Vol. 19 no. 1 2003 Pages 149–150



Genetic Power Calculator: design of linkage and association genetic mapping studies of complex traits

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Received on May 16, 2002; revised on July 11, 2002; accepted on July 15, 2002

ABSTRACT

Summary: A website for performing power calculations for the design of linkage and association genetic mapping studies of complex traits.

Availibility: The package is made available at http:// statgen.iop.kcl.ac.uk/gpc/

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An essential first step in the planning of any scientific study is to assess how many samples must be collected in order to achieve sufficient power to detect the hypothesized effect. In human genetics this requirement is particularly salient, given the costs involved in phenotyping and genotyping individuals. Futhermore, it is virtually impossible to submit a grant proposal to study a particular disease or trait without inclusion of detailed power calculations to show that the proposed research is likely to succeed, provided there is indeed a gene to be found. After conducting a study, power analysis can also shed light on negative results by indicating whether the study was underpowered, or what the smallest detectable effect size would be given the actual sample size. The power of a study is the probability of successfully detecting an effect of a particular size: if β is the probability of a false-negative (type II) error, then power is $1 - \beta$. Power depends on several factors: magnitude of effect, sample size, N, and required level of statistical significance, α (the false-positive, or type I, error rate). Although N and α are determined by the experimenter, many of the factors that contribute to the effect size are typically unknown. In order to compute power, we are therefore required to make assumptions regarding what we expect to find. For mapping loci, such factors include the proportion of variance explained by the trait locus, gene action, and marker heterozygosity and density. Although there is no shortage of statistical genetic literature to aid the

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researcher in performing such calculations (Cardon and Fulker, 1994; Carey and Williamson, 1991; Nance and Neale, 1989; Neale *et al.*, 1994; Schmitz *et al.*, 1998; Sham *et al.*, 2000; Suarez *et al.*, 1982) there are few software tools to make the task practical. The present paper describes an easy- to-use website which allows the researcher to quickly perform such necessary calculations.

Methods to map loci influencing complex traits fall into two broad classes: linkage and association. Linkage relies on correlating sharing of chromosomal segments among relatives with their similarity on a trait whereas association directly relates genotype to phenotype. Variance components models provide a powerful framework for both linkage and association mapping (Almasy and Blangero, 1998; Fulker and Cherny, 1996; Pratt et al., 2000; Fulker et al., 1999). It has been shown that maximum-likelihood variance components approaches to linkage mapping of quantitative trait loci (QTL), which utilize the full familial covariance structure, are more powerful than simple regression-based methods (Fulker and Cherny, 1996) Additionally, for association mapping, a powerful variance components approach has been presented (Fulker et al., 1999) which allows simultaneous modelling of linkage and association while controlling for population stratification effects in sibship data, by considering both between-sibship and within-sibship variation. This method has been made accessible by the release of QTDT (Abecasis et al., 2000a) and has been generalized to deal with extended families rather than just sibships (Abecasis et al., 2000b).

The computationally intensive approach to power calculation is to simulate hundreds or thousands of replicate samples under a specified set of population parameters. The proportion of replicates in which an effect is detected (the test statistic falling above a specified threshold) provides an estimate of power. Recently, however, closedform analytic power equations have been presented for

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variance components methods of linkage and association mapping (Sham *et al.*, 2000). The use of such equations greatly speeds up power calculation and allows a more comprehensive exploration of the parameter space (e.g. different models and sample types). The Genetic Power Calculator (GPC) implements these power equations and others, for both linkage and association methods using either qualitative (e.g. the presence or absence of a disease) or quantitative (e.g. a score on a personality inventory) traits.

Power for the variance components linkage test can be calculated for sibships of arbitrary size, under userdefinable levels of the proportion of variance explained by the trait locus, acting additively and/or via dominance. The background residuals sibling correlation can be varied, as can the polymorphism information content at the locus of interest, allowing accommodation of either twopoint or multipoint linkage. The output includes a table of power for various common α levels as well as a userselected α level, and required sample size to achieve the user-selected desired level of power, suitable for direct inclusion in a grant proposal. In addition the noncentrality parameter is provided to facilitate power calculation for samples of variable-sized sibships. For tests of association, GPC uses the variance components test described by Fulker et al. (1999). The user is presented with a similar set of options as for the linkage test, along with association-specific options such as the extent of trait locus-marker locus linkage disequilibrium and allele frequencies. Output is presented for between-sibship, within-sibship and combined tests of association.

For testing association in discrete (disease) traits, tools are available for both the TDT test, which employs parents and a single affected offspring (Spielman et al., 1993) and the case-control design (Sham, 1998). The user is able to explore power under conditions of varying disease allele frequency, disease prevalence, and genotype relative risk. Again, output is similar to that described above. In addition, calculation of power for the quantitative TDT and quantitative case-control designs is also available. These study designs assume that cases and controls are defined as scoring above or below specific thresholds. Additional utilities made available on the website include two-locus linkage power calculations and a facility for calculating the potential informativeness of sibships for linkage, conditional on observed trait values. This index of informativeness provides a basis for efficient selective genopyting (Purcell et al., 2001). Presently, there is no software widely available to employ such tests, but the situation is likely to improve in the near future. We will attempt to add additional tools for estimating power to this website as additional methods of analysis are developed and software distributed for their implementation.

ACKNOWLEDGEMENTS

Supported in part by National Institutes of Health (USA) grant EY-12562, MRC components grant G9700821 (UK) and the Wellcome Trust.

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