Simple haplotype analyses in R

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Statistical methods of varying complexity have been proposed to efficiently estimate haplotype effects and haplotype-environment interactions in case-control and prospective studies. We have proposed an alternate approach that is based on a non-iterative, two-step estimation process: first, an expectation-maximization algorithm is used to compute posterior estimates of the probability of all potential haplotypes consistent with the observed genotype for each subject; second, the estimated probabilities are used as weights in a regression model for the disease outcome, possibly including environmental factors. Standard error estimates are based on a robust variance estimator. We have shown that the two-step process provides valid tests for genetic associations and reliable estimates of modest genetic effects of common haplotypes for case-control studies (French et al, 2006). The two-step process has also been applied to prospective studies with a survival outcome subject to censoring (Neuhausen et al, 2009). An advantage of the two-step process is its straightforward implementation in software, so that analyses combining genetic and environmental information can be conducted by researchers expert in that subject matter using standard software, rather than by statisticians using specialized software. We illustrate the use of the two-step process for case-control studies using our R package haplo.ccs, which implements weighted logistic regression, and for prospective studies with a survival outcome using our working R package haplo.cph, which implements weighted Cox regression. We illustrate our method and software using data from a study of chronic heart failure patients (Cappola et al, 2011) to estimate the effect of CLCNKA haplotypes on time to death or cardiac transplantation.

References

