

## THEORY AND METHODS

# Size matters: just how big is BIG?

## Quantifying realistic sample size requirements for human genome epidemiology

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**Background** Despite earlier doubts, a string of recent successes indicates that if sample sizes are large enough, it is possible—both in theory and in practice—to identify and replicate genetic associations with common complex diseases. But human genome epidemiology is expensive and, from a strategic perspective, it is still unclear what ‘*large enough*’ really means. This question has critical implications for governments, funding agencies, bioscientists and the tax-paying public. Difficult strategic decisions with imposing price tags and important opportunity costs must be taken.

**Methods** Conventional power calculations for case–control studies disregard many basic elements of analytic complexity—e.g. errors in clinical assessment, and the impact of unmeasured aetiological determinants—and can seriously underestimate true sample size requirements. This article describes, and applies, a rigorous simulation-based approach to power calculation that deals more comprehensively with analytic complexity and has been implemented on the web as *ESPRESSO*: ([www.p3gobservatory.org/powercalculator.htm](http://www.p3gobservatory.org/powercalculator.htm)).

**Results** Using this approach, the article explores the *realistic* power profile of stand-alone and nested case–control studies in a variety of settings and provides a robust quantitative foundation for determining the required sample size both of individual biobanks and of large disease-based consortia. Despite universal acknowledgment of the importance of large sample sizes, our results suggest that contemporary initiatives are still, at best, at the lower end of the range of desirable sample size. Insufficient power remains particularly problematic for

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studies exploring gene–gene or gene–environment interactions.

**Discussion** Sample size calculation must be both accurate and *realistic*, and we must continue to strengthen national and international cooperation in the design, conduct, harmonization and integration of studies in human genome epidemiology.

**Keywords** Human genome epidemiology, biobank, sample size, statistical power, simulation studies, measurement error, reliability, aetiological heterogeneity

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## Introduction

By 2020, common chronic diseases will account for almost three-quarters of deaths worldwide.<sup>1</sup> The quest to find genetic variants underlying these disorders is in a ‘fast-moving, exciting and highly productive phase’.<sup>2</sup> If the common disease common variant hypothesis<sup>3–8</sup> is true for at least some genetic determinants of chronic diseases, their aetiological effects will typically be weak<sup>9,10</sup> and they will be identified more readily by association rather than linkage studies.<sup>11</sup> In consequence, although alternative strategies could have been adopted,<sup>12</sup> the majority of contemporary gene discovery studies are based on association studies in unrelated individuals.

A series of recent publications has convincingly identified or replicated genetic associations for a range of chronic diseases including: type 1 diabetes;<sup>13,14</sup> type 2 diabetes;<sup>14–18</sup> coronary artery disease;<sup>14,19–21</sup> breast cancer;<sup>22,23</sup> colorectal cancer;<sup>24–26</sup> prostate cancer;<sup>27,28</sup> age-related macular degeneration<sup>29–31</sup> and Crohns disease.<sup>14,32</sup> But, prior to these recent contributions, genetic association studies were strikingly inconsistent.<sup>7–9,33–42</sup> Although numerous scientific and technical issues were blamed,<sup>10,33,35,37,40,43–47</sup> perhaps the most fundamental problem was a serious lack of statistical power.<sup>10,33,35,37,40,43–47</sup> This raises important questions: how large must stand-alone and nested case–control studies really be if they are to power contemporary gene discovery? And specifically, will the current generation of ‘large’ initiatives,<sup>14,48,49</sup> <http://www.genome.gov/17516722>, <http://www.p3gob.servatory.org>, generate enough power to study the *joint* effects of genes and environment?<sup>50</sup>

These questions are crucial. Governments and funding agencies worldwide are deciding whether, and how much, to invest in population genomics. Difficult strategic decisions, with imposing price tags and substantial opportunity costs have to be taken. In Europe, for example, national governments must decide whether to adopt the European Union’s regional road map for research infrastructures in ‘big science’. This proposes development of a harmonized pan-European network of biobanks. But, are pre-existing projects, like UK Biobank,<sup>48</sup> the Wellcome Trust case control consortium (WTCCC),<sup>14</sup> EPIC (European prospective investigation into cancer and

nutrition)<sup>51</sup> and BioHealth Norway<sup>52</sup> already large enough to service all foreseeable needs, or is further investment required to facilitate larger pooled analyses and more powerful replication studies?

Rigorous power calculations are needed, but conventional approaches disregard key elements of analytic complexity, including the bioclinical complexity of causal pathways leading to disease and the inferential complexity that arises from key aspects of study design, conduct and analysis. For example, errors in assessing disease status and aetiological determinants are known to dramatically reduce statistical power if the primary outcome is a quantitative disease-related phenotype.<sup>53</sup> But, although their importance has been emphasized,<sup>54</sup> they are typically ignored by conventional power calculations for case–control studies.

This article describes and applies a simulation-based approach to power calculation for case–control studies in population genomics, generating a *realistic* power profile across a range of meaningful bioclinical scenarios. It also explores the incidence of common chronic diseases in a typical population-based cohort study recruiting middle-aged adults. Taken together, these data provide a logical basis for deciding the appropriate size of major new initiatives in population genomics, including the construction of disease-based and population-based biobanks and the pulling together of consortia based on case series, population controls, case–control projects and/or cohort studies.

## Materials and methods

All simulations were carried out in the statistical programming environment ‘R’.<sup>55</sup>

### The required size of case–control analyses

Simulation-based power calculation involves two steps—simulation and analysis. Here, both steps were based on logistic regression. All analyses in the main paper utilize an unmatched case–control design enrolling unrelated individuals with four controls per case (alternative case: control ratios are considered in Supplementary materials, and Supplementary Figure S1). Unless otherwise stated, genetic and environmental determinants are all dichotomous

(as explained in Supplementary Box S1). Supplementary methods include: (i) full specification of the mathematical models used (equations A–K); (ii) an annotated version of the R code for the primary simulation programme and (iii) discussion of the key assumptions invoked in the analysis and the effect of modifying them. Formal estimates of type 1 error were all nominal (Supplementary Table S1).

Genetic variants were modelled as having two levels: ‘at risk’ and ‘not at risk’. This would apply, for example, under a dominant genetic model (Supplementary box S1): one detrimental allele puts you ‘at risk’, but that risk is increased no further by a second copy. Under such a model, 9.75% of the general population would be ‘at risk’ if the minor allele frequency (MAF) was 5%, and at-risk prevalences of 19, 51 and 75% would correspond to MAFs of 10, 30 and 50%, respectively. A dichotomous genetic variant represents the setting of least power that is commonly encountered and it was for this reason that it was used as the default in the main paper. When it is mathematically valid, more power and a smaller sample size requirement, may be obtained if an additive genetic model (in contrast to a dichotomous model) is used. This model, which is used widely—e.g.<sup>14</sup>—is considered further in the discussion.

Interaction terms reflect departures from a multiplicative model—i.e. from additivity on the scale of log odds—[Supplementary methods, equations (A–C)]. In simulation studies, where a gene–environment interaction is of primary interest, the main effect ORs (odds ratios) associated with the genetic and environmental determinant is fixed at 1.5 while the magnitude of the interaction term is varied: results are insensitive to changing this fixed magnitude (Supplementary Table S2).

### Step 1 (simulation)

The parameters characterizing each scenario (a series of simulations all using the same bioclinical parameters) were set, and varied, under the following assumptions: (i) Prevalence of the ‘at risk’ genotypic and environmental determinants [0.0975 (MAF = 5%) or 0.51 (MAF = 30%), and 0.1 or 0.50, respectively]; (ii) ORs associated with genotypic and environmental main effects (1.10–3.0), and gene–environment interactions (1.20–10.00); (iii) Sensitivity and specificity of disease assessment appropriate to the particular disease under consideration (e.g.<sup>56</sup>, Supplementary box S2); Supplementary materials and Supplementary Figure S2 explore the impact of changing sensitivity and specificity; (iv) Controls assessed clinically in the same way as cases (except in the real WTCCC data); (v) Errors in classifying genotypes modelled as if arising primarily from incomplete linkage disequilibrium (LD) between an observed marker and a causative variant [ $R^2 = 1.00$  (no error), 0.80<sup>57</sup> or 0.50]; (vi) Lifestyle/environmental exposure status determined by dichotomization of an underlying quantitative variable measured with

error equivalent to a test–retest reliability of (a) 100%, (b) 90%, (c) 70%, (d) 50% or (e) 30%; (vii) Heterogeneity of underlying disease risk modelled using a random effect<sup>58</sup> with a variance reflecting a 10-fold ratio in baseline risk between individuals on ‘high’ (95%) and ‘low’ (5%) population centiles; Supplementary materials and Supplementary Figure S3 investigate changing the heterogeneity of risk; (viii) Disease prevalence appropriate to the particular disease under consideration<sup>56</sup>; Supplementary materials and Supplementary Figure S4 explore changing disease prevalence; (ix) Statistical significance defined at  $10^{-7}$  for a genome wide association (GWA) study and  $10^{-4}$  for a candidate gene study or for gene–environment interactions (Supplementary methods) and (x) No correction was made for substructure in population ancestry.<sup>14</sup>

Having set the required parameters, a dataset ( $D_1$ ) was simulated (Supplementary methods, equations A–F), containing cases and controls each associated with a set of aetiological determinants (e.g. a gene and an environmental determinant) distributed as would be expected for a case–control study given the particular bioclinical parameters specified.

### Step 2 (analysis)

Dataset  $D_1$  was analysed using unconditional logistic regression (Supplementary methods, equations G–J), as if it were a real case–control study. This generated estimates (and associated standard errors and  $P$ -values) for the regression coefficients reflecting the genetic and environmental main effects and, where incorporated, a gene–environment interaction. On the basis of the pre-specified type 1 error,  $D_1$  was categorized as either ‘significant’ or ‘non-significant’ for each of the genetic, environmental and interaction effects.

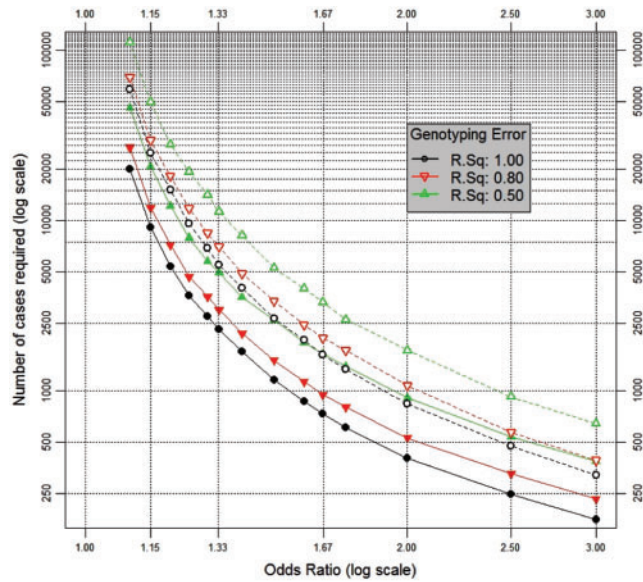
Under each scenario, steps 1 and 2 were repeated many times ( $\geq 1000$ ), generating and analysing datasets  $D_2, D_3, \dots, etc.$  The empirical statistical power of the test for each effect was then estimated as the proportion of the simulated datasets for which step 2 generated a ‘statistically significant’ result. Given the estimated power of a study based on whatever number of cases and controls had actually been specified under the particular scenario being considered, the sample size for an equivalent study (including the same ratio of controls to cases) that would generate a power of 80% was estimated as described in Supplementary methods (equation K). In exploring the power profile across a range of ORs, the required sample size for each OR was calculated, tabulated and plotted (Figures 1 and 2).

This approach is very flexible and can easily be extended by adding additional terms (Supplementary methods).

### The expected incidence of chronic disease in population-based cohorts

The number of incident cases of selected chronic diseases of public health relevance expected to



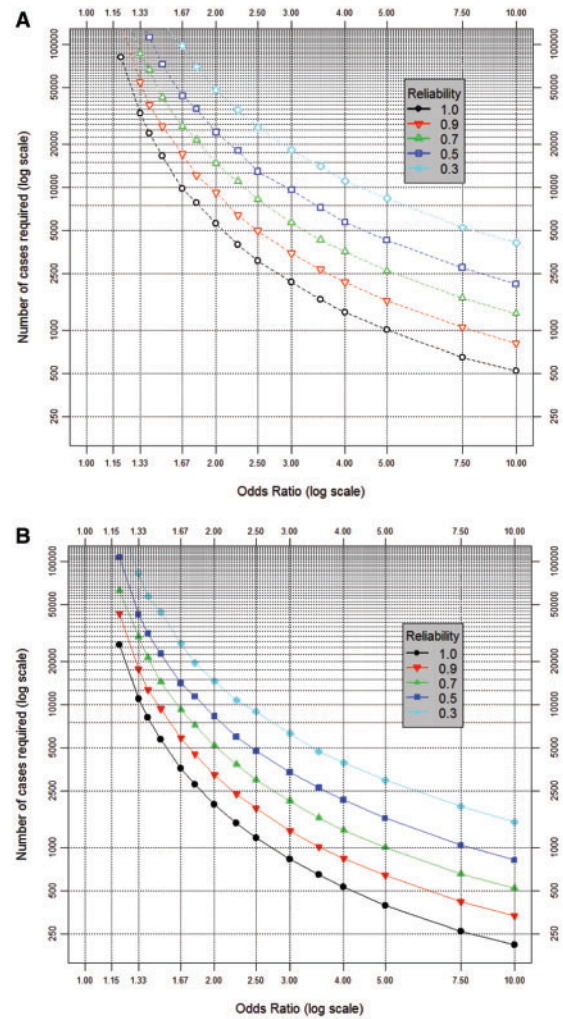


**Figure 1** A genetic main effect, in a candidate gene study. The number of cases required to detect ORs from 1.1 to 3.0 for a genetic main effect with a power of 80% (at  $P < 0.0001$ )—assuming a vague candidate gene) in a study with four controls per case. Assumptions (see Materials and methods section for details): (i) population genotypic prevalence = 9.75% [dashed lines] or 51% [solid lines], corresponding to dominant SNP effects with MAFs (minor allele frequencies) of 5 and 30%, respectively; (ii) genotypic ‘error’ corresponding to:  $R^2 = 1.0$ , 0.8 or 0.5; (iii) case-status determined with sensitivity 89.1% and specificity 97.4%: as for a study of diabetes diagnosed by a composite test (GP diagnosis or HbA1C  $\geq 2$  SD above the population mean<sup>56</sup>); (iv) controls phenotypically assessed in the same way as cases; (v) incorporation of heterogeneity in the baseline risk of disease arising from unmeasured determinants, corresponding in magnitude to a 10-fold risk ratio between individuals on the high (95%) and low (5%) centiles of population risk

accumulate over time was estimated in a simulated cohort of 500 000 individuals, recruited over 5 years, with equal numbers enrolled in all 5 year age bands between 40 and 69 years at entry (Figure 3). This simulated cohort corresponds closely to the design of UK Biobank<sup>48</sup> and provides important parallels to other cohorts worldwide ([www.p3gobserver.org](http://www.p3gobserver.org)). Each recruit was simulated in ‘R’ and his/her subsequent life-course was simulated taking appropriate account of the chance occurrence of disease, migrations, loss to follow-up and deaths (sources for the vital statistics are detailed in Supplementary Table S3). Adjustment was also made for the ‘healthy cohort effect’, whereby subjects recruited to cohort studies typically experience lower rates of morbidity than the general population.

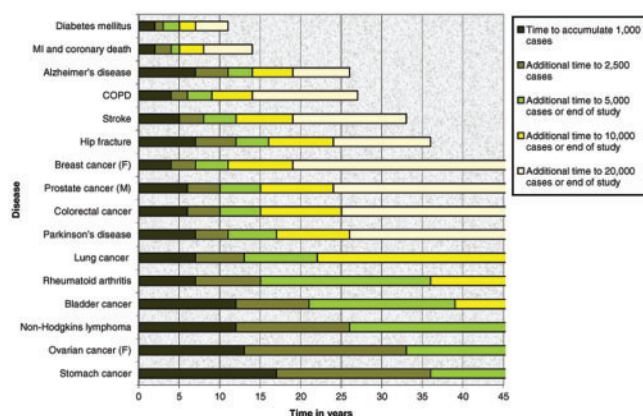
## Results

Figure 1 presents the sample size needed for 80% power to detect (at  $P < 0.0001$ ) the main effect of a



**Figure 2** (A) An uncommon interaction. The number of cases required to detect ORs from 1.2 to 10.0 for a gene–environment interaction with a power of 80% (at  $P < 10^{-4}$ ) in a study with four controls per case. Assumptions (see Materials and methods section for details): (i) population genotypic prevalence = 9.75%, corresponding to a dominant SNP effect with a MAF of 5%; (ii) population prevalence of binary environmental determinant = 20%; (iii) genotypic ‘error’ corresponding to  $r^2 = 0.8$ ; (iv) environmental error corresponding to dichotomization of an underlying normally distributed latent quantitative variable measured with a reliability of 100, 90, 70, 50 or 30%; (v) case–control status determined with sensitivity 89.1% and specificity 97.4%: as for a study of diabetes diagnosed by a composite test (GP diagnosis or HbA1C  $\geq 2$  SD above the population mean<sup>56</sup>); (vi) controls phenotypically assessed in the same way as cases; (vii) incorporation of heterogeneity in the baseline risk of disease arising from unmeasured determinants, corresponding in magnitude to a 10-fold risk ratio between individuals on the high (95%) and low (5%) centiles of population risk. The prevalences of the ‘at risk’ genotype and the ‘at risk’ environmental determinant imply a prevalence of  $\sim 2\%$  for the doubly ‘at risk’ interaction. (B) A common interaction. As (A), but assuming: population genotypic prevalence = 51% (corresponding to MAF = 30%) and prevalence of the environmental determinant = 50%, implying prevalence of the doubly ‘at risk’ interaction  $\sim 25\%$

dichotomous (binary) genetic variant in a vague candidate gene, using an unmatched case–control design enrolling four controls per case. As detailed in the Materials and methods section, and summarized in the figure legend, these calculations all incorporate cardinal elements of realistic analytic complexity. Table 1 details the multiplicative factor by which the sample sizes in the figure should be scaled if a  $P$ -value other than  $P < 0.0001$  is to be used or if one requires a power of 50 or 90% rather than 80%.



**Figure 3** Time to achieve required numbers of cases. The expected rate of generation of incident cases of 16 common complex diseases (MI = myocardial infarction; COPD = chronic obstructive pulmonary disease) in a cohort of 500 000 men and women recruited over 5 years, aged 40–69 years at baseline, assuming rates of mortality and morbidity as in the UK and that drop-out mirrors that of the British 1958 birth cohort and with adjustment for the ‘healthy cohort’ effect (for full details see<sup>68</sup>)

Figures 2A and B present sample size requirements for studies of gene–environment interaction (see Materials and Methods section, with details in Supplementary methods). Phenotypic and genotypic characteristics are detailed in the figure legend. Figure 2A considers an uncommon interaction where ‘doubly-at-risk’ individuals (i.e. subjects exposed to the at-risk level of both the genetic and the lifestyle determinant) represent ~2% of the general population. Figure 2B addresses a common interaction with ~25% of individuals being doubly-at-risk. Each figure details the sample size profile for a range of errors in assessing the environmental factor (see Materials and methods section). As a benchmark, Table 2 presents bioclinical exemplars that are typically measured with a corresponding reliability. Significance testing is at  $P < 0.0001$ : i.e. it is assumed that research involving the joint effects of genes and environment will focus on specific interactions with at least some vague basis for candidature. If a more rigorous threshold is required, the sample size multipliers in Table 1 may be used.

In light of the daunting sample size requirements implied by Figures 1 and 2, Figure 3 demonstrates that a cohort of 500 000 middle-aged recruits may be expected to generate 10 000 incident cases of very common conditions (e.g. diabetes and coronary artery disease) within 7–8 years, and 20 000 cases within 15 years. But even for the commonest cancers it will take ~20 years or more to generate 10 000 cases and >40 years to generate 20 000 cases. Such targets will never be attained for rarer conditions. However, population-based cohort studies also recruit prevalent cases of chronic disease (Supplementary Table S4)

**Table 1** The multiplicative change in required sample size, for a statistical power of 50, 80 or 90% using different levels of statistical significance, relative to the sample size indicated by Figures 1 and 2 (i.e. when  $P < 0.0001$  and power = 80%)

<i>P</i> -value defining ‘statistical significance’	Sample size multiplier for specified power		
	Power = 50%	Power = 80%	Power = 90%
Significance thresholds conventionally used in medical statistics			
$P < 0.05$	0.172	0.350	0.469
$P < 0.01$	0.296	0.522	0.664
Significance thresholds widely used in candidate gene studies			
$P < 0.005$	0.352	0.594	0.746
$P < 0.001$	0.484	0.762	0.933
$P < 0.0005$	0.541	0.834	1.013
$P < 0.0001$	0.676	as in figures	1.195
Significance thresholds widely used in GWA studies			
$P < 5 \times 10^{-7}$	1.128	1.538	1.777
$P < 10^{-7}$	1.267	1.699	1.950
$P < 5 \times 10^{-8}$	1.327	1.768	2.024
$P < 10^{-8}$	1.467	1.929	2.196

Sample size multipliers more than 1 indicate that the required sample size is larger than implied by the figures, values less than 1 that it is smaller.

**Table 2** Formal estimates of test–retest reliability for a number of exemplar lifestyle/environmental determinants that are widely studied

Reliability of measurement	Lifestyle/environmental factor
≥0.95	Body mass index (BMI) calculated from measured height and weight in various studies <sup>76</sup>
~0.9	Measured hip or waist circumference <sup>76,77</sup>
~0.7	Blood pressure measurement in the Intersalt Study <sup>78</sup>
~0.5	Many nutritional components in a dietary recall study, mean of four 24 h assessments <sup>79</sup>
~0.3	Many nutritional components in a dietary recall study, a single 24 h assessment <sup>79</sup>

and, if it is appropriate, these can be used to supplement statistical power.

## Discussion

'Big' bioscience is critically poised. It is now known that genetic associations with complex diseases can reliably be detected and replicated if sample sizes are large enough. This will fuel international investment in biobanking. But, how far should that investment go?

It is essential to close the 'reality-gap' that currently exists between the sample sizes really required to detect determinants of scientific interest that have plausible bioclinical effects, and the sample sizes that are typically used when studies are being designed. Extensive theoretical work has been undertaken to explore statistical power and sample size in human genomics. This includes studies of the effect of genotype misclassification on power,<sup>59–62</sup> and of strategies for power optimization for genetic main effects<sup>63–64</sup> and for gene–gene<sup>65</sup> and gene–environment interactions.<sup>66</sup> Furthermore, the effect of measurement error in both outcomes and exposures on statistical power of gene–environment interaction studies has been explored thoroughly for *quantitative traits*.<sup>53</sup> But, previous work on the power of case–control analyses (i.e. *binary traits*) has not addressed the impact of realistic assessment errors in both exposures and outcomes and the impact of unmeasured aetiological determinants. The important original contributions of the current article are 3-fold, therefore: (i) to extend the classes of analytic complexity addressed in a straightforward simulation-based power calculation engine; (ii) to use this calculator to undertake *realistic* sample size calculations for a class of analyses (case–control analyses with unavoidable assessment errors in both exposures and outcomes) that will be utilized widely over the next few decades—analyses in this

class are the least powerful that are likely to be applied commonly, and the resultant calculations, therefore, provide a valuable guide to study design in the many large-scale biobanks that are currently being conceived and launched; (iii) to alert readers, particularly those setting up new biobanks, to a web-based implementation—ESPRESSO (Estimating Sample-size and Power in R by Exploring Simulated Study Outcomes)—of the power calculator used in this article and to provide detailed information (in Supplementary methods) about the mathematical models on which it is based.

Studies enrolling several hundred subjects are commonplace in human genome epidemiology. But, even conventional power calculations<sup>67</sup> indicate that 400 cases and 400 controls provide <1% power to detect (at  $P < 0.0001$ ) an OR of 1.4 for a binary 'at risk' genetic variant with a general population frequency of 0.0975 (e.g. a dominant risk-determining allele with MAF = 0.05). There can be no doubt that a study involving several hundred cases and controls demands hard work and is large by historical comparison; nevertheless, the *reality* is that to generate a power of 80%, such a study would actually require 4000 cases and 4000 controls.

But even these figures substantially understate the challenge that really faces us. Conventional power calculations ignore many aspects of analytic complexity. Using ESPRESSO, the R-based<sup>55</sup> simulation-based power calculator<sup>68</sup> jointly developed by P<sup>3</sup>G, PHOEBE and UK Biobank ([www.p3gobservatory.org/powercalculator.htm](http://www.p3gobservatory.org/powercalculator.htm)), such complexities can be taken into proper account. Using this approach to mimic the conventional power calculation (above)—i.e. assuming disease and genotype to be assessed without error and no heterogeneity in disease risk—confirms a requirement for approximately 4000 cases and 4000 controls. But the sensitivity and specificity of the diagnostic test ought to be taken into proper account: e.g. 0.891 and 0.974, respectively, for a published screening test for type 2 diabetes based on glycosylated haemoglobin<sup>56</sup> (see Supplementary box 2). Genotyping error must also be considered. It may, for example, be reasonable to assume that this corresponds to incomplete LD with an  $R^2$  of 0.8.<sup>57</sup> Finally, heterogeneity in disease risk might be reflected in an assumed 10-fold ratio in the risk between subjects on high (95%) and low (5%) centiles of population risk. Having built in these assumptions, the required sample size *more than doubles* to 8500 cases and 8500 controls.

It might be argued that substantial power could be gained if a multiplicative model based on additive allelic effects [Supplementary methods, equations (H and I)], as in WTCCC,<sup>14</sup> were used instead of a binary genetic model (Supplementary box S1). Statistical power will be increased if there is a systematic gradation in the strength of association across the three genotypes defined by two alleles. This may reflect biological



reality, or it may arise as an artefact of the decay of incomplete LD when working with a linked marker rather than a causative variant. But (Supplementary Figure S5), the reduction in required sample size (typically, 5–50%) is only substantial for SNPs with a common minor allele. This is because when the minor allele is rare, subjects homozygous for the minor allele will be *very rare* and the genetic determinant will effectively act as if it were a binary exposure. But, power limitation is less of a problem for SNPs with a common minor allele and so the impact of moving to a valid multiplicative genetic model is less dramatic than might otherwise be assumed.

One of the landmark genomic studies of 2007 was the WTCCC that reported robust ‘hits’ in seven of eight complex diseases in its main experiment.<sup>14</sup> But the basic design—involving 2000 cases and 3000 controls for each disease—seems, at first sight, to be at the lower limit of required sample size as implied by our calculations (Figure 1). Therefore, it is tempting to conclude either that the WTCCC was lucky or that our calculations are overly conservative. But, the main experiment of WTCCC had a number of design features that contrast with the assumptions of the primary power calculations reported in our article.<sup>14</sup> The most relevant of these are: (i) use of a model invoking an additive genetic effect rather than a binary ‘at-risk’ genotype; (ii) cases rigorously phenotyped so that few, if any, non-diseased subjects will have appeared as cases; (iii) a *P*-value threshold of  $5 \times 10^{-7}$ ; (iv) a case:control ratio of 2:3; (v) no phenotyping of controls so diseased subjects will have contaminated the controls to an extent determined solely by general population prevalence. On the basis of simulations that invoke all of these assumptions, Supplementary Figure S6 presents the precise equivalent to Figure 1, but uses the design parameters of the WTCCC. On the basis of their own simulation-based power calculations (incorporating errors consequent upon incomplete LD),<sup>14</sup> the design team of the WTCCC estimated that its power would be ‘43% for alleles with a relative risk of 1.3, increasing to 80% for a relative risk of 1.5’.<sup>14</sup> These power calculations were based on averaging across all MAFs  $> 0.05$ ,<sup>14</sup> and the design should, therefore, be underpowered for SNPs with an uncommon minor allele and to have more power than required for common SNPs.<sup>14</sup> Our methods (Supplementary Figure S6) concur that for SNPs with a MAF in the range 0.2–0.5, the WTCCC design was well powered to detect *heterozygote ORs*<sup>14</sup> of 1.3 or greater and that even ORs as low as 1.2 should have been detected with a non-negligible probability (power ~9%). On the other hand, the power to detect rarer SNPs (MAF = 0.05–0.1) with ORs  $< 1.5$  should have been low. Without knowing which SNPs are truly associated with which complex disease, or how strong those associations might be, it is impossible to use the empirical evidence to *precisely* quantify how accurately our approach predicts the power of the

WTCCC. But, the predicted power profile is certainly consistent with the results reported in Table 3 of the WTCCC paper.<sup>14</sup> Three of the 19 SNPs they identified as having a ‘significant heterozygous OR’, had a MAF between 0.05 and 0.1 and these *all* had  $OR > 1.5$ . In contrast, 13 had a MAF between 0.2 and 0.5 and of these, four exhibited an  $OR < 1.3$  (1.19–1.29), five an  $OR$  between 1.3 and 1.5 and four an  $OR > 1.5$ . SNPs with a rarer minor allele are typically most common,<sup>57</sup> and if power was not a substantial issue, one would have expected more ‘hits’ to arise in rare SNPs. It is true that the observed ORs would have been subject to the ‘winner’s curse’,<sup>69</sup> but this does not detract from the consistency of the overall pattern that was found. As a second test of its validity, the ESPRESSO model was then used to estimate the power of the WTCCC to reconfirm the effect of 12 (non-HLA) loci that had been ‘previously robustly replicated’.<sup>14</sup> On the basis of the published bioclinical characteristics of these 12 variants (Supplementary Table S5), our simulations predicted a 24% probability that all 12 would replicate and probabilities of 44, 26 and 6%, respectively, that 11, 10 or  $\leq 9$  would replicate. These predictions are closely consistent with the published WTCCC analysis in which 10 of the 12 actually replicated.<sup>14</sup> Of course, these analyses provide no more than a rudimentary check of the calibration of our approach, nevertheless, it is encouraging that the predictions appear sensible and it would, therefore, seem reasonable to apply the methods to new problems, including those involving environmental as well as genetic determinants.

The fact that our power estimates appear consistent with those of the WTCCC team itself suggests that any additional elements of analytic complexity that were addressed by our methods had a limited impact on required sample size in this particular setting. Therefore, we explored the relative contribution to increased sample size requirement that was consequent upon those specific elements of our model that are not included in a conventional power calculation. Across an arbitrary, but not atypical, set of models incorporating a gene–environment interaction (see Supplementary materials, and Supplementary Table S6), it was found that it is a realistic level of error in assessing the environmental determinant that was most influential in inflating the required sample size. But, the WTCCC analysis focused solely on genetic main effects and so this was irrelevant. Furthermore, all cases in WTCCC were carefully phenotyped. Specificity was, therefore, close to 100% and very few, if any, healthy subjects would have appeared as cases (Supplementary materials and Supplementary Figures S2a and S2b). Finally, the sophisticated power calculations undertaken by WTCCC took appropriate account of error arising from incomplete LD, and so the only additional factor that *did* come into play in the WTCCC was heterogeneity in underlying disease risk—but, on its own, this has little impact (Supplementary Table S6).

Can biobanks ever be *large enough*? Although our methods are in accord with the power calculations undertaken by the WTCCC and suggest that it was appropriately powered to detect the effects that it set out to study, larger—sometimes *much* larger—sample sizes will be required (Figures 1 and 2) to reliably detect: (i) ORs at the lower end of the plausible range; (ii) SNP effects associated with rarer minor alleles; (iii) genotypic effects that are binary rather than multiplicative in nature; (iv) gene–environment (or, gene–gene) interactions or (v) aetiological effects in case series subject to less exhaustive phenotyping. If bioscience aims to rigorously investigate such effects, it will be necessary to design studies enrolling not thousands, but tens of thousands of cases. But, studies of such a size should not be contemplated unless relative risks  $\leq 1.5$  are really worth investigating. A central aim of modern bioscience is to understand the causal mechanisms underlying complex disease<sup>49,70</sup> and each quantum of new knowledge has the potential to provide an important insight that may have a dramatic impact on disease prevention or management. This implies that scientific interest may logically focus on any causal association that can convincingly be identified and replicated—it need not be ‘strong’ by any statistical or epidemiological criterion. The fundamental need, therefore, is for research platforms to support analyses powered to detect *plausible* aetiological effects. But, what does this mean? The majority of genetic effects on chronic diseases that have so far been identified and replicated are characterized<sup>8,9,13–32</sup> by allelic or genotypic relative risks of 1.5 or less—many in the range 1.1–1.3. Effect sizes may be greater for causal variants than for markers in LD, but it would be unwise to assume that the gain will necessarily be substantial. Although the search for ‘low hanging fruit’ must continue, therefore, we agree with Easton *et al.*<sup>22</sup> that much of the future harvest will be rather higher up the tree. But, even if they are of scientific interest, can ORs  $\leq 1.5$  reliably be detected by *any* observational study? In 1995, Taubes argued that: ‘[observational epidemiological studies]... are so plagued with biases, uncertainties, and methodological weaknesses that they may be inherently incapable of accurately discerning... weak associations’.<sup>71</sup> Fortunately, several of the central arguments underlying this bleak assessment do not hold in human genome epidemiology. Randomization at gamete formation renders simple phenotype–genotype associations robust to life style confounding and to uncertainty in the direction of causality—in other words, enhanced inferential rigour is a direct, but wholly fortuitous, consequence of what is often called Mendelian randomization.<sup>70,72–74</sup> At the same time, the increasing accuracy and precision of measurements in genome epidemiology<sup>14,53,54</sup> mean that—in the absence of intractable confounding and reverse causality—sufficient statistical power can *realistically* be accrued to draw meaningful inferences for small effect sizes.

Despite important caveats,<sup>70,73,75</sup> therefore, small effects reflecting the direct impact of genetic determinants (main effects and gene–gene interactions) or the differential impact of genetic variants in diverse environmental backgrounds (gene–environment interactions) are more robust than their counterparts in traditional environmental epidemiology.

Finally, we note that the primary simulations that underpin our conclusions are all based on a case: control ratio of 1:4, while a 1:1 ratio was adopted in considering the ‘conventional’ power calculations (see above). Furthermore, most of the case–control studies that we reference (including the WTCCC) are based on ratios that are much closer to unity.<sup>13–32</sup> But this presentation was deliberate. Given access to a fixed number of cases and an unrestricted number of well-characterized controls, substantial additional power can be obtained using a design based on four or more controls per case (Supplementary Figure S1). In the future, the existence of massive population-based biobanks such as UK Biobank<sup>48</sup> and extensive sets of nationally representative controls (e.g. as in WTCCC<sup>14</sup>) will mean that designs based on multiple controls will be highly cost effective and will be widely used. It would, therefore, have been inappropriate to present power calculations based primarily on the 1:1 design as this would have increased the estimated sample size requirement, thereby strengthening our main message in a manner that could have been seen as misleading. On the other hand, in exploring the implications of conventional power calculations (see above), most contemporary work is based on designs with approximately equal number of cases and controls and it was, therefore, felt to be more intuitive for readers to focus on designs of this nature. For the sake of completeness, Supplementary Figures S7, S8a and S8b replicate Figures 1, 2A and B but use equal numbers of cases and controls.

To finish, we note that the basic conclusions we have reached are stark and may appear disheartening. But, pessimism is unwarranted. Disentangling the causal architecture of chronic diseases will be neither cheap nor easy and it would be unwise to assume otherwise. But it has the potential to return investment many-fold with future improvements in promoting health and combating disease. Therefore, it is encouraging that several international case–control consortia have already managed to amass sample sizes of the magnitude that is realistically required.<sup>16,19,22,26</sup> Furthermore, the largest contemporary cohort-based initiatives<sup>48,49,51,52</sup> will generate enough cases to study the commonest diseases in their own right (Figure 3). To take things further, three complementary strategies will markedly enhance the capacity to study plausible relative risks right across the spectrum of complex diseases: (i) improve the accuracy and precision of measurements and assessments;<sup>14,53,54</sup> (ii) increase the size of individual studies and biobanks<sup>8</sup> and (iii) harmonize protocols for information collection, processing



and sharing<sup>10,46–49,70</sup> (<http://www.p3g.org>). Taken together, these actions will provide for a powerful global research platform to drive forward our understanding of the causal architecture of the common chronic diseases. But, such a platform will be of little value unless power calculations are both *accurate* and *realistic*. It is our hope that this article and access to ESPRESSO will be viewed as providing valuable guidance to those setting up individual biobanks and designing the case-control analyses to be based upon them.

## Supplementary data

Supplementary data are available on the P3G Observatory <http://www.p3gobservatory.org>.

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### KEY MESSAGES

- Biobanking is very expensive and the effect sizes to be investigated are often very small—accurate sample size estimation is vital, therefore.
- Conventional power calculations for case-control comparisons ignore key aspects of analytic complexity and can substantially understate sample size requirements—often by a factor of two or more.
- Power profiles for stand-alone and nested case-control studies are presented that are based on a simulation-based approach to calculation that takes robust account of analytic complexity (including several forms of assessment error) and has been implemented as the web-based utility ESPRESSO (<http://www.p3gobservatory.org/powercalculator.htm>).
- Taking appropriate account of realistic constraints on statistical power, any research infrastructure aimed at providing a robust platform for exploring genomic association will typically require several thousands of cases to study main effects and several tens of thousands of cases to properly support the investigation of gene-gene or gene-environment interaction.
- In order to enhance scientific return from the massive international investment in biobanking, power calculations must be both accurate and realistic and individual biobanks must be designed so as to enhance the quality of the data and samples that are collected, and harmonized to facilitate data sharing and pooled analysis.

## References

- 1 World Health Organization (WHO). *Diet, Nutrition and the Prevention of Chronic Diseases. The Aethiology of these Diseases*. Geneva: Joint WHO/FAO Expert Consultation, 2005.
- 2 O’Rahilly S, Wareham NJ. Genetic variants and common diseases—better late than never. *N Engl J Med* 2006; **355**:306–8.
- 3 Lander ES. The new genomics: global views of biology. *Science* 1996; **274**:536–39.
- 4 Cargill M, Daley GQ. Mining for SNPs: putting the common variants-common disease hypothesis to the test. *Pharmacogenomics* 2000; **1**:27–37.
- 5 Reich DE, Lander ES. On the allelic spectrum of human disease. *Trends Genet* 2001; **17**:502–10.
- 6 Pritchard JK, Cox NJ. The allelic architecture of human disease genes: common disease-common variant... or not? *Hum Mol Genet* 2002; **11**:2417–23.

- <sup>7</sup> Hirschhorn JN, Lohmueller K, Byrne E, Hirschhorn K. A comprehensive review of genetic association studies. *Genet Med* 2002;**4**:45–61.
- <sup>8</sup> Lohmueller KE, Pearce CL, Pike M, Lander ES, Hirschhorn JN. Meta-analysis of genetic association studies supports a contribution of common variants to susceptibility to common disease. *Nat Genet* 2003;**33**:177–82.
- <sup>9</sup> Hattersley AT, McCarthy MI. What makes a good genetic association study? *Lancet* 2005;**366**:1315–23.
- <sup>10</sup> Khoury MJ, Little J, Gwinn M, Ioannidis JP. On the synthesis and interpretation of consistent but weak gene-disease associations in the era of genome-wide association studies. *Int J Epidemiol* 2007;**36**:439–45.
- <sup>11</sup> Risch N, Merikangas K. The future of genetic studies of complex human diseases. *Science* 1996;**273**:1516–17.
- <sup>12</sup> Terwilliger JD, Weiss KM. Confounding, ascertainment bias, and the blind quest for a genetic 'fountain of youth'. *Ann Med* 2003;**35**:532–44.
- <sup>13</sup> Todd JA, Walker NM, Cooper JD *et al*. Robust associations of four new chromosome regions from genome-wide analyses of type 1 diabetes. *Nat Genet* 2007;**39**:857–64.
- <sup>14</sup> Wellcome\_Trust\_Case\_Control\_Consortium. Genome-wide association study of 14,000 cases of seven common diseases and 3,000 shared controls. *Nature* 2007;**447**:661–78.
- <sup>15</sup> Grant SF, Thorleifsson G, Reynisdottir I *et al*. Variant of transcription factor 7-like 2 (TCF7L2) gene confers risk of type 2 diabetes. *Nat Genet* 2006;**38**:320–23.
- <sup>16</sup> Zeggini E, Weedon MN, Lindgren CM *et al*. Replication of genome-wide association signals in U.K. samples reveals risk loci for type 2 diabetes. *Science* 2007;**316**:1336–39.
- <sup>17</sup> Saxena R, Voight BF, Lyssenko V *et al*. Genome-wide association analysis identifies loci for type 2 diabetes and triglyceride levels. *Science* 2007;**316**:1331–36.
- <sup>18</sup> Scott LJ, Mohlke KL, Bonnycastle LL *et al*. A genome-wide association study of type 2 diabetes in Finns detects multiple susceptibility variants. *Science* 2007;**316**:1341–45.
- <sup>19</sup> Samani NJ, Erdmann J, Hall AS, Hengstenberg C, Mangino M, Mayer B. Genome-wide association analysis of coronary artery disease. *N Engl J Med* 2007;**357**:443–53.
- <sup>20</sup> Helgadóttir A, Thorleifsson G, Manolescu A *et al*. A common variant on chromosome 9p21 affects the risk of myocardial infarction. *Science* 2007;**316**:1491–93.
- <sup>21</sup> McPherson R, Pertsemlidis A, Kavaslar N *et al*. A common allele on chromosome 9 associated with coronary heart disease. *Science* 2007;**316**:1488–91.
- <sup>22</sup> Easton DF, Pooley KA, Dunning AM, Pharoah PDP, Thompson D, Ballinger DG. Genome-wide association study identifies novel breast cancer susceptibility loci. *Nature* 2007;**447**:1087–93.
- <sup>23</sup> Stacey SN, Manolescu A, Sulem P *et al*. Common variants on chromosomes 2q35 and 16q12 confer susceptibility to estrogen receptor-positive breast cancer. *Nat Genet* 2007;**39**:865–69.
- <sup>24</sup> Haiman CA, Le Marchand L, Yamamoto J, Stram DO, Sheng X, Kolonel LN. A common genetic risk factor for colorectal and prostate cancer. *Nat Genet* 2007;**39**:954–56.
- <sup>25</sup> Tomlinson I, Webb E, Carvajal-Carmona L, Broderick P, Kemp Z, Spain S. A genome-wide association scan of tag SNPs identifies a susceptibility variant for colorectal cancer at 8q24.21. *Nat Genet* 2007;**39**:984–88.
- <sup>26</sup> Zanke BW, Greenwood CM, Rangrej J, Kustra R, Tenesa A, Farrington SM. Genome-wide association scan identifies a colorectal cancer susceptibility locus on chromosome 8q24. *Nat Genet* 2007;**39**:989–94.
- <sup>27</sup> Gudmundsson J, Sulem P, Steinthorsdóttir V, Bergthorsson JT, Thorleifsson G, Manolescu A. Two variants on chromosome 17 confer prostate cancer risk, and the one in TCF2 protects against type 2 diabetes. *Nat Genet* 2007;**39**:977–83.
- <sup>28</sup> Gudmundsson J, Sulem P, Manolescu A *et al*. Genome-wide association study identifies a second prostate cancer susceptibility variant at 8q24. *Nat Genet* 2007;**39**:631–37.
- <sup>29</sup> Klein RJ, Zeiss C, Chew EY *et al*. Complement factor H polymorphism in age-related macular degeneration. *Science* 2005;**308**:385–89.
- <sup>30</sup> Haines JL, Hauser MA, Schmidt S *et al*. Complement factor H variant increases the risk of age-related macular degeneration. *Science* 2005;**308**:419–21.
- <sup>31</sup> Edwards AO, Ritter R 3rd, Abel KJ, Manning A, Panhuysen C, Farrer LA. Complement factor H polymorphism and age-related macular degeneration. *Science* 2005;**308**:421–24.
- <sup>32</sup> Rioux JD, Xavier RJ, Taylor KD *et al*. Genome-wide association study identifies new susceptibility loci for Crohn disease and implicates autophagy in disease pathogenesis. *Nat Genet* 2007;**39**:596–604.
- <sup>33</sup> Ioannidis JP, Ntzani EE, Trikalinos TA, Contopoulos-Ioannidis DG. Replication validity of genetic association studies. *Nat Genet* 2001;**29**:306–9.
- <sup>34</sup> Tabor HK, Risch NJ, Myers RM. Opinion: candidate-gene approaches for studying complex genetic traits: practical considerations. *Nat Rev Genet* 2002;**3**:391–97.
- <sup>35</sup> Weiss KM, Terwilliger JD. How many diseases does it take to map a gene with SNPs? *Nat Genet* 2000;**26**:151–57.
- <sup>36</sup> Goldstein DB, Ahmadi KR, Weale ME, Wood NW. Genome scans and candidate gene approaches in the study of common diseases and variable drug responses. *Trends Genet* 2003;**19**:615–22.
- <sup>37</sup> Cardon LR, Bell JI. Association study designs for complex diseases. *Nat Rev Genet* 2001;**2**:91–99.
- <sup>38</sup> Terwilliger JD, Goring HH. Gene mapping in the 20th and 21st centuries: statistical methods, data analysis, and experimental design. *Hum Biol* 2000;**72**:63–132.
- <sup>39</sup> Palmer LJ, Cardon LR. Shaking the tree: mapping complex disease genes with linkage disequilibrium. *Lancet* 2005;**366**:1223–34.
- <sup>40</sup> Buchanan AV, Weiss KM, Fullerton SM. Dissecting complex disease: the quest for the Philosopher's Stone? *Int J Epidemiol* 2006;**35**:562–71.
- <sup>41</sup> Moonesinghe R, Khoury MJ, Janssens AC. Most published research findings are false-but a little replication goes a long way. *PLoS Med* 2007;**4**:e28.
- <sup>42</sup> Chanock SJ, Manolio T, Boehnke M *et al*. Replicating genotype-phenotype associations. *Nature* 2007;**447**:655–60.
- <sup>43</sup> Risch NJ. Searching for genetic determinants in the new millennium. *Nature* 2000;**405**:847–56.
- <sup>44</sup> Little J, Khoury MJ, Bradley L *et al*. The human genome project is complete. How do we develop a handle for the pump? *Am J Epidemiol* 2003;**157**:667–73.
- <sup>45</sup> Ioannidis JP, Gwinn M, Little J *et al*. A road map for efficient and reliable human genome epidemiology. *Nat Genet* 2006;**38**:3–5.

- <sup>46</sup> Khoury MJ, Millikan R, Little J, Gwinn M. The emergence of epidemiology in the genomics age. *Int J Epidemiol* 2004;**33**:936–44.
- <sup>47</sup> Little J, Bradley L, Bray MS *et al*. Reporting, appraising, and integrating data on genotype prevalence and gene-disease associations. *Am J Epidemiol* 2002;**156**:300–10.
- <sup>48</sup> Collins R and UK Biobank Steering Committee. *UK Biobank: Protocol for a Large-scale Prospective Epidemiological Resource*. Manchester: UK Biobank Coordinating Centre, 2007.
- <sup>49</sup> Burton P, Fortier I, Deschenes M, Hansell A, Palmer L. Biobanks and biobank harmonization. In: Davey SG, Burton P, Palmer L (eds). *An Introduction to Genetic Epidemiology*. Bristol: Policy Press (In press).
- <sup>50</sup> Hunter DJ. Gene-environment interactions in human diseases. *Nat Rev Genet* 2005;**6**:287–98.
- <sup>51</sup> Weikert S, Boeing H, Pischon T *et al*. Fruits and vegetables and renal cell carcinoma: findings from the European prospective investigation into cancer and nutrition (EPIC). *Int J Cancer* 2006;**118**:3133–39.
- <sup>52</sup> Husebekk A, Iversen O-J, Langmark F, Laerum OD, Ottersen OP, Stoltenberg C. *Biobanks for Health - Report and Recommendations from an EU workshop*. Oslo: Technical report to EU Commission, 2003.
- <sup>53</sup> Wong MY, Day NE, Luan JA, Chan KP, Wareham NJ. The detection of gene-environment interaction for continuous traits: should we deal with measurement error by bigger studies or better measurement? *Int J Epidemiol* 2003;**32**:51–57.
- <sup>54</sup> Schwartz D, Collins F. Medicine: environmental biology and human disease. *Science* 2007;**316**:695–96.
- <sup>55</sup> R Development Core Team. *R: A Language and Environment for Statistical Computing*. Vienna, Austria: R Foundation for Statistical Computing, 2008.
- <sup>56</sup> Rohlfing CL, Little RR, Wiedmeyer HM *et al*. Use of GHb (HbA1c) in screening for undiagnosed diabetes in the U.S. population. *Diabetes Care* 2000;**23**:187–91.
- <sup>57</sup> Barrett JC, Cardon LR. Evaluating coverage of genome-wide association studies. *Nat Genet* 2006;**38**:659–62.
- <sup>58</sup> Burton P, Gurrin L, Sly P. Extending the simple linear regression model to account for correlated responses: an introduction to generalized estimating equations and multi-level mixed modelling. *Stat Med* 1998;**17**:1261–91.
- <sup>59</sup> de Bakker PI, Burt NP, Graham RR *et al*. Transferability of tag SNPs in genetic association studies in multiple populations. *Nat Genet* 2006;**38**:1298–303.
- <sup>60</sup> Jorgenson E, Witte JS. A gene-centric approach to genome-wide association studies. *Nat Rev Genet* 2006;**7**:885–91.
- <sup>61</sup> Gordon D, Finch SJ, Nothnagel M, Ott J. Power and sample size calculations for case-control genetic association tests when errors are present: application to single nucleotide polymorphisms. *Hum Hered* 2002;**54**:22–33.
- <sup>62</sup> Gordon D, Finch SJ. Factors affecting statistical power in the detection of genetic association. *J Clin Invest* 2005;**115**:1408–18.
- <sup>63</sup> Gordon D, Leal SM, Heath SC, Ott J. An analytic solution to single nucleotide polymorphism error-detection rates in nuclear families: implications for study design. *Pac Symp Biocomput* 2000;**5**:660–71.
- <sup>64</sup> Gordon D, Yang Y, Haynes C, Finch SJ, Mendell NR, Brown AM. Increasing power for tests of genetic association in the presence of phenotype and/or genotype error by use of double-sampling. *Stat Appl Genet Mol Biol* 2004;**3**:26.
- <sup>65</sup> Marchini J, Donnelly P, Cardon LR. Genome-wide strategies for detecting multiple loci that influence complex diseases. *Nat Genet* 2005;**37**:413–17.
- <sup>66</sup> Bermejo JL, Hemminki K. Gene-environment studies: any advantage over environmental studies? *Carcinogenesis* 2007;**28**:1526–32.
- <sup>67</sup> Armitage P, Berry G. *Statistical Methods in Medical Research*. 3rd edn. Oxford: Blackwell Scientific Publications, 1994.
- <sup>68</sup> Burton PR, Hansell A. *UK Biobank: The Expected Distribution of Incident and Prevalent Cases of Chronic Disease and the Statistical Power of Nested Casecontrol Studies*. Manchester, UK: UK Biobank Technical Reports, 2005.
- <sup>69</sup> Iles MM. What can genome-wide association studies tell us about the genetics of common disease? *PLoS Genetics* 2008;**4**:e33.
- <sup>70</sup> Davey Smith G, Ebrahim S, Lewis S, Hansell AL, Palmer LJ, Burton PR. Genetic epidemiology and public health: hope, hype, and future prospects. *Lancet* 2005;**366**:1484–98.
- <sup>71</sup> Taubes G. Epidemiology faces its limits. *Science* 1995;**269**:164–69.
- <sup>72</sup> Clayton D, McKeigue PM. Epidemiological methods for studying genes and environmental factors in complex diseases. *Lancet* 2001;**358**:1356–60.
- <sup>73</sup> Davey Smith G, Ebrahim S. ‘Mendelian randomization’: can genetic epidemiology contribute to understanding environmental determinants of disease? *Int J Epidemiol* 2003;**32**:1–22.
- <sup>74</sup> Tobin MD, Minelli C, Burton PR, Thompson JR. Commentary: development of Mendelian randomization: from hypothesis test to ‘Mendelian deconfounding’. *Int J Epidemiol* 2004;**33**:26–29.
- <sup>75</sup> Didelez V, Sheehan N. Mendelian randomization as an instrumental variable approach to causal inference. *Stat Methods Med Res* 2007;**16**:309–30.
- <sup>76</sup> Klipstein-Grobusch K, Georg T, Boeing H. Interviewer variability in anthropometric measurements and estimates of body composition. *Int J Epidemiol* 1997;**26** (Suppl 1):S174–80.
- <sup>77</sup> Rimm EB, Stampfer MJ, Colditz GA, Chute CG, Litin LB, Willett WC. Validity of self-reported waist and hip circumferences in men and women. *Epidemiology* 1990;**1**:466–73.
- <sup>78</sup> Dyer AR, Shipley M, Elliott P, for the Intersalt Cooperative Research Group. Urinary electrolyte excretion in 24 hours and blood pressure in the INTERSALT study: I. Estimates of Reliability. *Am J Epidemiol* 1994;**139**:927–39.
- <sup>79</sup> Grandits GA, Bartsch GE, Stamler J. Chapter 4. Method issues in dietary data analyses in the multiple risk factor intervention trial. *Am J Clin Nutr* 1997;**65** (Suppl):211S–27S.