t) \exp(\beta'x)$, where $h_0(t)$ is the baseline hazard. This is equivalent to say that $\log(\text{group hazard}/\text{baseline hazard}) = \log((h(t)/h_0(t)) = \sum_i \beta_i x_i$. Then, a unit increase in the independent variable i is associated with β_i increase in the log hazard rate. The regression coefficient allow thus to quantify the log of the hazard in the treatment group (compared to the control or placebo group), accounting for the covariates included in the model; it is interpreted as a relative risk (assuming no time-varying coefficients).

In the case of logistic regression, the regression coefficient reflects the log of the **odds-ratio**, hence the interpretation as an k -fold increase in risk. So yes, the interpretation of hazard ratios shares some resemblance with the interpretation of odds ratios.

Be sure to check Dave Garson's website where there is some good material on **Cox Regression** with SPSS.

234 Doing correlation on one variable vs many

To help you get started with the visualization, here is a snippet of R code with simulated data (a matrix with age and counts for 20 words, arranged in columns, for 100 subjects). The computation are done as proposed my @mbq (correlation).

```
n <- 100 # No. subjects
k <- 20 # No. words
words <- paste("word", 1:k, sep="")
df <- data.frame(age=rnorm(n, mean=25, sd=5),
                 replicate(k, sample(1:10, n, rep=T)))
colnames(df)[2:(k+1)] <- words
robs <- sort(cor(as.matrix(df))[-1,1])

library(lattice)
my.cols <- colorRampPalette(c("red", "blue"))
res <- data.frame(robs=robs, x=seq(1,20), y=rep(1,20))

trellis.par.set(clip=list(panel="off"), axis.line=list(col="transparent"))
levelplot(robs~y*x, data=res, col.regions=my.cols,
```

```

colorkey=F, xlab="", ylab="", scales=list(draw=F), in a matrix or data.frame, this is easily done as
panel=function(...) {
  panel.levelplot(...)
  panel.text(x=rep(1, k), y=seq(1, k), lab=rownames(sesnorm(100))
})
apply(X, 2, function(x) lm(y ~ x))

```



The above picture was saved as PDF, setting the margins to 1, and cropped with `pdfcrop` from my TeXLive distribution.

```

pdf("1.pdf")
op <- par(mar=c(1,1,1,1))
(...)
par(op)
dev.off()

```

I guess it would not be too difficult to make a similar looking chart with `barchart()` from `lattice`, or `ggfluctuation()` or any other `qplot()` from `ggplot2`.

235 Simple and multiple logistic regression

If I understand you correctly, you want to fit two successive simple logistic regression model. I don't know if there's a specific instruction in SPSS that allows to switch the covariate of interest or cycle through them, but I guess you can run the two models in succession. In R, if your data are organized

About your second question, models like this are generally estimated using listwise deletion: any individuals having one or more missing observations on the covariates are deleted before estimating model parameters. Again, in R:

```

X[2,2] <- NA
summary(lm(y ~ X))

```

shows that one observation has been deleted, yielding 96 DF (instead of 97).

236 Graph theory — analysis and visualization

There are various packages for representing directed and undirected graphs, incidence/adjacency matrix, etc. in addition to `graph`; look for example at the `gR` Task view.

For visualization and basic computation, I think the `igraph` package is the reliable one, in addition to `Rgraphviz` (on BioC as pointed out by @Rob). Be aware that for the latter to be working properly, `graphviz` must be installed too. The `igraph` package has nice algorithms for creating good layouts, much like `graphviz`.

Here is an example of use, starting from a fake adjacency matrix:

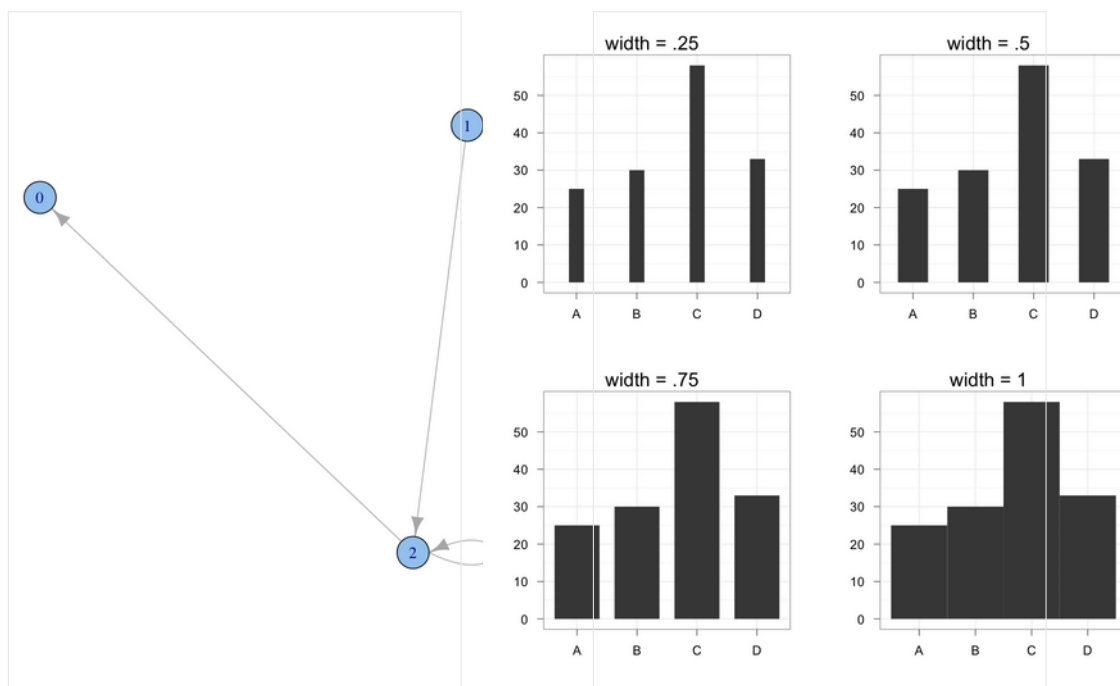
```

adj.mat <- matrix(sample(c(0,1), 9, replace=TRUE), nr=3)
g <- graph.adjacency(adj.mat)
plot(g)

```

237 Proc Calis (or TCalis) and p-values

@mpiktas is right and knowing the value of the test statistic (t or z) allows you to know which parameter estimate is significant at the desired α level. In practice, the t -statistic is equivalent to a z -score for large samples (which is often the case in SEM), and the significance thresholds are 1.96 and 2.58 for the .05 and .01 α levels. Most of the time, p -values are interesting when comparing models; as shown



in this nice tutorial on [Structural equation modeling](#) using SAS, by Y H Chan, giving t - or z -statistic with associated critical values at 5% should be enough, IMO.

238 How to increase the space between the bars in a bar plot in ggplot2?

You can always play with the `width` parameter, as shown below:

```
df <- data.frame(x=factor(LETTERS[1:4]), y=sample(1:100, 4))
library(ggplot2)
ggplot(data=df, aes(x=x, y=y, width=.5)) +
  geom_bar(stat="identity", position="identity") +
  opts(title="width = .5") + labs(x="", y="") +
  theme_bw()
```

Compare with the following other settings for `width`:

So far, so good. Now, suppose we have two factors. In case you would like to play with evenly spaced juxtaposed bars (like when using `space` together with `beside=TRUE` in `barplot()`), it's not so easy using `geom_bar(position="dodge")`: you can change bar width, but not add space in between adjacent bars (and I didn't find a convenient solution on Google). I ended up with something like that:

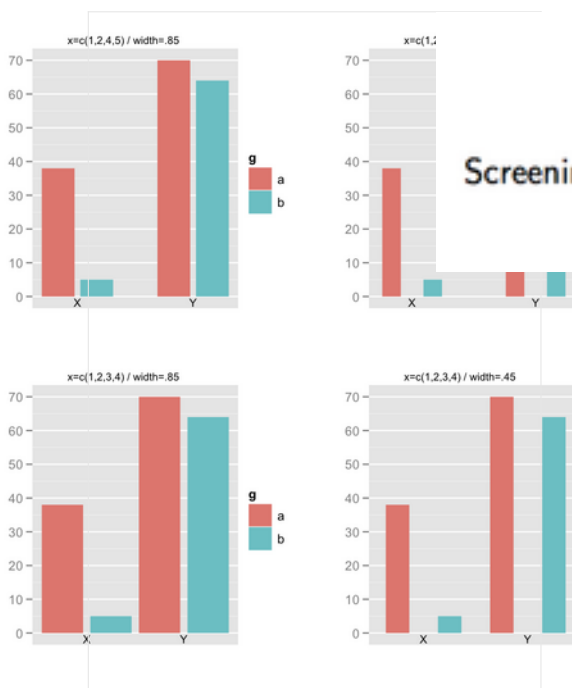
```
df <- data.frame(g=gl(2, 1, labels=letters[1:2]), y=sample(1:100, 4))
x.seq <- c(1,2,4,5)
ggplot(data=transform(df, x=x.seq), aes(x=x, y=y, width=.85)) +
  +
  geom_bar(stat="identity", aes(fill=g)) + labs(x="", y="") +
  +
  scale_x_discrete(breaks = NA) +
  geom_text(aes(x=c(sum(x.seq[1:2])/2, sum(x.seq[3:4])/2),
    y=0,
    label=c("X","Y")), vjust=1.2, size=8)
```

The vector used for the x -axis is “injected” in the data.frame, so that so you change the outer spacing if you want, while `width` allows to control for inner spacing. Labels for the x -axis might be enhanced by using `scale_x_discrete()`.

239 Analysing questionnaire data

In the spirit of an [earlier response](#), you might be interested in David A Kenny's webpage on [dyadic analysis](#), and models for matched pairs (See Agresti, [Categorical Data Analysis](#), Chapter 10, or this nice [handout](#), by C J Anderson). There's also a lot of literature on sib-pair study design, in genetics, epidemiology, and psychometrics.

As you may know, studies on sibling rivalry also



		True diagnosis		Total
		Positive	Negative	
Positive	Screening test	<i>a</i>	<i>b</i>	<i>a + b</i>
Negative		<i>c</i>	<i>d</i>	<i>c + d</i>
Total		<i>a + c</i>	<i>b + d</i>	<i>N</i>

and predictive power:

- *Sensitivity* (se), $a/(a + c)$, i.e. the probability of the screen providing a positive result given that disease is present;
- *Specificity* (sp), $d/(b + d)$, i.e. the probability of the screen providing a negative result given that disease is absent;
- *Positive predictive value* (PPV), $a/(a+b)$, i.e. the probability of patients with positive test results who are correctly diagnosed (as positive);
- *Negative predictive value* (NPV), $d/(c+d)$, i.e. the probability of patients with negative test results who are correctly diagnosed (as negative).

suggest that parents' attitude might play a role, but also that generally sibling relationships in early adulthood might be characterized by independent dimensions (warmth, conflict, and rivalry, according to Stocker et al., 1997). So it may be interesting for you to look at what has been done in psychometrics, especially whether your items share some similarity with previous studies or not. The very first hit on Google with [siblings rivalry scale statistical analysis](#) was a study on [The Effects of Working Mothers on Sibling Rivalry](#) which offers some clues on how to handle such data (although I still think that model for matched pairs are better than the χ^2 -based approach used in this study).

References

Stocker, CM, Lanthier, RP, Furman, W (1997). [Sibling Relationships in Early Adulthood](#). *Journal of Family Psychology*, 11(2), 210-221.

240 Statistical test for Positive and Negative Predictive Value

Assuming a cross-classification like the one shown below (here, for a screening instrument)

we can define four measures of screening accuracy

Each four measures are simple proportions computed from the observed data. A suitable statistical test would thus be a **binomial (exact) test**, which should be available in most statistical packages, or many online calculators. The tested hypothesis is whether the observed proportions significantly differ from 0.5 or not. I found, however, more interesting to provide confidence intervals rather than a single significance test, since it gives an information about the precision of measurement. Anyway, for reproducing the results you shown, you need to know the total margins of your two-way table (you only gave the PPV and NPV as %).

As an example, suppose that we observe the following data (the CAGE questionnaire is a screening questionnaire for alcohol):

then in R the PPV would be computed as follows:

```
> binom.test(99, 142)

Exact binomial test

data: 99 and 142
number of successes = 99, number of trials = 142, p-value
```


		Alcohol abuse	
		Positive	Negative
CAGE	Positive	99	43
	Negative	5	97
		104	140

```
= 2.958e-06
alternative hypothesis: true probability of success is not
equal to 0.5
95 percent confidence interval:
 0.6145213 0.7714116
sample estimates:
probability of success
      0.6971831
```

If you are using SAS, then you can look at the Usage Note 24170: [How can I estimate sensitivity, specificity, positive and negative predictive values, false positive and negative probabilities, and the likelihood ratios?](#).

To compute confidence intervals, the gaussian approximation, $p \pm 1.96 \times \sqrt{p(1-p)/n}$ (1.96 being the quantile of the standard normal distribution at $p = 0.975$ or $1 - \alpha/2$ with $\alpha = 5\%$), is used in practice, especially when the proportions are quite small or large (which is often the case here).

For further reference, you can look at

Newcombe, RG. [Two-Sided Confidence Intervals for the Single Proportion: Comparison of Seven Methods](#). *Statistics in Medicine*, 17, 857-872 (1998).

241 Resources for learning to use (/create) dynamic (/interactive) statistical visualization

Apart from [Protovis](#) (HTML+JS) or [Mayavi](#) (Python), I would recommend [Processing](#) which is

an open source programming language and environment for people who want to create images, animations, and interactions. Initially developed to serve as a software sketchbook and to teach fundamentals of computer programming within a visual context.

There are a lot of open-source scripts on <http://www.openprocessing.org/>, and a lot of [related books](#) that deal with Processing

but also data visualization.

I know there is a project to provide an R interface, [rprocessing](#), but I don't know how it goes. There's also an interface with clojure/incanter (see [fig. 2 Creating Processing Visualizations with Clojure and Incanter](#)).

There are many online resources, among which Stanford class notes, e.g. [CS448B](#), or [7 Classic Foundational Vis Papers You Might not Want to Publicly Confess you Don't Know](#).

242 How to present the gain in explained variance thanks to the correlation of Y and X?

Here are some suggestions (about your plot, not about how I would illustrate correlation/regression analysis):

- The two univariate plots you show in the right and left margins may be simplified with a call to [rug\(\)](#);
- I find more informative to show a density plot of X and Y or a boxplot, at risk of being evocative of the idea of a bi-normality assumption which makes no sense in this context;
- In addition to the regression line, it is worth showing a non-parametric estimate of the trend, like a loess (this is good practice and highly informative about possible local non linearities);
- Points might be highlighted (with varying color or size) according to Leverage effect or Cook distances, i.e. any of those measures that show how influential individual values are on the estimated regression line. I'll second @DWin's comment and I think it is better to highlight how individual points "degrade" goodness-of-fit or induce some kind of departure from the linearity assumption.

Of note, this graph assumes X and Y are non-paired data, otherwise I would stick to a [Bland-Altman plot](#) ($(X - Y)$ against $(X + Y)/2$), in addition to scatterplot.

243 How to fix the threshold for statistical validity of p-values produced by ANOVAs?

Hey, but it seems you already looked at the results!

Usually, the risk of falsely rejecting the null (Type I error, or α) should be decided before starting the analysis. Power might also be fixed to a given value (e.g., 0.80). At least, this is the “Neyman-Pearson” approach. For example, you might consider a risk of 5% ($\alpha = 0.05$) for all your hypotheses, and if the tests are not independent you should consider correcting for multiple comparisons, using any single-step or step-down methods you like.

When reporting your results, you should indicate the Type I (and II, if applicable) error you considered (before seeing the results!), corrected or not for multiple comparisons, and give your p-values as $p < .001$ or $p = .0047$ for example.

Finally, I would say that your tests allow you to reject a given null hypothesis not to prove Hypothesis A or B. Moreover, what you describe as 0.001 being a somewhat stronger indication of an interesting deviation from the null than 0.05 is more in light with the Fisher approach to [statistical hypothesis testing](#).

244 What software is used for maps of the US (or other arbitrary areas)

In addition to the package that @robin pointed too, you should look at the [Spatial](#) Task View on CRAN. What you are describing is known as a [choropleth map](#), as illustrated here: [Choropleth Maps of Presidential Voting](#), or [U.S. Unemployment Data: Animated Choropleth Maps](#).

In R, they can be handled using

- the base graphics routines (with [maps](#)),
- the lattice way (see [mapplot\(\)](#) in [latticeExtra](#)),
- the ggplot2 way (see e.g., [Choropleth Challenge Results](#) for example code).

Other softwares that allow to deal with geographical maps include [Mondrian](#), [Quantum GIS](#) (and I guess many other GIS programs), but data format may

vary from one software to the other.

245 How to measure/rank “variable importance” when using CART? (specifically using {rpart} from R)

Variable importance might generally be computed based on the corresponding reduction of predictive accuracy when the predictor of interest is removed (with a permutation technique, like in Random Forest) or some measure of decrease of node impurity, but see (1) for an overview of available methods. An obvious alternative to CART is RF of course ([randomForest](#), but see also [party](#)). With RF, the Gini importance index is defined as the averaged Gini decrease in node impurities over all trees in the forest (it follows from the fact that the Gini impurity index for a given parent node is larger than the value of that measure for its two daughter nodes, see e.g. (2)).

I know that Carolin Strobl and coll. have contributed a lot of simulation and experimental studies on (conditional) variable importance in RFs and CARTs (e.g., (3-4)), but there are many other ones, or her thesis, [Statistical Issues in Machine Learning – Towards Reliable Split Selection and Variable Importance Measures](#).

To my knowledge, the [caret](#) package (5) only considers a loss function for the regression case (i.e., mean squared error). Maybe it will be added in the near future (anyway, an example with a classification case by k-NN is available in the on-line help for [dotPlot](#)).

However, Noel M O’Boyle seems to have some R code for [Variable importance in CART](#).

References

1. Sandri and Zuccolotto. [A bias correction algorithm for the Gini variable importance measure in classification trees](#). 2008
2. Izenman. *Modern Multivariate Statistical Techniques*. Springer 2008
3. Strobl, Hothorn, and Zeileis. [Party on!](#). *R Journal* 2009 1/2
4. Strobl, Boulesteix, Kneib, Augustin, and Zeileis.

Conditional variable importance for random forests.
BMC Bioinformatics 2008, 9:307

5. Kuhn. *Building Predictive Models in R Using the caret Package*. *JSS* 2008 28(5)

246 likelihood ratio test in R

Basically, yes, provided you use the correct difference in log-likelihood:

```
> library(epicalc)
> model0 <- glm(case ~ induced + spontaneous, family=binomial, data=infert)
> model1 <- glm(case ~ induced, family=binomial, data=infert)
> lrtest (model0, model1)
Likelihood ratio test for MLE method
Chi-squared 1 d.f. = 36.48675 , P value = 0
> model1$deviance-model0$deviance
[1] 36.48675
```

and *not the deviance for the null model* which is the same in both cases. The number of df is the number of parameters that differ between the two nested models, here $df=1$. BTW, you can look at the source code for `lrtest()` by just typing

```
> lrtest
```

at the R prompt.

247 What is Deviance? (specifically in CART/rpart)

Deviance and GLM

Formally, one can view deviance as a sort of distance between two probabilistic models; in an GLM context, it amounts to two times the ratio of log-likelihood between two nested models ℓ_1/ℓ_0 where ℓ_0 is the “smaller” model, that is a linear restriction on model parameters (cf. the **Neyman–Pearson lemma**), as @suncoolsu said. As such, it can be used to perform *model comparison*. It can also be seen it as a generalization of the RSS used in OLS estimation (ANOVA, regression), for it provides a measure of *goodness-of-fit* of the model being evaluated when compared to the null model (intercept only). It works with LM too:

```
> x <- rnorm(100)
> y <- 0.8*x+rnorm(100)
```

```
> lm.res <- lm(y ~ x)
```

The residuals SS (RSS) is computed as $\hat{\varepsilon}^t \hat{\varepsilon}$, which is readily obtained as:

```
> t(residuals(lm.res))%*%residuals(lm.res)
[1,]
[1,] 98.66754
```

or from the (unadjusted) R^2

```
> summary(lm.res)
```

```
Call:
lm(formula = y ~ x)

(....)

Residual standard error: 1.003 on 98 degrees of freedom
Multiple R-squared: 0.4234, Adjusted R-squared: 0.4175
F-statistic: 71.97 on 1 and 98 DF, p-value: 2.334e-13
```

since $R^2 = 1 - \text{RSS}/\text{TSS}$ where TSS is the total variance. Note that it is directly available in an ANOVA table, like

```
> summary.aov(lm.res)
          Df Sum Sq Mean Sq F value    Pr(>F)
x           1  72.459   72.459   71.969 2.334e-13 ***
Residuals   98  98.668    1.007
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Now, look at the deviance:

```
> deviance(lm.res)
[1] 98.66754
```

In fact, for linear models the deviance equals the RSS (you may recall that OLS and ML estimates coincide in such a case).

Deviance and CART

We can see CART as a way to allocate already n labeled individuals into arbitrary classes (in a classification context). Trees can be viewed as providing a probability model for individuals class membership. So, at each node i , we have a probability distribution p_{ik} over the classes. What is important here is that the leaves of the tree give us a random sample n_{ik} from a multinomial distribution specified by p_{ik} . We can thus define the deviance of a tree, D , as the sum over all leaves of

$$D_i = -2 \sum_k n_{ik} \log(p_{ik}),$$

following Venables and Ripley's notations ([MASS](#), Springer 2002, 4th ed.). If you have access to this essential reference for R users (IMHO), you can check by yourself how such an approach is used for splitting nodes and fitting a tree to observed data (p. 255 ff.); basically, the idea is to minimize, by pruning the tree, $D + \alpha \#(T)$ where $\#(T)$ is the number of nodes in the tree T . Here we recognize the *cost-complexity trade-off*. Here, D is equivalent to the concept of node impurity (i.e., the heterogeneity of the distribution at a given node) which are based on a measure of entropy or information gain, or the well-known Gini index, defined as $1 - \sum_k p_{ik}^2$ (the unknown proportions are estimated from node proportions).

With a regression tree, the idea is quite similar, and we can conceptualize the deviance as sum of squares defined for individuals j by

$$D_i = \sum_j (y_j - \mu_i)^2,$$

summed over all leaves. Here, the probability model that is considered within each leaf is a gaussian $\mathcal{N}(\mu_i, \sigma^2)$. Quoting Venables and Ripley (p. 256), “ D is the usual scaled deviance for a gaussian GLM. However, the distribution at internal nodes of the tree is then a mixture of normal distributions, and so D_i is only appropriate at the leaves. The tree-construction process has to be seen as a *hierarchical refinement of probability models, very similar to forward variable selection in regression*.” Section 9.2 provides further detailed information about [rpart](#) implementation, but you can already look at the [residuals\(\)](#) function for [rpart](#) object, where “deviance residuals” are computed as the square root of minus twice the logarithm of the fitted model.

An introduction to recursive partitioning using the [rpart](#) routines, by Atkinson and Therneau, is also a good start. For more general review (including bagging), I would recommend

- Moissen, G.G. (2008). [Classification and Regression Trees](#). *Ecological Informatics*, pp. 582-588.
- Sutton, C.D. (2005). [Classification and Regression Trees, Bagging, and Boosting](#), in *Handbook of Statistics*, Vol. 24, pp. 303-329, Elsevier.

248 Correcting for multiple comparisons when running a bivariate correlation in SPSS

This first part of my response won't address your two questions directly since what I am suggesting departs from your correlational approach. If I understand you correctly, you have two blocks of variables, and they play an asymmetrical role in the sense that one of them is composed of response variables (performance on four cognitive tests) whereas the other includes explanatory variables (measures of blood flow at several locations). So, a nice way to answer your question of interest would be to look at [PLS regression](#). As detailed in an earlier response of mine, [Regression with multiple dependent variables?](#), the correlation between factor scores on the first dimension will reflect the overall link between these two blocks, and a closer look at the weighted combination of variables in each block (i.e., loadings) would help interpreting the contribution of each variable of the X block in predicting the Y block. The [SPSS implementation](#) is detailed on Dave Garson's website. This prevents from using any correction for multiple comparisons.

Back to your specific questions, yes the Bonferroni correction is known to be conservative and step-down methods are to be preferred (instead of correcting the p-values or the test statistic in one shot for all the tests, we adapt the threshold depending on the previous HT outcomes, in a sequential manner). Look into SPSS documentation (or [Pairwise Comparisons in SAS and SPSS](#)) to find a suitable one, e.g. Bonferroni-Holm.

249 How can one plot continuous by continuous interactions in ggplot2?

Here's my version with your simulated data set:

```
x1 <- rnorm(100,2,10)
x2 <- rnorm(100,2,10)
y <- x1+x2+x1*x2+rnorm(100,1,2)
dat <- data.frame(y=y,x1=x1,x2=x2)
res <- lm(y~x1*x2,data=dat)
z1 <- z2 <- seq(-1,1)
newdf <- expand.grid(x1=z1,x2=z2)

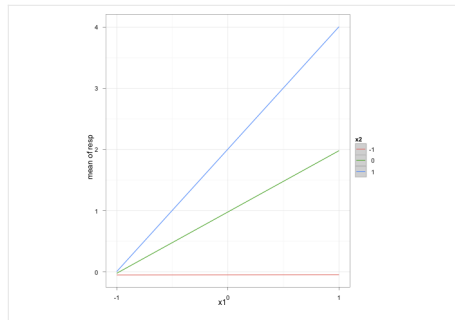
library(ggplot2)
p <- ggplot(data=transform(newdf, yp=predict(res, newdf)),
```

```

aes(y=yp, x=x1, color=factor(x2))) + stat_smooth(method="lm") +
p + scale_colour_discrete(name="x2") +
  labs(x="x1", y="mean of resp") +
  scale_x_continuous(breaks=seq(-1,1)) + theme_bw()

```

I let you manage the details about x/y-axis labels and legend positioning.



250 Assessing conditional independence of genes in Trans-eQTL cluster

I would recommend looking at the `snpMatrix` R package. Within the `snp.lhs.tests()` function, height will be the phenotype (or outcome), your SNP data will be stored in a vector (`snp.data`), and your gene expression levels will enter the model as covariates. I never used this kind of covariates (in the GWAS I am familiar with we adjust for population stratification and subject-specific covariates), so I can't give an example of use. Just give it a try to see if it suits your needs.

Otherwise, a larger modeling framework is available in the `GGtools` package, from the `Bioconductor` project too. It was developed by Vince J. Carey for dealing specifically with SNP and gene expression data, in GWAS or eQTL studies. There are numerous examples of use on the web (look up for tutorial by VJ Carey with `GGtools`). However, it's mainly to model gene expression as a function of observed genotypes (chromosome- or genome-wide).

Caution. Please note that `GGtools` package has a lot of dependencies, in particular `GGBase` (which provides specific environments for holding genetic data) and `snpMatrix` (which provides an efficient storage of SNP data). You will also need to install the base packages from the `Bioconductor` repository (this is not part of CRAN); detailed instructions are available [here](#).

251 Plotting a heatmap given a dendrogram and a distance matrix in R

I don't know a specific function for that. The ones I used generally take raw data or a distance matrix. However, it would not be very difficult to hack already existing code, without knowing more than basic R. Look at the source code for the `cim()` function in the `mixOmics` package for example (I choose this one because source code is very easy to read; you will find other functions on the `Bioconductor` project). The interesting parts of the code are l. 92-113, where they assign the result of HC to `ddc`, and around l. 193-246 where they devised the plotting regions (you should input the values of your distance matrix in place of `mat` when they call `image()`). HTH

252 Bayesian additive regression trees (BART) for classification analysis of gene expression data

I would suggest looking at the `BayesTree` package, from CRAN. I have no experience with it, so I cannot say if there are better option from there. Try looking at the `Machine Learning` Task View, or directly through www.rseek.org.

I don't know anything approaching in `Bioconductor`, but if the above package suits your needs I guess you won't have any problem with gene expression data. I know the `CMA` package offer a full-featured pipeline for supervised classification (see e.g., `CMA - A comprehensive Bioconductor package for supervised classification with high dimensional data`, by Slawski et al.). Maybe you can plug BART method in addition to the available methods?

253 Validating an existing questionnaire into another language

I don't know what your questionnaire aims to assess. In Health-related Quality-of-Life studies, for example, there are a certain number of recommendations for translation issues that were discussed in the following papers (among others):

1. Marquis et al., Translating and evaluating ques-

tionnaires: Cultural issues for international research, in Fayers & Hays (eds.), *Assessing Quality of Life in Clinical Trials (2nd ed.)*, Oxford, pp. 77-93 (2005).

2. Hui and Triandis, Measurement in cross-cultural psychology: A review and comparison of strategies. *Journal of Cross-Cultural Psychology*, 16(2), 131-152 (1985).
3. Mathias et al., Rapid translation of quality of life measures for international clinical trials: Avoiding errors in the minimalist approach. *Quality of Life Research*, 3, 403-412 (1994).

Translation can be done simultaneously in several languages, as was the case for the WHOQOL questionnaire, or in a primary language (e.g., English) for the SF-36 followed by translation in other target languages. There were many papers related to translation issues in either case that you will probably find on [Pubmed](#). Various procedures have been developed to ensure consistent translation, but forward/backward translation is most commonly found in the above studies.

In any case, most common issues when translating items (as single entities, and as a whole, that is at the level of the questionnaire) have to do with the equivalence of the hypothetical concept(s) that is/are supposed to be covered.

Otherwise, the very first things I would look at would be:

- Am I measuring the same concepts? – this is a question related to *validity*;
- Are the scores delivered in the foreign language reliable enough? – this has to do with *scores reliability*;
- Are the items behaving as expected for everybody, i.e. without differential effects depending on country or native language? – this is merely related to *differential item functioning* (DIF), which is said to occur when the probability of endorsing a particular item differs according to a subject-specific covariate (e.g., age, gender, country), when holding subject trait constant.

Some of the common techniques used to assess those properties were discussed in an earlier related question, [Validating questionnaires](#).

About DIF specifically, here are some examples of

subtle variation across subject-specific characteristics:

- In psychiatric studies, “I feel sad” / “Able to enjoy life” have been shown to highlight gender-related DIF (Crane et al., 2007).
- In personality assessment, there are well-known age and gender-effect on the NEO-PI questionnaire (reviewed in Kulas et al., 2008).
- In Health-related Quality-of-Life, items like “Did you worry?” / “Did you feel depressed?” have been shown to exhibit country-related DIF effect (Petersen et al. 2003).

A good starting point is this review paper by Jeanne Teresi in 2004: [Differential Item Functioning and Health Assessment](#).

254 Is there an anova procedure that doesn't assume equal variance?

There is a function named `oneway.test()` in the base `stats` package, which implements Welch correction for a one-way ANOVA. Its use is similar to the standard `t.test()` function. It is also referred to as O'Brien transformation (Biometrics 40 (1984), 1079–1087) and might be applied with two or more independent samples (here is my [implementation](#), check if it is not too buggy!).

255 How to graphically compare predicted and actual values from multivariate regression in R?

In addition to @mpiktas's comment, you can also have a look at the `rms` package from Frank Harrell. The advantage is that it handles both LM and GLM for model fitting and prediction; see for example the `plot.Predict()` function. If you're planning to do serious job in regression modeling, this package and its companion `Hmisc` are really good.

256 What are the text-mining packages for R and are there other open source text-mining programs?

Here are two further integrated projects:

- Python **Natural Language Toolkit** (easy installation, good documentation)
- Java [MALLET][2] (no experience with it, but looks promising; included in the link given by @Nick)

Both are open-source software.

[2]: <http://mallet.cs.umass.edu/>

257 Visualizing the intersections of many sets

This won't compete with @Shane's answer because circular displays are really well suited for displaying complex relationships with high-dimensional datasets.

For Venn diagrams, I've been using the **venneuler** R package. It has a simple yet intuitive interface and produce nifty diagrams with transparency, compared to the basic **venn()** function described in the *Journal of Statistical Software*. It does not handle more than 3 categories, though. Another project is **eVenn** and it deals with $K = 4$ sets.

More recently, I came across a new package that deal with higher-order relation sets, and probably allow to reproduce some of the Venn diagrams shown on Wikipedia or on this webpage, **What is a Venn Diagram?**, but it is also limited to $K = 4$ sets. It is called **VennDiagram**, but see the reference paper: **VennDiagram: a package for the generation of highly-customizable Venn and Euler diagrams in R** (Chen and Boutros, *BMC Bioinformatics* 2011, 12:35).

For further reference, you might be interested in

Kestler et al., **Generalized Venn diagrams: a new method of visualizing complex genetic set relations**, *Bioinformatics*, 21(8), 1592-1595 (2004).

Venn diagrams have their limitations, though. In this respect, I like the approach taken by Robert Kosara in **Sightings: A Vennerable Challenge**, or with **Parallel Sets** (but see also [this discussion](#) on Andrew Gelman weblog).

258 When is *interactive* data visualization useful to use?

In addition to linking quantitative or qualitative data to spatial patterns, as illustrated by @whuber, I would like to mention the use of EDA, with brushing and the various of linking plots together, for *longitudinal* and *high-dimensional* data analysis.

Both are discussed in the excellent book, **Interactive and Dynamic Graphics for Data Analysis With R and GGobi**, by Dianne Cook and Deborah F. Swayne (Springer UseR!, 2007), that you surely know. The authors have a nice discussion on EDA in Chapter 1, justifying the need for EDA to “force the unexpected upon us”, quoting John Tukey (p. 13): The use of interactive and dynamic displays is neither **data snooping**, nor preliminary data inspection (e.g., purely graphical summaries of the data), but it is merely seen as an interactive investigation of the data which might precede or complement pure hypothesis-based statistical modeling.

Using GGobi together with its R interface (**rggobi**) also solves the problem of how to generate static graphics for intermediate report or final publication, even with **Projection Pursuit** (pp. 26-34), thanks to the **DescribeDisplay** or **ggplot2** packages.

In the same line, **Michael Friendly** has long advocated the use of data visualization in Categorical Data Analysis, which has been largely exemplified in the **vcd** package, but also in the more recent **vcdExtra** package (including dynamic viz. through the **rgl** package), which acts as a glue between the **vcd** and **gnm** packages for extending log-linear models. He recently gave a nice summary of that work during the 6th CARME conference, **Advances in Visualizing Categorical Data Using the vcd, gnm and vcdExtra Packages in R**.

Hence, EDA can also be thought of as providing a visual explanation of data (in the sense that it may account for unexpected patterns in the observed data), prior to a purely statistical modeling approach, or in parallel to it. That is, EDA not only provides useful ways for studying the internal structure of the data at hand, but it may also help to

refine and/or summarize statistical models applied on it. It is in essence what **biplots** allow to do, for example. Although they are not multidimensional analysis techniques *per se*, they are tools for visualizing results from multidimensional analysis (by giving an *approximation* of the relationships when considering all individuals together, or all variables together, or both). Factor scores can be used in subsequent modeling in place of the original metric to either reduce the dimensionality or to provide intermediate levels of representation.

Sidenote

At risk of being old-fashioned, I'm still using **xlispstat** (Luke Tierney) from time to time. It has simple yet effective functionalities for interactive displays, currently not available in base R graphics. I'm not aware of similar capabilities in Clojure+Incanter (+Processing).

259 How to use variables derived from factor analysis as predictors in logistic regression?

If I understand you correctly, you are using FA to extract two subscales from your 11-item questionnaire. They are supposed to reflect some specific dimensions of self-efficacy (for example, self-regulatory vs. self-assertive efficacy).

Then, you are free to use *individual* mean (or sum) scores computed on the two subscales as predictors in a regression model. In other words, instead of considering 11 item scores, you are now working with 2 subscores, computed as described above for each individual. The only assumption that is made is that those scores reflect one's location on an "hypothetical construct" or latent variable, defined as a continuous scale.

As @JMS said, there are other issues that you might further clarify, especially which kind of FA was done. A subtle issue is that measurement error will not be accounted for by a standard regression approach. An alternative is to use **Structural Equation Models** or any latent variables model (e.g. those coming from the **IRT** literature), but here the regression approach should provide a good approximation. The analysis of ordinal variables (Likert-type item) has been discussed elsewhere on this site.

However, in current practice, your approach is what

is commonly found when validating a questionnaire or constructing scoring rules: We use weighted or unweighted combination of item scores (hence, they are treated as numeric variables) to report individual location on the latent trait(s) under consideration.

260 Calculating predicted values from categorical predictors in logistic regression

My initial thought would have been to display the probability of acceptance as a function of relative GPA for each of your four schools, using some kind of **trellis displays**. In this case, faceting should do the job well as the number of schools is not so large. This is very easy to do with **lattice** (`y ~ gpa | school`) or **ggplot2** (`facet_grid(. ~ school)`). In fact, you can choose the conditioning variable you want: this can be school, but also situation at undergrad institution. In the latter case, you'll have 4 curves for each plot, and three three plot of `Prob(admitting) ~ GPA`.

Now, if you are looking for effective displays of effects in GLM, I would recommend the **effects** package, from John Fox. Currently, it works with binomial and multinomial link, and ordinal logistic model. Marginalizing over other covariates is handled internally, so you don't have to bother with that. There are a lot of illustrations in the on-line help, see `help(effect)`. But, for a more thorough overview of effects displays in GLM, please refer to

1. Fox (2003). **Effect Displays in R for Generalised Linear Models**. JSS 8(15).
2. Fox and Andersen (2004). **Effect displays for multinomial and proportional-odds logit models**. ASA Methodology Conference – Here is the corresponding **JSS paper**

261 Clinical reasoning

I like @nico's response because it makes clear that statistical and pragmatic thinking shall come hand in hand; this also has the merit to bring out issues like statistical vs. clinical significance. But about your specific question, I would say this is clearly detailed in the two sections that directly follow your quote (p. 10).

Rereading Piantadosi's textbook, it appears that the author means that **clinical thinking** applies to the situation where a physician has to interpret the results of RCTs or other studies in order to decide of the best treatment to apply to a *new patient*. This has to do with the extent to which (population-based) conclusions drawn from previous RCT might generalize to new, unobserved, samples. In a certain sense, such decision or judgment call for some form of clinical experience, which is not necessarily of the resort of a consistent statistical framework. Then, the author said that "the solution offered by **statistical reasoning** is to control the signal-to-noise ratio by design." In other words, this is a way to reduce uncertainty, and "the chance of drawing incorrect conclusions from either good or bad data." In sum, both lines of reasoning are required in order to draw valid conclusions from previous (and 'localized') studies, and choose the right treatment to administer to a new individual, given his history, his current medication, etc. – treatment efficacy follows from a good balance between statistical facts and clinical experience.

I like to think of a statistician as someone who is able to mark off the extent to which we can draw firm inferences from the observed data, whereas the clinician is the one that will have a more profound insight onto the implications or consequences of the results at the individual or population level.

262 Spam filtering using naive Bayesian classifiers with the e1071/klaR package on R

The `NaiveBayes()` function in the `klaR` package obeys the classical `formula` R interface whereby you express your outcome as a function of its predictors, e.g. `spam ~ x1+x2+x3`. If your data are stored in a `data.frame`, you can input all predictors in the rhs of the formula using dot notation: `spam ~ ., data=df` means "`spam` as a function of all other variables present in the `data.frame` called `df`."

Here is a toy example, using the `spam` dataset discussed in the *Elements of Statistical Learning* (Hastie et al., Springer 2009, 2nd ed.), available on-line. This really is to get you started with the use of the R function, not the methodological aspects for using NB classifier.

```
data(spam, package="ElemStatLearn")
library(klaR)
```

```
# set up a test sample
train.ind <- sample(1:nrow(spam), ceiling(nrow(spam)*2/3),
replace=FALSE)

# apply NB classifier
nb.res <- NaiveBayes(spam ~ ., data=spam[train.ind,])

# show the results
opar <- par(mfrow=c(2,4))
plot(nb.res)
par(opar)

# predict on holdout units
nb.pred <- predict(nb.res, spam[-train.ind,])

# raw accuracy
confusion.mat <- table(nb.pred$class, spam[-train.ind,"spam"])
sum(diag(confusion.mat))/sum(confusion.mat)
```

A recommended add-on package for such ML task is the `caret` package. It offers a lot of useful tools for preprocessing data, handling training/test samples, running different classifiers on the same data, and summarizing the results. It is available from CRAN and has a lot of vignettes that describe common tasks.

263 Getting started with biclustering

I never used it directly, so I can only share some papers I had and general thoughts about that technique (which mainly address your questions 1 and 3).

My general understanding of biclustering mainly comes from genetic studies (2-6) where we seek to account for clusters of genes and grouping of individuals: in short, we are looking to groups samples sharing similar profile of gene expression together (this might be related to disease state, for instance) and genes that contribute to this pattern of gene profiling. A survey of the state of the art for biological "massive" datasets is available in Pardalos's slides, *Biclustering*. Note that there is an R package, `biclust`, with applications to microarray data.

In fact, my initial idea was to apply this methodology to clinical diagnosis, because it allows to put features or variables in more than one cluster, which is interesting from a semeiological perspective because symptoms that cluster together allow to define *syndrome*, but some symptoms can overlap in different diseases. A good discussion may be found

in Cramer et al., [Comorbidity: A network perspective](#) (Behavioral and Brain Sciences 2010, 33, 137-193).

A somewhat related technique is [collaborative filtering](#). A good review was made available by Su and Khoshgoftaar (*Advances in Artificial Intelligence*, 2009): [A Survey of Collaborative Filtering Techniques](#). Other references are listed at the end. Maybe analysis of [frequent itemset](#), as exemplified in the [market-basket problem](#), is also linked to it, but I never investigated this. Another example of co-clustering is when we want to simultaneously cluster words and documents, as in text mining, e.g. Dhillon (2001). [Co-clustering documents and words using bipartite spectral graph partitioning](#). *Proc. KDD*, pp. 269–274.

About some general references, here is a not very exhaustive list that I hope you may find useful:

1. Jain, A.K. (2010). [Data clustering: 50 years beyond K-means](#). *Pattern Recognition Letters*, 31, 651–666
2. Carmona-Saez et al. (2006). [Biclustering of gene expression data by non-smooth non-negative matrix factorization](#). *BMC Bioinformatics*, 7, 78.
3. Prelic et al. (2006). [A systematic comparison and evaluation of biclustering methods for gene expression data](#). *Bioinformatics*, 22(9), 1122–1129. www.tik.ee.ethz.ch/sop/bimax
4. DiMaggio et al. (2008). [Biclustering via optimal re-ordering of data matrices in systems biology: rigorous methods and comparative studies](#). *BMC Bioinformatics*, 9, 458.
5. Santamaria et al. (2008). [BicOverlapper: A tool for bicluster visualization](#). *Bioinformatics*, 24(9), 1212–1213.
6. Madeira, S.C. and Oliveira, A.L. (2004) [Bicluster algorithms for biological data analysis: a survey](#). *IEEE Trans. Comput. Biol. Bioinform.*, 1, 24–45.
7. Badea, L. (2009). [Generalized Clustergrams for Overlapping Biclusters](#). *IJCAI*
8. Symeonidis, P. (2006). [Nearest-Biclusters Collaborative Filtering](#). *WEBKDD*

264 Books with good coverage on Joint Distributions, Multivariate Statistics etc?

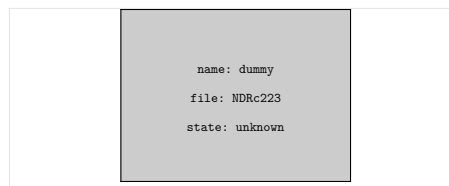
Despite @whuber's sound comment—covering all advances in MV analysis for the last 30 years is also outside the scope of e.g. the famous [Handbook of Statistics](#) series—, I would like to recommend

Izenman, [Modern Multivariate Statistical Techniques](#), Springer 2008.

Although it has pretty much the same coverage than the [Elements of Statistical Learning](#), from Hastie and coll., it has some different applications and covers extra topic, like Correspondence Analysis. There is a [short review](#) by John Maindonald in the JSS.

265 Complex regression plot in R

Does the picture below look like what you want to achieve?



Here's the **updated** R code, following your comments:

```
do.it <- function(df, type="confidence", ...) {
  require(ellipse)
  lm0 <- lm(y ~ x, data=df)
  xc <- with(df, xyTable(x, y))
  df.new <- data.frame(x=seq(min(df$x), max(df$x), 0.1))
  pred.ulb <- predict(lm0, df.new, interval=type)
  pred.lo <- predict(loess(y ~ x, data=df), df.new)
  plot(xc$x, xc$y, cex=xc$number*2/3, xlab="x", ylab="y",
  ...)
  abline(lm0, col="red")
  lines(df.new$x, pred.lo, col="green", lwd=1.5)
  lines(df.new$x, pred.ulb[, "lwr"], lty=2, col="red")
  lines(df.new$x, pred.ulb[, "upr"], lty=2, col="red")
  lines(ellipse(cor(df$x, df$y), scale=c(sd(df$x), sd(df$y))),
  centre=c(mean(df$x), mean(df$y)), lwd=1.5, col="green")
  invisible(lm0)
}

set.seed(101)
```

```
n <- 1000
x <- rnorm(n, mean=
y <- 1.5 + 0.4*x +
df <- data.frame(x=
```

```
# take a bootstrap
df <- df[sample(nro
```

```
do.it(df, pch=19, c
```

And here is the g
produced with th

```
xc <- with(df, xyTa
df2 <- cbind.data.f
df.ell <- as.data.f
```

```
library(ggplot2)
```

```
ggplot(data=df2, ae
  geom_point(aes(si
  stat_smooth(data=
+
  stat_smooth(data=
  geom_path(data=df
```

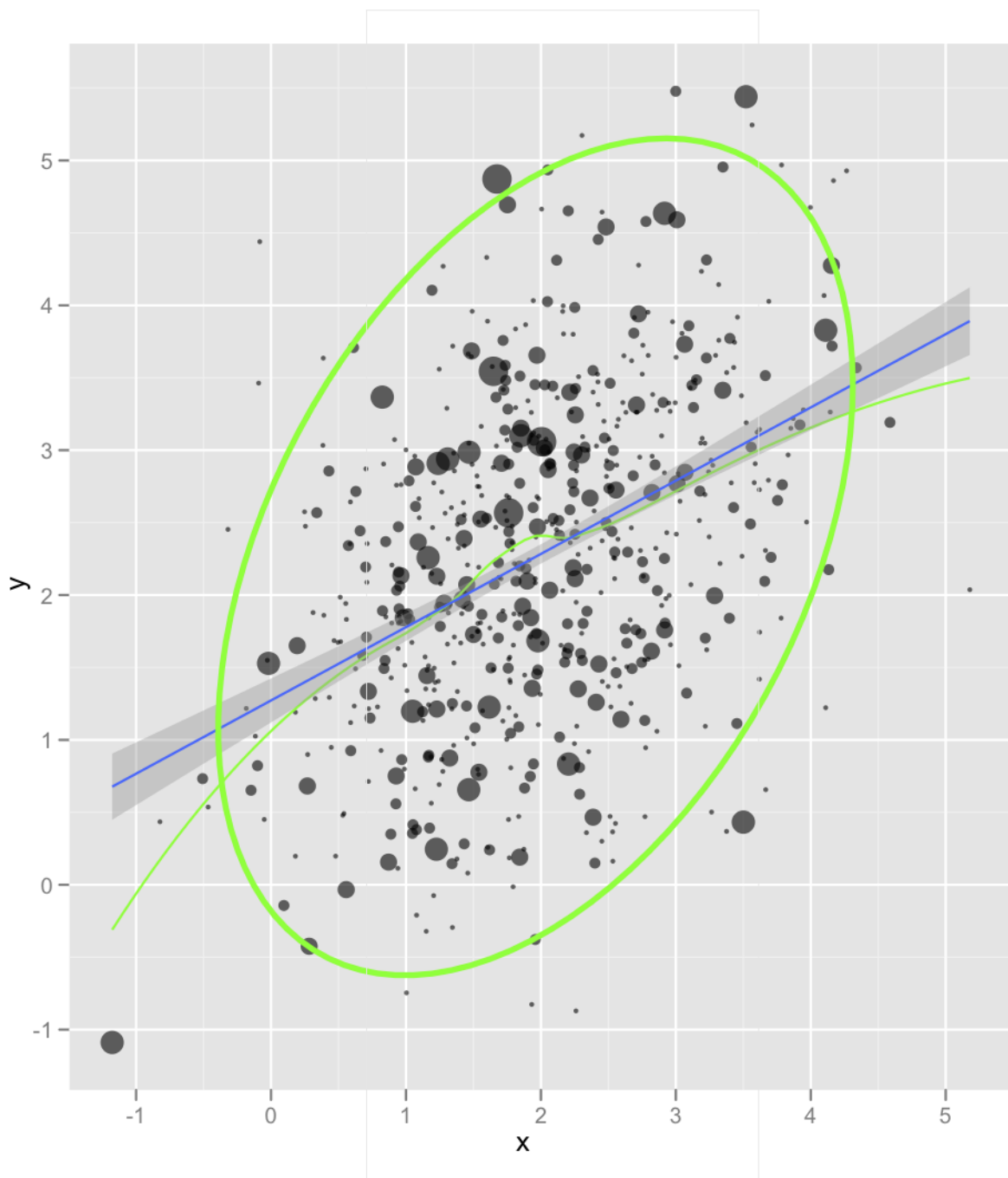
It could be custo
model fit indices,
shading effect.

266 How to with S

Although I didn
R (I usually post
rectly with `\rowc`
package), I thin
by playing with
`print.xtable()`.
nents (passed as
L^AT_EX command
added at the end

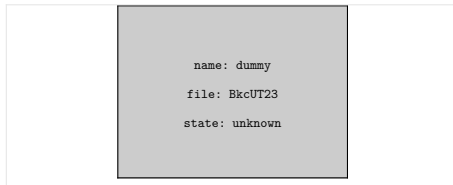
It seems to work, with the `colortbl` package. So,
something like this

```
<<result=tex>>
library(xtable)
m <- matrix(sample(1:10,10), nr=2)
print(xtable(m), add.to.row=list(list(1),"\rowcolor{gray}{.83}"))
@
```



gives me

(This is a customized Beamer template, but this should work with a standard document. With Beamer, you'll probably want to add the `table` option when loading the package.)



267 How to do a ‘beer and diapers’ correlation analysis

In addition to the links that were given in comments, here are some further pointers:

- [Association rules and frequent itemsets](#)
- [Survey on Frequent Pattern Mining](#) – look around Table 1, p. 4

About Python, I guess now you have an idea of what you should be looking for, but the [Orange](#) data mining package features a package on [Association rules](#) and [Itemsets](#) (although for the latter I cannot find any reference on the website).

Edit:

I recently came across [pysuggest](#) which is

a Top-N recommendation engine that implements a variety of recommendation algorithms. Top-N recommender systems, a personalized information filtering technology, are used to identify a set of N items that will be of interest to a certain user. In recent years, top-N recommender systems have been used in a number of different applications such to recommend products a customer will most likely buy; recommend movies, TV programs, or music a user will find enjoyable; identify web-pages that will be of interest; or even suggest alternate ways of searching for information.

268 Binary classification when many binary features are missing

Assuming data are considering missing completely at random (cf. @whuber’s comment), using an ensemble learning technique as described in the following paper might be interesting to try:

Polikar, R. et al. (2010). [Learn++.MF: A random subspace approach for the missing](#)

[feature problem](#). *Pattern Recognition*, 43(11), 3817-3832.

The general idea is to train multiple classifiers on a subset of the variables that composed your dataset (like in Random Forests), but to use only the classifiers trained with the observed features for building the classification rule. Be sure to check what the authors call the “distributed redundancy” assumption (p. 3 in the preprint linked above), that is there must some evenly spaced redundancy among your features set.

269 Labeling boxplots in R

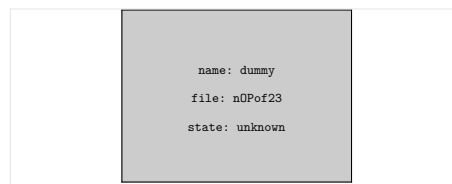
Try something like this for a standalone version:

```
bxp <- boxplot(rnorm(100), horizontal=TRUE, axes=FALSE)
mtext(c("Min", "Max"), side=3, at=bxp$stats[c(1,5)], line=-3)
```

Note that you can get some information when calling [boxplot](#), in particular the “five numbers”.

If you want it to be superimposed onto another graphic, use `add=T` but replace `mtext` by `text`; you will need to set a y value (which depend on the way you plot the other graphic).

A more complete example was given by [John Maindonald](#) (code should be on his website):











270 Plotting sparklines in R

I initially managed to produce something approaching your original picture with some quick and dirty R code (see this [gist](#)), until I discovered that the [sparkTable](#) package should do this very much better, provided you are willing to use $LaTeX$. (In the meantime, it has also been pointed out by @Bernd!)

Here is an example, from [help\(sparkEPS\)](#):

It should not be too difficult to arrange this the way you want.

	line	gdp - bar	gdp - min	gdp - max	gdp - last
			100	100	100
			123	131.6	123.5
			115.2	124.3	115.6
			32.5	45.6	45.6

be referred to an $F_{p-1, n-p}$ distribution for a critical or p -value. It applies for the simple regression model as well, and obviously bears some analogy with the classical ANOVA framework.

Sidenote. When you have more than one predictor, then you may wonder whether considering only a subset of those predictors “reduces” the quality of model fit. This corresponds to a situation where we consider *nested models*. This is exactly the same situation as the above ones, where we compare a given regression model with a *null* model (no predictors included). In order to assess the reduction in explained variance, we can compare the residual sum of squares (RSS) from both model (that is, what is left unexplained once you account for the effect of predictors present in the model). Let \mathcal{M}_0 and \mathcal{M}_1 denote the base model (with p parameters) and a model with an additional predictor ($q = p + 1$ parameters), then if $\text{RSS}_{\mathcal{M}_1} - \text{RSS}_{\mathcal{M}_0}$ is small, we would consider that the smaller model performs as good as the larger one. A good statistic to use would be the ratio of such SS, $(\text{RSS}_{\mathcal{M}_1} - \text{RSS}_{\mathcal{M}_0})/\text{RSS}_{\mathcal{M}_0}$, weighted by their degrees of freedom ($p - q$ for the numerator, and $n - p$ for the denominator). As already said, it can be shown that this quantity follows an F (or Fisher-Snedecor) distribution with $p - q$ and $n - p$ degrees of freedom. If the observed F is larger than the corresponding F quantile at a given α (typically, $\alpha = 0.05$), then we would conclude that the larger model makes a “better job”. (This by no means implies that the model is correct, from a practical point of view!)

271 Logic behind the ANOVA F -test in simple linear regression

In the simplest case, when you have only one predictor (simple regression), say X_1 , the F -test tells you whether including X_1 does explain a larger part of the variance observed in Y compared to the null model (intercept only). The idea is then to test if the added explained variance (total variance, TSS, minus residual variance, RSS) is large enough to be considered as a “significant quantity”. We are here comparing a model with one predictor, or explanatory variable, to a baseline which is just “noise” (nothing except the grand mean).

Likewise, you can compute an F statistic in a multiple regression setting: In this case, it amounts to a test of *all predictors* included in the model, which under the HT framework means that we wonder whether any of them is useful in predicting the response variable. This is the reason why you may encounter situations where the F -test for the whole model is significant whereas some of the t or z -tests associated to each regression coefficient are not.

The F statistic looks like

$$F = \frac{(\text{TSS} - \text{RSS})/(p - 1)}{\text{RSS}/(n - p)},$$

where p is the number of model parameters and n the number of observations. This quantity should

A generalization of the above idea is the **likelihood ratio test**.

If you are using R, you can play with the above concepts like this:

```
df <- transform(X <- as.data.frame(replicate(2, rnorm(100))),
                y = V1+V2+rnorm(100))

## simple regression
anova(lm(y ~ V1, df))           # "ANOVA view"
summary(lm(y ~ V1, df))         # "Regression view"

## multiple regression
summary(lm0 <- lm(y ~ ., df))
lm1 <- update(lm0, . ~ . -V2) # reduced model
anova(lm1, lm0)                # test of V2
```

272 R: Choose factor level as dummy base in `lm()`

You can use `relevel()` to change the baseline level of your factor. For instance,

```
> g <- gl(3, 2, labels=letters[1:3])
> g
[1] a a b b c c
Levels: a b c
> relevel(g, "b")
[1] a a b b c c
Levels: b a c
```

273 How to visualize/summarize a matrix with number of rows \gg number of columns?

I was about to suggest something along @whuber's answer (I used this reordering technique but in a context of feature selection, so I was mainly concerned with the "variables view"). So let me suggest two other directions (but the first one is close to the already proposed one).

As far as **heatmaps** are concerned, you can display them after a slight rearrangement of rows (samples) and/or columns (genes) through hierarchical clustering (yet another aggregation method based on a (dis)similarity measure). There're a lot of R packages that can do this, for example the `cim()` function in **mixOmics**. Another package that might be of interest is **MADE4**; it relies on the very good **ade4** package for multivariate data analysis and visualization.

If you are concerned with the rather large number of variables, you might also consider some reduction method for **genes clustering**. One that I've heard about is *PCA-gene shaving* (Hastie et al., 2000), that is largely described in Izenman (2008). In essence, this is a two-stage iterative procedure where (a) for *feature selection*, we single out genes whose correlation with the first principal component is below a distribution-based threshold (say, the 10% of genes having the lowest correlation at each step), and (b) for *clustering*, we seek to maximize an R^2 measure (between-cluster/within-cluster variances) for j successive clusters of size k_j , where the optimal k_j is obtained by a permutation technique and the use of the *gap statistic* (after effects of the former cluster has been removed by residualization). More detailed informations can be

found in the paper referenced below, but the general idea is to cluster genes into *small and potentially overlapping subsets of correlated genes that vary as much as possible across individuals*.

References

1. Hastie, T., Tibshirani, R., Eisen, M.B., Alzadeh, A., Levy, R., Staudt, L., Chan, W.C., Botstein, D., and Brown, P.O. (2000). '**Gene shaving**' as a method for identifying distinct sets of genes with similar expression patterns. *Genome Biology*, 1(2).
2. Izenman, A.J. (2008). *Modern Multivariate Statistical Techniques*. Springer.

274 How can I specify a level of a factor while in an `lme()`

If you just want to fit separate models for each level of your factor, then probably the easiest way is to use the `subset=` argument to `lme` (or any other GLM in R, btw). For example,

```
lme(y ~ x + factor(repeatedmeasures) + fact,
    random = ~1 | z, data=mydata, subset=as.numeric(fact) == 1)
```

should subset on the first level of your factor. I used `as.numeric()` because you didn't provide labels to your newly created factor. To do so, you can fill the `labels=` argument when calling `cut()`, such that the above code could read

```
lme(..., subset=fact == "low")
```

for example (assuming the first level of `fact` is named `low`). In both cases, you can easily set up a loop (using something like `for i in 1:nlevels(fact)` or `for i in seq_along(levels(fact))`) to iterate over your different models.

275 Examples of studies using $p < 0.001$, $p < 0.0001$ or even lower p-values?

My opinion is that it does (and should) not depend on the field of study. For example, you may well work at a lower significance level than $p < 0.001$ if, for example, you are trying to replicate a study with

historical or well-established results (I can think of several studies on the **Stroop effect**, which had led to some controversies in the past few years). That amounts to consider a lower “threshold” within the classical Neyman-Pearson framework for testing hypothesis. However, statistical and practical (or substantive) significance is another matter.

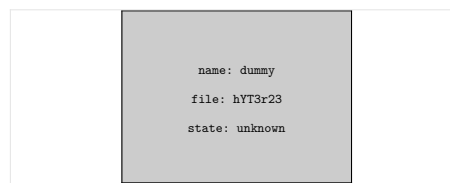
Sidenote. The “star system” seems to have dominated scientific inquiries as early as the 70’s, but see *The Earth Is Round* ($p < .05$), by J. Cohen (*American Psychologist*, 1994, 49(12), 997-1003), despite the fact that what we often want to know is given the data I have observed, what is the probability that H_0 is true? Anyway, there’s also a nice discussion on “**Why $P=0.05$?**”, by Jerry Dallal.

276 References for using networks to display correlations?

Do you know the **qgraph** project (and the related **R package**)? It aims at providing various displays for psychometric models, especially those relying on correlations. I discovered this approach for displaying correlation measures when I was reading a very nice and revolutionary article on diagnostic medicine by Denny Borsboom and coll.: **Comorbidity: A network perspective**, BBS (2010) 33: 137-193.

An oversimplified summary of their *network approach* of comorbidity is that it is “hypothesized to arise from direct relations between symptoms of multiple disorders”, contrary to the more classical view where these are comorbid disorders themselves that causes their associated symptoms to correlate (as reflected in a latent variable model, like factor or item response models, where a given symptom would allow to measure a particular disorder). In fact, symptoms are part of disorder, but they don’t measure it (and this is a mereological relationship). Their figure 5 describes such a “comorbidity network” and is particularly interesting as it embeds the frequency of symptoms and magnitude of their bivariate association in the same picture. They were using **Cytoscape** at that time, but the **qgraph** project has now reached a mature state.

Here are some examples from the on-line R help; basically, these are (1) an association graph with circular or (2) spring layout, (3) a concentration graph with spring layout, and (4) a factorial graph with spring layout (but see **help(qgraph.panel)**):



(See also **help(qgraph.pca)** for nice circular displays of an observed correlation matrix for the NEO-FFI, which is a 60-item personality inventory.)

277 In genome wide association studies, what are Principal Components?

In this particular context, PCA is mainly used to account for population-specific variations in alleles distribution on the SNPs (or other DNA markers, although I’m only familiar with the SNP case) under investigation. Such “population substructure” mainly arises as a consequence of varying frequencies of minor alleles in genetically distant ancestries (e.g. japanese and black-african or european-american). The general idea is well explained in **Population Structure and Eigenanalysis**, by Patterson et al. (*PLoS Genetics* 2006, 2(12)), or the *Lancet*’s special issue on genetic epidemiology (2005, 366; most articles can be found on the web, start with Cordell & Clayton, **Genetic Association Studies**).

The construction of principal axes follows from the classical approach to PCA, which is applied to the scaled matrix (individuals by SNPs) of observed genotypes (AA, AB, BB; say B is the minor allele in all cases), to the exception that an additional normalization to account for population drift might be applied. It all assumes that the frequency of the minor allele (taking value in $\{0,1,2\}$) can be considered as numeric, that is we work under an *additive model* (also called allelic dosage) or any equivalent one that would make sense. As the successive orthogonal PCs will account for the maximum variance, this provides a way to highlight groups of individuals differing at the level of minor allele frequency. The software used for this is known as **Eigenstrat**. It is worth to note that other methods to detect population substructure were proposed, in particular model-based cluster reconstruction (see references at the end). More information can be found by browsing the **Hapmap** project, and available tutorial coming from the **Bioconductor** project.

(Search for Vince J Carey or David Clayton’s nice tutorials on Google).

Considering that eigenanalysis allows to uncover some structure at the level of the individuals, we can use this information when trying to explain observed variations in a given phenotype (or any distribution that might be defined according to a binary criterion, e.g. disease or case-control situation). Specifically, we can adjust our analysis with those PCs (i.e., the factor scores of individuals), as illustrated in [Principal components analysis corrects for stratification in genome-wide association studies](#), by Price et al. (*Nature Genetics* 2006, 38(8)), and later work (there was a nice picture showing axes of genetic variation in Europe, but I can’t find it actually). Note also that another solution is to carry out a stratified analysis (by including ethnicity in an GLM)—this is readily available in the [snpMatrix](#) package, for example.

References

1. Daniel Falush, Matthew Stephens, and Jonathan K Pritchard (2003). [Inference of population structure using multilocus genotype data: linked loci and correlated allele frequencies](#). *Genetics*, 164(4): 1567–1587.
2. B Devlin and K Roeder (1999). [Genomic control for association studies](#). *Biometrics*, 55(4): 997–1004.
3. JK Pritchard, M Stephens, and P Donnelly (2000). [Inference of population structure using multilocus genotype data](#). *Genetics*, 155(2): 945–959.
4. Gang Zheng, Boris Freidlin, Zhaohai Li, and Joseph L Gastwirth (2005). [Genomic control for association studies under various genetic models](#). *Biometrics*, 61(1): 186–92.
5. Chao Tian, Peter K. Gregersen, and Michael F. Seldin (2008). [Accounting for ancestry: population substructure and genome-wide association studies](#). *Human Molecular Genetics*, 17(R2): R143–R150.
6. Kai Yu, [Population Substructure and Control Selection in Genome-wide Association Studies](#).
7. Alkes L. Price, Noah A. Zaitlen, David Reich and Nick Patterson (2010). [New approaches to population stratification in genome-wide association studies](#), *Nature Reviews Genetics*
8. Chao Tian, et al. (2009). [European Population Genetic Substructure: Further Definition of Ancestry Informative Markers for Distinguishing among Diverse European Ethnic Groups](#), *Molecular Medicine*, 15(11-12): 371–383.

278 How to customize axis labels in a boxplot?

Here’s a reproducible example, that you might adapt to fit with what you want to achieve with your data.

```
opar <- par(las=1)
df <- data.frame(y=rnorm(100), x=gl(2, 50, labels=letters[1:2]))
with(df, plot(y ~ x, axes=FALSE))
axis(1, at=1:2, labels=levels(df$x))
axis(2, at=seq(-3, 3, by=1),
      labels=paste(seq(-3, 3, by=1), "hr", sep=""))
box()
par(opar)
```

279 Problems with pie charts

I wouldn’t say there’s an increasing interest or debate about the use of pie charts. They are just found everywhere on the web and in so-called “predictive analytic” solutions.

I guess you know Tufte’s work (he also discussed the use of [multiple pie charts](#)), but more funny is the fact that the second chapter of Wilkinson’s *Grammar of Graphics* starts with “How to make a pie chart?”. You’re probably also aware that Cleveland’s [dotplot](#), or even a barchart, will convey much more precise information. The problem seems to really stem from the way our visual system is able to deal with spatial information. It is even quoted in the R software; from the on-line help for [pie](#),

Cleveland (1985), page 264: “Data that can be shown by pie charts always can be shown by a dot chart. This means that judgements of position along a common scale can be made instead of the less accurate angle judgements.” This statement is based on the empirical investigations of Cleveland and McGill as well as investigations by perceptual psychologists.

Cleveland, W. S. (1985) *The elements of graphing data*. Wadsworth: Monterey, CA, USA.

There are variations of pie charts (e.g., donut-like charts) that all raise the same problems: We are not good at evaluating angle and area. Even the ones used in “corrgram”, as described in Friendly, *Corrgrams: Exploratory displays for correlation matrices*, *American Statistician* (2002) 56:316, are hard to read, IMHO.

At some point, however, I wondered whether they might still be useful, for example (1) displaying two classes is fine but increasing the number of categories generally worsen the reading (especially with strong imbalance between %), (2) relative judgments are better than absolute ones, that is displaying two pie charts side by side should favor a better appreciation of the results than a simple estimate from, say a pie chart mixing all results (e.g. a two-way cross-classification table). Incidentally, I asked a similar question to Hadley Wickham who kindly pointed me to the following articles:

1. Spence, I. (2005). *No Humble Pie: The Origins and Usage of a Statistical Chart*. *Journal of Educational and Behavioral Statistics*, 30(4), 353–368.
2. Heer, J. and Bostock, M. (2010). *Crowdsourcing Graphical Perception: Using Mechanical Turk to Assess Visualization Design*. *CHI 2010*, April 10–15, 2010, Atlanta, Georgia, USA.

In sum, I think they are just good for grossly depicting the distribution of 2 to 3 classes (I use them, from time to time, to show the distribution of males and females in a sample on top of an histogram of ages), but they must be accompanied by relative frequencies or counts for being really informative. A table would still do a better job since you can add margins, and go beyond 2-way classifications.

Finally, there are alternative displays that are built upon the idea of pie chart. I can think of square pie or *waffle chart*, described by Robert Kosara in *Understanding Pie Charts*.

280 What is the difference between `independence.test` in R and COTT?

As a follow-up to my comment, if `independence.test`

refers to `coin::independence_test`, then you can reproduce a Cochran and Armitage trend test, as it is used in GWAS analysis, as follows:

```
> library(SNPassoc)
> library(coin)
> data(SNPs)
> datSNP <- setupSNP(SNPs,6:40,sep="")
> ( tab <- xtabs(~ casco + snp10001, data=datSNP) )
      snp10001
casco T/T C/T C/C
      0  24  21   2
      1  68  32  10
> independence_test(casco~snp10001, data=datSNP, teststat="quad",
                    scores=list(snp10001=c(0,1,2)))
```

Asymptotic General Independence Test

```
data: casco by snp10001 (T/T < C/T < C/C)
chi-squared = 0.2846, df = 1, p-value = 0.5937
```

This is a conditional version of the CATT. About scoring of the ordinal variable (here, the frequency of the minor allele denoted by the letter C), you can play with the `scores=` arguments of `independence_test()` in order to reflect the model you want to test (the above result is for a log-additive model).

There are five different genetic models that are generally considered in GWAS, and they reflect how genotypes might be collapsed: codominant (T/T (92) C/T (53) C/C (12), yielding the usual $\chi^2(2)$ association test), dominant (T/T (92) vs. C/T-C/C (65)), recessive (T/T-C/T (145) vs. C/C (12)), overdominant (T/T-C/C (104) vs. C/T (53)) and log-additive (0 (92) < 1 (53) < 2 (12)). Note that genotype recoding is readily available in inheritance functions from the *SNPassoc* package. The “scores” should reflect these collapsing schemes.

Following Agresti (CDA, 2002, p. 182), CATT is computed as $n \cdot r^2$, where r stands for the linear correlation between the numerical scores and the binary outcome (case/control), that is

```
z.catt <- sum(tab)*cor(datSNP$casco, as.numeric(datSNP$snp10001))^2
1 - pchisq(z.catt, df = 1) # p=0.5925
```

(There’s even a `prop.trend.test()` function in base R.)

There also exist various built-in CATT functions in R/Bioconductor ecosystem for GWAS, e.g.

- `CATT()` from *Rassoc*, e.g. with(`datSNP`, `CATT(table(casco,`

snp10001), 0.5)) # p=0.5925 (additive/multiplicative displays).

- in **snpMatrix**, there are headed as 1-df χ^2 -test when you call `single.snp.tests()` (see the **vignette**); please note that the default mode of inheritance is the codominant/additive effect.

Finally, here are two references that discuss the choice of scoring scheme depending on the genetic model under consideration, and some issues with power/robustness

1. Zheng, G, Freidlin, B, Li, Z and Gastwirth, JL (2003). Choice of scores in trend tests for case-control studies of candidate-gene associations. *Biometrical Journal*, **45**: 335-348.
2. Freidlin, B, Zheng, G, Li, Z, and Gastwirth, JL (2002). **Trend Tests for Case-Control Studies of Genetic Markers: Power, Sample Size and Robustness**. *Human Heredity*, **53**: 146-152.

See also the **GeneticsDesign** (bioc) package for power calculation with linear trend tests.

281 In R, fisher.test returns different results if I use vectors or contingency table

The first test should read

```
> fisher.test(expData[,1], expData[,2])

Fisher's Exact Test for Count Data

data: expData[, 1] and expData[, 2]
p-value = 0.4857
alternative hypothesis: true odds ratio is not equal to 1
95 percent confidence interval:
 0.001607888 4.722931239
sample estimates:
odds ratio
 0.156047
```

as per the doc: **x** is the outcome and **y** is the factor (or vice-versa).

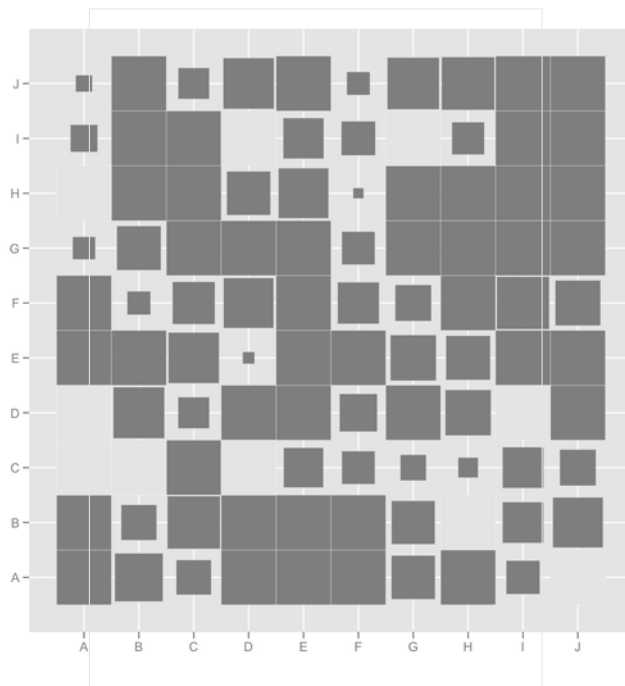
282 How to display a matrix of correlations with missing entries?

Building upon @GaBorgulya's response, I would suggest trying fluctuation or level plot (aka heatmap)

For example, using **ggplot2**:

```
library(ggplot2, quietly=TRUE)
k <- 100
rvals <- sample(seq(-1,1,by=.001), k, replace=TRUE)
rvals[sample(1:k, 10)] <- NA
cc <- matrix(rvals, nr=10)
ggfluctuation(as.table(cc)) + opts(legend.position="none")
+
  labs(x="", y="")
```

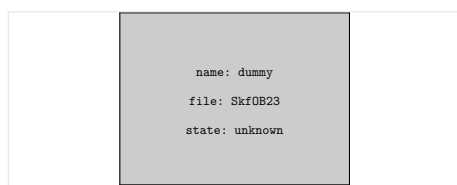
(Here, missing entry are displayed in plain gray, but the default color scheme can be changed, and you can also put "NA" in the legend.)



or

```
ggfluctuation(as.table(cc), type="color") + labs(x="", y="")
+
  scale_fill_gradient(low = "red", high = "blue")
```

(Here, missing values are simply not displayed. However, you can add a **geom_text()** and display something like "NA" in the empty cell.)



283 Two-Way Joining in R

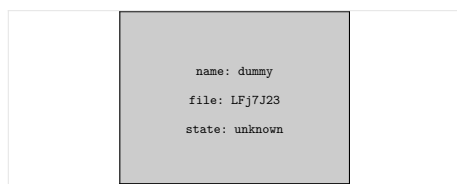
Generally speaking, you should always find useful pointers by looking at the relevant CRAN Task Views, in this case the one that deals with **Cluster** packages, or maybe **Quick-R**.

It's not clear to me whether the link you gave referenced standard clustering techniques for n (individuals) by k (variables) matrix of measures where we impose constraints on the resulting heatmap displays, or two-mode clustering or **biclustering**.

In the first approach, we could, for example,

1. compute a measure of (dis)similarity between individuals, or correlation between variables, and show the resulting $n \times n$ or $k \times k$ matrix where rows and columns are rearranged by some kind of partitioning or ordering technique – this help highlighting possible substructures in the association matrix, and you will find more information in this [related question](#);
2. compute the correlation between two blocks of data observed on the same individuals, and reorder the pattern of correlations following an external ordination technique (e.g., hierarchical clustering) – it amounts to show a heatmap of the observed statistics reordered by rows and columns.

As proposed in an [earlier response](#), the latter is readily available in the `cim()` function from the **mixOmics** package. From the on-line help, we can end up with something like that:



Please, note that this is just a two-step process to conveniently display summary measures of association: clustering of rows (individuals or variables)

and columns (individuals or variables) is done separately.

In the second approach (biclustering), that I'm inclined to favour, I only know one R package, **biclust**, that is greatly inspired for research in bioinformatics. Some pointers were also given in an [earlier thread](#). (But there's even some papers in the psychometrics literature.) In this case, we need to put some constraints during clustering because we want to cluster both individuals *and* variables *at the same time*.

Again, you can display the resulting structure as heatmaps (see `help(heatmapBC)`), as shown below

284 Software for easy-yet-robust data exploration

As far as exploratory (possibly interactive) data analysis is concerned, I would suggest to take a look at:

- **Weka**, originally targets data-mining applications, but can be used for data summaries.
- **Mondrian**, for interactive data visualization.
- **KNIME**, which relies on the idea of building data flows and is compatible with Weka and R.

All three accept data in **arff** or **csv** format.

In my view, Stata does not require so much programming expertise. This is even part of its attractiveness, in fact: Most of basic analysis can be done by point-and-click user actions, with dialog boxes for customizing specific parameters, say, for prediction in a linear model. The same applies, albeit to a lesser extent, to R when you use external GUIs like **Rcmdr**, **Deducer**, etc. as said by @gsk3.

285 Measure of association for 2x3 contingency table

Linear or monotonic trend tests— M^2 association measure, WMW test cited by @GaBorgulya, or the Cochran-Armitage trend test—can also be used, and they are well explained in Agresti (**CDA**, 2002, §3.4.6, p. 90).

The latter is actually equivalent to a score test for

but it can be computed from the M^2 statistic, defined as $(n-1)r^2$ ($\sim \chi^2(1)$ for large sample), where r is the sample correlation coefficient between the two variables (the ordinal measure being recoded as numerical scores), by replacing $n-1$ with n (ibid., p. 182). It is easy to compute in any statistical software, but you can also use the `coin` package in R (I provided an example of use for a [related question](#)).

Sidenote

If you are using R, you will find useful resources in either Laura Thompson's *R (and S-PLUS) Manual to Accompany Agresti's Categorical Data Analysis (2002)*, which shows how to replicate Agresti's results with R, or the `gnm` package (and its companion packages, `vcd` and `vcdExtra`) which allows to fit row-column association models (see the vignette, *Generalized nonlinear models in R: An overview of the gnm package*).

286 Estimates and C.I. of percentiles for a survival function

The `bootkm()` function in `Hmisc` provides bootstrapped estimate of the probability of survival, as well as the estimate of the quantile of the survival distribution (through either `describe` or `quantile` applied onto the result of `bootkm`).

287 Graphics encyclopedia

For an online summary, check out A [Periodic Table of Visualization Methods](#).

288 How to get generalisation performance from nnet in R using k-fold cross-validation?

Implementing k-fold CV (with or without nesting) is relatively straightforward in R; and stratified sampling (wrt. class membership or subjects' characteristics, e.g. age or gender) is not that difficult.

About the way to assess one's classifier performance, you can directly look at the R code for the `tune()` function. (Just type `tune` at the R prompt.) For a classification problem, this is the class agreement (between predicted and observed class membership) that is computed.

testing $H_0 : \beta = 0$ in a logistic regression model,

However, if you are looking for a complete R framework where data preprocessing (feature elimination, scaling, etc.), training/test resampling, and comparative measures of classifiers accuracy are provided in few commands, I would definitely recommend to have a look at the [caret](#) package, which also includes a lot of useful vignettes (see also the [JSS paper](#)).

Of note, although NNs are part of the methods callable from within [caret](#), you may probably have to look at other methods that perform as well and most of the times better than NNs (e.g., Random Forests, SVMs, etc.)

289 Multidimensional scaling pseudo-code

There are different kind of MDS (e.g., see this [brief review](#)). Here are two pointers:

- the [smacof](#) R package, developed by Jan de Leeuw and Patrick Mair has a nice vignette, [Multidimensional Scaling Using Majorization: SMACOF in R](#) (or see, the *Journal of Statistical Software* (2009) 31(3)) – R code is available, of course.
- there are some handouts on [Multidimensional Scaling](#), by Forrest Young, where several algorithms are discussed (including INDSCAL (Individual Difference Scaling, or weighted MDS) and ALSCAL, with Fortran [source code](#) by the same author) – this two keywords should help you to find other source code (mostly Fortran, C, or Lisp).

You can also look for “Manifold learning” which should give you a lot of techniques for dimension reduction (Isomap, PCA, MDS, etc.); the term was coined by the Machine Learning community, among others, and they probably have a different view on MDS compared to psychometricians.

290 How to compute the standard error of measurement (SEM) from a reliability estimate?

You should use the point estimate of the reliability, not the lower bound or whatsoever. I guess by lb/up you mean the 95% CI for the ICC (I don’t have SPSS, so I cannot check myself)? It’s unfortunate

that we also talk of Cronbach’s alpha as a “lower bound for reliability” since this might have confused you.

It should be noted that this formula is not restricted to the use of an estimate of ICC; in fact, you can plug in any “valid” measure of reliability (most of the times, it is Cronbach’s alpha that is being used). Apart from the NCME tutorial that I linked to in my comment, you might be interested in this recent article:

Tighe et al. [The standard error of measurement is a more appropriate measure of quality for postgraduate medical assessments than is reliability: an analysis of MRCP\(UK\) examinations](#). *BMC Medical Education* 2010, 10:40

Although it might seem to barely address your question at first sight, it has some additional material showing how to compute SEM (here with Cronbach’s α , but it is straightforward to adapt it with ICC); and, anyway, it’s always interesting to look around to see how people use SEM.

291 What are some interesting and well-written applied statistics papers?

It’s a bit difficult for me to see what paper might be of interest to you, so let me try and suggest the following ones, from the psychometric literature:

Borsboom, D. (2006). [The attack of the psychometricians](#). *Psychometrika*, 71, 425-440.

for dressing the scene (Why do we need to use statistical models that better reflect the underlying hypotheses commonly found in psychological research?), and

Borsboom, D. (2008). [Psychometric perspectives on diagnostic systems](#). *Journal of Clinical Psychology*, 64, 1089-1108.

for an applied perspective on diagnostic medicine (transition from yes/no assessment as used in the DSM-IV to the “dimensional” approach intended for the DSM-V). A larger review of latent variable models in biomedical research that I like is:

Rabe-Hesketh, S. and Skrondal, A. (2008). [Classical latent variable models for medical](#)

research. *Statistical Methods in Medical Research*, 17(1), 5-32.

292 What are some interesting and well-written applied statistics papers?

From a genetic epidemiology perspective, I would now recommend the following series of papers about **genome-wide association studies**:

1. Cordell, H.J. and Clayton, D.G. (2005). **Genetic association studies**. *Lancet* 366, 1121-1131.
2. Cantor, R.M., Lange, K., and Sinsheimer, J.S. (2010). **Prioritizing GWAS results: A review of statistical methods and recommendations for their application**. *The American Journal of Human Genetics* 86, 6-22.
3. Ioannidis, J.P.A., Thomas, G., Daly, M. [†] (2009). **Validating, augmenting and refining genome-wide association signals**. *Nature Reviews Genetics* 10, 318-329.
4. Balding, D.J. (2006). **A tutorial on statistical methods for population association studies**. *Nature Reviews Genetics* 7, 781-791.
5. Green, A.E. et al. (2008). **Using genetic data in cognitive neuroscience: from growing pains to genuine insights**. *Nature Reviews Neuroscience* 9, 710-720.
6. McCarthy, M.I. et al. (2008). **Genome-wide association studies for complex traits: consensus, uncertainty and challenges**. *Nature Reviews Genetics* 9, 356-369.
7. Psychiatric GWAS Consortium Coordinating Committee (2009). **Genomewide Association Studies: History, Rationale, and Prospects for Psychiatric Disorders**. *American Journal of Psychiatry* 166(5), 540-556.
8. Sebastiani, P. et al. (2009). **Genome-wide association studies and the genetic dissection of complex traits**. *American Journal of Hematology* 84(8), 504-15.
9. The Wellcome Trust Case Control Consortium (2007). **Genome-wide association study of 14,000 cases of seven common diseases and 3,000 shared**

controls. *Nature* 447, 661-678.

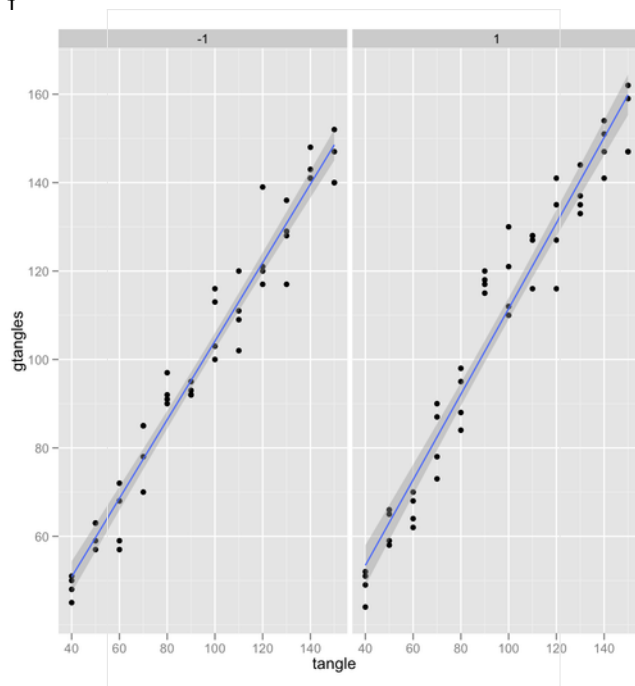
10. The Wellcome Trust Case Control Consortium (2010). **Genome-wide association study of CNVs in 16,000 cases of eight common diseases and 3,000 shared controls**. *Nature* 464, 713-720.

293 R - interaction plot with confidence intervals?

If you're willing to use **ggplot**, you can try the following code.

```
library(ggplot2)
gp <- ggplot(data=br, aes(x=tangle, y=gtangles))
gp + geom_point() + stat_smooth(method="lm", fullrange=T)
+ facet_grid(. ~ up)
```

for a faceted interaction plot



For a standard interaction plot (like the one produced by **interaction.plot()**), you just have to remove the facetting.

```
gp <- ggplot(data=br, aes(x=tangle, y=gtangles, colour=factor(up)))
gp + geom_point() + stat_smooth(method="lm")
```

name: dummy
file: UUHnu23
state: unknown

294 Multiple regression with no origin and mix of directly entered and stepwise entered variables using R

I think you can set up your base model, that is the one with your 12 IVs and then use `add1()` with the remaining predictors. So, say you have a model `mod1` defined like `mod1 <- lm(y ~ 0+x1+x2+x3)` (0+ means *no intercept*), then

```
add1(mod1, ~ .+x4+x5+x6, test="F")
```

will add and test one predictor after the other on top of the base model.

More generally, if you know in advance that a set of variables should be included in the model (this might result from prior knowledge, or whatsoever), you can use `step()` or `stepAIC()` (in the **MASS** package) and look at the `scope=` argument.

Here is an illustration, where we specify a priori the functional relationship between the outcome, y , and the predictors, x_1, x_2, \dots, x_{10} . We want the model to include the first three predictors, but let the selection of other predictors be done by stepwise regression:

```
set.seed(101)
X <- replicate(10, rnorm(100))
colnames(X) <- paste("x", 1:10, sep="")
y <- 1.1*X[,1] + 0.8*X[,2] - 0.7*X[,5] + 1.4*X[,6] + rnorm(100)
df <- data.frame(y=y, X)

# say this is one of the base model we think of
fm0 <- lm(y ~ 0+x1+x2+x3+x4, data=df)

# build a semi-constrained stepwise regression
fm.step <- step(fm0, scope=list(upper = ~ 0+x1+x2+x3+x4+x5+x6+x7+x8+x9+x10,
                                lower = ~ 0+x1+x2+x3), trace=FALSE)

summary(fm.step)
```

The results are shown below:

```
Coefficients:
      Estimate Std. Error t value Pr(>|t|)
```

```
x1    1.0831      0.1095   9.888 2.87e-16 ***
x2    0.6704      0.1026   6.533 3.17e-09 ***
x3   -0.1844      0.1183  -1.558  0.123
x6    1.6024      0.1142  14.035 < 2e-16 ***
x5   -0.6528      0.1029  -6.342 7.63e-09 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
Residual standard error: 1.004 on 95 degrees of freedom
Multiple R-squared:  0.814,    Adjusted R-squared:  0.8042
F-statistic: 83.17 on 5 and 95 DF,  p-value: < 2.2e-16
```

You can see that x_3 has been retained in the model, even if it proves to be non-significant (well, the usual caveats with univariate tests in multiple regression setting and model selection apply here – at least, its relationship with y was not specified).

295 Graphics encyclopedia

[A Tour through the Visualization Zoo](#) (Heer et al., Visualization 8(5) 2010) offers a particularly interesting overview of “innovative” and interactive techniques for displaying data.

On a related point, a good software for data visualization, including the aforementioned gallery, is **Protovis**, which comes with a lot of [examples](#).

296 Calculating the MacKinnon empirical distribution test to test mediation

Tests for full and partial mediation are well explained on the [Mediation FAQ](#) webpage by David P. MacKinnon, and they can be implemented in any statistical package offering tools for regression modeling. Bootstrapping can be used to derive standard errors and confidence intervals for the estimated coefficients.

If you are using R, there’s even a **mediation** package that helps you to estimate those effects, but also to conduct a sensitivity analysis on mediation effect for violations of sequential ignorability assumption. I’ve been previously using the **QuantPsyc** package, from [Thomas Fletcher](#), which implements methods proposed by MacKinnon and coll. With Stata, it can be done as described on the [UCLA Stata FAQ](#), [How can I do moderated mediation in Stata?](#)

A good overview is also available in

Preacher, K.J., Rucker, D.D., and Hayes, A.F. (2007). [Addressing Moderated Mediation Hypotheses: Theory, Methods, and Prescriptions](#). *Multivariate Behavioral Research*, 42(1), 185-227.

297 Prediction in simple and multiple ANOVA

You can use `lm()` instead of `aov()` in this case (the latter is a wrapper of the former).

Here is an illustration:

```
n <- 100
A <- gl(2, n/2, n, labels=paste("a", 1:2, sep=""))
B <- gl(2, n/4, n, labels=paste("b", 1:2, sep=""))
# generate fake data for a balanced two-way ANOVA
df <- data.frame(y=rnorm(n), A, B)
summary(lm1 <- lm(y~A+B, data=df)) # compare with summary.aov(...)
predict(lm1, expand.grid(A=levels(A), B=levels(B)), interval=
```

The latter command gives you predictions for each combination of the A and B factor levels (here, I didn't include the interaction), in the following order:

```
  A B
1 a1 b1
2 a2 b1
3 a1 b2
4 a2 b2
```

Another option is to use the [effects](#) package.

298 Using R online - without installing it

Yes, there are some Rweb interface, like [this one](#).

Note: Installation of the R software is pretty straightforward and quick, on any platform.

299 Cox model with LASSO

Here are two suggestions. First, you can take a look at the [glmnet](#) package, from Friedman, Hastie and Tibshirani, but see their JSS 2010 (33) paper, [Regularization Paths for Generalized Linear Models via](#)

[Coordinate Descent](#).

Second, although I've never used this kind of penalized model, I know that the [penalized](#) package implements L1/L2 penalties on GLM and the Cox model. What I found interesting in this package (this was with ordinary regression) was that you can include a set of unpenalized variables in the model.

The associated publication is now:

Goeman J.J. (2010). [L-1 Penalized Estimation in the Cox Proportional Hazards Model](#). *Biometrical Journal* 52 (1) 70-84.

300 How to calculate pseudo-R² from R's logistic regression?

Don't forget the [rms](#) package, by Frank Harrell. You'll find everything you need for fitting and validating GLMs.

Here is a toy example (with only one predictor):

```
set.seed(101)
n <- 200
x <- rnorm(n)
a <- 1
b <- -2
p <- exp(a+b*x)/(1+exp(a+b*x))
y <- factor(ifelse(runif(n)<p, 1, 0), levels=0:1)
mod1 <- glm(y ~ x, family=binomial)
summary(mod1)
```

This yields:

```
Coefficients:
              Estimate Std. Error z value Pr(>|z|)
(Intercept)   0.8959      0.1969   4.55 5.36e-06 ***
x             -1.8720      0.2807  -6.67 2.56e-11 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 258.98  on 199  degrees of freedom
Residual deviance: 181.02  on 198  degrees of freedom
AIC: 185.02
```

Now, using the `lrm` function,

```
require(rms)
mod1b <- lrm(y ~ x)
```


You soon get a lot of model fit indices, including Nagelkerke R^2 , with `print(mod1b)`:

Logistic Regression Model

```
lrm(formula = y ~ x)
```

Rank Discrim.		Model Likelihood		Discrimination	
		Ratio Test		Indexes	
Obs	200	LR chi2	77.96	R2	0.445
C	0.852				
0	70	d.f.	1	g	2.054
Dxy	0.705				
1	130	Pr(> chi2)	<0.0001	gr	7.801
gamma	0.705				
max deriv	2e-08			gp	0.319
tau-a	0.322				
				Brier	0.150

	Coef	S.E.	Wald Z	Pr(> Z)
Intercept	0.8959	0.1969	4.55	<0.0001
x	-1.8720	0.2807	-6.67	<0.0001

Here, $R^2 = 0.445$ and it is computed as $(1 - \exp(-LR/n)) / (1 - \exp(-\chi^2/n))$ where LR is the χ^2 stat (comparing the two nested models you described), whereas the denominator is just the max value for R^2 . For a perfect model, we would expect $LR = 2L_0$, that is $R^2 = 1$.

By hand,

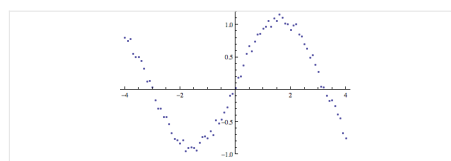
```
> mod0 <- update(mod1, .~.-x)
> lr.stat <- lrtest(mod0, mod1)
> (1-exp(-as.numeric(lr.stat$stats[1])/n))/(1-exp(-as.numeric(logLik(mod0)/n)))
[1] 0.4445742
> mod1b$stats["R2"]
R2
0.4445742
```

Ewout W. Steyerberg discussed the use of R^2 with GLM, in his book *Clinical Prediction Models* (Springer, 2009, § 4.2.2 pp. 58-60). Basically, the relationship between the LR statistic and Nagelkerke's R^2 is approximately linear (it will be more linear with low incidence). Now, as discussed on the earlier thread I linked to in my comment, you can use other measures like the c statistic which is equivalent to the AUC statistic (there's also a nice illustration in the above reference, see Figure 4.6).

301 How can I calculate the auto-correlation of a signal in Mathematica environment?

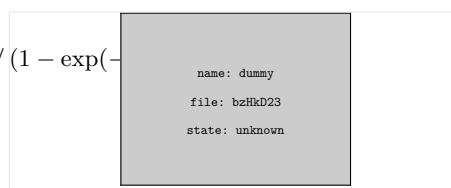
It's been a long since I didn't play with Mathematica, and I just had a quick look on Google, but can't you just use (here with some fake data)

```
x = Table[Sin[x] + 0.2 RandomReal[], {x, -4, 4, .1}];
ListPlot[x, DataRange -> {-4, 4}]
```



the function `ListCorrelate`?

```
acf = ListCorrelate[x, x, {1, 1}, 0]
ListPlot[acf, Filling -> Axis]
```



302 Problem with pvclust in R

It's difficult to answer without seeing the data itself, but my best guess is that you have some non numerical entries in the matrix/dataframe (which is what is expected by `pvclust`). For example,

```
> as.numeric(c(1,2,"NA"))
[1] 1 2 NA
```

or

```
> dist(c(1,2,"NA"))
1 2
2 1
3 NA NA
```

will produce the same warning message ('NAs introduced by coercion'). I deliberately used "NA", but any element that is not numerical will result in the same warning message.

So,

1. A warning message is issued when trying to compute a distance matrix from the non-numerical input
2. Then, `hclust` failed when it is called within `pvclust`. Again,

```
> hclust(dist(c(1,2,"NA")))
```

will throw the same error message.

In your first try, you called `hclust` by using a matrix. Can't you just check that you use the same variables in both cases, or that there is no strange values in your data (e.g., `summary(transpose)`), or no missing values coded as the character "NA" instead of `NA`, as below:

```
> xx <- data.frame(replicate(3, sample(c(1,2,3), 3)))
> xx[2,3] <- "NA"
> is.na(xx[2,3])
[1] FALSE
> sapply(xx, is.character)
  X1  X2  X3
FALSE FALSE TRUE
> apply(xx, 2, function(x) sum(is.na(x)))
X1 X2 X3
 0  0  0
# Now if we had a true NA value, we would see
> xx[2,3] <- NA
> apply(xx, 2, function(x) sum(is.na(x)))
X1 X2 X3
 0  0  1
```

303 How to print a single column in the output of forecast function in R forecast package?

Although I never used the `forecast` package, looking at the output of

```
ts1 <- ets(USAccDeaths)
fc <- forecast(ts1, h=48)
str(fc)
```

it seems that what you are looking for is just stored as

```
fc$mean
```

while `fc$lower` (resp. `fc$upper`) will give you 80

and 95% lower (resp. upper) limits for prediction. Note that you'll have to replace `fc` with the name of your variable, `fit`. Note also that this is *fully documented* in the on-line help for `forecast` (see returned value):

```
An object of class "forecast" is a list containing
at least the
following elements:

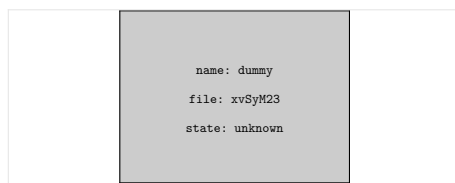
model: A list containing information about the fitted model
method: The name of the forecasting method as a character
string
mean: Point forecasts as a time series
lower: Lower limits for prediction intervals
upper: Upper limits for prediction intervals
```

304 How to compute correlation between/within groups of variables?

What @rolando suggested looks like a good start, if not the whole response (IMO). Let me continue with the correlational approach, following the Classical Test Theory (CTT) framework. Here, as noted by @Jeromy, a summary measure for your group of characteristics might be considered as the totalled (or sum) score of all items (a characteristic, in your words) belonging to what I will now refer to as a scale. Under CTT, this allows us to formalize individual "trait" propensity or liability as one's location on a continuous scale reflecting an underlying construct (a latent trait), although here it is merely an ordinal scale (but this another debate in the psychometrics literature).

What you described has to do with what is known as *convergent* (to what extent items belonging to the same scale do correlate one with each other) and *discriminant* (items belonging to different scales should not correlate to a great extent) validity in psychometrics. Classical techniques include multi-trait multi-method (MTMM) analysis (Campbell & Fiske, 1959). An illustration of how it works is shown below (three methods or instruments, three constructs or traits):

In this MTMM matrix, the diagonal elements might be Cronbach's alpha or test-retest intraclass correlation; these are indicators of the *reliability* of each measurement scale. The *validity* of the hypothesized (shared) constructs is assessed by the correlation of scales scores when different instruments



are used to assess the same trait; if these instrument were developed independently, high correlation (> 0.7) would support the idea that the traits are defined in a consistent and objective manner. The remaining cells in this MTMM matrix summarize relations *between traits within method*, and *between traits across methods*, and are indicative of the way unique constructs are measured with different scales and what are the relations between each trait in a given scale. Assuming independent traits, we generally don't expect them to be high (a recommended threshold is $< .3$), but more formal test of hypothesis (on correlation point estimates) can be carried out. A subtlety is that we use so-called "rest correlation", that is we compute correlation between an item (or trait) and its scale (or method) after removing the contribution of this item to the sum score of this scale (correction for overlap).

Even if this method was initially developed to assess convergent and discriminant validity of a certain number of traits as studied by different measurement instruments, it can be applied for a single multi-scale instrument. The traits then becomes the items, and the methods are just the different scales. A generalization of this method to a single instrument is also known as *multitrait scaling*. Items correlating as expected (i.e., with their own scale rather than a different scale) are counted as *scaling success*. We generally assume, however, that the different scales are not correlated, that is they are targeting different hypothetical constructs. But averaging the within and between-scale correlations provide a quick way of summarizing the internal structure of your instrument. Another convenient way of doing so is to apply a cluster analysis on the matrix of pairwise correlations and see how your variables do hang together.

Of note, in both cases, the usual caveats of working with correlation measures apply, that is you cannot account for measurement error, you need a large sample, instruments or tests are assumed to be "parallel" (tau-equivalence, uncorrelated errors, equal error variances).

The second part addressed by @rolando is also interesting: If there's no theoretical or substantive

indication that the already established grouping of items makes sense, then you'll have to find a way to highlight the structure of your data with e.g., exploratory factor analysis. But even if you trust those "characteristics within a group", you can check that this is a valid assumption. Now, you might be using confirmatory factor analysis model to check that the pattern of items loadings (correlation of an item with its own scale) behaves as expected.

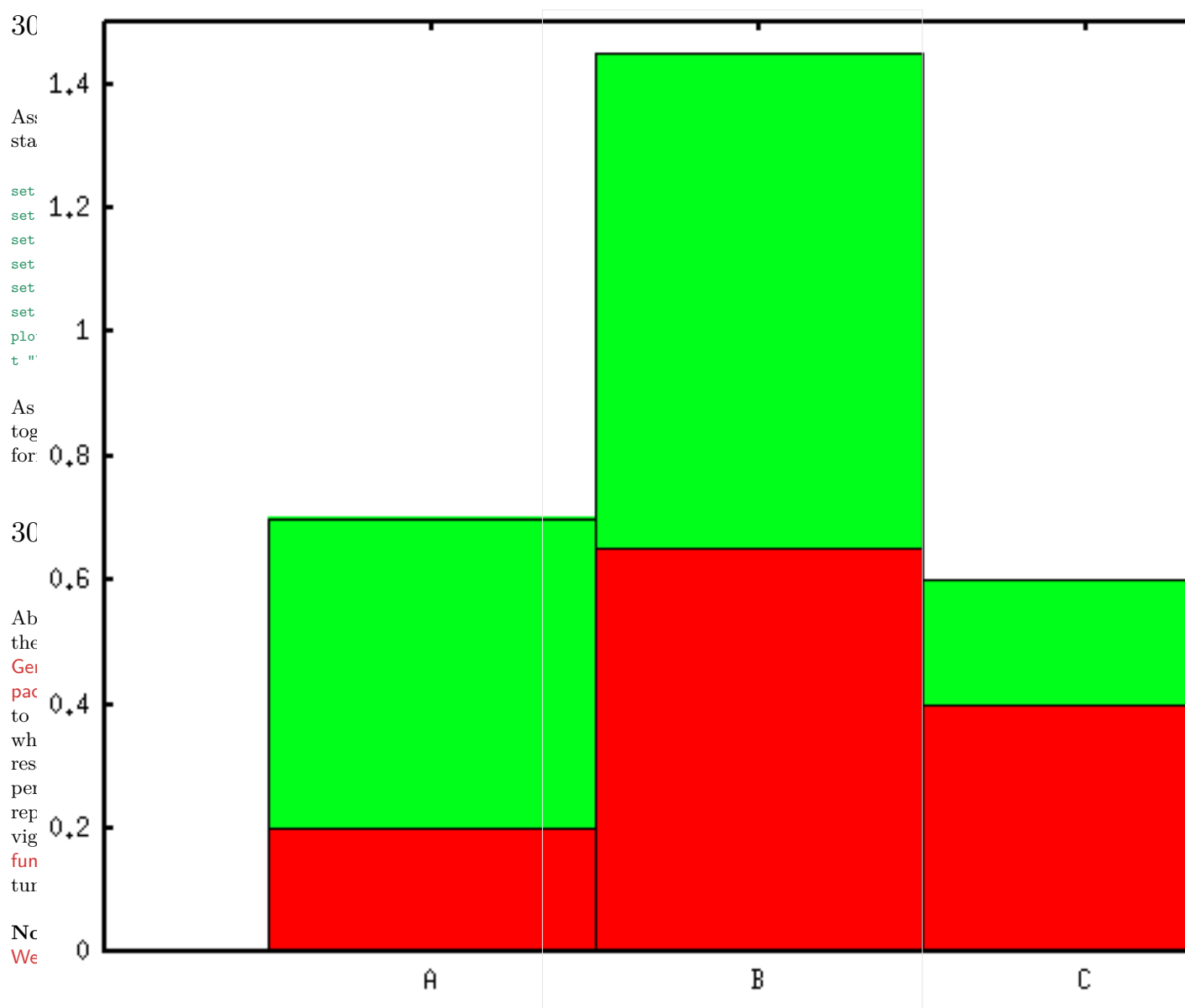
Instead of traditional factor analytic methods, you can also take a look at items clustering (Revelle, 1979) which relies on a Cronbach's alpha-based split-rule to group together items into homogeneous scales.

A final word: If you are using R, there are two very nice packages that will ease the aforementioned steps:

- **psych**, provides you with everything you need for getting started with psychometrics methods, including factor analysis (`fa`, `fa.parallel`, `principal`), items clustering (ICLUST and related methods), Cronbach's alpha (`alpha`); there's a nice overview available on William Revelle's website, especially [An introduction to psychometric theory with applications in R](#).
- **psy**, also includes scree plot (via PCA + simulated datasets) visualization (`scree.plot`) and MTMM (`mtmm`).

References

1. Campbell, D.T. and Fiske, D.W. (1959). Convergent and discriminant validation by the multitrait-multimethod matrix. *Psychological Bulletin*, 56: 81–105.
2. Hays, R.D. and Fayers, P. (2005). Evaluating multi-item scales. In *Assessing quality of life in clinical trials*, (Fayers, P. and Hays, R., Eds.), pp. 41-53. Oxford.
3. Revelle, W. (1979). Hierarchical Cluster Analysis and the Internal Structure of Tests. *Multivariate Behavioral Research*, 14: 57-74.



307 Implementations of the Random Forest algorithm

The [ELSI](#) used [randomForest](#) (see e.g., footnote 3 p.591), which is an R implementation of the Breiman and Cutler's [Fortran code](#) from Salford. Andy Liaw's code is in C.

There's another implementation of RFs proposed in the [party](#) package (in C), which relies on R/Lapack, which has some dependencies on BLAS (see [include/R_ext/Lapack.h](#) in your base R di-

rectory).

As far as bagging is concerned, it should not be too hard to parallelize it, but I'll let more specialized users answer on this aspect.

308 How to get a redundancy index when performing canonical correlation analysis in SPSS?

As it is not really difficult to import SAV dataset in R nowadays, with e.g.,

```
library(foreign)
df <- read.spss("yourfilename", to.data.frame=TRUE)
```

you can check your SPSS results against one of the R packages that allow to perform CCA (see the CRAN Task View on [Multivariate](#) or [Psychometrics](#) analysis). In particular, the [vegan](#) package offers an handy way to apply CCA and has nice graphical and numerical summary through the [CCorA\(\)](#) function.

Also, note that redundancy indexes apply onto one block of variables, conditional on the other block (hence the distinction you'll find in the aforementioned function between $Y|X$ and $X|Y$); they are intended to provide a measure of the variance of one set of variables predicted from the linear combination of the other set of variables. However, in essence CCA consider that you have two sets of measures that play a symmetrical role. They are both descriptions of the same individuals or statistical units. If your blocks really play an asymmetric role—that is you have a block of predictors and a block of response variable—then you're better using [PLS regression](#).

309 Two-stage clustering in R

The closest package that I can think of is [birch](#), but it is not available on CRAN anymore so you have to get the source and install it yourself ([R CMD install birch_1.1-3.tar.gz](#) works fine for me, OS X 10.6 with R version 2.13.0 (2011-04-13)). It implements the original algorithm described in

Zhang, T. and Ramakrishnan, R. and Livny, M. (1997). [BIRCH: A New Data Clustering Algorithm and Its Applications](#). *Data Mining and Knowledge Discovery*, 1, 141-182.

which relies on cluster feature tree, as does SPSS TwoStep (I cannot check, though). There's a possibility of using the k-means algorithm to perform clustering on [birch](#) object ([kmeans.birch\(\)](#)), that is partition the subclusters into k groups such that the sum of squares of all the points in each subclus-

ter to the assigned cluster centers is minimized.

310 Assumption of additivity for intra-class correlation

What you describes about Tukey's nonadditivity test sounds good to me. In effect, it allows to test for an item by rater interaction. Some words of caution, though:

- Tukey's nonadditivity test effectively allows to test for a *linear-by-linear* product of two factor main effects.
- The possibility of deriving a total score is irrelevant here, as this particular Tukey's test can be applied in any randomized block design, as described on [Stata FAQ](#), for example.
- It applies in situation where you have a *single observation per cell*, that is each rater assess only one item (no replicates).

You might recall that the interaction term is confounded with the error term when there're no replicates in an ANOVA design; in inter-rater studies, it means we have only one rating for each rater x item cell. Tukey's test in this case provide a 1-DF test for assessing any deviation from additivity, which is a common assumption to interpret a main effect in two-factor models. Here is a [tutorial](#) describing how it works.

I must admit I never used it when computing ICC, and I spent some times trying to reproduce Dave Garson's results with R. This led me to the following two papers that showed that Tukey's nonadditivity test might not be the "best" test to use as it will fail to recover a true interaction effect (e.g., where some raters exhibit an opposite rating behavior compared to the rest of the raters) when there's no main effect of the target of the ratings (e.g., marks given to items):

1. Lahey, M.A., Downey, R.G., and Saal, F.E. (1983). Intraclass Correlations: There's More There Than Meets the Eye. *Psychological Bulletin*, 93(3), 586-595.
2. Johnson, D.E. and Graybill, F.A. (1972). An analysis of a two-way model with interaction and no replication. *Journal of the American Statistical Association*, 67, 862-868.

3. Hegemann, V. and Johnson, D.E. (1976). The power of two tests for nonadditivity. *Journal of the American Statistical Association*, 71(356), 945-948.

(I'm very sorry but I couldn't find ungated PDF version of those papers. The first one is really a must-read one.)

About your particular design, you considered raters as fixed effects (hence the use of Shrout and Fleiss's type 3 ICC, i.e. mixed model approach). In this case, Lahey et al. (1) stated that you face a situation of nonorthogonal interaction components (i.e., the interaction is not independent of other effect) and a biased estimate of the rating effect – but, this for the case where you have a single observation per cell (ICC(3,1)). With multiple ratings per items, estimating ICC(3,k) requires the “assumption of nonsignificance of the interaction. In this case, the ANOVA effects are neither theoretically nor mathematically independent, and without adequate justification, the assumption of no interaction is very tenuous.”

In other words, such an interaction test aims at offering you diagnostic information. My opinion is that you can go on with you ICC, but be sure to check that (a) there's a significant effect for the target of ratings (otherwise, it would mean the reliability of measurements is low), (b) no rater systematically deviates from others' ratings (this can be done graphically, or based on the residuals of your ANOVA model).

More technical details are given below.

The alternative test that is proposed is called the *characteristic root test of the interaction* (2,3). Consider a multiplicative interaction model of the form (here, as an effect model, that is we use parameters that summarize deviations from the grand mean):

$$\mu_{ij} = \mu + \tau_i + \beta_j + \lambda\alpha_i\gamma_j + \varepsilon_{ij}$$

with τ ($i = 1, \dots, t$) the effect due to targets/items, β ($j = 1, \dots, b$) the effect of raters, $\alpha\gamma$ the interaction targets x raters, and the usual assumptions for the distribution of errors and parameters constraints. We can compute the largest characteristic root of $Z'Z$ or ZZ' , where $Z = z_{ij} = y_{ij} - y_{i.} - y_{.j} + y_{..}$ is the $t \times b$ matrix of residuals from an additive model.

The test then relies on the idea of using λ_1/RSS as a test statistic ($H_0 : \lambda = 0$) where λ_1 is the largest nonzero characteristic root of ZZ' (or $Z'Z$), and RSS equals the residual sum of squares from an additive model (2).

311 References on numerical optimization for statisticians

Optimization, by Kenneth Lange (Springer, 2004), reviewed in JASA by Russell Steele. It's a good textbook with Gentle's *Matrix algebra* for an introductory course on Matrix Calculus and Optimization, like the one by Jan de Leeuw (courses/202B).

312 Making a heatmap with a pre-computed distance matrix and data matrix in R

Ok, so you can just look at the code by typing the name of the function at the R prompt, or use `edit(pheatmap)` to see it in your default editor.

Around line 14 and 23, you'll see that another function is called for computing the distance matrices (for rows and columns), given a distance function (R `dist`) and a method (compatible with `hclust` for hierarchical clustering in R). What does this function do? Use `getAnywhere("cluster_mat")` to print it on screen, and you soon notice that it does nothing more than returning an `hclust` object, that is your dendrogram computed from the specified distance and linkage options.

So, if you already have your distance matrix, change line 14 (rows) or 23 (columns) so that it reads, e.g.

```
tree_row = hclust(my.dist.mat, method="complete")
```

where `my.dist.mat` is your own distance function, and `complete` is one of the many methods available in `hclust` (see `help(hclust)`). Here, it is important to use `fix(pheatmap)` and not `edit(pheatmap)`; otherwise, the edited function will not be callable in the correct environment/namespaces.

This is a quick and dirty hack that I would not recommend with larger package. It seems to work for me at least, that is I can use a custom distance matrix with complete linkage for the rows.

In sum, assuming your distance matrix is stored in a variable named `dd`,

```
library(pheatmap)
fix(pheatmap)
# 1. change the function as you see fit
# 2. save and go back to R
# 3. if your custom distance matrix was simply read as a matrix,
make sure
# it is read as a distance matrix
my.dist.map <- dd # or as.dist(dd)
```

Then, you can call `pheatmap` as you did but now it will use the results of `hclust` applied to `my.dist.map` with `complete` linkage. Please note that you just have to ensure that `cluster_rows=TRUE` (which is the default). Now, you may be able to change

- the linkage method
- choose between rows or columns

by editing the package function appropriately.

313 How to get Cooks distance and carry out residual analysis for non `lm()` and non `glm()` with R

As described in the on-line help, the `cooks.distance()` function expects an object of class `lm` or `glm` so it is not possible to get it work with other type of models. It is defined in `src/library/stats/R/lm.influence.R`, from R source, so you can browse the code directly and build your own function if nothing exists in other places. A quick way of seeing what it does is to type `stats:::cooks.distance.lm` at the R prompt, though.

Also, as `tobit` is nothing more than a wrapper for `survreg`, all attached methods to the latter kind of R object might be used. For example, there's a `residuals.survreg` (in the `survival` package) S3 method for extracting residuals from objects inheriting from class `survreg`.

314 Using LaTeX expression in gnuplot

If your penultimate goal is to embed your plot in

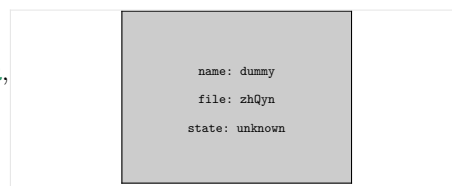
a Latex document, you might consider using the `gnuplottex` package (as an alternative to `pgfplots` which is an awesome package). The idea is rather simple: you write your code chunk in your tex document directly (like you would do with Sweave), et voilà!

Here is an example (grabbed from gnuplot demos):

```
\documentclass{standalone}
\usepackage{gnuplottex}

\begin{document}
\begin{gnuplot}[scale=0.95, terminal=epslatex]
set style fill transparent pattern 4 bo
set style function filledcurves y1=0
set clip two
Gauss(x,mu,sigma) = 1./(sigma*sqrt(2*pi)) * exp( -(x-mu)**2
/ (2*sigma**2) )
d1(x) = Gauss(x, 0.5, 0.5)
d2(x) = Gauss(x, 2., 1.)
d3(x) = Gauss(x, -1., 2.)
set xrange [-5:5]
set yrange [0:1]
set xlabel "$\bar{x}$ values"
unset colorbox
plot d1(x) fs solid 1.0 lc rgb "forest-green", \
      d2(x) lc rgb "gold", d3(x) lc rgb "red"
\end{gnuplot}
\end{document}
```

You'll need to compile with the `-shell-escape` option to `pdflatex`.



315 Calculating AUPR in R

A little googling returns one bioc package, `qpgraph` (`qpPrecisionRecall`), and a cran one, `minet` (`auc.pr`). I have no experience with them, though. Both have been devised to deal with biological networks.

316 Reference or book on simulation of experimental design data in R

Statistical models in S, by Chambers and Hastie (Chapmann and Hall, 1991; or the so-called *White Book*), and to a lesser extent *Modern Applied Statistics with S*, by Venables and Ripley (Springer, 2002, 4th ed.), include some material about DoE and the analysis of common designs in S and R. Vikneswaran wrote [An R companion to “Experimental Design”](#), although it is not very complete (IMHO), but there are a lot other textbooks in the [Contributed](#) section on CRAN that might help you get started.

Apart from textbook, the CRAN Task View on [Design of Experiments \(DoE\) & Analysis of Experimental Data](#) has some good packages that ease the creation and analysis of various experimental designs; I can think of [dae](#), [agricolae](#), or [AlgDesign](#) (which comes with a nice [vignette](#)), to name a few.

317 Using an SVM for feature selection

As I understand them, SVMs have built-in *regularization* because they tend to penalize large weights of predictors which amounts to favor simpler models. They’re often used with [recursive feature elimination](#) (in neuroimaging paradigms, at least).

About R specifically, there’s the [kernlab](#) package, by [Alex Smola](#) who co-authored *Learning with Kernels* (2002, MIT Press), which implements SVM (in addition to [e1071](#)). However, if you are after a dedicated framework, I would warmly recommend the [caret](#) package.

318 Multinomial choice with binary observations

Unless I misunderstood the question, this refer to paired preference (1) or [pair comparison](#) data. A well-known example of such a model is the Bradley-Terry model (2), which shares some connections with item scaling in psychometrics (3). There is an R package, [BradleyTerry2](#), described in the JSS (2005) 12(1), [Bradley-Terry Models in R](#), and a detailed overview in Agresti’s CDA, pp. 436-439, with

R code available in Laura Thompson’s textbook, [R \(and S-PLUS\) Manual to Accompany Agresti’s Categorical Data Analysis \(2002\) 2nd edition](#).

References

1. Thurstone, L.L. (1927). A law of comparative judgment. *Psychological Review*, 3, 273-286.
2. Bradley, R.A. and Terry, M.E. (1952). Rank analysis of incomplete block designs I: The methods of paired comparisons. *Biometrika*, 39, 324-345.
3. Andrich, D. (1978). Relationships between the Thurstone and Rasch approaches to item scaling. *Applied Psychological Measurement*, 2(3), 451-462.

319 Friedman’s test and post-hoc analysis

As @caracal’s said, this script implements a permutation-based approach to Friedman’s test with the [coin](#) package. The maxT procedure is rather complex and there is no relation with the traditional χ^2 statistic you’re probably used to get after a Friedman ANOVA. The general idea is to control the [FWER](#). Let’s say you perform 1000 permutations, for every variable of interest, then you can derive not only pointwise empirical p-values for each variable (as you would do with a single permutation test) but also a value that accounts for the fact that you tested all those variables at the same time. The latter is achieved by comparing each observed test statistic against the maximum of permuted statistics over all variables. In other words, this p-value reflects the chance of seeing a test statistic as large as the one you observed, given you’ve performed as many tests. More information (in a genomic context, and with algorithmic considerations) can be found in

Dudoit, S., Shaffer, J.P., and Boldrick, J.C. (2003). [Multiple Hypothesis Testing in Microarray Experiments](#). *Statistical Science*, 18(1), 71–103.

(Here are some [slides](#) from the same author with applications in R with the [multtest](#) package.)

Another good reference is *Multiple Testing Procedures with Applications to Genomics*, by Dudoit and van der Laan (Springer, 2008).

Now, if you want to get more “traditional” statistic, you can use the `agricolae` package which has a `friedman()` function that performs the overall Friedman’s test followed by post-hoc comparisons.

The permutation method yields a $\max T=3.24$, $p=0.003394$, suggesting an overall effect of the target when accounting for the blocking factor. The post-hoc tests basically indicate that only results for Wine A vs. Wine C ($p=0.003400$) are statistically different at the 5% level.

Using the non-parametric test, we have

```
> library(agricolae)
> with(WineTasting, friedman(Taster, Wine, Taste, group=FALSE))
Friedman's Test
=====
Adjusted for ties
Value: 11.14286
Pvalue chisq : 0.003805041
F value : 7.121739
Pvalue F: 0.002171298

Alpha      : 0.05
t-Student  : 2.018082

Comparison between treatments
Sum of the ranks
```

	Difference	pvalue	sig	LCL	UCL
Wine A - Wine B	6	0.301210		-5.57	17.57
Wine A - Wine C	21	0.000692	***	9.43	32.57
Wine B - Wine C	15	0.012282	*	3.43	26.57

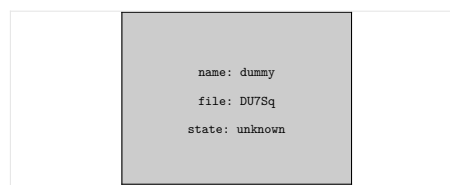
The two global tests agree and basically say there is a significant effect of Wine type. We would, however, reach different conclusions about the pairwise difference. It should be noted that the above pairwise tests (Fisher’s LSD) are not really corrected for multiple comparisons, although the difference B-C would remain significant even after Holm’s correction (which also provides strong control of the FWER).

320 How can I rank observations within groups in Stata?

The following works for me:

```
bysort group_id: egen desired_rank=rank(var_to_rank)
```

321 Elbow criteria to determine number of cluster



The idea underlying the k-means algorithm is to try to find clusters that minimize the within-cluster variance (or up to a constant the corresponding sum of squares or SS), which amounts to maximize the between-cluster SS because the total variance is fixed. As mentioned on the wiki, you can directly use the within SS and look at its variation when increasing the number of clusters (like we would do in Factor Analysis with a screeplot): an abrupt change in how SS evolve is suggestive of an optimal solution, although this merely stands from visual appreciation. As the total variance is fixed, it is equivalent to study how the ratio of the between and total SS, also called the percentage of variance explained, evolves, because in this case it will present a large gap from one k to the next $k+1$. (Note that the between/within ratio is not distributed as an F-distribution because k is not fixed; so, test are meaningless.)

In sum, you just have to compute the squared distance between each data point and their respective center (or centroid), for each cluster—this gives you the within SS, and the total within SS is just the sum of the cluster-specific WSS (transforming them to variance is just a matter of dividing by the corresponding degrees of freedom); the between SS is obtained by subtracting the total WSS from the total SS, the latter being obtained by considering $k=1$ for example.

By the way, with $k=1$, $WSS=TSS$ and $BSS=0$.

If you’re after determining the number of clusters or where to stop with the k-means, you might consider the Gap statistic as an alternative to the elbow criteria:

Tibshirani, R., Walther, G., and Hastie, T. (2001). **Estimating the numbers of clusters in a data set via the gap statistic.** *J. R. Statist. Soc. B*, 63(2): 411-423.

322 Calculating p-value for a two-way ANOVA

The eta-square (η^2) value you are describing is intended to be used as a measure of *effect size* in the observed data (i.e., your sample), as it amounts to quantify how much of the total variance can be explained by the factor considered in the analysis (that is what you wrote in fact, BSS/TSS). With more than one factor, you can also compute partial η^2 that reflect the percentage of variance explained by one factor when holding constant the remaining ones.

The F-ratio (BSS/WSS) is the right *test statistic* to use if you want to test the null hypothesis (H_0) that there is no effect of your factor (all group means are equal), that is your factor of interest doesn't account for a sufficient amount of variance compared to the residual (unexplained) variance. In other words, we test whether the added explained variance (BSS=TSS-RSS) is large enough to be considered as a "significant quantity". The distribution of the ratio of these two sum of squares (scaled by their corresponding degrees of freedom—this answers one of your question, about why we don't use directly SSs), which individually follow a χ^2 distribution, is known as the **Fisher-Snedecor** distribution.

I don't know which software you are using, but

- If you have R, everything you need for basic modeling is given in the `aoV()` base function (η^2 might be computed with `etasq` from the **heplots** package; and there's a lot more to see for diagnostics and plotting in other packages).
- If you're more versed into C programming, you may have a look at the **apophenia** library which features a nice set of statistical functions with bindings for MySQL and Python.

323 How can I assess how descriptive feature vectors are?

One generally consider that a "good partitioning" must satisfy one or more of the following criteria: (a) *compactness* (small within-cluster variation), *connectedness* (neighbouring data belong to the same cluster), and *spatial separation* (must be combined with other criteria like compactness or balance of cluster sizes). As part of a large battery of internal measures of cluster validity (where

we do not use additional knowledge about the data, like some a priori on class labeling), they can be complemented with so-called *combination measures* (for example, assessing intra-cluster homogeneity and inter-cluster separation), like Dunn or Davies–Bouldin index, silhouette width, SD-validity index, etc., but also estimates of predictive power (self-consistency and stability of a partitioning), how well distance information are reproduced in the resulting partitions (e.g., cophenetic correlation and Hubert's Gamma statistic). A more complete review, and simulation results, are available in

Handl, J., Knowles, J., and Kell, D.B. (2005). **Computational cluster validation in post-genomic data analysis**. *Bioinformatics*, 21(15): 3201-3212.

I guess you could rely on some of them for comparing your different cluster solutions and choose the features set that yields the better indices. You can even use bootstrap to get an estimate of the variability of those indices (e.g., cophenetic correlation, Dunn's index, silhouette width), as was done by Tom Nichols and coll. in a neuroimaging study, **Finding Distinct Genetic Factors that Influence Cortical Thickness**.

If you are using R, I warmly recommend taking a look at the **fpc** package, by **Christian Hennig**, which provides almost all statistical indices described above (`cluster.stats()`) as well as a bootstrap procedure (`clusterboot()`).

About the use of mutual information in clustering, I have no experience with it but here is a paper that discusses its use in a genomic context (with comparison to k-means):

Priness, I., Maimon, O., and Ben-Gal, I. (2007). **Evaluation of gene-expression clustering via mutual information distance measure**. *BMC Bioinformatics*, 8: 111.

324 Effect size of Cochran's Q

I found this paper with Google but I cannot access it, so I don't really know what it is about really:

Berry KJ, Johnston JE, Mielke PW Jr. **An alternative measure of effect size for Cochran's Q test for related proportions**. *Percept Mot Skills*. 2007 Jun;104(3 Pt 2):1236-42.

I initially thought that using pairwise multiple comparisons with Cochran or McNemar test* (if the overall test is significant) would give you further indication of where the differences lie, while reporting simple difference for your binary outcome would help asserting the magnitude of the observed difference.

* I found an [online tutorial with R](#).

325 The effect of the number of replicates in different cells on the results of ANOVA

I don't have Matlab but from what I've read in the on-line help for [N-way analysis of variance](#) it's not clear to me whether Matlab would automatically adapt the [type](#) (1–3) depending on your design. My best guess is that yes you got different results because the tests were not designed in the same way.

Generally, with an imbalanced design it is recommended to use Type III sum of squares (SS), where each term is tested after all other (the difference with Type II sum of squares is only apparent when an interaction term is present), while with an incomplete design it might be interesting to compare Type III and Type IV SS. Note that the use of type III vs. Type II in the case of unbalanced data is subject to discussion in the literature.

(The following is based on a French tutorial that I cannot found anymore on the original website. Here is a [personal copy](#), and here is another paper that discussed the different ways to compute SS in factorial ANOVAs: [Which Sums of Squares Are Best In Unbalanced Analysis of Variance?](#))

The difference between Type I/II and Type III (also called Yates's weighted squares of means) lies in the model that serves as a reference model when computing SS, and whether factors are treated in the order they enter the model or not. Let's say we have two factors, A and B, and their interaction A*B, and a model like $y \sim A + B + A:B$ (Wilkinson's notation).

With Type I SS, we first compute SS associated to A, then B, and finally A*B. Those SS are computed as the difference in residual SS (RSS) between the largest model omitting the term of interest and the smallest one including it.

For Type II and III, SS are computed in a sequan-

tial manner, starting with those associated to AB, then B, and finally A. For AB, it is simply the difference between the RSS in the full model and the RSS in the model without interaction. The SS associated to B is computed as the difference between RSS for a model where B is omitted and a model where B is included (reference model); with Type III SS, the reference model is the full model (A+B+AB), whereas for Type I and II SS, it is the additive model (A+B). This explains why Type II and III will be identical when no interaction is present in the full model. However, to obtain the first SS, we need to use dummy variables to code the levels of the factor, or more precisely difference between those dummy-coded levels (which also means that the reference level considered for a given factor matters; e.g., SAS consider the last level, whereas R consider the first one, in a lexicographic order). To compute SS for the A term, we follow the same idea: we consider the difference between the RSS for the model A+B+AB and that for the reduced model B+A*B (A omitted), in case of Type III SS; with Type II SS, we consider A+B vs. B.

Note that in a complete balanced design, all SS will be equal. Moreover, with Type I SS, the sum of all SS will equal that of the full model, whatever the order of the terms in the model is. (This is not true for Type II and Type III SS.)

A detailed and concrete overview of the different methods is available in one of Howell's handout: [Computing Type I, Type II, and Type III Sums of Squares directly using the general linear model](#). That might help you check your code. You can also use R with the [car](#) package, by John Fox who discussed the use of incremental sum of squares in his textbook, *Applied Regression Analysis, Linear Models, and Related Methods* (Sage Publications, 1997, § 8.2.4–8.2.6). An example of use can be found on [Daniel Wollschläger](#) website.

Finally, the following paper offers a good discussion on the use of Type III SS (§ 5.1):

Venables, W.N. (2000). [Exegeses on Linear Models](#). Paper presented to the S-PLUS User's Conference Washington, DC, 8-9th October, 1998.

(See also this [R-help thread](#), references therein, and the following post [Anova – Type I/II/III SS explained](#).)

326 How to analyse a study looking at relationship between one set of five items (predictors) and a second set of five items (outcomes)

About your first question (using Spearman rank correlation with ordinal scales), I think you will find useful responses on this site (search for spearman, likert, ordinal or scale).

About your second question: As I understand the situation, for each dimension (what you call a “section”), you have a set of five questions scored on a 7-point Likert-type scale. If those five questions all define a single construct, that is if we can consider they form an unidimensional scale (such an assumption might be checked, anyway), why don’t you use a summated scale score (add up the individuals response to the five responses)? This way, your problem would vanish because then you only have one single estimate for the correlation between, say Perceived ease of use and Weblog publishing. Another option is to use [Canonical Correlation Analysis \(CCA\)](#) which allows to build maximally correlated linear combinations of two-block data structure, as described in [this response](#). The pattern of loadings on the two blocks will help you to summarize which item contribute the most information in each block and how they relate each other (under the constraint imposed by CCA). The canonical correlation itself will give you a single number to summarize the association between any two section (again, when considering a linear combination of the 5 questions that compose a section).

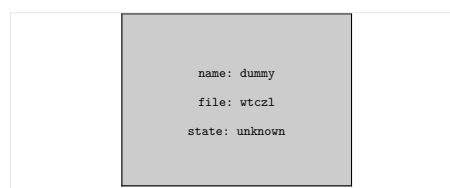
For your third question, I would suggest considering [PLS regression](#), where you define one block of variables as outcomes and the other one as predictors. The idea of PLS regression is to build successive linear combinations of the variables belonging to each block such that their covariance (instead of the correlation like in CCA) is maximal, within a regression approach (because there’s an asymmetric deflation process when constructing the next linear combination). In other words, you build “latent variables” that account for a maximum of information included in the block of predictors while allowing to predict the block of outcomes with minimal error. As you are working with ordinal data, you can even preprocess each variable with optimal scaling if you want, see for example the [homals](#) package in R, and the papers referenced in the documentation.

327 What do you call adding zeros to a table of frequency counts of consecutive integers where the given integer does not occur

Mapping a set of observed value onto the observable values expected for a given variable?

That is, a variable is characterized by all hypothetical values that can be observed when using it, but observed values may not reflect the full range of possible values. For example, when collecting $n=100$ discrete scores on a 0-20 point scale, you might end up with some scores that were observed more often than other, while some never occurred.

And if the number of observable (distinct) value is so small, I would suggest some kind of bar graph (or a dot chart) rather than an histogram, which for a random sample might look like this:



328 What is a propensity weighting sampling / RIM?

You may know that weighting generally aims at ensuring that a given sample is representative of its target population. If in your sample some attributes (e.g., gender, SES, type of medication) are less well represented than in the population from which the sample comes from, then we may adjust the weights of the incriminated statistical units to better reflect the hypothetical target population.

RIM weighting (or raking) means that we will equate the sample marginal distribution to the theoretical marginal distribution. It bears some idea with post-stratification, but allows to account for many covariates. I found a good overview in this handout about [Weighting Methods](#), and here is an example of its use in a real study: [Raking Fire Data](#).

Propensity weighting is used to compensate for unit non-response in a survey, for example, by increas-

ing the sampling weights of the respondents in the sample using estimates of the probabilities that they responded to the survey. This is in spirit the same idea than the use of propensity scores to adjust for treatment selection bias in observational clinical studies: based on external information, we estimate the probability of patients being included in a given treatment group and compute weights based on factors hypothesized to influence treatment selection. Here are some pointers I found to go further:

- The propensity score and estimation in nonrandom surveys - an overview
- A Simulation Study to Compare Weighting Methods for Nonresponses in the National Survey of Recent College Graduates
- A Comparison of Propensity Score and Linear Regression Analysis of Complex Survey Data.

As for a general reference, I would suggest

Kalton G, Flores-Cervantes I. Weighting Methods. J. Off. Stat. (2003) 19: 81-97.
Available on <http://www.jos.nu/>

329 Inspect generators and defining relations of a fractional factorial design

Disclaimer: Not really a positive answer...

Take a look at the **FrF2** package, for example:

```
des.24 <- FrF2(16,8)
design.info(des.24)$aliased # look at the alias structure
```

create a randomized fractional design with 8 factors, 16 runs. To print all designs,

```
print(catlg, nfactor=8, nruns=16)
```

For example, we have design 8-4.1 for a 2^{8-4}_{IV} design, whose generators are $E = ABC$, $F = ABD$, $G = ACD$, and $H = BCD$ (with defining relations in e.g., Montgomery 5ed Appendix X p. 629):

```
summary(FrF2(design="8-4.1"))
FrF2:::generators.from.design(FrF2(design="8-4.1"))
```

Yet I found no way to update the design matrix.

It seems we can update a response vector (there's an example of use with `design()/undesign()`), but AFAIK there's no function that would import a matrix of contrasts and allow to match it in the catalog or find the generators.

P.S. Apart from Minitab and StatGraphics, **DOE++** seems to offer many facilities to work with two-level fractional designs, but I cannot test it unfortunately.

330 Visual representations of the p-value in ANOVA to assist intuitive understanding

Here is a toy example for simulating a one-way ANOVA in R.

First, I just defined a general function that expect an effect size (**es**), which is simply the ratio MSB/MSW (between/within mean squares), a value for the MSB , the number of groups, which might or not be of equal sizes:

```
sim.exp <- function(es=0.25, msb=10, groups=5, n=NULL, verbose=FALSE)
{
  msw <- msb/es
  N <- ifelse(is.null(n), sample(10:40, groups), groups*n)
  means <- rnorm(n=groups, mean=0, sd=sqrt(msb))
  my.df <- data.frame(grp=gl(groups, 1, N),
                     y=rnorm(N, means, sqrt(msw)))
  aov.res <- aov(y ~ grp, my.df)
  if (verbose) print(summary(aov.res))
  ave <- with(my.df, tapply(y, grp, function(x) c(mean(x),
sd(x))))
  invisible(list(ave=ave, p.value=summary(aov.res)[[1]][1,5]))
}
```

This function returns the p-value associated to the F-test, as well as the sample means and SDs. We can use it as follows:

```
> sim.exp(verbose=TRUE)
      Df Sum Sq Mean Sq F value Pr(>F)
grp      4  32.71   8.176  0.1875 0.9418
Residuals 18 784.93  43.607
> sim.exp(es=2, verbose=TRUE)
      Df Sum Sq Mean Sq F value   Pr(>F)
grp      4 555.66 138.915  33.567 1.653e-09 ***
Residuals 24  99.32   4.138
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
> sim.exp(es=.5, n=30, groups=3, verbose=TRUE)
      Df Sum Sq Mean Sq F value    Pr(>F)
grp      2   639.12   319.56  16.498 8.42e-07 ***
Residuals  87 1685.13    19.37
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
for (i in 1:n.sim)
  my.design$p[i] <- mean(unlist(replicate(500,
    sim.exp(my.design[i,1], my.design[i,2],
    n=20))[2,]))
```

and the results are:

Then, I created a grid of values for **es** and **msb**, that is I want to check whether varying these parameters has an effect on the estimated p-value.

```
my.design <- expand.grid(es=seq(.2, 2.4, by=.2), msb=seq(2,
10, by=2))
n.sim <- nrow(my.design)
```

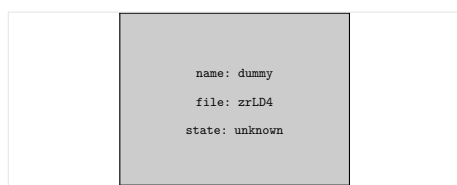
Finally, let's use it. First, with a single replicate of each condition:

```
for (i in 1:n.sim)
  my.design$p[i] <- sim.exp(my.design[i,1], my.design[i,2],
n=20)$p.value
```

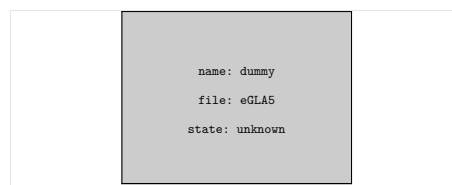
As can be seen, when increasing the effect size we are more likely to reject the null (averaged over MSB):

```
> with(my.design, aggregate(p, list(es=es), mean))
      es      x
1  0.2 1.178042e-01
2  0.4 1.315028e-02
3  0.6 5.765548e-02
4  0.8 5.742882e-02
5  1.0 8.940993e-05
6  1.2 9.199611e-09
7  1.4 9.115640e-06
8  1.6 8.100427e-10
9  1.8 2.656848e-07
10 2.0 3.577391e-05
11 2.2 5.477981e-14
12 2.4 1.219156e-04
```

The results are shown below, although for clarity I took the log of the p-value. The horizontal dashed line shows the 5% limit for type I risk.

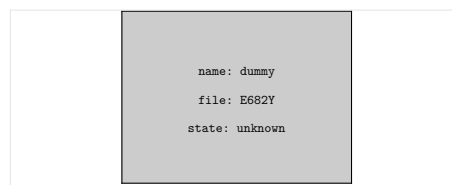


Ok, it's somewhat noisy. So, let's try to average p-values for 500 replicates in each conditions:



We can play with **es** only as follows:

```
k <- 10000
es1 <- sample(seq(.1, 5, by=.1), k, rep=T)
pp <- numeric(k)
for (i in 1:k)
  pp[i] <- sim.exp(groups=3, es=es1[i])$p.value
plot(es1, -log10(pp), pch=19, col="#FF737350", cex=.6, xlab="Effect
size (MSB=10)")
xx <- seq(.1, 5, by=.1)
lines(xx, predict(loess(-log10(pp) ~ es1), data.frame(es1=xx)),
col="green", lwd=2)
```



Many other experiments are possible, and probably a better code too.

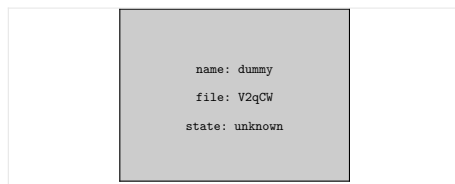
331 Internet statistics resources suitable for psychology students doing research

The **UCLA** server has a lot of resources for statistical computing, including annotated output from various statistical packages.

332 Individuals standard deviations and/or standard errors for groups after implementing ANOVA?

My personal view on this is that

- For *descriptive purpose*, we usually want to show the within-group (i.e., individual) variations (barplot + SD, or better boxplot).
- Within the *inferential context* of the ANOVA, we might rather want to show the SE, 95% CIs, or LSD intervals, for example. Showing 95% CIs has the merit of visually conveying the precision of the estimates, and they are easier to interpret, IMO. In this context, what we really want to show is how good is our estimate of the mean, not so much individual fluctuations on a single sample. Note that the question arises then as to whether we display pooled (when the homoscedasticity assumption holds) or group-specific SEs. We can combine any of the above estimates, of course. E.g., for a one-way ANOVA, we can show 95% CIs associated to each group mean on a barplot and show the overall mean $\pm 1SE$ next to it. The figure below illustrates this idea by showing the SE for an interaction effect, centered on the overall mean (without the 95% CI):



Finally, the following paper offers an interesting discussion of the use of error bars when presenting experimental results (and gauging significant difference from non overlapping error bars):

Cumming, G., Fidler, F., and Vaux, D.L. (2007). **Error bars in experimental biology**. J Cell Biol, 177(1): 7-11.

333 Bayesian two-factor ANOVA

Simon Jackman has some working code for fitting ANOVA and regression models with JAGS (which is pretty like BUGS), e.g. **two-way ANOVA via JAGS** (R code) or maybe among his handouts on **bayesian analysis for the social sciences**.

A lot of WinBUGS code, including one- and two-way ANOVA, seem to be available on the companion website for **Bayesian Modeling Using WinBUGS: An introduction**.

334 Assumptions of cluster analysis

Well, clustering techniques are not limited to *distance-based* methods where we seek groups of statistical units that are unusually close to each other, in a geometrical sense. There're also a range of techniques relying on *density* (clusters are seen as "regions" in the feature space) or *probability distribution*.

The latter case is also known as *model-based clustering*; psychometricians use the term **Latent Profile Analysis** to denote this specific case of **Finite Mixture Model**, where we assume that the population is composed of different unobserved groups, or latent classes, and that the joint density of all manifest variables is a mixture of this class-specific density. Good implementations are available in the **Mclust** package or **Mplus** software. Different class-invariant covariance matrices can be used (in fact, Mclust uses the BIC criterion to select the optimal one while varying the number of clusters).

The standard **Latent Class Model** also makes the assumption that observed data come from a mixture of g multivariate multinomial distributions. A good overview is available in **Model-based cluster analysis: a Defence**, by Gilles Celeux.

Inasmuch these methods rely on distributional assumptions, this also renders possible to use formal tests or goodness-of-fit indices to decide about the number of clusters or classes, which remains a difficult problem in distance-based cluster analysis, but see the following articles that discussed this issue:

1. Handl, J., Knowles, J., and Kell, D.B. (2005). Computational cluster validation in post-genomic data analysis. *Bioinformatics*, 21(15), 3201-3212.
2. Hennig, C. (2007) Cluster-wise assessment of cluster stability. *Computational Statistics and Data Analysis*, 52, 258-271.
3. Hennig, C. (2008) Dissolution point and isolation robustness: robustness criteria for general cluster analysis methods. *Journal of Multivariate Analysis*.

ate Analysis, 99, 1154-1176.

335 Uncorrected pairwise p-values for one-way ANOVA?

You can use `pairwise.t.test()` with one of the available options for multiple comparison correction in the `p.adjust.method=` argument; see `help(p.adjust)` for more information on the available option for single-step and step-down methods (e.g., `BH` for FDR or `bonf` for Bonferroni). Of note, you can directly give `p.adjust()` a vector of raw p-values and it will give you the corrected p-values.

So, I would suggest to run something like

```
pairwise.t.test(time, breed, p.adjust.method = "none") # uncorrected
p-value
pairwise.t.test(time, breed, p.adjust.method = "bonf") # Bonferroni
p-value
```

The first command gives you t-test-based p-values without controlling for FWER or FDR. You can then use whatever command you like to get corrected p-values.

336 Stepwise model selection, Hosmer-Lemeshow statistics and prediction success of model in nested logistic regression in R

Everything is already available in Frank Harrell's `rms` package (which model to choose, how to evaluate its predictive performance or how to validate it, how not to fall into the trap of overfitting or stepwise approach, etc.), with formal discussion in his textbook, *Regression Modeling Strategies* (Springer, 2001), and a nice set of handouts on his [website](#).

Also, I would recommend the following papers if you're interested in predictive modeling:

1. Aliferis, C.F., Statnikov, A., Tsamardinos, I., Schildcrout, J.S., Shepherd, B.E., and Harrell, F.E. Jr (2009). *Factors Influencing the Statistical Power of Complex Data Analysis Protocols for Molecular Signature Development from Microarray Data*. *PLoS ONE* 4(3): e4922.

2. Harrell, F.E. Jr, Margolis, P.A., Gove, S., Mason, K.E., Mulholland, E.K., Lehmann, D., Muhe, L., Gatchalian, S., and Eichenwald, H.F.. (1998). *Development of a clinical prediction model for an ordinal outcome: the World Health Organization Multicentre Study of Clinical Signs and Etiological agents of Pneumonia, Sepsis and Meningitis in Young Infants*. WHO/ARI Young Infant Multicentre Study Group. *Statistics in Medicine*, 17(8): 909-44.
3. Harrell, F.E. Jr, Lee, K.L., and Mark, D.B. (1996). *Multivariable prognostic models: issues in developing models, evaluating assumptions and adequacy, and measuring and reducing errors*. *Statistics in Medicine*, 15(4): 361-87.

337 Importing time series from SQL base into R

Unless I missed something, you want to convert your `data.frame` into a suitable time-indexed series of measurement. In this case, you can use the `zoo` package as follows:

```
> library(zoo)
> memdata.ts <- with(memdata, zoo(vsize, date))
> str(memdata.ts)
'zoo' series from 2011-04-22 to 2011-04-30
Data: Factor w/ 9 levels "3535.178","4403.515",...: 1 9 8
4 2 3 5 6 7
Index: Factor w/ 9 levels "2011-04-22","2011-04-23",...:
1 2 3 4 5 6 7 8 9
```

338 Textbook with list of hypothesis tests and practical guidance on use

Statistical Rules of Thumb (Wiley, 2002), by van Belle, has a lot of useful rules of thumb for applied statistics.

339 Data transposition from 'clustered rows' into columns

340 Load data

Assuming `fd.txt` contains the following

```
N: toto
Y: 2000
S: tata

N: titi
Y: 2004
S: tutu

N: toto
Y: 2000
S: tata2

N: toto
Y: 2000
S: tata3

N: tete
Y: 2002
S: tyty

N: tete
Y: 2002
S: tyty2
```

here is one solution in R:

```
tmp <- scan("fd.txt", what="character")
res <- data.frame(matrix(tmp[seq(2, length(tmp), by=2)], nc=
byrow=TRUE))
```

The first command read everything a single vector of character, skipping blank lines; then we remove every odd elements (“N:”, “S:”, “Y:”); finally, we arrange them in a data.frame (this a convenient way to make each column a factor).

The output is

```
   X1  X2  X3
1 toto 2000 tata
2 titi 2004 tutu
3 toto 2000 tata2
4 toto 2000 tata3
5 tete 2002 tyty
6 tete 2002 tyty2
```

Please note that if you have some GNU utilities on your machine, you can use `awk`

```
sed 's/[NYS]: //' fd.txt | awk 'ORS=(FNR%4)?FS:RS' > res.txt
```

The first command uses `sed` to filter the descriptor (replace by blank); then `awk` will produce its output (Output Record Separator) as follows: arrange each record using default Field Separator (space), and put a new Record Separator (new line) every 4 fields. Of note, we could filter the data using `awk` directly, but I like separating tasks a little.

The result is written in `res.txt` and can be imported into R using `read.table()`:

```
toto 2000 tata
titi 2004 tutu
toto 2000 tata2
toto 2000 tata3
tete 2002 tyty
tete 2002 tyty2
```

341 Process and transform data

I didn’t find a very elegant solution in R, but the following works:

```
library(plyr)
tmp <- ddply(res, .(X1,X2), mutate, S=list(X3))[, -3]
resf <- tmp[!duplicated(tmp[,1:2]),]
```

Then, `resf` has three columns, where column `S` list the levels of the `X3` factor (siblings’ name). So, instead of putting siblings in different columns, I concatenated them in a list. In other words,

```
as.character(resf$S[[1]])
```

gives you the name of `tete`’s siblings, which are `tyty` and `tyty2`.

I’m pretty sure there’s a better way to do this with `plyr`, but I didn’t manage to get a nice solution for the moment.

With repeated “S:” motif, here is one possible quick and dirty solution. Say `fd.txt` now reads

```
N: toto
Y: 2000
S: tata
S: tata2
S: tata3
```

```
Y: 2004
S: tutu
```

```
N: tete
Y: 2002
S: tyty
S: tyty2
```

then,

```
tmp <- read.table("fd.txt")
tmp$V1 <- gsub(":", "", tmp$V1)
start <- c(which(tmp$V1=="N"), nrow(tmp)+1)
max.col <- max(diff(start))
res <- matrix(nr=length(start)-1, nc=max.col)
for (i in 1:(length(start)-1))
  res[i,1:diff(start[i:(i+1)])] <- t(tmp[start[i]:(start[i+1]-1),])
res <- data.frame(res)
colnames(res) <- c("name", "year", paste("S", 1:(max.col-2), sep=""))
```

will produce

```
  name year  S1   S2   S3
1 toto 2000 tata tata2 tata3
2 titi 2004 tutu <NA> <NA>
3 tete 2002 tyty tyty2 <NA>
```

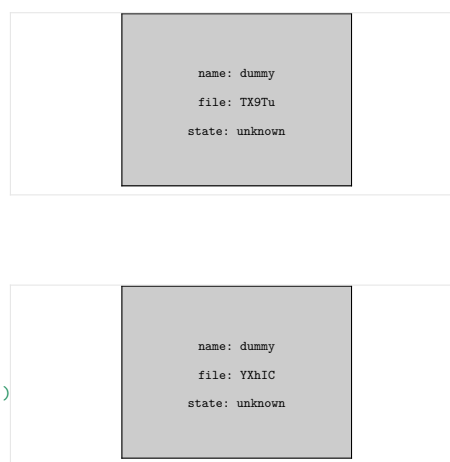
342 Is there a good browser/viewer to see an R dataset (.rda file)

Here are some other thoughts (although I am always reluctant to leave Emacs):

- **Deducer** (with **JGR**) allows to view a data.frame with a combined variable/data view (à la SPSS).
- J Fox's **Rcmdr** also offers edit/viewing facilities, although in an X11 environment.
- J Verzani's Poor Man Gui (**pmg**) only allows for quick preview for data.frame and other R objects. Don't know much about **Rattle** capabilities.

Below are two screenshots when viewing a 704 by 348 data.frame (loaded as an RData) with Deducer (top) and Rcmdr (bottom).

343 Pairwise comparisons after significant interaction results: parametric or non?



If I understand your question correctly, you are wondering why you got different p-values from your t-tests when they are carried out as post-hoc tests or as separate tests. But did you control the **FWER** in the second case (because this is what is done with the step down Sidak-Holm method)? Because, in case of simple t-tests, the t-values won't change, unless you use a different pooling method for computing variance at the denominator, but the p-value of the unprotected tests will be lower than the corrected one.

This is easily seen with Bonferroni adjustment, since we multiply the observed p-value by the number of tests. With step-down methods like **Holm-Sidak**, the idea is rather to sort the null hypothesis tests by increasing p-values and correct the alpha value with Sidak correction factor in a stepwise manner ($\alpha' = 1 - (1 - \alpha)^k$, with k the number of possible comparisons, updated after each step). Note that, in contrast to Bonferroni-Holm's method, control of the FWER is only guaranteed when comparisons are independent. A more detailed description of the different kind of correction for multiple comparisons is available here: **Pairwise Comparisons in SAS and SPSS**.

344 R only alternatives to BUGS

You can take a look at the **MCMCglmm** package

that comes with very nice vignettes. There's also a `bayesglm()` function for fitting Bayesian generalized linear models in the `arm` package, by Andrew Gelman. I've also heard of a [future release](#) `blmer/bglmer` functions for hierarchical modeling in the same package.

345 Is there an R package with a pretty function that can deal effectively with outliers?

If you're importing your data with a command like, say,

```
read.table('yourfile.txt', header=TRUE, ...)
```

you can indicate what values are to be considered as "null" or NA values, by specifying `na.strings = "999999999"`. We can also consider different values for indicating NA values. Consider the following file (`fake.txt`) where we want to treat "." and "999999999" as NA values:

```
1 2 .
3 999999999 4
5 6 7
```

then in R we would do:

```
> a <- read.table("fake.txt", na.strings=c(".", "999999999"))
> a
  V1 V2 V3
1  1  2 NA
2  3 NA  4
3  5  6  7
```

Otherwise, you can always filter your data as indicated by @Sacha in his comment. Here, it could be something like

```
a[a=="." | a==999999999] <- NA
```

Edit

In case there are multiple abnormal values that can possibly be observed in different columns with different values, *but you know the likely range of admissible values*, you can apply a function to each column. For example, define the following filter:

```
my.filter <- function(x, threshold=100) ifelse(x > threshold,
NA, x)
```

then

```
a.filt <- apply(a, 2, my.filter)
```

will replace every value > 100 with NA in the matrix `a`.

Example:

```
> a <- replicate(10, rnorm(10))
> a[1,3] <- 999999999
> a[5,6] <- 999999999
> a[8,10] <- 999999990
> summary(a[,3])
   Min. 1st Qu.  Median    Mean 3rd Qu.    Max.
-1e+00  0e+00  0e+00  1e+07  1e+00  1e+08
> af <- apply(a, 2, my.filter)
> summary(af[,3])
   Min. 1st Qu.  Median    Mean 3rd Qu.    Max.   NA's
-1.4640 -0.2680  0.4671 -0.0418  0.4981  0.7444  1.0000
```

It can be vector-based of course:

```
> summary(my.filter(a[,3], 500))
   Min. 1st Qu.  Median    Mean 3rd Qu.    Max.   NA's
-1.4640 -0.2680  0.4671 -0.0418  0.4981  0.7444  1.0000
```

346 What kind of residuals and Cook's distance are used for GLM?

If you take a look at the code (simple type `plot.lm`, without parenthesis, or `edit(plot.lm)` at the R prompt), you'll see that **Cook's distances** are defined line 44, with the `cooks.distance()` function. To see what it does, type `stats:::cooks.distance.glm` at the R prompt. There you see that it is defined as

```
(res/(1 - hat))^2 * hat/(dispersion * p)
```

where `res` are Pearson residuals (as returned by the `influence()` function), `hat` is the **hat matrix**, `p` is the number of parameters in the model, and `dispersion` is the dispersion considered for the current model (fixed at one for logistic and Poisson regression, see `help(glm)`). In sum, it is computed as a function of the leverage of the observations and their standardized residuals. (Compare with `stats:::cooks.distance.lm`.)

For a more formal reference you can follow references in the `plot.lm()` function, namely

Belsley, D. A., Kuh, E. and Welsch, R. E.

(1980). *Regression Diagnostics*. New York: Wiley.

Moreover, about the additional information displayed in the graphics, we can look further and see that R uses

```
plot(xx, rsp, ...                # line 230
panel(xx, rsp, ...)             # line 233
cl.h <- sqrt(crit * p * (1 - hh)/hh) # line 243
lines(hh, cl.h, lty = 2, col = 2)  #
lines(hh, -cl.h, lty = 2, col = 2) #
```

where `rsp` is labeled as Std. Pearson resid. in case of a GLM, Std. residuals otherwise (line 172); in both cases, however, the formula used by R is (lines 175 and 178)

```
residuals(x, "pearson") / s * sqrt(1 - hii)
```

where `hii` is the hat matrix returned by the generic function `lm.influence()`. This is the usual formula for std. residuals:

$$rs_j = \frac{r_j}{\sqrt{1 - \hat{h}_j}}$$

where j here denotes the j th covariate of interest. See e.g., Agresti *Categorical Data Analysis*, §4.5.5.

The next lines of R code draw a smoother for Cook's distance (`add.smooth=TRUE` in `plot.lm()` by default, see `getOption("add.smooth")`) and contour lines (not visible in your plot) for critical standardized residuals (see the `cook.levels=` option).

347 Recommended procedure for factor analysis on dichotomous data with R

To sum up, with $n=45$ subjects you're left with correlation-based and multivariate descriptive approaches. However, since this questionnaire is supposed to be unidimensional, this always is a good start.

What I would do:

- Compute pairwise correlations for your 22 items; report the range and the median – this will give an indication of the relative consistency of observed items responses (correlations above 0.3 are generally thought of as indicative of good

convergent validity, but of course the precision of this estimate depends on the sample size); an alternative way to study the internal consistency of the questionnaire would be to compute **Cronbach's alpha**, although with $n=45$ the associated confidence interval (use bootstrap for that) will be relatively large.

- Compute **point-biserial correlation** between items and the summated scale score; it will give you an idea of the discriminative power of each item (like loadings in FA), where values above 0.3 are indicative of a satisfactory relationship between each item and their corresponding scale.
- Use a PCA to summarize the correlation matrix (it yields an equivalent interpretation to what would be obtained from a **multiple correspondence analysis** in case of dichotomously scored items). If your instrument behaves as a unidimensional scale for your sample, you should observe a dominant axis of variation (as reflected by the first eigenvalue).

Should you want to use R, you will find useful function in the **ltm** and **psych** package; browse the CRAN **Psychometrics** Task View for more packages. In case you get 100 subjects, you can try some CFA or SEM analysis with bootstrap confidence interval. (Bear in mind that loadings should be very large to consider there's a significant correlation between any item and its factor, since it should be at least two times the standard error of a reliable correlation coefficient, $2(1 - r^2)/\sqrt{(n)}$.)

348 How to extract residuals from function `cv.lm` in R?

Looking at the R code, computation for individual fold are done in the inner loop, starting with

```
for (i in sort(unique(rand))) { # line 37
```

but results are just returned with a `print` statement (line 67-68), if `printit=TRUE` (which is the default). So, you can use what I suggested for a **related question** and edit the function in place so that it returns the SS for each fold in a list. That is, use

```
fix(cv.lm)
```

at the R prompt, then add the following three lines

in the code

```
...
sumss <- 0
sumdf <- 0
ssl <- list()          # (*)
...
  ms <- ss/num
  ssl[[i]] <- ss        # (*)
  if (printit)
    cat("\nSum of squares =", round(ss, 2), "   Mean square = ",
    round(ss/num, 2), "\n")
...
invisible(c(ss = sumss, df = sumdf,
            ss.fold=ssl)) # (*)
}
```

To check that it worked, try

```
> res <- cv.lm(printit=FALSE, plotit=FALSE)
> str(res)
List of 5
 ss      : num 59008
 df      : num 15
 ss.fold1: num 24351
 ss.fold2: num 20416
 ss.fold3: num 14241
```

You can also return a list of the fold SS by replacing `ss.fold=ssl` with `ss.fold=list(ssl)`, so that the output would look like

```
List of 3
 ss      : num 59008
 df      : num 15
 ss.fold:List of 3
 ..$ : num 24351
 ..$ : num 20416
 ..$ : num 14241
```

349 How to tell if data is “clustered” enough for clustering algorithms to produce meaningful results?

About k-means specifically, you can use the Gap statistics. Basically, the idea is to compute a goodness of clustering measure based on average dispersion compared to a reference distribution for an increasing number of clusters. More information can be found in the original paper:

Tibshirani, R., Walther, G., and Hastie, T.

(2001). [Estimating the numbers of clusters in a data set via the gap statistic](#). *J. R. Statist. Soc. B*, 63(2): 411-423.

The answer that I provided to a [related question](#) highlights other general validity indices that might be used to check whether a given dataset exhibits some kind of a structure.

When you don’t have any idea of what you would expect to find if there was noise only, a good approach is to use resampling and study clusters stability. In other words, resample your data (via bootstrap or by adding small noise to it) and compute the “closeness” of the resulting partitions, as measured by [Jaccard](#) similarities. In short, it allows to estimate the frequency with which similar clusters were recovered in the data. This method is readily available in the [fpc](#) R package as `clusterboot()`. It takes as input either raw data or a distance matrix, and allows to apply a wide range of clustering methods (hierarchical, k-means, fuzzy methods). The method is discussed in the linked references:

Hennig, C. (2007) [Cluster-wise assessment of cluster stability](#). *Computational Statistics and Data Analysis*, 52, 258-271.

Hennig, C. (2008) [Dissolution point and isolation robustness: robustness criteria for general cluster analysis methods](#). *Journal of Multivariate Analysis*, 99, 1154-1176.

Below is a small demonstration with the k-means algorithm.

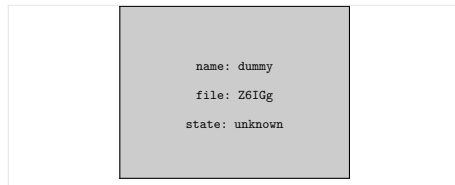
```
sim.xy <- function(n, mean, sd) cbind(rnorm(n, mean[1], sd[1]),
rnorm(n, mean[2], sd[2]))
xy <- rbind(sim.xy(100, c(0,0), c(.2,.2)),
            sim.xy(100, c(2.5,0), c(.4,.2)),
            sim.xy(100, c(1.25,.5), c(.3,.2)))

library(fpc)
km.boot <- clusterboot(xy, B=20, bootmethod="boot",
                      clustermethod=kmeansCBI,
                      krange=3, seed=15555)
```

The results are quite positive in this artificial (and well structured) dataset since none of the three clusters ([krange](#)) were dissolved across the samples, and the average clusterwise Jaccard similarity is > 0.95 for all clusters.

Below are the results on the 20 bootstrap samples. As can be seen, statistical units tend to stay grouped into the same cluster, with few exceptions

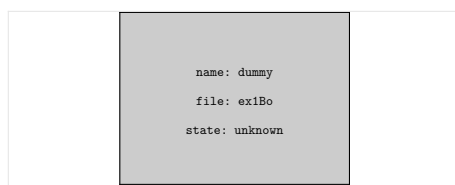
for those observations lying in between.



You can extend this idea to any validity index, of course: choose a new series of observations by bootstrap (with replacement), compute your statistic (e.g., silhouette width, cophenetic correlation, Hubert's gamma, within sum of squares) for a range of cluster numbers (e.g., 2 to 10), repeat 100 or 500 times, and look at the boxplot of your statistic as a function of the number of cluster.

Here is what I get with the same simulated dataset, but using Ward's hierarchical clustering and considering the cophenetic correlation (which assess how well distance information are reproduced in the resulting partitions) and silhouette width (a combination measure assessing intra-cluster homogeneity and inter-cluster separation).

The cophenetic correlation ranges from 0.6267 to 0.7511 with a median value of 0.7031 (500 bootstrap samples). Silhouette width appears to be maximal when we consider 3 clusters (median 0.8408, range 0.7371-0.8769).



350 Testing the importance of an item among a finite set of items

The naive approach would be to compute the marginal distribution of rankings (e.g., mean score for each item), but it would throw away a lot of information as it does not account for the within-person relationship between ranks.

As an extension to **paired preference model** (e.g., the Bradley-Terry model, described in Agresti's CDA pp. 436-439), there exist model for ordinal or likert-type comparison data with or without subject covariates, as well as model for ranking data (bais-

cally, it relies on the use of log-linear model). Here is a **short intro** to the package, and a mathematical explanation in this technical report: **Fitting Paired Comparison Models in R**. You will find everything you need in the **prefmod** R package, see the `pattR.fit()` function which expects data in the form you described:

The responses have to be coded as consecutive integers starting with 1. The value of 1 means highest rank according to the underlying scale. Each column in the data file corresponds to one of the ranked objects. For example, if we have 3 objects denoted by 'A', 'B', and 'C', with corresponding columns in the data matrix, the response pattern '(3,1,2)' represents: object 'B' ranked highest, 'C' ranked second, and 'A' ranked lowest. Missing values are coded as 'NA', ties are not allowed (in that case use 'pattL.fit'). Rows with less than 2 ranked objects are removed from the fit and a message is printed.

For additional information (about and beyond your particular study), you might find useful the following papers:

1. Böckenholt, U. and Dillon, W.R. (1997). Modelling within-subject dependencies in ordinal paired comparison data. *Psychometrika*, 62, p.411-434
2. Dittrich, R., Francis, B., Hatzinger, R., and Katzenbeisser, W. (2006). **Modelling dependency in multivariate paired comparisons: A log-linear approach**. *Mathematical Social Sciences*, 52, 197-209.
3. Maydeu-Olivares, A. (2004). **Thurstone's Case V model: A structural equations modeling perspective**. In K.van Montfort et al. (eds), *Recent Developments on Structural Equation Models*, 41-67.

351 Intraclass correlation coefficient vs. F-test (one-way ANOVA)?

Both methods rely on the same idea, that of decomposing the observed variance into different parts or components. However, there are subtle differences in whether we consider items and/or raters as fixed or random effects. Apart from saying what part of the total variability is explained by the between factor (or how much the between variance departs from the residual variance), the F-test doesn't say much.

At least this holds for a one-way ANOVA where we assume a fixed effect (and which corresponds to the ICC(1,1) described below). On the other hand, the ICC provides a bounded index when assessing rating reliability for several “exchangeable” raters, or homogeneity among analytical units.

We usually make the following distinction between the different kind of ICCs. This follows from the seminal work of Shrout and Fleiss (1979):

- *One-way random effects model*, ICC(1,1): each item is rated by different raters who are considered as sampled from a larger pool of potential raters, hence they are treated as random effects; the ICC is then interpreted as the % of total variance accounted for by subjects/items variance. This is called the consistency ICC.
- *Two-way random effects model*, ICC(2,1): both factors – raters and items/subjects – are viewed as random effects, and we have two variance components (or mean squares) in addition to the residual variance; we further assume that raters assess all items/subjects; the ICC gives in this case the % of variance attributable to raters + items/subjects.
- *Two-way mixed model*, ICC(3,1): contrary to the one-way approach, here raters are considered as fixed effects (no generalization beyond the sample at hand) but items/subjects are treated as random effects; the unit of analysis may be the individual or the average ratings.

This corresponds to cases 1 to 3 in their Table 1. An additional distinction can be made depending on whether we consider that observed ratings are the average of several ratings (they are called ICC(1,k), ICC(2,k), and ICC(3,k)) or not.

In sum, you have to choose the right model (one-way vs. two-way), and this is largely discussed in Shrout and Fleiss’s paper. A one-way model tend to yield smaller values than the two-way model; likewise, a random-effects model generally yields lower values than a fixed-effects model. An ICC derived from a fixed-effects model is considered as a way to assess *raters consistency* (because we ignore rater variance), while for a random-effects model we talk of an estimate of *raters agreement* (whether raters are interchangeable or not). Only the two-way models incorporate the rater x subject interaction, which might be of interest when trying to unravel untypical rating patterns.

The following illustration is readily a copy/paste of the example from `ICC()` in the `psych` package (data come from Shrout and Fleiss, 1979). Data consists in 4 judges (J) assessing 6 subjects or targets (S) and are summarized below (I will assume that it is stored as an R matrix named `sf`)

```

      J1 J2 J3 J4
S1  9  2  5  8
S2  6  1  3  2
S3  8  4  6  8
S4  7  1  2  6
S5 10  5  6  9
S6  6  2  4  7

```

This example is interesting because it shows how the choice of the model might influence the results, therefore the interpretation of the reliability study. All 6 ICC models are as follows (this is Table 4 in Shrout and Fleiss’s paper)

```

Intraclass correlation coefficients
                                type  ICC    F df1 df2      p lower
bound upper bound
Single_raters_absolute  ICC1 0.17  1.8    5 18 0.16477  -
0.133      0.72
Single_random_raters    ICC2 0.29 11.0    5 15 0.00013  0.019
0.76
Single_fixed_raters     ICC3 0.71 11.0    5 15 0.00013  0.342
0.95
Average_raters_absolute ICC1k 0.44  1.8    5 18 0.16477  -
0.884      0.91
Average_random_raters   ICC2k 0.62 11.0    5 15 0.00013  0.071
0.93
Average_fixed_raters    ICC3k 0.91 11.0    5 15 0.00013  0.676
0.99

```

As can be seen, considering raters as fixed effects (hence not trying to generalize to a wider pool of raters) would yield a much higher value for the homogeneity of the measurement. (Similar results could be obtained with the `irr` package (`icc()`), although we must play with the different option for model type and unit of analysis.)

What do the ANOVA approach tell us? We need to fit two models to get the relevant mean squares:

- a one-way model that considers subject only; this allows to separate the targets being rated (between-group MS, BMS) and get an estimate of the within-error term (WMS)
- a two-way model that considers subject + rater + their interaction (when there’s no replications, this last term will be confounded with the resid-

No need to look at the F-test, only MSs are of interest here.

Now, we can assemble the different pieces in an extended ANOVA Table which looks like the one shown below (this is Table 3 in Shrout and Fleiss's paper):

where the first two rows come from the one-way model, whereas the next two ones come from the two-way ANOVA.

- In case of $ICC(1,1)=(BMS-WMS)/(BMS+(k-1) \bullet WMS)$, the overall reliability is computed as $(BMS-WMS)/BMS=0.443$.
- For the $ICC(2,1)=(BMS-EMS)/(BMS+(k-1) \bullet EMS)/N$, the overall reliability is $(N \bullet (BMS-EMS))/(N \bullet BMS + JMS-EMS)=0.620$.
- Finally, for the $ICC(3,1)=(BMS-EMS)/(BMS+(k-1) \bullet EMS)$, we have a reliability of $(BMS-EMS)/B$

352 References

1. Shrout, P.E. and Fleiss, J.L. (1979). **Intraclass correlations: Uses in assessing rater reliability.** *Psychological Bulletin*, 86, 420-3428.
2. Hays, R.D. and Revicki, D. (2005). Reliability and validity (including responsiveness). In Fayers, P. and Hays, R.D. (eds.), *Assessing Quality of Life in Clinical Trials*, 2nd ed., pp. 25-39. Oxford University Press.

353 Assessing reliability of a questionnaire: dimensionality, problematic items, and whether to use alpha, lambda6 or some other index?

The Cronbach's alpha is a sample-dependent index used to ascertain a lower-bound of the reliability of an instrument. It is no more than an indicator of variance shared by all items considered in the computation of a scale score. Therefore, it should not be confused with an absolute measure of reliability, nor does it apply to a multidimensional instrument as a whole. In effect, the following assumptions are made: (a) no residual correlations, (b) items have identical loadings, and (c) the scale is unidimensional. This means that the sole case where alpha will be essentially the same as *reliability* is the case of uniformly high factor loadings, no error covariances, and unidimensional instrument (1). As its precision depends on the standard error of items intercorrelations it depends on the spread of item correlations, which means that alpha will reflect this range of correlations regardless of the source or sources of this particular range (e.g., measurement error or multidimensionality). This point is largely discussed in (2). It is worth noting that when alpha is 0.50, a widely referred reliability threshold for group comparison purpose (3,4), the standard error of measurement will be over half (0.55) a standard deviation. Moreover, Cronbach alpha is a measure of *internal consistency*, it is not a measure of unidimensionality and can't be used to infer unidimensionality (5). Finally, we can quote L.J. Cronbach himself.

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bring to the surface many subtleties implied by variance components. In particular, the interpretations being made in current assessments are best evaluated through use of a standard error of measurement. — Cronbach & Shavelson, (6)

There are many other pitfalls that were largely discussed in several papers in the last 10 years (e.g., 7-10).

Guttman (1945) proposed a series of 6 so-called lambda indices to assess a similar lower bound for reliability, and Guttman's λ_3 lowest bound is strictly equivalent to Cronbach's alpha. If instead of estimating the true variance of each item as the average covariance between items we consider the amount of variance in each item that can be accounted for by the linear regression of all other items (aka, the squared multiple correlation), we get the λ_6 estimate, which might be computed for multi-scale instrument as well. More details can be found in William Revelle's forthcoming textbook, *An introduction to psychometric theory with applications in R* (chapter 7). (He is also the author of the *psych* R package.) You might be interested in reading section 7.2.5 and 7.3, in particular, as it gives an overview of alternative measures, like McDonald's ω_{t} or ω_h (instead of using the squared multiple correlation, we use item uniqueness as determined from an FA model) or Revelle's β (replace FA with hierarchical cluster analysis, for a more general discussion see (12,13)), and provide simulation-based comparison of all indices.

354 References

1. Raykov, T. (1997). Scale reliability, Cronbach's coefficient alpha, and violations of essential tau-equivalence for fixed congeneric components. *Multivariate Behavioral Research*, 32, 329-354.
2. Cortina, J.M. (1993). *What Is Coefficient Alpha? An Examination of Theory and Applications*. *Journal of Applied Psychology*, 78(1), 98-104.
3. Nunnally, J.C. and Bernstein, I.H. (1994). *Psychometric Theory*. McGraw-Hill Series in Psychology, Third edition.
4. De Vaus, D. (2002). *Analyzing social science data*. London: Sage Publications.
5. Danes, J.E. and Mann, O.K.. (1984). Unidimensional measurement and structural equation models with latent variables. *Journal of Business Research*, 12, 337-352.
6. Cronbach, L.J. and Shavelson, R.J. (2004). *My current thoughts on coefficient alpha and successor procedures*. *Educational and Psychological Measurement*, 64(3), 391-418.
7. Schmitt, N. (1996). *Uses and Abuses of Coefficient Alpha*. *Psychological Assessment*, 8(4), 350-353.
8. Iacobucci, D. and Duhachek, A. (2003). *Advancing Alpha: Measuring Reliability With Confidence*. *Journal of Consumer Psychology*, 13(4), 478-487.
9. Shevlin, M., Miles, J.N.V., Davies, M.N.O., and Walker, S. (2000). *Coefficient alpha: a useful indicator of reliability?* *Personality and Individual Differences*, 28, 229-237.
10. Fong, D.Y.T., Ho, S.Y., and Lam, T.H. (2010). *Evaluation of internal reliability in the presence of inconsistent responses*. *Health and Quality of Life Outcomes*, 8, 27.
11. Guttman, L. (1945). A basis for analyzing test-retest reliability. *Psychometrika*, 10(4), 255-282.
12. Zinbarg, R.E., Revelle, W., Yovel, I., and Li, W. (2005). *Cronbach's alpha, Revelle's beta, and McDonald's ω_h : Their relations with each other and two alternative conceptualizations of reliability*. *Psychometrika*, 70(1), 123-133.
13. Revelle, W. and Zinbarg, R.E. (2009) *Coefficients alpha, beta, omega and the glb: comments on Sijtsma*. *Psychometrika*, 74(1), 145-154.

355 Confidence intervals for repeatability

I would go for bootstrap to compute 95% CIs. This is what is generally done with coefficient of heritability or intraclass correlation. (I found no other indication in Falconer's book.) There is an example in the *gap* package of an handmade bootstrap (see *help(h2)*) in case of the correlation-based heritability coefficient, h^2 . IMO, you're better off comput-

ing the variance components yourself, and using the **boot** package. Briefly, the idea is to write a small function that returns your MSs ratio and then call the **boot()** function, e.g.

```
library(boot)
repeat.boot <- function(data, x) { foo(data[x,])$ratio }
res.boot <- boot(yourdata, repeat.boot, 500)
boot.ci(res.boot, type="bca")
```

where **foo(x)** is a function that take a data.frame, compute the variance ratio, and return it as **ratio**.

Sidenote: I just checked on <http://rseek.org> and found this project, **rptR: Repeatability estimation for Gaussian and non-Gaussian data**. I don't know if the above is not simpler.

356 Computing percentile rank in R

Given a vector of raw data values, a simple function might look like

```
perc.rank <- function(x, xo) length(x[x <= xo])/length(x)*100
```

where **x0** is the value for which we want the percentile rank, given the vector **x**, as suggested on **R-bloggers**.

However, it might easily be vectorized as

```
perc.rank <- function(x) trunc(rank(x))/length(x)
```

which has the advantage of not having to pass each value. So, here is an example of use:

```
my.df <- data.frame(x=rnorm(200))
my.df <- within(my.df, xr <- perc.rank(x))
```

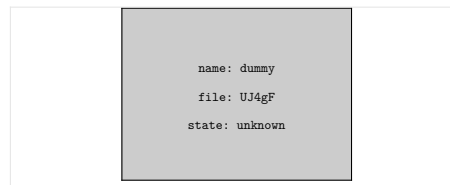
357 Multiple histograms

Without seeing the data, I would suggest trying **trellis displays**. If you are using R, this is very easy to do with the **lattice** (even **latticeExtra**) or **ggplot2** packages.

```
> my.df <- data.frame(x=rnorm(300), year=gl(3, 100, 300, labels=c(2000, 2002, 2004)))
> head(my.df)
      x year
1 -0.3260365 2000
2  0.5524619 2000
```

```
3 -0.6749438 2000
4  0.2143595 2000
5  0.3107692 2000
6  1.1739663 2000
> library(lattice)
> densityplot(~ x, data=my.df, groups=year)
```

which gives



Compare to **densityplot(~ x | year, data=my.df, layout=c(3,1))** (for a faceted display).

358 Explanatory power of a variable

The **relaimpo** R package does exactly what you want to do, and it also provides bootstrap CIs when assessing relative contribution of individual predictor to the overall R^2 .

An example of use can be found at the end of this tutorial: **Getting Started with a Modern Approach to Regression**.

359 Organizing a classification tree (in rpart) into a set of rules?

Such a functionality (or a close one) seems to be available in the **rattle** package, as described in **RJournal 1/2 2009** (p. 50), although I only checked it from the command-line.

For your example, it yields the following output:

```
Rule number: 3 [Kyphosis=present cover=19 (23%) prob=0.58]
  Start< 8.5

Rule number: 23 [Kyphosis=present cover=7 (9%) prob=0.57]
  Start>=8.5
  Age<=2000
  Age<=2002
  Age>=55
  Age< 111
```

```

Rule number: 22 [Kyphosis=absent cover=14 (17%) prob=0.14] 4300]';
Start>=8.5
Start< 14.5
Age>=55
Age>=111

Rule number: 10 [Kyphosis=absent cover=12 (15%) prob=0.00]
Start>=8.5
Start< 14.5
Age< 55

Rule number: 4 [Kyphosis=absent cover=29 (36%) prob=0.00]
Start>=8.5
Start>=14.5

```

```

g = [1 1 1 1 2 2 2 2 3 3 3 3]';
gcat = dummyvar(g);
gcat = gcat(:,2:3); % remove the first column
X = [x gcat x.*gcat(:,1) x.*gcat(:,2)];
n = [48 42 31 34 31 21 23 23 21 16 17 21]';
y = [1 2 0 3 8 8 14 17 19 15 17 21]';
[b, dev, stats] = glmfit(X, [y n], 'binomial', 'link', 'probit');

```

I didn't include a column of ones for the intercept as it is included by default. The design matrix looks like

2100	0	0	0	0
2300	0	0	0	0
2500	0	0	0	0
2700	0	0	0	0
2900	1	0	2900	0
3100	1	0	3100	0
3300	1	0	3300	0
3500	1	0	3500	0
3700	0	1	0	3700
3900	0	1	0	3900
4100	0	1	0	4100
4300	0	1	0	4300

To get this output, I source the `rattle/R/rpart.R` source file (from the source package) in my workspace, after having removed the two calls to `Rtxt()` in the `asRules.rpart()` function (you can also replace it with `print`). Then, I just type

```
> asRules(fit)
```

360 Coding an interaction between a nominal and a continuous predictor for multinomial regression in MATLAB

The easiest way, IMO, is to build the design matrix yourself, as `glmfit` accepts either a matrix of raw (observed) values or a design matrix. Coding an interaction term isn't that much difficult once you wrote the full model. Let's say we have two predictors, x (continuous) and g (categorical, with three unordered levels, say $g = \{1, 2, 3\}$). Using Wilkinson's notation, we would write this model as $y \sim x + g + x:g$, neglecting the left-hand side (for a binomial outcome, we would use a logit link function). We only need two dummy vectors to code the g levels (as present/absent for a particular observation), so we will have 5 regression coefficients, plus an intercept term. This can be summarized as

$$\beta_0 + \beta_1 \cdot x + \beta_2 \cdot \mathbb{I}_{g=2} + \beta_3 \cdot \mathbb{I}_{g=3} + \beta_4 \cdot x \times \mathbb{I}_{g=2} + \beta_5 \cdot x \times \mathbb{I}_{g=3},$$

where \mathbb{I} stands for an indicator matrix coding the level of g .

In Matlab, using the online example, I would do as follows:

```
x = [2100 2300 2500 2700 2900 3100 3300 3500 3700 3900 4100
```

and you can see that the interaction terms are just coded as the product of x with the corresponding column of g ($g=2$ and $g=3$, since we don't need the first level).

The results are given below, as coefficients, standard errors, statistic and p-value (from `stats` structure):

int.	-3.8929	2.0251	-1.9223	0.0546
x	0.0009	0.0008	1.0663	0.2863
g2	-3.2125	2.7622	-1.1630	0.2448
g3	-5.7745	7.5542	-0.7644	0.4446
x:g2	0.0013	0.0010	1.3122	0.1894
x:g3	0.0021	0.0021	0.9882	0.3230

Now, testing the interaction can be done by computing the difference in deviance from the full model above and a reduced model (omitting the interaction term, that is the last two columns of the design matrix). This can be done manually, or using the `lratiotest` function which provides Likelihood ratio hypothesis test. The deviance for the full model is 4.3122 (`dev`), while for the model without interaction it is 6.4200 (I used `glmfit(X(:,1:3), [y n], 'binomial', 'link', 'probit')`), and the associated LR test has two degrees of freedom (the difference in the number of parameters between the two models). As the scaled deviance is just two times the log-likelihood for GLMs, we can use

```
[H, pValue, Ratio, CriticalValue] = lratiotest(4.3122/2, 6.4200/2, 2)
```

where the statistic is distributed as a χ^2 with 2 df (the critical value is then 5.9915, see `chi2inv(0.95, 2)`). The output indicates a non-significant result: We cannot conclude to the existence of an interaction between `x` and `g` in the observed sample.

I guess you can wrap up the above steps in a convenient function of your choice. (Note that the LR test might be done by hand in very few commands!)

I checked those results against R output, which is given next.

Here is the R code:

```
x <- c(2100,2300,2500,2700,2900,3100,3300,3500,3700,3900,4100,4300)
g <- gl(3, 4)
n <- c(48,42,31,34,31,21,23,23,21,16,17,21)
y <- c(1,2,0,3,8,8,14,17,19,15,17,21)
f <- cbind(y, n-y) ~ x*g
model.matrix(f) # will be model.frame() for glm()
m1 <- glm(f, family=binomial("probit"))
summary(m1)
```

Here are the results, for the coefficients in the full model,

Call:

```
glm(formula = f, family = binomial("probit"))
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-1.7124	-0.1192	0.1494	0.3036	0.5585

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	-3.892859	2.025096	-1.922	0.0546
x	0.000884	0.000829	1.066	0.2863
g2	-3.212494	2.762155	-1.163	0.2448
g3	-5.774400	7.553615	-0.764	0.4446
x:g2	0.001335	0.001017	1.312	0.1894
x:g3	0.002061	0.002086	0.988	0.3230

For the comparison of the two nested models, I used the following commands:

```
m0 <- update(m1, . ~ . -x:g)
anova(m1,m0)
```

which yields the following “deviance table”:

```
Model 1: cbind(y, n - y) ~ x + g
Model 2: cbind(y, n - y) ~ x * g
Resid. Df Resid. Dev Df Deviance
1      8      6.4200
2      6      4.3122  2    2.1078
```

361 What is a null conjunction analysis in an fMRI study?

The original paradigm originates from the work of Price and Friston (1997, 1999), and it has been criticized in a more recent paper by Tom Nichols et al. (2005), but see Friston et al. (2005) for a reply.

The idea behind *conjunction analysis* is to determine whether two tasks activate the same region(s) of the brain. Quoting **Towards Evidence of Absence: Conjunction Analyses in fMRI**, contrary to the *subtraction approach* ((a) sum all of the activation maps (AM) from the baseline condition, (b) sum all of the AMs from the stimulation condition, (c) rescale them by converting these summated AMs to average images, (d) subtract the baseline AM from the stimulation AM in order to reveal the activation locations (voxels)), the idea is that

- each voxel should be significantly activated by the two tasks;
- each voxel should not be significantly modulated by an interaction effect between tasks;
- the estimated relationships between each voxel and each task are not significantly different.

The rest of the blog post is worth reading, IMO.

362 References

1. Price, C.J. and Friston, K.J. (1997). **Cognitive conjunction: a new approach to brain activation experiments**. *Neuroimage*, 5, 261-70.
2. Friston, K.J., Holmes, A.P., Price, C.J., Büchel, C., and Worsley, K.J. (1999). Multisubject fMRI Studies and Conjunction Analyses. *NeuroImage*, 10(4), 385-396.
3. Nichols, T., Brett, M., Andersson, J., Wa-

ger, T., Poline, J.-B. (2005). [Valid conjunction inference with the minimum statistic](#). *Neuroimage*, 15;25(3): 653-60.

4. Friston, K.J., Penny, W.D., and Glaser, D.E. (2005). [Conjunction revisited](#). *NeuroImage*, 25, 661-667.

363 Reporting χ^2 test results in APA format

Given the data you posted in your comment, here are the results I get from R:

```
> x <- matrix(c(141,29,43,26,5,10,26,12,10), nc=3)
> x
      [,1] [,2] [,3]
[1,]  141   26   26
[2,]   29    5   12
[3,]   43   10   10
> chisq.test(x)

Pearson's Chi-squared test

data:  x
X-squared = 4.8007, df = 4, p-value = 0.3084
```

So, the χ^2 statistic is actually 4.80. (The expected values are the ones you gave in your comment.)

364 R package for identifying relationships between variables

AFAIK, no. To be more precise, I don't know of a single R package that would do part of what is called *Exploratory Data Analysis* (EDA) for you through a single function call – I'm thinking of the *re-expression* and *revelation* aspects discussed in Hoaglin, Mosteller and Tukey, *Understanding Robust and Exploratory Data Analysis*. Wiley-Interscience, 1983, in particular.

However, there exist some nifty alternatives in R, especially regarding interactive exploration of data (Look here for interesting discussion: [When is interactive data visualization useful to use?](#)). I can think of

- [iplots](#), or its successor [Acynonix](#), for interactive visualization (allowing for brushing, linked plots, and the like) (Some of these functionalities can be found in the [latticist](#) package; finally,

[rgl](#) is great for 3D interactive visualization.)

- [ggobi](#) for interactive and dynamic displays, including data reduction (Multidimensional scaling) and [Projection Pursuit](#)

This is only for interactive data exploration, but I would say this is the essence of EDA. Anyway, the above techniques might help when exploring bivariate or higher-order relationships between numerical variables. For categorical data, the [vcd](#) package is a good option (visualization and summary tables). Then, I would say than the [vegan](#) and [ade4](#) packages come first for exploring relationships between variables of mixed data types.

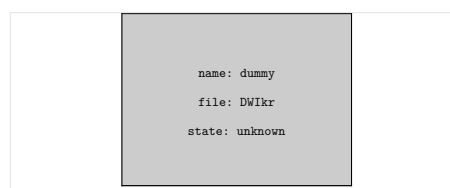
Finally, what about *data mining* in R? (Try this keyword on [Rseek](#))

365 How to best display graphically type II (beta) error, power & sample size

A few thoughts: (a) Use transparency, and (b) Allow for some interactivity.

Here is my take, largely inspired by a Java applet on [Type I and Type II Errors - Making Mistakes in the Justice System](#). As this is rather pure drawing code, I pasted it as [gist #1139310](#).

Here is how it looks:



It relies on the [aplpack](#) package (slider and push button). So, basically, you can vary the deviation from the mean under H_0 (fixed at 0) and the location of the distribution under the alternative. Please note that there's no consideration of sample size.

366 Degrees of freedom for Chi-squared test

How many variables are present in your cross-classification will determine the degrees of freedom

of your χ^2 -test. In your case, you are actually cross-classifying two variables (period and country) in a 2-by-3 table.

So the dof are $(2-1) \times (3-1) = 2$ (see e.g., [Pearson's chi-square test](#) for justification of its computation). I don't see where you got the 6 in your first formula, and your expected frequencies are not correct, unless I misunderstood your dataset.

A quick check in R gives me:

```
> my.tab <- matrix(c(100, 59, 150, 160, 20, 50), nc=3)
> my.tab
      [,1] [,2] [,3]
[1,]  100  150  200
[2,]   59  160   50
> chisq.test(my.tab)

Pearson's Chi-squared test

data:  my.tab
X-squared = 23.7503, df = 2, p-value = 6.961e-06

> chisq.test(my.tab)$expected
      [,1] [,2] [,3]
[1,]  79.6475 155.2876 35.06494
[2,]  79.3525 154.7124 34.93506
```

367 Determining the variables when training a/classifying with a random forest

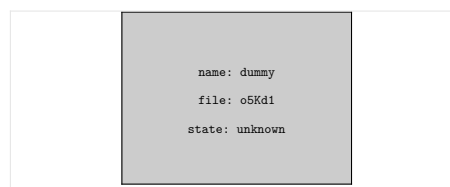
As requested, I'll turn my comment to an answer, although I doubt it will provide you with a working solution.

The Random Forest™ algorithm is well described on [Breiman's webpage](#). See references therein. Basically, it relies on the idea of *bagging* (bootstrap aggregation) where we “average” the results of one (or more) classifier run over different subsamples of the original dataset (1). This is part of what we called more generally *ensemble learning*. The advantage of this approach is that it allows to keep a “training” sample (approx. 2/3 when sampling with replacement) to build the classifier, and a “validation” sample to evaluate its performance; the latter is also known as the out-of-bag (OOB) sample. Classification and regression trees (CART, e.g. (2)) work generally well within this framework, but it is worth noting that in this particular case we do not want to prune the tree (to avoid *overfitting*; on

the contrary, we deliberately choose to work with “weak” classifiers). By constructing several models and combining their individual estimates (most of the times, by a majority vote), we introduce some kind of variety, or fluctuations, in model outputs, while at the same time we circumvent the decision of a single weak classifier by averaging over a collection of such decisions. The [ESLII textbook](#) (3) has a full chapter devoted to such techniques.

However, in RF there is a second level of randomization, this time at the level of the variables: only random subsets of the original variables are considered when growing a tree, typically \sqrt{p} , where p is the number of predictors. This means that RF can work with a lot more variables than other traditional methods, especially those where we expect to have more observations than variables. In addition, this technique will be less impacted by collinearity (correlation between the predictors). Finally, subsampling the predictors has for a consequence that the fitted values across trees are more independent.

As depicted in the next figure, let's consider a block of predictors X ($n \times p$) and a response vector Y , of length n , which stands for the outcome of interest (it might be a binary or numerical variable).



Here, we'll be considering a classification task (Y is categorical). The three main steps are as follows:

1. Take a random sample of the n individuals; they will be used to build a single tree.
2. Select $k < p$ variables and find the first split; *iterate* until the full tree is grown (do not prune); drop the OOB data down the tree and record the outcome assigned to each observation.

Do steps 1–2 b times (say, $b = 500$), and each time count the number of times each individual from the OOB sample is classified into one category or the other. Then assign each case a category by using a majority vote over the b trees. You have your predicted outcomes.

In order to assess *variable importance*, that is how much a variable contributes to classification accu-

racy, we could estimate the extent to which the predictions would be affected if there were no association between the outcome and the given variable. An obvious way to do this is to shuffle the individual labels (this follows from the principle of [re-randomization](#)), and compute the difference in predictive accuracy with the original model. This is step 3. The more predictive a variable is (when trying to predict a future outcome—this really is forecasting, because remember that we only use our OOB sample for assessing the predictive ability of our model), the more likely the classification accuracy will decrease when breaking the link between its observed values and the outcome.

Now, about your question,

- Reliable ways to assess the importance of your predictors have been proposed in the past (this might be the one described above, or additional resampling strategy related to backward elimination (4,5)); so there's no need to reinvent the wheel.
- The hard task, IMO, is to choose your descriptors or features, that is the characteristics of your records. Note that it should not be specific to a subset of your units, but rather encompasses a general description of possible events, e.g. “One Shot Game” vs. other (I'm not very good at game :-).
- Do an extensive literature search on what has been carried out in this direction on this specific field.
- Do not break the control on overfitting that is guaranteed by RF by running additional feature selection + new classification tasks *on the same sample*.

If you're interested in knowing more about RF, there's probably a lot of [additional information](#) on this site, and currently available implementations of RF have been listed [here](#).

368 References

1. Sutton, C.D. (2005). [Classification and Regression Trees, Bagging, and Boosting](#), in *Handbook of Statistics*, Vol. 24, pp. 303-329, Elsevier.

2. Moissen, G.G. (2008). [Classification and Regression Trees](#). *Ecological Informatics*, pp. 582-588.
3. Hastie, T., Tibshirani, R., and Friedman, J. (2009). [The Elements of Statistical Learning: Data Mining, Inference, and Prediction \(2nd ed.\)](#). Springer.
4. Díaz-Uriarte, R., Alvarez de Andrés, S. (2006). [Gene selection and classification of microarray data using Random Forest](#). *BMC Bioinformatics*, 7:3.
5. Díaz-Uriarte, R. (2007). [GeneSrf and varSelRF: a web-based tool and R package for gene selection and classification using Random Forest](#). *BMC Bioinformatics*, 8:328.

369 How to group-center / standardize variables in R?

Here is a possible [plyr](#) solution. Note that it relies on the base [transform\(\)](#) function.

```
my.df <- data.frame(x=rnorm(100, mean=10),
                    sex=sample(c("M","F"), 100, rep=T),
                    group=gl(5, 20, labels=LETTERS[1:5]))

library(plyr)
ddply(my.df, c("sex", "group"), transform, x.std = scale(x))
```

(We can check whether it works as expected with e.g., [with\(subset\(my.df, sex=="F" & group=="A"\), scale\(x\)\)](#))

Basically, the 2nd argument describes how to “split” the data, the 3rd argument what function to apply to each chunk. The above will append a variable `x.std` to the data.frame. Use `x` if you want to replace your original variable by the scaled one.

370 Online calculator for power analysis: what value to give?

For power/sample size analysis, you have to fix either one or the other: You're generally interested in determining the sample size to achieve a given power, or you want to know the power of a test given a certain sample size. In both cases, the type I risk (α) is also fixed at a given value (typically, 5%), and we can accommodate group imbalance, dropouts, etc.

Given the way statistical test of null hypothesis are

framed (definition of a null hypothesis, H_0 , and the alternative, H_1 , yielding the acceptance and rejection regions), the calculator is asking you the expected correlation, ρ , under the alternative.

Now, be aware that computing power “after the fact” (so-called [post-hoc power analysis](#)) is clearly not a definitive solution if you are working with a planned design.

371 Calculate prediction interval for ridge regression?

This has been partly discussed on this [related thread](#). The problem is that this technique introduces bias while trying to decrease the variance of parameter estimates, which works well in situations where multicollinearity does exist. However, the nice properties of the OLS estimators are lost and one has to resort to approximations in order to compute confidence intervals. While I think the bootstrap might offer a good solution to this, here are two references that might be useful:

1. Crivelli, A., Firinguetti, L., Montano, R., and Munoz, M. (1995). Confidence intervals in ridge regression by bootstrapping the dependent variable: a simulation study. *Communications in statistics. Simulation and computation*, 24(3), 631-652.
2. Firinguetti, L. and Bobadilla, G. (2011). [Asymptotic confidence intervals in ridge regression based on the Edgeworth expansion](#). *Statistical Papers*, 52(2), 287-307.

As a good starter to IRT, I always recommend reading [A visual guide to item response theory](#).

The [FreeIRT project](#) features a list of dedicated macros or packages to be used with SAS, Stata, R, among others. Another survey of available software is available on www.rasch.org.

From my experience, I found the [Raschtest](#) (and associated) Stata command(s) very handy in most cases where one is interested in fitting one-parameter model. For more complex design, one can resort on [GLLAMM](#); there’s a nice [working example](#) based on De Boeck and Wilson’s book, *Explanatory Item and Response Models* (Springer, 2004).

About R specifically, there are plenty of packages that have become available in the past five years, see for instance the related [CRAN Task View](#). Most of them are discussed in a [special issue](#) of the *Journal of Statistical Software* (vol. 20, 2007). As discussed in another response, the [ltm](#) and [eRm](#) allow to fit a wide range of IRT models. As they rely on different method of estimation—[ltm](#) used the marginal approach while [eRm](#) use the conditional approach—choosing one or the other is mainly a matter of the model you want to fit ([eRm](#) won’t fit 2- or 3-parameter models) and the measurement objective you follow: conditional estimation of person parameters have some nice psychometric properties while a marginal approach let you easily switch to mixed-effects model, as discussed in the following two papers:

- Doran, H., Bates, D., Bliese, P. and Dowling, M. (2007). [Estimating the Multilevel Rasch Model: With the lme4 Package](#). *Journal of Statistical Software*, 20(2). See also Doug Bates’s [slides on R-forge](#)
- De Boeck, P., Bakker, M., Zwitser, R., Nivard, M., Hofman, A., Tuerlinckx, F., and Partchev, I. (2011). [The Estimation of Item Response Models with the lmer Function from the lme4 Package in R](#). *Journal of Statistical Software*, 39(12). See also the aforementioned De Boeck’s handbook and this [handout](#)

There are also some possibilities to fit Rasch models using MCMC method, see e.g. the [MCMCpack](#) package (or [WinBUGS/JAGS](#), but see [BUGS Code for Item Response Theory](#), JSS (2010) 36).

I have no experience with SAS for IRT modeling, so I’ll let that to someone who is more versed into SAS programming.

Other dedicated software (mostly used in educational assessment) include: RUMM, Conquest, Winsteps, BILOG/MULTILOG, Mplus (not citing the list already available on [wikipedia](#)). None are free to use, but time-limited demonstration version are proposed for some of them. I found [jMetrik](#) very limited when I tried it (one year ago), and all functionalities are already available in R. Likewise, [ConstructMap](#) can be safely replaced by [lme4](#), as illustrated in the [handout](#) linked above. I should also mention [mdlrm](#) (Multidimensional Discrete Latent Trait Models) for mixture Rasch models, by von Davier and coll, which is supposed to accompany the book *Multivariate and Mixture Distribution Rasch Models* (Springer, 2007).

372 How to perform factor and canonical correlation analysis on correlation matrices in R?

For factor analysis, the `psych` package accepts either raw data or a correlation matrix (see e.g., `factor.pa()`). About CCA, I'm not aware of a package that would take correlation matrices as input instead of row data tables.

373 Visualize sets and their connections

I think you may be interested in circular displays for tabular data (in your case, a two-way table denoting the co-occurrence of every binary features), as proposed through `Circos`; see example and on-line demo [here](#).

Sidenote: As an alternative, you can also take a look at `Parallel Sets` that were developed by Robert Kosara. See also,

Robert Kosara, [Turning a Table into a Tree: Growing Parallel Sets into a Purposeful Project](#), in Steele, Iliinsky (eds), *Beautiful Visualization*, pp. 193–204, O'Reilly Media, 2010.

374 How would you explain covariance to someone who understands only the mean?

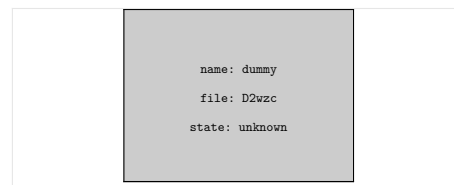
To elaborate on my comment, I used to teach the covariance as a measure of the (average) co-variation between two variables, say x and y .

It is useful to recall the basic formula (simple to explain, no need to talk about mathematical expectancies for an introductory course):

$$\text{cov}(x, y) = \frac{1}{n} \sum_{i=1}^n (x_i - \bar{x})(y_i - \bar{y})$$

so that we clearly see that each observation, (x_i, y_i) , might contribute positively or negatively to the covariance, depending on the product of their deviation from the mean of the two variables, \bar{x} and \bar{y} . Note that I do not speak of magnitude here, but simply of the sign of the contribution of the i th observation.

This is what I've depicted in the following diagrams. Artificial data were generated using a linear model (left, $y = 1.2x + \varepsilon$; right, $y = 0.1x + \varepsilon$, where ε were drawn from a gaussian distribution with zero mean and $\text{SD} = 2$, and x from an uniform distribution on the interval $[0, 20]$).



The vertical and horizontal bars represent the mean of x and y , respectively. That mean that instead of “looking at individual observations” from the origin $(0, 0)$, we can do it from (\bar{x}, \bar{y}) . This just amounts to a translation on the x - and y -axis. In this new coordinate system, every observation that is located in the upper-right or lower-left quadrant contributes positively to the covariance, whereas observations located in the two other quadrants contribute negatively to it. In the first case (left), the covariance equals 30.11 and the distribution in the four quadrants is given below:

```

+   -
+ 30  2
-   0 28

```

Clearly, when the x_i 's are above their mean, so do the corresponding y_i 's (wrt. \bar{y}). Eye-balling the shape of the 2D cloud of points, when x values increase y values tend to increase too. (But remember we could also use the fact that there is a clear relationship between the covariance and the slope of the regression line, i.e. $b = \text{Cov}(x, y) / \text{Var}(x)$.)

In the second case (right, same x_i), the covariance equals 3.54 and the distribution across quadrants is more “homogeneous” as shown below:

```

+   -
+ 18 14
- 12 16

```

In other words, there is an increased number of case where the x_i 's and y_i 's do not covary in the same direction wrt. their means.

Note that we could reduce the covariance by scaling either x or y . In the left panel, the covariance of $(x/10, y)$ (or $(x, y/10)$) is reduced by a ten fold amount (3.01). Since the units of measurement and

the spread of x and y (relative to their means) make it difficult to interpret the value of the covariance in absolute terms, we generally scale both variables by their standard deviations and get the correlation coefficient. This means that in addition to re-centering our (x, y) scatterplot to (\bar{x}, \bar{y}) we also scale the x- and y-unit in terms of standard deviation, which leads to a more interpretable measure of the linear covariation between x and y .

375 How to compute simulated differences of means repetitively?

Let's go for the one-line solution:

```
replicate(1000, mean(rnorm(100, 69.5, 2.9)))-replicate(1000,
mean(rnorm(100, 63.9, 2.7)))
```

376 Predicted by residual plot in R

A plot of residuals versus predicted response is essentially used to spot possible heteroskedasticity (non-constant variance across the range of the predicted values), as well as influential observations (possible outliers). Usually, we expect such plot to exhibit no particular pattern (a funnel-like plot would indicate that variance increase with mean). Plotting residuals against one predictor can be used to check the linearity assumption. Again, we do not expect any systematic structure in this plot, which would otherwise suggest some transformation (of the response variable or the predictor) or the addition of higher-order (e.g., quadratic) terms in the initial model.

More information can be found in any textbook on regression or on-line, e.g. [Graphical Residual Analysis](#) or [Using Plots to Check Model Assumptions](#).

As for the case where you have to deal with multiple predictors, you can use [partial residual plot](#), available in R in the [car](#) ([crPlot](#)) or [faraway](#) ([prplot](#)) package. However, if you are willing to spend some time reading on-line documentation, I highly recommend installing the [rms](#) package and its ecosystem of goodies for regression modeling.

377 Displaying multiple conditional distributions using lattice

Here is a (not so elegant) solution using [lattice](#), where I consider quartiles in case the variables have

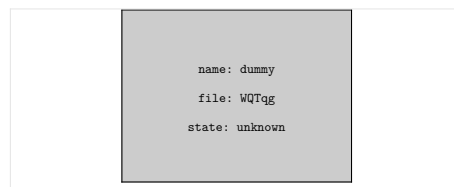
numeric or integer values. Note that I assume that your classification factor is always in the latest position in your dataframe.

```
mydata <- data.frame(age=rnorm(100, 25, 4),
                     sugar=sample(0:10, 100, rep=T),
                     spinach=sample(c("true","false"), 100,
rep=T),
                     meat=sample(c("true","false"), 100, rep=T),
                     milk=sample(c("true","false"), 100, rep=T),
                     class=sample(c("true","false"), 100,
rep=T))

library(lattice)
library(gridExtra)
library(Hmisc)

plt <- list()
for (i in 1:(ncol(mydata)-1)) {
  if (is.numeric(mydata[,i])) vv <- cut2(mydata[,i], g=4)
  else vv <- mydata[,i]
  plt[[i]] <- barchart(xtabs(~ vv + mydata[, "class"]), horizontal=F,
                       main=colnames(mydata)[i],
                       col=c("red", "blue"), xlab="", ylab="",
box.width=1,
                       lattice.options=list(axis.padding=list(factor=0.5)),
                       scales=list(x=list(rot=ifelse(is.numeric(mydata[,i]),45,0))))
}
plt[[i+1]] <- barchart(xtabs(~ class, mydata), col=c("red", "blue"),
                       xlab="", ylab="", box.width=1,
                       lattice.options=list(axis.padding=list(factor=0.5)),
                       horizontal=F, main="class")

do.call(grid.arrange, plt)
```



Using the same dataset with 10 more variables

```
mydata <- data.frame(age=rnorm(100, 25, 4),
                     sugar=sample(0:10, 100, rep=T),
                     spinach=sample(c("true","false"), 100,
rep=T),
                     meat=sample(c("true","false"), 100, rep=T),
                     milk=sample(c("true","false"), 100, rep=T),
                     replicate(10, sample(c("true","false"),
100, rep=T)),
                     class=sample(c("true","false"), 100,
```

```
rep=T))
```

this is the **base** version (you will still need the **Hmisc** library):

```
opar <- par(mfrow=c(4,4))
for (i in 1:15) {
  if (is.numeric(mydata[,i])) vv <- cut2(mydata[,i], g=4)
  else vv <- mydata[,i]
  barplot(xtabs(~ mydata[, "class"] + vv), col=c("red", "blue"),
    main=colnames(mydata)[i],
    las=ifelse(is.numeric(mydata[,i]), 2, 1))
}
barplot(xtabs(~ class, mydata), col=c("red", "blue"), main="class",
  las=ifelse(is.numeric(mydata[,i]), 2, 1))
par(opar)
```

378 What is a good source to learn about multidimensional scaling?

A good textbook on multivariate data analysis, mixing introductory material and more advanced theory, is *Modern Multivariate Statistical Techniques*, by Alan J. Izenman (Springer, 2008). A [review](#) by John Maindonald was published in the JSS.

It features a complete chapter dedicated to MDS (chapter 13), with a lot of illustration using the open-source **R** statistical software. More on R packages can be found on CRAN **Multivariate** Task View, among others.

As an alternative, I would suggest the *Handbook of Applied Multivariate Statistics and Mathematical Modeling*, by Howard E. A. Tinsley and Steven D. Brown (Academic Press, 2000). Again, a complete chapter is devoted to MDS. Less mathematical background is required.

As for online reference, I can also recommend Forrest W. Young's course on [Multidimensional Scaling](#).

379 Tukey-Kramer test

It is called the *LSD Threshold Matrix*, according to the *JMP Statistics and Graphics Guide* (it is headed as **Abs(Dif)-LSD**), and those numbers reflect absolute difference between group means minus Fisher's least significant difference (LSD). They are displayed as off diagonal entries with the great-

est difference in the upper-right or lower-left corner. The diagonal entries represent the comparison of each group mean with itself and are just the opposite of the LSD value. (Note that the matrix is symmetric so you just need to look at the upper or lower diagonal entries.)

Hence, as stated in the documentation, a high positive value would indicate a large departure from the LSD, whereas negative values would mean that the observed difference of means is less than it. See also Ramirez and Ramirez, *Analyzing and Interpreting Continuous Data Using JMP: A Step-by-Step Guide*, SAS Publishing 2009 (pp. 314-316), for more information on how JMP handles the computation of LSD and HSD. Gerard E. Dallal also offers a good overview of [Multiple Comparison Procedures](#).

Note: Given that you only have two groups, a t-test would be sufficient.

380 Can you recommend a book to read before Elements of Statistical Learning?

Introduction to Machine Learning, by E. Alpaydin (MIT Press, 2010, 2nd ed.), covers a lot of topics with nice illustrations (much like Bishop's *Pattern Recognition and Machine Learning*).

In addition, Andrew W. Moore has some nice tutorials on [Statistical Data Mining](#).

381 How to add symbols on smooth lines in a R's lattice xyplot?

Here is a way to achieve this:

```
library(lattice)
library(gridExtra)

my.df <- data.frame(y=c(y1, y2, y3, z1, z2, z3),
  g=gl(6, 50), x=rep(x, 6))

my.panel.loess <- function(x, y, span=2/3, ...) {
  loess.fit <- loess.smooth(x, y, span=span)
  panel.lines(loess.fit$x, loess.fit$y, ...)
}

pp <- list()
pp[[1]] <- xyplot(y ~ x, data=my.df, groups=g, type="b",
```

```
col=rep(c("blue","red"), each=3), cex=.6,
panel=function(...)
  panel.superpose(panel.groups=my.panel.lines,
...))

pp[[2]] <- update(pp[[1]], span=1/5)
pp[[3]] <- update(pp[[1]], col=1:6, type=c("p", "g"))
pp[[4]] <- update(pp[[1]], pch=rep(c(2,6), each=3))

do.call(grid.arrange, pp)
```

You will notice that I had to briefly define a custom panel function for the smoother. This is because I need to pass an extra parameter to `panel.lines` ("`p`"+"`l`"="`b`"); note that it cannot be used directly when calling `panel.lines` since this formal parameter is already defined in `xyplot`.

The rest of the code is pure drawing with varying parameters (all digested through the `...` argument). Although I arranged your data in a more convenient `data.frame`, I believe your original formulation will work with slight adaptation around the call to `panel.groups`.

![[enter image description here]]

However, you can annotate the curves directly with the `directlabels` package (a lot of `examples` are available on R-forge). Something along those lines should work fine:

```
library(directlabels)
pp0 <- xyplot(y ~ x, data=my.df, groups=g, type="smooth")
direct.label(pp0, "follow.points")
```

I should note that there is a nice function, `labcurve()`, in the `Hmisc` package that does pretty much the same job. [1]: <http://i.stack.imgur.com/H9poh.png>

382 Possible extensions to the default diagnostic plots for lm (in R and in general)?

This answer focus on what's available in base R, rather than external packages, although I agree that Fox's package is worth to adopt.

The function `influence()` (or its wrapper, `influence.measures()`) returns most of what we need for model diagnostic, including jackknifed statistics. As stated in Chambers and Hastie's *Statistical Models in S*

(Wadsworth & Brooks, 1992), it can be used in combination to `summary.lm()`. One of the example provided in the so-called "white book" (pp. 130-131) allows to compute standardized (residuals with equal variance) and studentized (the same with a different estimate for SE) residuals, DFBETAS (change in the coefficients scaled by the SE for the regression coefficients), DFFIT (change in the fitted value when observation is dropped), and DFFITS (the same, with unit variance) measures without much difficulty.

Based on your example, and defining the following objects:

```
lms <- summary(fit)
lmi <- influence(fit)
e <- residuals(fit)
s <- lms$sigma
xxi <- diag(lms$cov.unscaled)
si <- lmi$sigma
h <- lmi$hat
bi <- coef(fit) - coef(lmi)
```

we can compute the above quantities as follows:

```
std. residuals  e / (s * (1-h)^.5)
stud. residuals e / (si * (1-h)^.5)
dfbetas        bi / (si %o% xxi^.5)
dffit          h * e / (1-h)
dffits         h^.5 * e / (si * (1-h))
```

(This is *Table 4.1*, p. 131.)

Chambers and Hastie give the following S/R code for computing DFBETAS:

```
dfbetas <- function(fit, lms = summary(fit), lmi = lm.influence(fit))
{
  xxi <- diag(lms$cov.unscaled)
  si <- lmi$sigma
  bi <- coef(fit) - coef(lmi)
  bi / (si %o% xxi^0.5)
}
```

Why do I mention that approach? Because, first, I find this is interesting from a pedagogical perspective (that's what I am using when teaching introductory statistics courses) as it allows to illustrate what can be computed from the output of a fitted linear model fitted in R (but the same would apply with any other statistical package). Second, as the above quantities will be returned as simple vectors or matrices in R, that also means that we can choose the graphics device we want—lattice or ggplot—to

display those statistics, or use them to enhance an existing plot (e.g., highlight DFFITS values in a scatterplot by varying point size `cex`).

383 Functions for regression diagnostics on mer objects in R

Not really about partial residuals plots, but I came across the `influence.ME` package which seems to address the last point, if we consider diagnostic measures to extend to influential observations.

I also found some illustrations that were apparently done in R in the following paper:

Nobre, JS and da Motta Singer, J (2007).
Residual Analysis for Linear Mixed Models.
Biometrical Journal, 49(6), 863–875.

(But check out the following slides, [Residual Analysis for Linear Mixed Models](#), by one of the authors.)

384 Creating a plot with boxplots ranked by quantiles in R

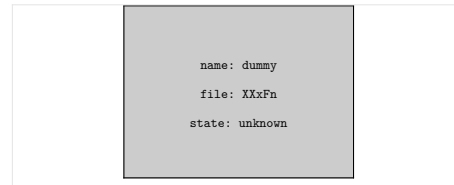
Here is a possible solution using base R graphics:

```
n <- 1000
x <- runif(n, 0, 100)
y <- 1.1*x + rnorm(n)
library(Hmisc)
xq <- cut2(x, g=10, levels.mean=TRUE)
ym <- tapply(y, xq, mean)
# display the mean for each decile
plot(as.numeric(levels(xq)), ym, pch="x", xlab="x", ylab="y")
# add the boxplots
boxplot(y ~ xq, add=TRUE, at=as.numeric(levels(xq)), axes=FALSE)
abline(v=cut2(x, g=10, onlycuts=TRUE))
```

If data are in a `data.frame`, just add a `data=` argument when calling `boxplot()`. You can play with the `boxwex` argument to increase box plots widths. If you prefer to stick on the default `cut()` function, you can probably parse right values of the deciles as in the code below (surely there's a cleaner way to do that!):

```
xq <- cut(x, quantile(x, seq(0, 1, by=.1)))
vx <- gsub("\\(", "", unlist(strsplit(levels(xq), ",")))[seq(1,
18, by=2)]
```

A simple `ggplot` solution might look like this:



```
xy <- data.frame(x=x, y=y)
ggplot(xy, aes(x, y, group=xq)) + geom_boxplot() + xlim(0, 100)
```

I don't know of any package for "decile plots", but I would like to recommend the `bpplt()` and `panel.bpplot()` from the `Hmisc` package. E.g., try this

```
library(lattice)
bwplot(xq ~ y, panel=panel.bpplot, probs=.25, datadensity=TRUE)
```

385 Multiple comparisons with ANOVA including one between and one within-subject effect

The `lme()` function, from the `nlme` package (or see the [companion website](#)), allows to fit mixed-effects models with gaussian errors. For an alternative framework, see this question: [How to choose nlme or lme4 R library for mixed effects models?](#) Your formula reads as follows: your model includes a between- by within-subject interaction assuming **compound symmetry** for the variance-covariance structure; subjects are considered random effects (in other words, this is a varying intercept model, as symbolized by the `~1|ID` formula for the **random** term). The `type="marginal"` argument indicates that you want to compute Type III sum of squares for fixed effects. (See [this](#) or [this](#) question for issues with the different types of SSs, and discussion on the UCLA server on [How can I get Type III tests of fixed effects in R?](#)) This corresponds to what would be obtained using `aoV()` with the appropriate error term (see the corresponding section in [Notes on the use of R for psychology experiments and questionnaires](#)). More details about R's formulas can be found on our [R's lmer cheat-sheet](#).

For more information, you can refer to:

- John Fox's appendix on [Linear Mixed Models](#) (2002).
- A nice guide to [R: Analysis of variance \(ANOVA\)](#), full of illustrations for various experimental de-

signs, which uses the `ez` package.

- Another tutorial on [Multilevel Linear Model](#)
- The [GLMM wiki](#), which summarizes the main issues raised on [r-sig-mixed-models](#) mailing-list.

(And if you have some background in psychology, I'd recommend *Practical Data Analysis for the Language Sciences with R*, by Baayen, which has a lot of illustrations on the analysis of crossed effects in psycholinguistic; there's also a R package, [languageR](#).)

Finally, as you can see a lot has been discussed in related threads so it's worth browsing the `[tag:repeated-measures]` or `[tag:mixed-model]` tags. I'd also like to put a direct link to a related thread that might be of interest: [Why do lme and aov return different results for repeated measures ANOVA in R?](#)

386 Power calculation for a case-control study with a continuous outcome

As I said in my comment, too much information (e.g., expected % of dropouts or missing values, type of matching, one-sided or two-sided hypothesis) is missing to propose a 'magic' formula. So, for general references (apart from classical epidemiological textbook), I would recommend:

- Donner, A. [Approaches to sample size estimation in the design of clinical trials—A review](#), *Statistics in Medicine* (1984) 3:199.
- Edwards, MD. [Sample size requirements for case-control study designs](#), *BMC Medical Research Methodology* (2001) 1:11.

387 Calculating predicted probabilities for ordinal logistic regression

A [similar issue](#) was raised on Stack Overflow more than one year ago. I don't know if re-installing Zelig and its dependencies will solve your problem (especially because I would prefer to understand why this error message came up before reinstalling).

Anyway, you can use the `lrm()` function from the

`rms` package, as it allows to fit several models for categorical outcomes including proportional odds model. There is a `predict()` (but also `Predict()`) function to get the desired predicted values. As an alternative, you may want to look at the `ordinal` package (see the `clm()` function).

388 What programming language do you recommend to prototype a machine learning problem?

The `scikit-learn` (now `sklearn`) should meet several of the criteria you described (speed, well-designed classes for handling data, models, and results), including targeted applications (L1/L2 penalized regression, SVM, etc.). It comes with a rich [documentation set](#) and a lot of [examples](#). See also its description in a [paper](#) published in the JMLR.

An alternative framework in Python is [Orange](#), which can be used through a gentle GUI or on the command line directly. For collaborative filtering, `pyrsvd` might be interesting but I've never tried it. However, [Apache Mahout](#) might certainly be used for [collaborative filtering](#).

389 Obtaining random number from a mixture of two normal distributions

If you want to sample unequally (with probability 0.7 and 0.3) from two gaussians with parameters (μ_1, σ_1^2) and (μ_2, σ_2^2) , then you can probably try something like that:

```
n <- 100
yn <- rbinom(n, 1, .7)
# draw n units from a mixture of N(0,1) and N(100,3^2)
s <- rnorm(n, 0 + 100*yn, 1 + 2*yn)
```

In fact, this is one of the illustrations provided in *Modern Applied Statistics with S*, by Venables and Ripley (Springer, 2002; §5.2, pp. 110-111).

With different parameters, you can use an `ifelse` expression to select the mean and SD according to the binomial sequence given in `yn`, e.g. `rnorm(n, mean=ifelse(yn, 21, 26), sd=ifelse(yn, 3.3, 4))`. (No need to cast `yn` to a logical with `as.logical()`.)

390 Highlighting significant results from non-parametric multiple comparisons on boxplots

The simplest code that comes to my mind is shown below. I'm pretty certain there's some already existing function(s) to do that on CRAN but I'm too lazy to search for them, even on [R-seek][1].

```
dd <- data.frame(y=as.vector(unlist(junk)),
                 g=rep(paste("g", 1:4, sep=""), unlist(lapply(junk,
length))))

aov.res <- kruskal.test(y ~ g, data=dd)
alpha.level <- .05/nlevels(dd$g) # Bonferroni correction,
but use
# whatever you want using
p.adjust()

# generate all pairwise comparisons
idx <- combn(nlevels(dd$g), 2)

# compute p-values from Wilcoxon test for all comparisons
pval.res <- numeric(ncol(idx))
for (i in 1:ncol(idx))
  # test all group, pairwise
  pval.res[i] <- wilcox.test(with(dd, y[as.numeric(g)==idx[1,i]],
                                y[as.numeric(g)==idx[2,i]]))$p.value

# which groups are significantly different (arranged by column)
signif.pairs <- idx[,which(pval.res<alpha.level)]

boxplot(y ~ g, data=dd, ylim=c(min(dd$y)-1, max(dd$y)+1))
# use offset= to increment space between labels, thanks to
vectorization
for (i in 1:ncol(signif.pairs))
  text(signif.pairs[,i], max(dd$y)+1, letters[i], pos=4,
offset=i*.8-1)
```

![enter image description here][2]

Also, be sure to check Rudolf Cardinal's R tips about R: [basic graphs 2](#) (see in particular, *Another bar graph, with annotations*). [1]: <http://www.rseek.org/> [2]: <http://i.stack.imgur.com/Fmg8p.png>

391 Calculating the p-value of an F- statistic

About Scheme libraries specifically, here are two

GSL bindings that you might be interested in:

- Noel Welsh's [fork of mzgsl](#).
- The [Science collection](#), by [Doug Williams](#), provides a collection of modules for numerical computing; it includes [random number distributions](#), among others.

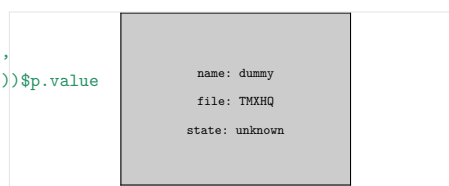
The second project is readily available on [PLaneT](#) if you use [Racket](#).

Here is an example that returns the p -value for the quantile $x = 4.2$ of an $\mathcal{F}(2, 10)$ distribution ($p = 0.047$):

```
(require (planet williams/science/random-distributions/f-
distribution))
(- 1 (f-distribution-cdf 4.2 2 10))
```

with the corresponding CDF

```
(require (planet williams/science/random-distributions/f-
distribution-graphics))
(f-distribution-plot 2 10)
```



There are also some [statistical functions](#) available for [Chicken Scheme](#) (release branch 4). After having installed the required dependencies, e.g.

```
$ sudo chicken-install statistics
```

you will be able to do something like

```
(use statistics)
(f-significance 4.2 2 10 #:one-tailed? #t)
```

in the interactive Chicken shell ([csi](#)). The most commonly used **correlational approach** is called *multi-trait scaling* (MTS). The idea behind MTS is that items belonging to the same dimension are supposed to be more correlated to each other, as well as to their own scale, and that their correlations with items from other dimensions should be zero or at least of a lower magnitude. *Scaling success* then refers to the proportion of items having a larger correlation with their own dimension than with any other dimensions, with higher values indi-

cating good convergent validity. Basically, we compute interitem correlations within each scale and between scales; that is, for each subscale, every item score is correlated with every other item from the same subscale (convergent validity) and from other scales (divergent validity). Such a method is readily available in the `psy` as `mtmm()`; it gives numerical results and graphical output when requested.

Some authors have suggested to examine one-tailed correlation tests, with appropriate correction for multiple comparisons (most of the times, a conservative approach like Bonferroni method). When assessing convergent validity, the usefulness of such tests might be discussed because the null hypothesis that is being considered is simply $H_0 : \rho = 0$ while we generally consider that a correlation of 0.3 (or 0.4) or above is indicative of a ‘meaningful’ correlation for two items belonging to the same subscale. This approach further allows to consider *scaling success with significance tests* as the number of significant correlations (i.e., only non-significant one-tailed correlation tests are counted as scaling errors). Finally, one generally report the average *homogeneity index* (within-scale interitem correlations) and an indicator of internal consistency like Cronbach’s alpha. The above terminology comes from Fayers and Machin (2007). The `psy::mtmm()` function does not perform such tests, but it is not really hard to code.

Another approach relies on factor analytical methods, either exploratory or confirmatory **factor analysis** (CFA). In the CFA framework, the pattern matrix (correlations between items and factor, or loadings) of the hypothesized measurement model is specified in advance, whereas in exploratory factor analysis loadings, communalities and uniquenesses are estimated from the data without imposing constraints. Two critical points that are worth to remember are that (a) if the same sample is used to extract the factors (and construct scoring rules for each subscale) and assess goodness of fit of a factor model, then the generalizability of CFA findings will obviously be limited, and (b) if the model does not fit the data, nothing tell us what could be the correct model (even if so-called modification indices are used to refine the initial model, because in this case it is easy to fall into the trap of data snooping). In R, there are three packages that might be used: `sem`, `lavaan`, `OpenMx`. All three packages provide numerous example of CFA applications.

The following are rough guidelines for assessing model fit: a Comparative Fit (CFI) and Tucker-

Lewis Index (TLI) greater than .90 are indicative of adequate model fit, with values near .95 being preferable; a Standardised Root Mean Square Residual (SRMR) below .10 (resp. .08) and a Root Mean Square Error of Approximation (RMSEA) below .08 (resp. .06) are indicative of acceptable (resp. good) model fit (Hu and Bentler, 1999).

As a sidenote, I would recommend having a look at the `psych` package as well. It features a lot of useful methods for psychometrics (item and reliability analysis, factor-related methods). William Revelle has a book in progress on [applied psychometrics with R](#).

References

1. Fayers, P. M. and Machin, D. (2007). *Quality of Life: The assessment, analysis and interpretation of patient-reported outcomes*. Wiley.
2. Hu, L. T. and Bentler, P. M. (1999). Cutoff criteria for fit indexes in covariance structure analysis: Conventional criteria versus new alternatives. *Structural Equation Modeling*, 6, 1-55.

392 Getting the variance-covariance matrix of regression coefficients in GEE

The naive VC matrix of the parameter estimates is stored in the `vbeta.naiv` component of the GEE fit.

Here is an illustration with the Ohio data:

```
library(geepack)
data(ohio)
fm <- resp ~ age*smoke
gee.fit <- geese(fm, id=id, data=ohio, family=binomial,
                 corstr="exch", scale.fix=TRUE)
summary(gee.fit)
```

This is a basic model where the working correlation is considered symmetric with correlation ρ . Here, the within-cluster correlation is estimated at $\hat{\rho} = 0.355$. Estimates with robust standard error (computed from a sandwich estimator, as detailed in the [JSS paper](#), pp. 4-5) are shown below:

	estimate	san.se	wald	p
(Intercept)	-1.90050	0.11909	254.6860	0.00000
age	-0.14124	0.05820	5.8889	0.01524
smoke	0.31383	0.18784	2.7912	0.09478
age:smoke	0.07083	0.08828	0.6438	0.42234

The robust and naive VC matrices are obtained as follows:

```
> gee.fit$vbeta
      [,1]      [,2]      [,3]      [,4]
[1,] 0.014182 0.002677 -0.014182 -0.002677
[2,] 0.002677 0.003387 -0.002677 -0.003387
[3,] -0.014182 -0.002677 0.035285 0.005348
[4,] -0.002677 -0.003387 0.005348 0.007793
> gee.fit$vbeta.naiv
      [,1]      [,2]      [,3]      [,4]
[1,] 0.01407 0.002400 -0.014072 -0.002400
[2,] 0.00240 0.003139 -0.002400 -0.003139
[3,] -0.01407 -0.002400 0.034991 0.005373
[4,] -0.00240 -0.003139 0.005373 0.007938
```

We can check that the Wald statistics computed using those values (as the ratio of the estimates to their standard errors, which are the diagonal entries of the VC matrix) match the ones displayed in the summary table:

```
> (gee.fit$beta/sqrt(diag(gee.fit$vbeta)))^2
(Intercept)      age      smoke  age:smoke
    254.6860     5.8889     2.7912     0.6438
```

(If you use `geeglm` instead, coefficients are available through the accessor `coef()`, and the robust VC matrix is stored in `gee.fit$geese$vbeta` where `gee.fit` now holds the results of the call to `geeglm`.)

A more detailed account of GEE computing is available in this excellent tutorial, [Generalized Estimating Equations \(GEE\)](#), by Søren Højsgaard and Ulrich Halekoh.

393 Standard formula for quick calculation of scores

There are different options for rescaling your summed scale scores (or **scaled scores**):

- Express every score on a 0-100 points scale, with higher scores reflecting higher locations on the latent trait each scale purports to assess;
- Standardize scores (T - or z -score) such that scores are deviations from the mean, expressed in standard deviation (SD) units. For T -scores, the mean and SD that are considered are 50 and 10, respectively.

(Percentile-based or normalized scores are also com-

mon options. Note that for T -scores, we usually rely on the empirical mean and SD of a larger population who endorsed all items previously (e.g., during large-scale field study) and which might be considered as a “reference population”. Of course, more complex methods exist in the case of grading or **equating** raw scoring gathered throughout different measurement instruments.)

The use of a common scale makes more sense with sum scores (it doesn’t matter much if you consider the mean instead of the sum, which is what social scientists generally prefer compared to psychologists), as @Macro pointed out.

Simple formulae exist in this case (this is just a rescaling problem), but the general idea can be summarized as follows:

```
Scaled score = [(Raw score - Min response category score)
/
                Range of possible response category scores]
* 100
```

to get scores on a 100-point scale. If some items (or response categories) are negatively worded, you will need to reverse-score them first. Subtract the resulting score from 100 to get scores expressed in the reverse direction.

Once each scale score (A, B, C) have been expressed on a common scale, you can use the arithmetic (un-weighted) mean to compute your final score.

394 How can I draw a boxplot without boxes in R?

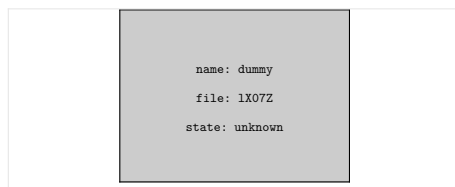
One interesting application of R’s `stripchart()` is that you can use jittering or stacking when there is some overlap in data points (see `method=`). With **lattice**, the corresponding function is `stripplot()`, but it lacks the above method argument to separate coincident points (but see below for one way to achieve stacking).

An alternative way of doing what you want is to use Cleveland’s dotchart. Here are some variations around this idea using **lattice**:

```
my.df <- data.frame(x=sample(rnorm(100), 100, replace=TRUE),
                    g=factor(sample(letters[1:2], 100, replace=TRUE)))
library(lattice)
dotplot(x ~ g, data=my.df) # g on the x-axis
```

```
dotplot(g ~ x, data=my.df, aspect="xy") # g on the y-axis
## add some vertical jittering (use 'factor=' to change its
amount in both case)
dotplot(g ~ x, data=my.df, jitter.y=TRUE)
stripplot(g ~ x, data=my.df, jitter.data=TRUE)
## use stacking (require the 'HH' package)
stripplot(g ~ x, data=my.df, panel=HH::panel.dotplot.tb, factor=
## using a custom sunflowers panel, available through
## http://r.789695.n4.nabble.com/ Grid- graphics- issues-
tp797307p797307.html
stripplot(as.numeric(g) ~ x, data=my.df, panel=panel.sunflowerp
col="black", seg.col="black", seg.lwd=1, size=.08)
## with overlapping data, it is also possible to use transparency
dotplot(g ~ x, data=my.df, aspect=1.5, alpha=.5, pch=19)
```

Some previews of the above commands:



395 What are good datasets to illustrate particular aspects of statistical analysis?

396 The low birth weight study

This is one of the datasets in Hosmer and Lemeshow's textbook on *Applied Logistic Regression* (2000, Wiley, 2nd ed.). The goal of this prospective study was to identify risk factors associated with giving birth to a low birth weight baby (weighing less than 2,500 grams). Data were collected on 189 women, 59 of which had low birth weight babies and 130 of which had normal birth weight babies. Four variables which were thought to be of importance were age, weight of the subject at her last menstrual period, race, and the number of physician visits during the first trimester of pregnancy.

It is available in R as `data(birthwt, package="MASS")` or in Stata with `webuse lbw`. A text version appears here: [lowbwt.dat](#) ([description](#)). Of note, there are several versions of this dataset because it was extended to a case-control study (1-1 or 1-3, matched on age), as illustrated by Hosmer and Lemeshow in ALR chapter 7.

I used to teach introductory courses based on this dataset for the following reasons:

- It is interesting from an historical and epidemiological perspective (data were collected in 1986); no prior background in medicine or statistics is required to understand the main ideas and what questions can be asked from that study.
- Several variables of mixed types (continuous, ordinal, and nominal) are available which makes it easy to present basic association tests (t-test, ANOVA, χ^2 -test for two-way tables, odds-ratio, Cochran and Armitage trend test, etc.). Moreover, birth weight is available as a continuous measure as well as a binary indicator (above or below 2.5 kg): We can start building simple linear models, followed by multiple regression (with predictors of interest selected from prior exploratory analysis), and then switch to GLM (logistic regression), possibly discussing the choice of a cutoff.
- It allows to discuss different modeling perspectives (explanatory or predictive approaches), and the implication of the sampling scheme when developing models (stratification/matched cases).

Other points that can be emphasized, depending on the audience and level of expertise with statistical software, or statistics in general.

1. As for the dataset available in R, categorical predictors are scored as integers (e.g., for mother's ethnicity we have '1' = white, '2' = black, '3' = other), notwithstanding the fact that natural ordering for some predictors (e.g., number of previous premature labors or number of physician visits) or the use of explicit labels (it is always a good idea to use 'yes'/'no' instead of 1/0 for binary variables, even if that doesn't change anything in the design matrix!) are simply absent. As such, it is easy to discuss what issues may be raised by ignoring levels or units of measurement in data analysis.
2. Variables of mixed types are interesting when it comes to do some exploratory analysis and discuss what kind of graphical displays are appropriate for summarizing univariate, bivariate or trivariate relationships. Likewise, producing nice summary tables, and more generally reporting, is another interesting aspect of this dataset (but the `Hmisc::summary.formula` com-

mand makes it so easy under R).

3. Hosmer and Lemeshow reported that actual data were modified to protect subject confidentiality (p. 25). It might be interesting to discuss data confidentiality issues, as was done in one of our earlier [Journal Club](#), but see its [transcript](#). (I must admit I never go into much details with that.)
4. It is easy to introduce some missing values or erroneous values (which are common issues in real life of a statistician), which lead to discuss (a) their detection through codebook (`Hmisc::describe` or Stata's `codebook`) or exploratory graphics (always plot your data first!), and (b) possible remedial (data imputation, list-wise deletion or pairwise measure of association, etc.).

397 Introduction to structural equation modeling

I would go for some papers by Muthén and Muthén, who authored the [Mplus](#) software, especially

1. Muthén, B.O. (1984). [A general structural equation model with dichotomous, ordered categorical and continuous latent indicators](#). *Psychometrika*, 49, 115–132.
2. Muthén, B., du Toit, S.H.C. & Spisic, D. (1997). Robust inference using weighted least squares and quadratic estimating equations in latent variable modeling with categorical and continuous outcomes. Unpublished technical report.

(Available as PDFs from here: [Weighted Least Squares for Categorical Variables](#).)

There is a lot more to see on Mplus wiki, e.g. [WLS vs. WLSMV results with ordinal data](#); the two authors are very responsive and always provide detailed answers with accompanying references when possible. Some comparisons of robust weighted least squares vs. ML-based methods of analyzing polychoric or polyserial correlation matrices can be found in:

Lei, P.W. (2009). [Evaluating estimation methods for ordinal data in structural equation modeling](#). *Quality & Quantity*, 43, 495–507.

For other mathematical development, you can have

a look at:

Jöreskog, K.G. (1994) [On the estimation of polychoric correlations and their asymptotic covariance matrix](#). *Psychometrika*, 59(3), 381–389. (See also [S-Y Lee's](#) papers.)

Sophia Rabe-Hesketh and her colleagues also have good papers on SEM. Some relevant references include:

1. Rabe-Hesketh, S. Skrondal, A., and Pickles, A. (2004b). [Generalized multilevel structural equation modeling](#). *Psychometrika*, 69, 167–190.
2. Skrondal, A. and Rabe-Hesketh, S. (2004). *Generalized Latent Variable Modeling: Multilevel, Longitudinal, and Structural Equation Models*. Chapman & Hall/CRC, Boca Raton, FL. (This is the reference textbook for understanding/working with Stata [gllamm](#).)

Other good resources are probably listed on John Uebersax's excellent website, in particular [Introduction to the Tetrachoric and Polychoric Correlation Coefficients](#).

Given that you are also interested in applied work, I would suggest taking a look at [OpenMx](#) (yet another software package for modeling covariance structure) and [lavaan](#) (which aims at delivering output similar to those of EQS or Mplus), both available under R.

398 How to plot decision boundary of a k-nearest neighbor classifier from Elements of Statistical Learning?

To reproduce this figure, you need to have the [ElemStatLearn](#) package installed on your system. The artificial dataset was generated with `mixture.example()` as pointed out by @StasK.

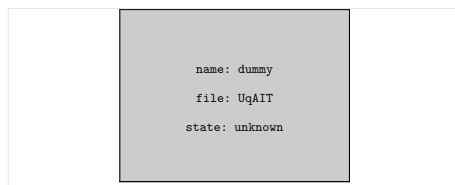
```
library(ElemStatLearn)
require(class)
x <- mixture.example$x
g <- mixture.example$y
xnew <- mixture.example$xnew
mod15 <- knn(x, xnew, g, k=15, prob=TRUE)
prob <- attr(mod15, "prob")
prob <- ifelse(mod15=="1", prob, 1-prob)
px1 <- mixture.example$px1
px2 <- mixture.example$px2
prob15 <- matrix(prob, length(px1), length(px2))
```

```

par(mar=rep(2,4))
contour(px1, px2, prob15, levels=0.5, labels="", xlab="",
ylab="", main=
      "15-nearest neighbour", axes=FALSE)
points(x, col=ifelse(g==1, "coral", "cornflowerblue"))
gd <- expand.grid(x=px1, y=px2)
points(gd, pch=".", cex=1.2, col=ifelse(prob15>0.5, "coral",
"cornflowerblue"))
box()

```

All but the last three commands come from the on-line help for `mixture.example`. Note that we used the fact that `expand.grid` will arrange its output by varying `x` first, which further allows to index (by column) colors in the `prob15` matrix (of dimension 69x99), which holds the proportion of the votes for the winning class for each lattice coordinates (`px1,px2`).



399 Where to find raw data about clinical trials?

You can have a look at the [Clinical Trials Network](#), where data are available in [CDISC](#) format. You must agree to their terms and conditions, although the following point might be of concern for teaching purpose:

To retain control over the received data, and not to transfer any portion of the received data, with or without charge, to any other entity or individual

(Anyway, I think that you can just send an email to the contact support to check that their data can be used for teaching.)

The [NIDKK Data Repository](#) is specifically concerned with studies on kidney and liver disease, and diabetes; however, you have to submit an application.

Otherwise, perhaps the [ADNI](#) project, which aims at characterizing change in cognitive functions and brain structures with age, with a particular emphasis on Alzheimer's disease and neuroimaging, might be interesting. This is not a clinical trial, but avail-

able data include: demographics, clinical and cognitive data, neuroimaging (MRI/PET) data. Details about protocols and data can be found under [ADNI Scientist's Home](#), and data are available on [ADCS](#) website.

There doesn't seem to be anything related to RCTs on <http://www.infochimps.com/>. However, I remember having seen some clinical data used with the [Weka](#) software, as e.g. on this page: [Data mining to predict patient outcome in a clinical trial of a lung cancer treatment](#).

400 Good source of information about AMOVA

There is a nice tutorial on [Analysis of Molecular Variance](#), by Peter Werner, which discusses the use of F - or ϕ -statistics. You may also want to take a closer look at the following paper, for a more complete overview:

Meirmans PG. Using the AMOVA framework to estimate a standardized genetic differentiation measure. *Evolution*. 2006 **60**(11):2399-402.

To add to Peter's response, there's a nice "add-on" to [ade4](#) for genetic data: [adegenet](#), and AMOVA is also available in the [pegas](#) R package.

401 How to convert molecular categorical variables to dummy variables for cluster analysis?

Yes, you can use dummy-coding since the 'representation' of SNP data for statistical analysis depends on the methods used and the underlying genetic model. When using PCA for unravelling population substructure or GWAS for modeling the association between SNPs and one or several phenotypes, each SNP is usually treated as a single integer-coded variable: under the "allelic dosage" model, with values in $\{0,1,2\}$ coding for the frequency of the minor allele; under dominant or recessive effect, with the two extreme categories aggregated yielding a 0/1 response, etc. If you want to use multiple correspondence analysis or a method expecting discrete variables, it would make sense to use dummy coded variables.

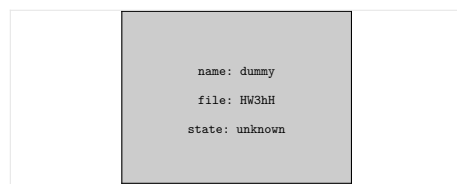
I am aware of two cases where different approaches were retained. Waaijenborg and Zwinderman (1) used *optimal scaling* to transform SNP into one continuous variable as an input into a penalized canonical correlation analysis framework. This allows to consider the three different genotypes (AA, AB, BB) under four different genetic model of inheritance (additive, dominant, recessive or constant). Wolf et al. (2) used dummy-coded SNP as input to Logic Forest, where, for each SNP, the first dummy takes a value of one for 1+ copy of the minor allele (dominant effect) while the second dummy variable takes a value of one if individual was homozygous on the minor allele (recessive effect). In the latter approach, you can use whatever approach you think might best represent the underlying genetic models you want to consider.

401.1 References

1. Waaijenborg and Zwinderman (2009). **Correlating multiple SNPs and multiple disease phenotypes: penalized non-linear canonical correlation analysis**. *Bioinformatics* **25**(21): 2764-2771.
2. Wolf, B.J., Hill, E.G., and Slate, E.H. (2010). **Logic Forest: an ensemble classifier for discovering logical combinations of binary markers**. *Bioinformatics* **26**(17): 2183-2189.

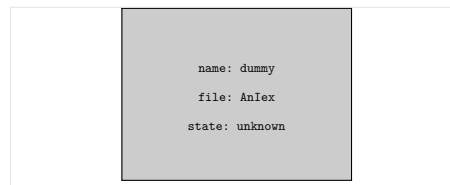
402 2D artificial data of different distributions and forms, existing datasets or generation code

R comes with a lot of datasets, and it looks like it would not be a big deal to reproduce most of the examples you cited with few lines of code. You may also find the **mlbench** package useful, in particular synthetic datasets starting with **mlbench.***. Some illustrations are given below.



You will find additional examples by looking at the **Cluster** Task View on CRAN. For example, the **fpc** package has a built-in generator for “face-shaped”

clustered benchmark datasets (**rFace**).



Similar considerations apply to Python, where you will find interesting benchmark tests and datasets for clustering with the **scikit-learn**.

The UCI Machine Learning Repository hosts a **lot of datasets** as well, but you’re better off simulating data yourself with the language of your choice.

403 Sequential testing

Here is some quick and dirty code for carrying out multiple t-tests over a fake dataset. There may be more elegant way of doing this, but that should let you started with R in a simple way. This solution *assumes* that the 20 samples are independent.

```
# Simulate some data
dfrm <- replicate(20, rnorm(1e6))
colnames(dfrm) <- paste("V", 1:20, sep="")

# Add random signal
inc <- sample(1:20, 3)
dfrm[,inc] <- dfrm[,inc]+.5

# Setup testing framework (each combination of variables)
idx <- combn(20, 2)
pval <- numeric(ncol(idx))
for (i in 1:ncol(idx))
  pval[i] <- t.test(dfrm[,idx[1,i]], dfrm[,idx[2,i]])$p.value

# No. significant test at 5%
sum(pval<.05)

# All variables that were found associated
colnames(dfrm)[unique(as.vector(idx[,pval<.05]))]

# Pretty print as table (well, a data.frame; but it can be
exported
# as LaTeX with the xtable package.)
res <- data.frame(t(idx[,pval<.05]), p=pval[pval<.05])
res$X1 <- colnames(dfrm)[res$X1]
res$X2 <- colnames(dfrm)[res$X2]

# After Bonferroni correction
```

```
res$padj <- p.adjust(res$p, method="bonf")
res[res$padj<.05,]
```

I only reported variable names and associated p-values (with and without correction for multiple tests), but you can add extra information in the `for` loop (e.g., t value, dof, etc.). Please note that R uses Welch's t-test as the default. See `help(t.test)` to use classical Student t-test instead. Sample results look like

```
> res
  X1 X2      p      padj
1 V1 V4 0.000000000 0.00000000
2 V1 V5 0.000000000 0.00000000
3 V1 V7 0.000000000 0.00000000
4 V2 V4 0.000000000 0.00000000
5 V2 V5 0.000000000 0.00000000
6 V2 V7 0.000000000 0.00000000
7 V3 V4 0.000000000 0.00000000
8 V3 V5 0.000000000 0.00000000
9 V3 V6 0.009132565 0.5296888
10 V3 V7 0.000000000 0.00000000
11 V3 V9 0.014075639 0.8163871
```

If all variables are numerical, and data were collected on the same sample, you need a paired t-test (`paired=TRUE`) but why not using simple correlation tests directly? (Simply replace `t.test()` with `cor.test()` in the testing loop.)

404 CCA/KCCA for more than two
views

If by more than two ‘views’ you actually mean extending the CCA framework to k-blocks data structure, then you might be interested in

Tenenhaus, A. and Tenenhaus, M. (2011). Regularized Generalized Canonical Correlation Analysis. *Psychometrika*, 76(2), 257-284.

The corresponding R package is called **RGCCA**.

405 How to apply coefficient term
for factors and interactive terms
in a linear equation?

This is just a comment but it won't fit as such in the limited edit boxes we have at our disposal.

I like seeing a regression equation clearly written in plain text, as @whuber did in his reply. Here is a quick way to this in R, with the **Hmisc** package. (I'll be using **rms** too, but that does not really matter.) Basically, it only assumes that a *L*^AT_EX typesetting system is available on your machine.

Let's simulate some data first,

```
n <- 200
x1 <- runif(n)
x2 <- runif(n)
x3 <- runif(n)
g1 <- gl(2, 100, n, labels=letters[1:2])
g2 <- cut2(runif(n), g=4)
v <- x1 + x2 + rnorm(200)
```

then fit a regression model.

```
f <- ols(y ~ x1 + x2 + x3 + g1 + g2 + x1:g1)
```

which yields the following results:

Linear Regression Model

```
ols(formula = y ~ x1 + x2 + x3 + g1 + g2 + x1:g1)
```

		Model Likelihood		Discrimination	
		Ratio Test		Indexes	
Obs	200	LR chi2	35.22	R2	0.161
sigma	0.9887	d.f.	8	R2 adj	0.126
d.f.	191	Pr(> chi2)	0.0000	g	0.487

Residuals

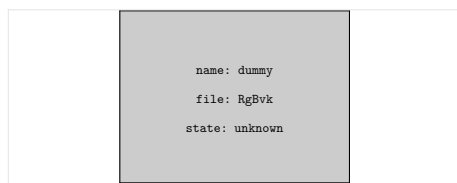
Min	1Q	Median	3Q	Max
-3.1642	-0.7109	0.1015	0.7363	2.7342

	Coef	S.E.	t	Pr(> t)
Intercept	0.0540	0.2932	0.18	0.8541
x1	1.1414	0.3642	3.13	0.0020
x2	0.8546	0.2331	3.67	0.0003
x3	-0.0048	0.2472	-0.02	0.9844
g1=b	0.2099	0.2895	0.73	0.4692
g2=[0.23278,0.553)	0.0609	0.1988	0.31	0.7598
g2=[0.55315,0.777)	-0.2615	0.1987	-1.32	0.1896
g2=[0.77742,0.985]	-0.2107	0.1986	-1.06	0.2901
x1 * g1=b	-0.2354	0.5020	-0.47	0.6396

Then, to print the corresponding regression equation, just use the generic `latex` function, like this:

`latex(f)`

Upon conversion of the dvi to png, you should get something like that



IMO, this has the merit of showing how to compute predicted values depending on actual or chosen values for numerical and categorical predictors. For the latter, factor levels are indicated in bracket near the corresponding coefficient.

406 PLS in R with the pls package

PLS regression relies on iterative algorithms (e.g., NIPALS, SIMPLS). Your description of the main ideas is correct: we seek one (PLS1, one response variable/multiple predictors) or two (PLS2, with different modes, multiple response variables/multiple predictors) vector(s) of weights, u (and v), say, to form linear combination(s) of the original variable(s) such that the covariance between Xu and Y (Yv , for PLS2) is maximal. Let us focus on extracting the first pair of weights associated to the first component. Formally, the criterion to optimize reads

$$\max \text{cov}(Xu, Yv). \quad (1)$$

In your case, Y is univariate, so it amounts to maximize

$$\text{cov}(Xu, y) \equiv \text{Var}(Xu)^{1/2} \times \text{cor}(Xu, y) \times \text{Var}(y)^{1/2},$$

Since $\text{Var}(y)$ does not depend on u , we have to maximise $\text{Var}(Xu)^{1/2} \times \text{cor}(Xu, y)$. Let's consider $X=[x_1; x_2]$, where data are individually standardized (I initially made the mistake of scaling your linear combination instead of x_1 and x_2 separately!), so that $\text{Var}(x_1) = \text{Var}(x_2) = 1$; however, $\text{Var}(Xu) \neq 1$ and depends on u . In conclusion, *maximizing the correlation between the latent component and the response variable will not yield the same results.*

I should thank [Arthur Tenenhaus](#) who pointed me in the right direction.

Using unit weight vectors is not restrictive and some packages (`pls`, `regression` in `pls.genomics`, based on code from Wehrens's earlier package `pls.pcr`) will return unstandardized weight vectors (but with latent components still of norm 1), if requested. But most of PLS packages will return standard-

ized u , including the one you used, notably those implementing the SIMPLS or NIPALS algorithm; I found a good overview of both approaches in Barry M. Wise's presentation, [Properties of Partial Least Squares \(PLS\) Regression, and differences between Algorithms](#), but the `chemometrics` vignette offers a good discussion too (pp. 26-29). Of particular importance as well is the fact that most PLS routines (at least the one I know in R) assume that you provide unstandardized variables because centering and/or scaling is handled internally (this is particularly important when doing cross-validation, for example).

Given the constraint $u'u = 1$, the vector u is found to be

$$u = \frac{X'y}{\|X'y\|}.$$

Using a little simulation, it can be obtained as follows:

```
set.seed(101)
X <- replicate(2, rnorm(100))
y <- 0.6*X[,1] + 0.7*X[,2] + rnorm(100)
X <- apply(X, 2, scale)
y <- scale(y)

# NIPALS (PLS1)
u <- crossprod(X, y)
u <- u/drop(sqrt(crossprod(u))) # X weights
t <- X%*%u
p <- crossprod(X, t)/drop(crossprod(t)) # X loadings
st.||u|| = 1.
```

You can compare the above results ($u=[0.5792043; 0.8151824]$, in particular) with what R packages would give. E.g., using NIPALS from the `chemometrics` package (another implementation that I know is available in the `mixOmics` package), we would obtain:

```
library(chemometrics)
pls1_nipals(X, y, 1)$W # X weights [0.5792043; 0.8151824]
pls1_nipals(X, y, 1)$P # X loadings
```

Similar results would be obtained with `pls` and its default kernel PLS algorithm:

```
> library(pls)
> as.numeric(loadings.weights(plsr(y ~ X, ncomp=1)))
[1] 0.5792043 0.8151824
```

In all cases, we can check that u is of length 1.

Provided you change your function to optimize to one that reads

```
f <- function(u) cov(y, X%*(u/sqrt(crossprod(u))))
```

and normalize `u` afterwards (`u <- u/sqrt(crossprod(u))`) you should be closer to the above solution.

Sidenote: As criterion (1) is equivalent to

$$\max u'X'Yv,$$

u can be found as the left singular vector from the SVD of $X'Y$ corresponding to the largest eigenvalue:

```
svd(crossprod(X, y))$u
```

In the more general case (PLS2), a way to summarize the above is to say that the first PLS canonical vectors are the best approximation of the covariance matrix of X and Y in both directions.

407 References

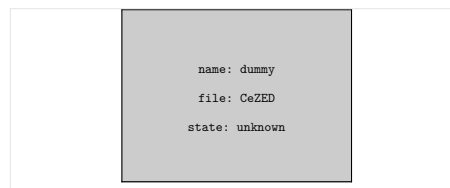
1. Tenenhaus, M (1999). **L'approche PLS**. *Revue de Statistique Appliquée*, 47(2), 5-40.
2. ter Braak, CJF and de Jong, S (1993). **The objective function of partial least squares regression**. *Journal of Chemometrics*, 12, 41–54.
3. Abdi, H (2010). **Partial least squares regression and projection on latent structure regression (PLS Regression)**. *Wiley Interdisciplinary Reviews: Computational Statistics*, 2, 97-106.
4. Boulesteix, A-L and Strimmer, K (2007). **Partial least squares: a versatile tool for the analysis of high-dimensional genomic data**. *Briefings in Bioinformatics*, 8(1), 32-44.

408 How to highlight predefined groups in PCA individual map?

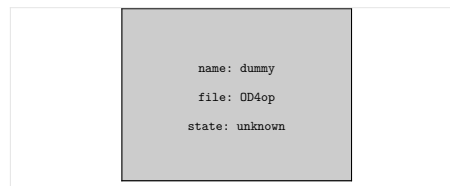
Let me continue my comment with an illustration for the case where you're interested in existing R packages. There are several package in the **Multivariate** Task View that will provide enhanced method for PCA-related methods (as compared to R base `prcomp` and `princomp`), e.g. **ade4** or **FactorMineR**. I personally like FactoMineR because of its simple syntax, and you check the **associated website** for more information on the available methods.

One can use supplementary categorical and/or numerical variables when applying a PCA. Those variables are not used to construct factor axes, but can be showed afterwards on the correlation circle (for numerical variables) or the individual map (for categorical variables). Here is a toy example of use (from the on-line help):

```
data(decathlon)
res.pca <- PCA(decathlon, quanti.sup = 11:12, quali.sup=13)
plotellipses(res.pca,13)
```



If you have multiple passive variables, you can select the one to display (with or without confidence ellipses) using the `keepvar=` argument. Here is another picture with two illustrative variables.



Be careful with arguments that are a little bit non-standard if you are used to default plotting functions in R. The `plotellipses()` function makes use of the helper function `ellipse::ellipse` that you can use (or not) in any plot (look for `monpanel.ellipse` subfunction in `plotellipses()` to see how confidence lines are computed). That's what I did to build specific individual map (B&W, different plotting symbol, etc.). For example, the following snippet just plot all individuals with two different symbols depending on the type of sporting event (2004 Olympic Game or 2004 Decastar):

```
labs <- paste(round(res.pca$eig[1:2, 2]), 2), "%", sep="")
plot(res.pca$ind$coord[,1:2], pch=as.numeric(decathlon$Competition),
      xlab=paste("Dim. 1 (", labs[1], ")", sep=""),
      ylab=paste("Dim. 2 (", labs[2], ")", sep=""))
abline(v=0, h=0, lty=2)
```

Besides, I would like to point you to @vqv's excellent **ggbiplot** package, available on GitHub, which follows from **one of his answer**. (It uses R base functions and `ggplot2`.)

409 A multitrait-multimethod matrix and data set

It looks like I forgot to link to the [original resource](#) I used to construct this picture, that was used as an illustration for an old course (I tend to prefer B&W pictures :-). I know nothing about the data, and that was not of primary interest at the time I used it (it was done with Omnigraffle for Mac).

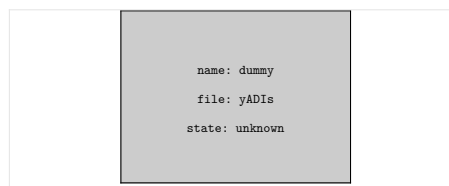
If the question is about how to reach such figures, you can try to generate correlation matrices on your own, using the excellent [psych](#) package. (Be sure to check William Revelle's [website](#).) However, for well-established data you could probably refer to

Brown, TA (2006). *Confirmatory Factor Analysis for Applied Research*. The Guilford Press.

See data for [Table 6.1](#). Some context (pp. 214-216):

In this illustration, the researcher wishes to examine the construct validity of the DSM-IV Cluster A personality disorders, which are enduring patterns of symptoms characterized by odd or eccentric behaviors (American Psychiatric Association, 1994). Cluster A is comprised of three personality disorder constructs: (1) paranoid (an enduring pattern of distrust and suspicion such that others' motives are interpreted as malevolent); (2) schizoid (an enduring pattern of detachment from social relationships and restricted range of emotional expression); and (3) schizotypal (an enduring pattern of acute discomfort in social relationships, cognitive and perceptual distortions, and behavioral eccentricities). In a sample of 500 patients, each of these three traits is measured by three assessment methods: (1) a self-report inventory of personality disorders; (2) dimensional ratings from a structured clinical interview of personality disorders; and (3) observational ratings made by paraprofessional staff. Thus, Table 6.1 is a 3 (T) x 3 (M) matrix, arranged such that the correlations among the different traits (personality disorders: paranoid, schizotypal, schizoid) are nested within each method (assessment type: inventory, clinical interview, observer ratings).

The result should look like this:



If you are using R, you might be interested in looking into the `mtmm()` function from the [psy](#) package (which can be used to assess convergent and discriminant validity within a single measurement instrument as well), as already mentioned in earlier replies of mine: [How to compute correlation between/within groups of variables?](#), [Which package to use for convergent and discriminant validity in R?](#)

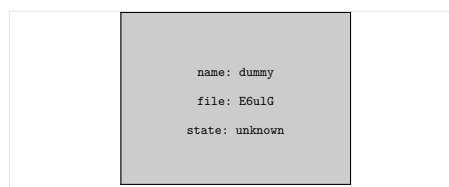
410 Tool to draw normal QQ-plot with 45 degree reference line

Here is one possible Octave solution to your question: (largely inspired from the corresponding Matlab function)

```
randn("state",255)
x = normrnd(10, 2, 200, 1);
[q, s] = qqplot(x);

% compute the y=x line
dx = prctile(q, 75) - prctile(q, 25);
dy = prctile(s, 75) - prctile(s, 25);
b = dy./dx; % slope
xc = (prctile(q, 25) + prctile(q, 75))/2; % center points
yc = (prctile(s, 25) + prctile(s, 75))/2; % ...
ymax = yc + b.*(max(q)-xc);
ymin = yc - b.*(xc-min(q));

plot(q, s, "LineStyle", "none", "Marker", "+")
line([min(q); max(q)], [ymin; ymax])
```



411 How to get an R-squared for a loess fit?

My first thought was to compute a [pseudo \$R^2\$](#) mea-

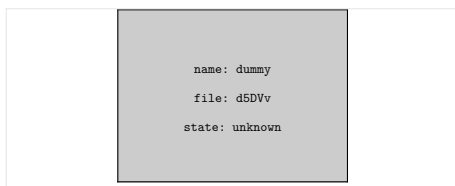
sure as follows:

```
ss.dist <- sum(scale(cars$dist, scale=FALSE)^2)
ss.resid <- sum(resid(cars.lo)^2)
1-ss.resid/ss.dist
```

Here, we get a value of 0.6814984 ($\approx \text{cor}(\text{cars}\$dist, \text{predict}(\text{cars.lo}))^2$), close to what would be obtained from a **GAM**:

```
library(mgcv)
summary(gam(dist ~ speed, data=cars))
```

This also seems to be in agreement with what **S loess** function would return (I don't have S so I can't check by myself) as **Multiple R-squared**. For example, using the **airquality** R dataset, which looks like the **air** data Chambers and Hastie used in the 'white book' (the one that is being referenced in the on-line help for **loess**; but that's not the exact same dataset), I got an R^2 of 0.8101377 using the above formula. That's pretty in agreement with what Chambers and Hastie reported.



I should note that I didn't find any paper dealing specifically with that (ok, that was just a quick googling), and William Cleveland doesn't speak about R^2 -like measure in **his paper**.

However, I wonder if the liberty with which you can choose the degree of smoothing (or window **span**) does not preclude any use of R^2 -based measure.

412 Visualizing Likert responses using R or SPSS

If you really want to use stacked barcharts with such a large number of items, here are two possible solutions.

413 Using **irutils**

I came across this package some months ago.

As of commit 0573195c07 on **GitHub**, the code won't

work with a **grouping=** argument. Let's go for Friday's debugging session.

Start by downloading a zipped version from Github. You'll need to hack the **R/likert.R** file, specifically the **likert** and **plot.likert** functions. First, in **likert**, **cast()** is used but the **reshape** package is never loaded (although there's an **import(reshape)** instruction in the **NAMESPACE** file). You can load this yourself beforehand. Second, there's an incorrect instruction to fetch items labels, where a **i** is dangling around line 175. This has to be fixed as well, e.g. by replacing all occurrences of **likert\$items[,i]** with **likert\$items[,1]**. Then you can install the package the way you are used to do on your machine. On my Mac, I did

```
% tar -czf irutils.tar.gz jbryer-irutils-0573195
% R CMD INSTALL irutils.tar.gz
```

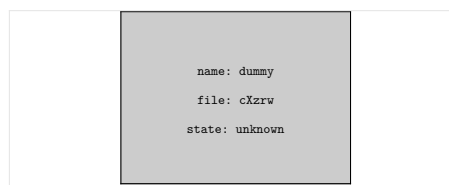
Then, with R, try the following:

```
library(irutils)
library(reshape)

# Simulate some data (82 respondents x 66 items)
resp <- data.frame(replicate(66, sample(1:5, 82, replace=TRUE)))
resp <- data.frame(lapply(resp, factor, ordered=TRUE,
                           levels=1:5,
                           labels=c("Strongly disagree","Disagree",
                                     "Neutral","Agree","Strongly
                                     Agree"))))
grp <- gl(2, 82/2, labels=LETTERS[1:2]) # say equal group
size for simplicity

# Summarize responses by group
resp.likert <- likert(resp, grouping=grp)
```

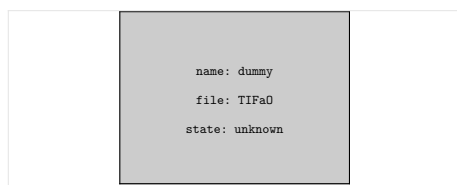
That should just work, but the visual rendering will be awful because of the high number of items. It works without grouping (e.g., **plot(likert(resp))**), though.



I would thus suggest to reduce your dataset to smaller subsets of items. E.g., using 12 items,

```
plot(likert(resp[,1:12], grouping=grp))
```

I get a ‘readable’ stacked barchart. You can probably process them afterwards. (Those are `ggplot2` objects, but you won’t be able to arrange them on a single page with `gridExtra::grid.arrange()` because of readability issue!)



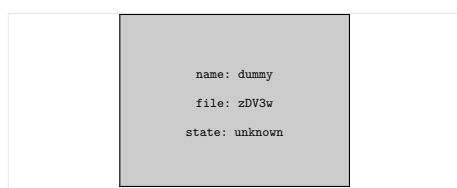
414 Alternative solution

I would like to draw your attention on another package, `HH`, that allows to plot Likert scales as diverging stacked barcharts. We could reuse the above code as shown below:

```
resp.likert <- likert(resp)
detach(package:irutils)
library(HH)
plot.likert(resp.likert$results[, -6]*82/100, main="")
```

but that will complicate things a bit because we need to convert frequencies to counts, subset the `likert` object produced by `irutils`, detach package, etc. So let’s start again with fresh (counts) statistics:

```
plot.likert(t(apply(resp, 2, table)), main="", as.percent=TRUE,
  rightAxisLabels=NULL, rightAxis=NULL, ylab.right="main",
  positive.order=TRUE)
```

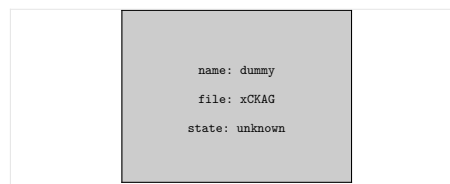


To use a grouping variable, you’ll need to work with an `array` of numerical values.

```
# compute responses frequencies separately by grp
resp.array <- array(NA, dim=c(66, 5, 2))
resp.array[, , 1] <- t(apply(subset(resp, grp=="A"), 2, table))
resp.array[, , 2] <- t(apply(subset(resp, grp=="B"), 2, table))
dimnames(resp.array) <- list(NULL, NULL, group=levels(grp))
plot.likert(resp.array, layout=c(2,1), main="")
```

This will produce two separate panels, but it fits on

a single page.



415 How do I compute class probabilities in caret package using ‘glmnet’ method?

I suspect your `y` is of class `numeric` and is not an R `factor`. You can look at the documentation for `glmnet` directly,

```
y: response variable. Quantitative for ‘family="gaussian"’
or
‘family="poisson"’ (non-negative counts). For
‘family="binomial"’ should be either a **factor with
two
levels, or a two-column matrix of counts or proportions**.
```

(emphasize is mine.)

or check it with the following toy example:

```
library(caret)
data(iris)
iris.sub <- subset(iris, Species %in% c("setosa", "versicolor"))
train(iris.sub[, 1:4], factor(iris.sub$Species), method='glmnet',
  trControl=trainControl(classProbs=TRUE)) # work
train(iris.sub[, 1:4], as.numeric(iris.sub$Species), method='glmnet',
  trControl=trainControl(classProbs=TRUE)) # 'cannot
compute class probabilities for regression'
```

416 Cluster quality measures

The `Statistics` toolbox provides `silhouette plot` which allows to gauge clusters *tightness* and *separation*. (There is also a function to compute cophenetic correlation (to assess how well distance information is reproduced in hierarchical clustering), but it won’t answer your question.)

It seems there is a dedicated `Clustering Toolbox` on Matlab Central, but I have no experience with it. I believe other utilities related to `cluster validity` are within easy reach with Google.

417 Using PCA for feature selection

The basic idea when using PCA as a tool for feature selection is to select variables according to the magnitude (from largest to smallest in absolute values) of their coefficients (*loadings*). You may recall that PCA seeks to replace p (more or less correlated) variables by $k < p$ uncorrelated linear combinations (projections) of the original variables. Let us ignore how to choose an optimal k for the problem at hand. Those k *principal components* are ranked by importance through their explained variance, and each variable contributes with varying degree to each component. Using the largest variance criteria would be akin to *feature extraction*, where principal component are used as new features, instead of the original variables. However, we can decide to keep only the first component and select the $j < p$ variables that have the highest absolute coefficient; the number j might be based on the proportion of the number of variables (e.g., keep only the top 10% of the p variables), or a fixed cutoff (e.g., considering a threshold on the normalized coefficients). This approach bears some resemblance with the **Lasso** operator in penalized regression (or **PLS** regression). Neither the value of j , nor the number of components to retain are obvious choices, though.

The problem with using PCA is that (1) measurements from all of the original variables are used in the projection to the lower dimensional space, (2) only linear relationships are considered, and (3) PCA or SVD-based methods, as well as univariate screening methods (t-test, correlation, etc.), do not take into account the potential multivariate nature of the data structure (e.g., higher order interaction between variables).

About point 1, some more elaborate screening methods have been proposed, for example **principal feature analysis** or stepwise method, like the one used for ‘**gene shaving**’ in gene expression studies. Also, **sparse PCA** might be used to perform dimension reduction and variable selection based on the resulting variable loadings. About point 2, it is possible to use kernel PCA (using the **kernel trick**) if one needs to embed nonlinear relationships into a lower dimensional space. **Decision trees**, or better the **random forest** algorithm, are probably better able to solve Point 3. The latter allows to derive Gini- or permutation-based measures of **variable importance**.

A last point: If you intend to perform feature se-

lection before applying a classification or regression model, be sure to cross-validate the whole process (see §7.10.2 of the **Elements of Statistical Learning**, or **Ambroise and McLachlan, 2002**).

As you seem to be interested in R solution, I would recommend taking a look at the **caret** package which includes a lot of handy functions for data preprocessing and variable selection in a classification or regression context.

418 Firth logistic regression in R

You can probably compute any predictions you want with little algebra. Let consider the example dataset,

```
data(sex2)
fm <- case ~ age+oc+vic+vic1+vis+dia
fit <- logistf(fm, data=sex2)
```

A design matrix is the only missing piece to compute predicted probabilities once we get the regression coefficients, given by

```
betas <- coef(fit)
```

So, let’s try to get prediction for the observed data, first:

```
X <- model.matrix(fm, data=sex2) # add a column of 1's
to sex2[, -1]
pi.obs <- 1 / (1 + exp(-X %*% betas)) # in case there's an
offset,  $\delta$ , it
# should be subtracted
as exp(-X $\beta$  -  $\delta$ )
```

We can check that we get the correct result

```
> pi.obs[1:5]
[1] 0.3389307 0.9159945 0.9159945 0.9159945 0.9159945
> fit$predict[1:5]
[1] 0.3389307 0.9159945 0.9159945 0.9159945 0.9159945
```

Now, you can put in the above design matrix, **X**, values you are interested in. For example, with all covariates set to one

```
new.x <- c(1, rep(1, 6))
1 / (1 + exp(-new.x %*% betas))
```

we get an individual probability of 0.804, while

when all covariates are set to 0 (`new.x <- c(1, rep(0, 6))`), the estimated probability is 0.530.

419 How to display multiple density or distribution functions on a single plot?

I like R too. Here is a more or less generic function to plot any probability distribution from the **base R functions**. It should not be difficult to extend the code with functions available in other packages, e.g. **SuppDists**.

```
plot.func <- function(distr=c("beta", "binom", "cauchy", "chisq",
                             "exp", "f", "gamma", "geom",
                             "hyper",
                             "logis", "lnorm", "nbinom",
                             "norm",
                             "pois", "t", "unif", "weibull"),
                      what=c("pdf", "cdf"), params=list(),
                      type="b",
                      xlim=c(0, 1), log=FALSE, n=101, add=FALSE,
                      ...) {
  what <- match.arg(what)
  d <- match.fun(paste(switch(what, pdf = "d", cdf = "p"),
                      distr, sep=""))
  # Define x-values (because we won't use 'curve') as last
  parameter
  # (with pdf, it should be 'x', while for cdf it is 'q').
  len <- length(params)
  params[[len+1]] <- seq(xlim[1], xlim[2], length=n)
  if (add) lines(params[[len+1]], do.call(d, params), type,
  ...)
  else plot(params[[len+1]], do.call(d, params), type, ...)
}
```

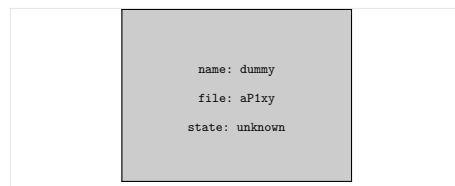
It's a bit crappy and I haven't tested it a lot. The `params` list must obey R's conventions for naming {C|P}DF parameters (e.g., `shape` and `scale` for the Weibull distribution, and not `a` or `b`). There's room for improvement, especially about the way it handles multiple plotting on the same graphic device (and, actually, passing vector of parameters only works as a side-effect when `type="p"`). Also, there's not much parameter checking!

Here are some examples of use:

```
# Normal CDF
xl <- c(-5, 5)
plot.func("norm", what="pdf", params=list(mean=1, sd=1.2),
          xlim=xl, ylim=c(0,.5), cex=.8, type="l", xlab="x",
          ylab="F(x)")
```

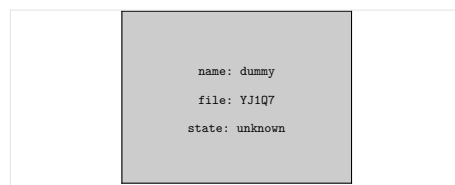
```
plot.func("norm", what="pdf", params=list(mean=3, sd=.8),
          xlim=xl, add=TRUE, pch=19, cex=.8)
plot.func("norm", what="pdf", params=list(mean=.5, sd=1.3),
          n=201,
          xlim=xl, add=TRUE, pch=19, cex=.4, type="p", col="steelblue")
title(main="Some gaussian PDFs")

# Standard normal PDF
plot.func("norm", "cdf", xlab="Quantile (x)", ylab="P(X<x)",
          xlim=c(-3,3), type="l",
          main="Some gaussian CDFs")
plot.func("norm", "cdf", list(sd=c(0.5,1.5)), xlim=c(-3,3),
          add=TRUE,
          type="p", pch=c("o", "+"), n=201, cex=.8)
legend("topleft", paste("N(0;", c(1,0.5,1.5), ")"), sep=""),
      col=1:3, pch=19)
```



420 Logistic regression: grouped and ungrouped variables (using R)

Table 3.1 is reproduced below:



Agresti considered the following numerical scores for snoring level: {0,2,4,5}.

There are two ways to fit a GLM with R: either your outcome is provided as a vector of 0/1 or a factor with two levels, with the predictors on the rhs of

your formula; or you can give a matrix with two columns of counts for success/failure as the lhs of the formula. The latter corresponds to what Agresti call ‘grouped’ data.

Data in matrix view would read:

```
snoring <- matrix(c(24,35,21,30,1355,603,192,224), nc=2)
```

From this, we can generate a `data.frame` in long format (2484 rows = `sum(snoring)` observations) as follows:

```
snoring.df <- data.frame(snoring=gl(4, 1, labels=c("Never",
"Occasional",
"Nearly
every night",
"Every
night")),
disease=gl(2, 4, labels=c("Yes",
"No")),
counts=as.vector(snoring))
snoring.df <- snoring.df[rep(seq_len(nrow(snoring.df)), snoring.df$counts),
1:2]
```

And the following two models will yield identical results:

```
levels(snoring.df$snoring) <- c(0, 2, 4, 5)
y <- abs(as.numeric(snoring.df$disease)-2)
x <- as.numeric(as.character(snoring.df$snoring))
fit.glm1 <- glm(y ~ x, family=binomial)

fit.glm2 <- glm(snoring ~ c(0, 2, 4, 5), family=binomial)
```

That is, $\text{logit}[\hat{\pi}(x)] = -3.87 + 0.40x$, using Agresti’s notation.

The second notation is frequently used on aggregated table with an instruction like `cbind(a, b)`, where `a` and `b` are columns of counts for a binary event (see e.g., [Generalized Linear Models](#)). It looks like it would also work when using table instead of matrix (as in your example), e.g.

```
glm(as.table(snoring) ~ c(0, 2, 4, 5), family=binomial)
```