

Likelihood Formulation of Parent-of-Origin Effects on Segregation Analysis, Including Ascertainment

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We developed a likelihood-based method for testing for parent-of-origin effect in complex diseases. The likelihood formulations model parent-of-origin effect and allow for incorporation of ascertainment, as well as differential male and female ascertainment probabilities. The results based on simulated data indicated that the estimates of parental effect (either maternal or paternal) were biased when ascertainment was ignored or when the wrong ascertainment model was used. The exception was single ascertainment, in which we proved that ignoring ascertainment does not bias the estimation of parental effect, in a simple parent-of-origin model. These results underscore the importance of considering ascertainment models when testing for parent-of-origin effect in complex diseases.

Introduction

Parent-of-origin effect refers to differential penetrance or expression of disease in the offspring, depending on the sex of the transmitting parent. Parent-of-origin effect may encompass several possible underlying biological phenomena, including genomic imprinting, trinucleotide-repeat expansion, or mitochondrial inheritance. Genomic imprinting (also referred to as “gametic” or “parental” imprinting) is the epigenetic marking of a gene, on the basis of its parental origin, that results in monoallelic expression. Genomic imprinting differs from classical Mendelian genetics in that the parental complements of imprinted genes are not equivalent with respect to their expression. In maternal imprinting, gene expression is inhibited after passage through the mother’s germline, whereas, in paternal imprinting, gene expression is inhibited after passage through the father’s germline. Prader-Willi syndrome (maternally imprinted) and Angelman syndrome (paternally imprinted) are two classic examples of the numerous human diseases in which the effects of imprinting are observed (Falls et al. 1999). Studies have revealed that genomic imprinting has the following intrinsic properties: silencing of gene expression, stable propagation in dividing somatic cells, possible reversal of the imprint pattern under certain conditions, and establishment of the imprint during gametogenesis. The mechanism(s) of genomic imprinting are complex and not well understood; however, evi-

dence suggests that methylation is a likely candidate, since it satisfies the aforementioned criteria (Tycko et al. 1997; Constanca et al. 1998).

Another biological phenomenon that can result in a parent-of-origin effect is trinucleotide-repeat expansion. Instability in expansion of trinucleotide repeats (e.g., CAG, CGG, CTG, and GAA) is observed during germline transmission when the length of the repeat exceeds a critical value (Reddy and Housman 1997). The instability is generally observed when the transmitted repeat size is 40–100 bp. Studies of individuals diagnosed with early-onset Huntington disease have revealed a significant increase in sperm trinucleotide repeat (CAG) lengths compared with the repeat lengths of the father. On the other hand, in Fragile X syndrome, expansion of trinucleotide repeats (CGG) in the *FMR1* gene is observed when the gene is transmitted maternally. Numerous models have been proposed to explain the triplet-repeat expansion leading to human disease (Pearson and Sinden 1998). For example, one model involves the formation of DNA hairpin structures that lead to errors in replication (e.g., replication slippage) and/or promote recombination via unequal sister-chromatid exchange.

A third phenomenon is mitochondrial inheritance, which manifests a transmission pattern consistent with parent-of-origin effect. mtDNA is almost exclusively maternally inherited (Lightowlers et al. 1997), but a mitochondrial disorder may exhibit either a maternal or a Mendelian inheritance pattern, depending on the site of the primary gene defect. Leber hereditary optic neuropathy was the first disease found to be caused by a point mutation in mtDNA (Wallace et al. 1988); since then, numerous other mutations that lead to diseases such as myoclonic epilepsy and ragged-red fibers, mitochondrial encephalomyopathy, lactic acidosis and

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stroke-like episodes, and progressive external ophthalmoplegia have been found. The clinical spectrum of possible mitochondrial defects has been expanded to include several common disorders. In disorders such as Parkinson disease and Alzheimer disease, mitochondrial defects may not be the primary cause but have been suggested to modify the outcome of disease (Suomalainen 1997).

The phenomenon of parent-of-origin effect has been investigated in the transmission of neurological and psychiatric diseases such as Tourette syndrome, bipolar disorder, and panic disorder (Lichter et al. 1995; McMahon et al. 1995; Stine et al. 1995; Gershon et al. 1996; Kato et al. 1996; Eapen et al. 1997; Battaglia et al. 1999; Haghighi et al. 1999). In previous studies, similar approaches have been adopted in the analysis of disease transmission. Some investigators have systematically ascertained two-generation pedigrees and have dichotomized their data into maternal- and paternal-transmission groups (McMahon et al. 1995). Others have utilized multigenerational pedigrees and have divided them into maternal- and paternal-transmission branches (Gershon et al. 1996; Haghighi et al. 1999). The latter approach discards valuable information by breaking down multigenerational pedigrees into maternal and paternal branches and by ignoring parental mating types (MTs) in which maternal or paternal transmission cannot be determined (i.e., unaffected \times unaffected and affected \times affected MTs). In the consideration of the maternal and paternal branches, these methods do not account for the potential transmission of the disease gene through the unaffected parent (via reduced penetrance). Also, these methods do not incorporate ascertainment models. Consequently, we developed a likelihood-based approach that utilizes all available data for testing for parent-of-origin effect, allowing for modeling of ascertainment (Haghighi and Hodge 1999).

The likelihood-based method presented here models parent-of-origin effect in nuclear families, for a single locus. Extension of this model to general pedigree structures is described in the "Discussion" section. The likelihood calculation handles all possible parental MTs and variable sibship sizes. For each family in the data set, the exact likelihood is computed, allowing for reduced penetrance. This entails consideration of possible disease-gene transmission from unaffected parent(s) to offspring who, in turn, may or may not express the disease phenotype. The likelihood is parameterized to model parental effect, by including the penetrances of maternal and paternal transmission. To assess the potential effect that ascertainment has on detection of parent-of-origin effect or on estimation of penetrances of maternal and paternal transmission, we also incorporated a general ascertainment model into our likelihood formulation. We demonstrated that, in the special case of single as-

certainment, no correction needs to be made for ascertainment (for this simple model).

The two goals of this paper are (1) to formulate the correct full likelihood for parent-of-origin effect in nuclear families, incorporating the " π "-based ascertainment model of Weinberg (1928) and Morton (1959) but also allowing for differential male and female ascertainment probabilities, and (2) to determine the effects that ascertainment has on our ability to detect parent-of-origin effect. We demonstrate the likelihood derivation and assess its utility, using simulated data generated under a range of inheritance and ascertainment models. The two principal likelihood models that are presented consist of the parental-effect model and the parental-effect-with-ascertainment model. These models were systematically studied by using simulated data generated under maternal or paternal parental effects with "complete" or "single" ascertainment. The likelihood models were evaluated by examining the estimated parental effect and the power to detect such an effect in the presence or absence of ascertainment.

Methods

Notation

To test for parent-of-origin effect, we have developed two likelihood models. Model I models parent-of-origin effect but does not incorporate ascertainment. This model would apply to situations of "random" ascertainment (see the "Discussion" section). Model II models parent-of-origin effect and also incorporates ascertainment, as well as allowing for differential male and female ascertainment probabilities. Before describing the models in detail, we will define the parameters used in the likelihood formulations:

- q = frequency of disease allele (denoted by " D ");
- p = frequency of nondisease allele (denoted by " d "), where $p = 1 - q$;
- f = disease penetrance, defined as the mean between the maternal- and paternal-transmission penetrances, $f = (f_m + f_p)/2$ (see below);
- δ = deviation of the maternal- and paternal-transmission penetrances from the mean penetrance, where δ may be either positive or negative—that is, $\delta = (f_m + f_p)/2$ so that
 - f_m = maternal-transmission penetrance, defined as $f + \delta$;
 - f_p = paternal-transmission penetrance, defined as $f - \delta$;
 - $f_{mp} = f_m + f_p - f_m f_p$;
 - (note that explanations for each of these penetrance probabilities are given in the subsection "Likelihood Model I," below);
- π_b = male ascertainment probability—that is, $P(\text{male is$

a proband|he is affected);
 π_g = female ascertainment probability—that is, $P(\text{female is a proband|she is affected})$;
 s_b = number of male offspring in a sibship;
 s_g = number of female offspring in a sibship;
 s = sibship size, defined as $s = s_b + s_g$;
 r_b = number of affected male offspring;
 r_g = number of affected female offspring;
 r = number of affected offspring, defined as $r = r_b + r_g$.

Note that “b” (for “boy”) and “g” (for “girl”) are used to denote male and female offspring, respectively. Also, in both likelihood models, I and II, δ is the parameter of interest; it is a nongenetic (i.e., “dummy”) parameter, which is used as an indicator of maternal or paternal transmission. The use of δ in this manner reduces model complexity, since the maternal- and the paternal-transmission penetrances are both defined with respect to one parameter (see the “Discussion” section).

Likelihood Model I

We began by calculating the exact likelihood for each of the four phenotypic parental MTs: (1) affected mother \times unaffected father, (2) unaffected mother \times affected father, (3) unaffected mother \times unaffected father, and (4) affected mother \times affected father. We assumed an autosomal dominant mode of inheritance. For each phenotypic MT, we enumerated the possible underlying parental genotypes consistent with the genetic model. These parental genotypes were then used to enumerate the possible offspring genotypes and to assign probabilities to them (see equation (1), below). A sample likelihood computation for the unaffected mother \times unaffected father MT is shown in table 1, with corresponding parental probabilities, as well as offspring penetrance and transmission probabilities. We chose to illustrate this particular MT because it illustrates the use of all possible underlying parental genotypes. The remaining parental MTs are calculated in the same fashion, except that, depending on the parental phenotypes, not all parental genotypes are possible. Thus, the likelihood tables for these other MTs will contain some empty cells.

The exact likelihood calculation described above can be formulated as a simple probability, which is $P(\phi_c, \phi_m, \phi_p)$, where ϕ_c denotes the vector $(\phi_{c_1}, \dots, \phi_{c_s})$ of the observed phenotypes of the s children, c_1, \dots, c_s ; and ϕ_m and ϕ_p denote the observed maternal and paternal phenotypes, respectively. Using the law of total probability, we rewrite this probability to allow for all underlying genotypes, where g_m and g_p denote maternal and paternal genotypes, respectively (the symbols “ ϕ_c ”

and “ g_c ,” not in boldface, denote the phenotype and the genotype, respectively, of a single child):

$$\begin{aligned}
 P(\phi_c, \phi_m, \phi_p) &= \sum_{g_m} \sum_{g_p} P(\phi_c | \phi_m, \phi_p, g_m, g_p) P(\phi_m, \phi_p | g_m, g_p) \\
 &= \sum_{g_m} \sum_{g_p} P(\phi_c | g_m, g_p) P(\phi_m | g_m) P(g_m) P(\phi_p | g_p) P(g_p) \\
 &= \sum_{g_m} \sum_{g_p} P(\phi_m | g_m) P(g_m) P(\phi_p | g_p) P(g_p) \\
 &\quad \times [P(\phi_c = \text{aff} | g_m, g_p)]^r [P(\phi_c = \text{unaff} | g_m, g_p)]^{s-r} \\
 &= \sum_{g_m} \sum_{g_p} P(\phi_m | g_m) P(g_m) P(\phi_p | g_p) P(g_p) \\
 &\quad \times \left[\sum_{g_c} P(\phi_c = \text{aff} | g_c, g_m, g_p) P(g_c | g_m, g_p) \right]^r \\
 &\quad \times \left[\sum_{g_c} P(\phi_c = \text{unaff} | g_c, g_m, g_p) P(g_c | g_m, g_p) \right]^{s-r}.
 \end{aligned} \tag{1}$$

This is essentially Elston and Stewart’s (1971) algorithm, except that, to model parent-of-origin effect, we condition the offspring phenotypes on the parental genotypes in addition to the usual offspring genotypes.

The probabilities in the last expression in equation (1) can be found in table 1. They are all derived from the usual functions of penetrance f and from Mendel’s first law, except the terms $P(\phi_c | g_c, g_m, g_p)$, which we now explain:

$P(\phi_c = \text{aff} | g_c = dd, g_m, g_p)$ is assumed to be 0, independent of g_m and g_p .

$P(\phi_c = \text{aff} | g_c = Dd, g_m, g_p)$ is taken to equal f_m (or f_p), if the Dd child clearly inherited the D allele from the mother (or father) (i.e., if one parent is Dd and the other parent is either DD or dd , or if one parent is DD and the other parent is dd). If either parent is equally likely to have contributed the D allele (i.e., if both parents are Dd), then $P(\phi_c = \text{aff} | g_c = Dd, g_m = Dd, g_p = Dd)$ is set to $f = (1/2)(f_m + f_p)$ —that is, it is the mean of the two parental-transmission penetrances.

For $P(\phi_c = \text{aff} | g_c = DD, g_m, g_p)$, the DD child must have received the D allele from both parents. This probability is found by first considering its complement, $P(\phi_c = \text{unaff} | g_c = DD, g_m, g_p)$. For a child with a DD genotype to be unaffected requires that the D allele not be “expressed” from either parent; thus, $P(\phi_c = \text{unaff} | g_c = DD, g_m, g_p) = (1 - f_m)(1 - f_p)$, under the assumption of the independence of the two parents. Therefore, $P(\phi_c = \text{aff} | g_c = DD, g_m, g_p) = 1 - (1 - f_m)(1 - f_p) = f_m + f_p - f_m f_p$, which we subsequently write as “ f_{mp} ” for short.

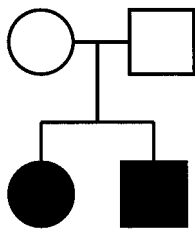


Figure 1 Pedigree used for illustration of likelihood calculation

We demonstrate an actual likelihood calculation using the family in figure 1. For the sake of simplicity, we assume that the disease allele is rare (i.e., q is very small), only for this example. This reduces the likely set of parental genotypes to the following ($g_m \times g_p$): $Dd \times dd$ and $dd \times Dd$. (This is a simplified example; in our actual likelihood calculations, all parental genotypes are considered.) We focus on the MT $Dd \times dd$, to illustrate the terms in equation (1). In this case, the parental penetrance terms $P(\phi_m|g_m)$ and $P(\phi_p|g_p)$ become $P(\phi_m = \text{unaff}|g_m = Dd) = 1 - f$ and $P(\phi_p = \text{unaff}|g_p = dd) = 1$, respectively. Similarly, the parental genotypic probabilities $P(g_m)$ and $P(g_p)$ become $P(Dd) = 2pq$ and $P(dd) = p^2$, respectively. The penetrance and transmission probabilities for the offspring are $P(\phi_c|g_c, g_m, g_p)$ and $P(g_c|g_m, g_p)$, which, for the affected children, become $P(\phi_c = \text{aff}|g_c = Dd, g_m = Dd, g_p = dd) = f_m$ and $P(g_c = Dd|g_m = Dd, g_p = dd) = 1/2$. Note that both children must have genotype Dd . The probabilities for the remaining MT are taken

from table 1. Thus, the (simplified) likelihood for this family is

$$L = 2p^3q(1 - f) \left(\frac{1}{2}f_m\right)^2 + 2p^3q(1 - f) \left(\frac{1}{2}f_p\right)^2,$$

where the two terms correspond to the $Dd \times dd$ and $dd \times Dd$ parental mating genotypes, respectively. If we (a) did not assume that q is small and (b) included all eight possible parental genotypic MTs, the complete likelihood for this family would be,

$$L = 2p^3q(1 - f) \left[\left(\frac{1}{2}f_m\right)^2 + \left(\frac{1}{2}f_p\right)^2 \right] + 4p^2q^2(1 - f)^2 \left[\frac{1}{4}(f_m + f_p) + \frac{1}{4}f_{mp} \right]^2 + p^2q^2(1 - f)(f_m^2 + f_p^2) + q^4(1 - f)^2(f_{mp})^2 + 2pq^3(1 - f)^2 \left[\left(\frac{1}{2}f_m + \frac{1}{2}f_{mp}\right)^2 + \left(\frac{1}{2}f_p + \frac{1}{2}f_{mp}\right)^2 \right],$$

and this is indeed what our program calculates.

Likelihood Model II

The likelihood model (i.e., model I, described above) was extended to incorporate ascertainment. This second model—model II—allows us to assess the potential influence that ascertainment has on detection of parent-of-origin effect. The potential for ascertainment bias, as is well known, is always a consideration in segregation analysis. In this case, the problem of ascertainment bias

Table 1

Transmission Probabilities (of Offspring Genotypes) and Penetrances (of Offspring Phenotypes), Conditioned on Indicated Parental Genotypes, for Parental Genotypes Compatible with Unaffected Mother \times Unaffected Father MT

UNAFFECTED FATHER'S GENOTYPE	TRANSMISSION PROBABILITIES WHEN UNAFFECTED MOTHER'S GENOTYPE IS		
	$dd; p^2$	$Dd; 2pq(1 - f)$	$DD; q^2(1 - f)$
$dd; p^2$	$dd; P(dd) = 1, P(\text{aff} dd) = 0, P(\text{unaff} dd) = 1$	$dd; P(dd) = .5, P(\text{aff} dd) = 0, P(\text{unaff} dd) = 1$ $Dd; P(Dd) = .5, P(\text{aff} Dd) = f_m, P(\text{unaff} Dd) = 1 - f_m$	$Dd; P(Dd) = 1, P(\text{aff} Dd) = f_m, P(\text{unaff} Dd) = 1 - f_m$
$Dd; 2pq(1 - f)$	$dd; P(dd) = .5, P(\text{aff} dd) = 0, P(\text{unaff} dd) = 1$ $Dd; P(Dd) = .5, P(\text{aff} Dd) = f_p, P(\text{unaff} Dd) = 1 - f_p$	$dd; P(dd) = .25, P(\text{aff} dd) = 0, P(\text{unaff} dd) = 1$ $Dd; P(Dd) = .5, P(\text{aff} Dd) = (1/2)(f_m + f_p), P(\text{unaff} Dd) = 1 - (1/2)(f_m + f_p)$ $DD; P(DD) = .25, P(\text{aff} DD) = f_{mp}, P(\text{unaff} DD) = 1 - f_{mp}$	$Dd; P(Dd) = .5, P(\text{aff} Dd) = f_m, P(\text{unaff} Dd) = 1 - f_m$ $DD; P(DD) = .5, P(\text{aff} DD) = f_{mp}, P(\text{unaff} DD) = 1 - f_{mp}$
$DD; q^2(1 - f)$	$Dd; P(Dd) = 1, P(\text{aff} Dd) = f_p, P(\text{unaff} Dd) = 1 - f_p$	$Dd; P(Dd) = .5, P(\text{aff} Dd) = f_p, P(\text{unaff} Dd) = 1 - f_p$ $DD; P(DD) = .5, P(\text{aff} DD) = f_{mp}, P(\text{unaff} DD) = 1 - f_{mp}$	$DD; P(DD) = 1, P(\text{aff} DD) = f_{mp}, P(\text{unaff} DD) = 1 - f_{mp}$

NOTE.—We define f_{mp} as $f_m + f_p - f_m f_p$ (see text).

is also a concern because estimation of parental effect falls under the rubric of segregation analysis.

The ascertainment model assumes that families are ascertained through children (Weinberg 1928; Morton 1959) and incorporates sex-based ascertainment in the likelihood. The likelihood with differential male and female ascertainment probabilities is as follows:

$$P(r_g, r_b, \phi_m, \phi_p | \text{family ascertained}) = \frac{P(\text{family ascertained} | r_g, r_b) P(r_g, r_b, \phi_m, \phi_p)}{P(\text{family ascertained})} \quad (2)$$

Note that $P(\text{family ascertained} | \text{parental and child phenotypes}) = P(\text{family ascertained} | \text{child phenotypes})$, since we assume that families are ascertained through the children. Hence, the first term in the numerator of equation (2) is simply $P(\text{family ascertained} | r_g, r_b)$. This ascertainment term, $P(\text{family ascertained} | r_g, r_b)$, allows for modeling of general ascertainment, by a variety of possible ascertainment criteria, such as sex (male vs. female), disease subtypes (e.g., early onset vs. late onset and mild vs. severe), and so on. In this situation, we were interested in modeling the differential male and female ascertainment probabilities, so, for any π_g and π_b , this ascertainment probability for a family can be expressed as $1 - (1 - \pi_g)^{r_g} (1 - \pi_b)^{r_b}$. The second term in the numerator, $P(r_g, r_b, \phi_m, \phi_p)$, corresponds to $P(\phi_c, \phi_m, \phi_p)$ derived in equation (1), here with $r = r_g + r_b$. Thus, the numerator of equation (2) becomes

$$[1 - (1 - \pi_g)^{r_g} (1 - \pi_b)^{r_b}] P(r_g, r_b, \phi_m, \phi_p) .$$

The denominator of equation (2) is found by summing the numerator over all possible configurations for sibship sizes and all possible parental phenotypes:

$$\sum_{s_b=0}^s \sum_{r=1}^s \sum_{r_b=\max(0, r-s_g)}^{\min(r, s_b)} [1 - (1 - \pi_g)^{r_g} (1 - \pi_b)^{r_b}] \times \sum_{\phi_m} \sum_{\phi_p} P(r_g, r_b, \phi_m, \phi_p) ,$$

where $r_g = r - r_b$ and $s_g = s - s_b$. Proceeding from left to right, the first summation traverses through all sibship configurations with respect to sex; the second summation enumerates all sibship configurations with at least $r = 1$ to, at most, $r = s$ affected children; and the third summation keeps track of the sex of the affected children, which is used in the ascertainment term. The last two (internal) summations traverse through all possible parental phenotypes. Thus, the expanded form of equa-

tion (2) for unequal male and female ascertainment probabilities is

$$P(r_g, r_b, \phi_m, \phi_p | \text{family ascertained}) = \{ [1 - (1 - \pi_g)^{r_g} (1 - \pi_b)^{r_b}] P(r_g, r_b, \phi_m, \phi_p) \} / \left\{ \sum_{s_b=0}^s \sum_{r=1}^s \sum_{r_b=\max(0, r-s_g)}^{\min(r, s_b)} [1 - (1 - \pi_g)^{r_g} (1 - \pi_b)^{r_b}] \times \sum_{\phi_m} \sum_{\phi_p} P(r_g, r_b, \phi_m, \phi_p) \right\} , \quad (3)$$

where each $P(r_g, r_b, \phi_m, \phi_p)$ is given by equation (1).

For the special case in which the male and female ascertainment probabilities are equal (i.e., $\pi_g = \pi_b = \pi$), equation (3) is simplified, such that

$$P(r, \phi_m, \phi_p | \text{family ascertained}) = \frac{P(\text{family ascertained} | r) P(r, \phi_m, \phi_p)}{P(\text{family ascertained})} .$$

In the numerator, the ascertainment term is now $P(\text{family ascertained} | r) = 1 - (1 - \pi)^r$, and $P(r, \phi_m, \phi_p)$ is as in equation (1). In the denominator, the sibship is traversed with respect to affection status but not sex. The final probability for equal male and female ascertainment probabilities becomes

$$P(r, \phi_m, \phi_p | \text{family ascertained}) = \frac{[1 - (1 - \pi)^r] P(r, \phi_m, \phi_p)}{\sum_{r=1}^s [1 - (1 - \pi)^r] \sum_{\phi_m} \sum_{\phi_p} P(r, \phi_m, \phi_p)} . \quad (4)$$

Models I and II described above give the full likelihood for a single nuclear family. The likelihood for a collection of families is found by multiplying the individual family likelihoods over all families:

$$\prod_{\text{all families}} L(\text{family}) ,$$

where the likelihood of the individual family is given by equation (1), for model I, and by equations (3) or (4), for model II.

Details of Simulation and Analysis

We designed and implemented a simulation program (`sim_poo.pl`) to simulate nuclear families under the likelihood models described above. The parental MTs (i.e., both genotypes and phenotypes) were randomly generated, on the basis of Hardy-Weinberg proportions and the user-defined disease penetrances and allele fre-

quencies. Next, offspring genotypes were generated assuming Mendelian laws of inheritance. The affection statuses of the offspring were randomly determined, given the user-defined maternal- and paternal-transmission penetrance probabilities and conditioning on offspring and parental genotypes.

We examined three situations (by “situation,” we mean a combination of generating model [GM], analysis model [AM], family structure, and data-set size).

Situation 1. Fixed family structure, rare disease ($q = 0.0001$). Data set includes 100 families (four sibs/sibship). GM includes complete and single ascertainment; selected values of f and of δ . AM includes complete, single, and random ascertainment.

Situation 2. Variable sibship sizes. Data set includes 50 families; otherwise same as situation 1.

Situation 3. Higher gene frequency ($q = 0.1$); otherwise, same as situation 2, except that AM includes only complete and random ascertainment.

For each situation, we evaluated some or all of the following: (a) bias in the estimate of δ , (b) power to detect parent-of-origin effect when $\delta > 0$, and (c) type I error rate when there is no parent-of-origin effect (i.e., when $\delta = 0$). To assess bias in the estimate of δ , we used the sample mean of the maximum-likelihood estimates (MLEs) of δ (henceforth denoted simply as “ $\hat{\delta}$ ”). For detection of power, we used the asymptotic approximation, $2\ln(LR) \sim \chi^2(1 \text{ df})$, where LR is the likelihood ratio of $L(\hat{\delta})$ versus $L(\delta = 0)$. For analysis of type I error rate (α), we set the nominal test size to 0.05 and then determined the actual test size from our simulations. Power was computed for a test size of $\alpha = 0.05$.

Each simulated family was then considered for inclusion in the sample, subject to a user-specified ascertainment criterion (i.e., “complete” ascertainment, single ascertainment, or “random” ascertainment). Under complete ascertainment, the family was ascertained with probability unity if there was at least one affected child in the sibship. (Thus, by “complete” ascertainment we mean the case in which $\pi = 1$. This situation was originally termed “truncate” ascertainment by Morton [1959], because the corresponding probability distribution is a truncated binomial distribution; however, since many investigators currently refer to this model as “complete” ascertainment, we use that terminology in this study as well.) Under single ascertainment, the probability that any one family will be ascertained is small and is proportional to the number of affected children in the family (Morton 1959; Stene 1979; Hodge and Vieland 1996) (also see Appendix A). Under random ascertainment, all families, including those with no affected children, are ascertained with a probability of unity.

The families were then analyzed with the `calc_poo.pl` program, which implements the aforementioned likeli-

hood-based algorithms. The program analyzes the data under the assumption of complete, single, or random ascertainment. It yields the logarithm of LR , $\ln(LR)$, by comparing the likelihood for a range of δ values and the likelihood of $\delta = 0$ (i.e., no parent-of-origin effect). The $\hat{\delta}$ value is recorded for each data set, and the sample mean of these estimates is calculated over all data sets. The number of data sets considered was 500–1,000, with 50–100 families per data set. The programs `sim_poo.pl` and `calc_poo.pl` were written in PERL and are available on request.

Results

We evaluated the likelihood models by simulation analyses, in which we incrementally increased the complexity of the data to emulate “real” data sets. The results are presented for a number of GMs and AMs. The models covered a range of f and δ parameter values and ascertainment criteria (i.e., complete ascertainment, single ascertainment, and “random” ascertainment). In the likelihood calculations, complete ascertainment corresponds to $\pi_g = \pi_b = 1.0$, whereas single ascertainment corresponds to $\pi_g = \pi_b = 0.01$. Random ascertainment means that the likelihoods were computed under model I, which assumes that all families are equally likely to be ascertained whether they have any affected children or not. For all analyses described below, values of f and of q in the AM match those in the GM.

Situation 1.—The fixed family structure and the rare-disease ($q = 0.0001$) model enabled us to easily confirm the simulation results analytically in any given data set. The data sets consisted of 100 families, each of which had exactly four sibs per sibship. The data were generated under complete ascertainment and then were analyzed assuming complete, single, and random ascertainment (table 2). We observed that, when the AM and GM were the same, $\hat{\delta}$ was approximately unbiased, except when the value of f was near the defined boundary limits (0 or 1) and/or δ was small (e.g., $f = 0.9$ and

Table 2
Observed $\hat{\delta}$ Values from Simulations of Situation 1, Generated under Complete Ascertainment

AM	f	δ				
		.1	.2	.3	.4	.5
Complete ascertainment	.9	.072
	.7	.101	.193	.280
	.5	.097	.198	.300	.399	.500
Single ascertainment	.9	.066
	.7	.062	.121	.195
	.5	.049	.105	.178	.269	.432
Random ascertainment	.9	.066
	.7	.061	.121	.194
	.5	.048	.105	.177	.268	.430

Table 3
Observed $\hat{\delta}$ Values from Simulations of Situation 1, Generated under Single Ascertainment

AM	<i>f</i>	δ				
		.1	.2	.3	.4	.5
Complete ascertainment	.9	.082
	.7	.160	.272	.300
	.5	.189	.316	.404	.466	.500
Single ascertainment	.9	.076
	.7	.100	.198	.280
	.5	.101	.201	.298	.400	.499
Random ascertainment	.9	.076
	.7	.100	.197	.279
	.5	.101	.200	.297	.399	.499

$\delta = 0.1$). However, when the data were incorrectly analyzed assuming single or random ascertainment, the $\hat{\delta}$ values were consistently lower than they were when the correct ascertainment model had been used. The $\hat{\delta}$ values analyzed assuming random ascertainment were approximately equal to those for single ascertainment. In addition, data were generated under single ascertainment and were analyzed assuming complete, single and random ascertainment (table 3). Similar to the previous results, $\hat{\delta}$ was approximately unbiased when the AM matched the underlying GM. Again, the $\hat{\delta}$ values for single and random ascertainment were virtually identical (see “Discussion” and Appendix A). However, when the data were analyzed assuming complete ascertainment the $\hat{\delta}$ values were inflated.

Situation 2.—We extended our data sets to include families with variable sibship sizes, to better emulate realistic data sets with different family configurations. Also, we chose to use data sets that contained 50 families, to evaluate the performance of the likelihood models, since this would be a reasonably attainable size for a real data set. These analyses were performed using data simulated under a rare-disease model with the same parameter values and the same ascertainment models as before (tables 4 and 5). Even with the smaller data-set size, the trends in the $\hat{\delta}$ values were consistent with those from the first set of analyses (tables 2 and 3). Specifically, for the analyses in which the AMs were the same as the GMs, $\hat{\delta}$ was generally unbiased. The $\hat{\delta}$ values were either lower (table 4) or higher (table 5) than expected, when the wrong ascertainment model was used in the analysis. Again, the exception occurs for single ascertainment, in which ignoring ascertainment does not bias $\hat{\delta}$ (see table 5, “Discussion,” and Appendix A). The $\hat{\delta}$ values that we observed when the data were analyzed assuming single or random ascertainment were always very similar.

In situation 2, we also investigated the *power* of the likelihood models to detect parent-of-origin effect. As expected, we observed higher power when the AM

matched the true GM (fig. 2A). In contrast, when the data were generated under single ascertainment, we found that power was higher when the data were analyzed assuming complete ascertainment (fig. 2B). Again, the observed power levels for analyses under single and random ascertainment were very close. Overall, the power increased with increasing parental effect in the data set (i.e., the power increased with increasing δ), as one would expect. For these analyses, power to detect a maternal effect converged to unity for $\delta > 0.3$.

Last, for situation 2, we examined the performance of our likelihood-based methods in the *absence* of parent-of-origin effect, to assess type I error. In this case, the data were simulated with no parental effect ($\delta = 0$), under both complete and single ascertainment, and were analyzed assuming complete, single, and random ascertainment. For these models, the $\hat{\delta}$ values did not appear to be influenced by potential ascertainment bias in which the resulting $\hat{\delta}$ values were approximately equal to 0. Furthermore, for the same models, we found that the type I error rate did not exceed the nominal size of the test (0.05), except in one analysis (table 6). This exception occurred when the data were generated under single ascertainment and were analyzed assuming complete ascertainment. Note that, in the presence of a parental effect, this particular GM/AM combination always gave inflated $\hat{\delta}$ values, yielding a higher proportion of individual data sets with inflated $\hat{\delta}$ values, thus resulting in a higher type I error.

Situation 3.—So far, all the analyses involved data generated under a rare disease model. However, our goal was to evaluate the likelihood models in the context of complex common diseases with potential parent-of-origin effect. Therefore, in situation 3 a higher disease-gene frequency ($q = 0.1$) was used. This situation is similar to situation 2 because the data sets were generated with 50 families of variable sibship sizes under complete ascertainment and were analyzed assuming complete or random ascertainment (table 7). We found

Table 4
Observed $\hat{\delta}$ Values from Simulations of Situation 2, Generated under Complete Ascertainment

AM	<i>f</i>	δ				
		.1	.2	.3	.4	.5
Complete ascertainment	.9	.061
	.7	.095	.193	.267
	.5	.095	.196	.295	.397	.499
Single ascertainment	.9	.057
	.7	.067	.136	.206
	.5	.056	.118	.192	.289	.447
Random ascertainment	.9	.057
	.7	.064	.135	.205
	.5	.055	.118	.191	.287	.445

Table 5
Observed $\hat{\delta}$ Values from Simulations of Situation 2, Generated under Single Ascertainment

AM	f	δ				
		.1	.2	.3	.4	.5
Complete ascertainment	.9	.073
	.7	.139	.252	.296
	.5	.160	.297	.396	.464	.500
Single ascertainment	.9	.062
	.7	.100	.198	.275
	.5	.097	.199	.297	.400	.498
Random ascertainment	.9	.069
	.7	.099	.197	.275
	.5	.096	.198	.296	.399	.498

that the trends in the estimates of parental effect were the same for both the common- and rare-disease models (tables 4 and 7). However, the $\hat{\delta}$ values were slightly lower for the common- versus the rare-disease model, for both AMs examined. This was most likely due to an increase in the proportion of uninformative parental MTs, because of the higher disease-gene frequency in the population.

We also performed power simulations in situation 3. For the two AMs, power to detect parental effect was consistently lower under the common-disease model. This is because, for common diseases, the proportion of informative parental MTs with distinct maternal or paternal disease-gene transmission is smaller than that for rare diseases. When the data were analyzed assuming complete ascertainment, in which the AM and GM were the same, the maximum drop in power was 22% at $\delta = 0.2$ (fig. 3A). When the same data were analyzed assuming the wrong ascertainment criterion (i.e., random ascertainment), there was a dramatic reduction in power, in which the observed drop was as high as 89% at $\delta = 0.1$ (fig. 3B). Similar to the previous observations' estimates of power for both disease models, the estimates of power increased as the magnitude of parental effect (i.e., δ) increased. Note that, for all the results presented, the data were simulated for maternal transmission (positive δ s); however, when the data were generated for paternal transmission (negative δ s) the results were symmetric (data not shown).

Discussion

Summary

In this study, we first formulated likelihood-based models for parent-of-origin effect in transmission of disease, allowing for ascertainment in nuclear families. This likelihood formulation was then used to test for the existence of parent-of-origin effect and to estimate the size of this effect (δ), in the presence of different ascertain-

ment models. Second, we evaluated the performance of the procedure under different circumstances, by means of simulation analyses. In this way, we assessed whether ascertainment bias would affect test results or estimates of δ . We showed that the estimates of δ were approximately unbiased when the data were analyzed for the same parametric GM, except when f and δ were near their defined boundary limits. The estimates of δ were biased when ascertainment was ignored or when the wrong ascertainment model was assumed. The only exception was for single ascertainment, in which the ignoring of ascertainment does not bias the estimates of δ (see below).

We examined only dominant modes of inheritance—because, in these models, the origin of the disease allele can often be determined on the basis of its maternal or paternal transmission, whereas, in recessive inheritance, both parents transmit the disease allele to the affected offspring. This may be problematic for studies of common complex diseases in which the disease gene(s) is frequent in the population, since a high proportion of the parents would be homozygous. Also, the lower the overall penetrance of the condition being studied, the more families there will be in which the transmitting parent is not phenotypically affected, so that it is not obvious whether the mother or father is the transmitting parent. An additional limitation in the model concerns our assumption of disease penetrances. The disease penetrance f is defined as the average of maternal- and paternal-transmission penetrances. In the model considered here, this average penetrance also equals the population-wide penetrance of the disease. In a more complex model (e.g., a model with differential penetrances for male and female *individuals*, in addition to the differential effects from male and female transmitting parents), that would no longer be the case. Also note that, in our study, the penetrance parameters are taken at a single time point and do not allow for potential age and environmental effects (e.g., drug exposure).

In our model, we assigned a disease risk of f_{mp} to homozygous DD children (i.e., children who have received a D allele from both parents), where f_{mp} is defined as $f_m + f_p - f_m f_p$. This definition of f_{mp} can be justified by a model such as the following: The child receives a “hit” from the maternal D allele with probability f_m and a hit from the paternal D allele with probability f_p ; if the child receives at least one hit, then the child is affected. Dr. Gary Chase (personal communication) has pointed out that, alternatively, one could formulate a regression model such as this: let f_m and f_p represent the regression coefficients in a binary model, where $Y = 1$ if the child is affected and $Y = 0$ otherwise; let f_m (or f_p) represent the lifetime-risk increase associated with maternal (or paternal) transmission. Thus, our formula for f_{mp} rep-

Table 6

Observed Type I Error Rates from Simulations of Situation 2, in Which Nominal Test Size is 0.05

AM ASSUMED	TYPE I ERROR RATE UNDER GM					
	Complete Ascertainment			Single Ascertainment		
	$f = .9$	$f = .7$	$f = .5$	$f = .9$	$f = .7$	$f = .5$
Complete ascertainment	.046	.045	.046	.060	.096	.144
Single ascertainment	.041	.045	.046	.041	.045	.046
Random ascertainment	.041	.045	.044	.041	.045	.044

resents one particular type of interaction, but other types could be modeled as well.

Importance of Ascertainment

It is well known that the ascertainment model plays a critical role in classical segregation analysis and that,

except when families (i.e., including those families with no affected members) are sampled completely randomly from the population, failure to allow for ascertainment can seriously bias a segregation analysis (Morton 1959; Stene 1979; Greenberg 1986). However, it may be less obvious that the ascertainment model would also affect analyses of parent-of-origin effect. After all, in estimating δ , we are assessing not segregation ratios per se but, rather, a quantity proportional to the *difference* between segregation ratios. Conceivably, biases in the segregation ratios themselves would be canceled out in the difference, δ . In fact, we have demonstrated that this is what *does* happen when families are ascertained under single ascertainment—both in our simulations (see tables) and in a proof (see below and Appendix A). However, this does not happen for other ascertainment models. Thus, our results are important, because they indicate that, in general, it *is* critical to allow for ascertainment when parent-of-origin effect is being assessed.

Note that we considered only two particular ascertainment models (i.e., those for single and complete as-

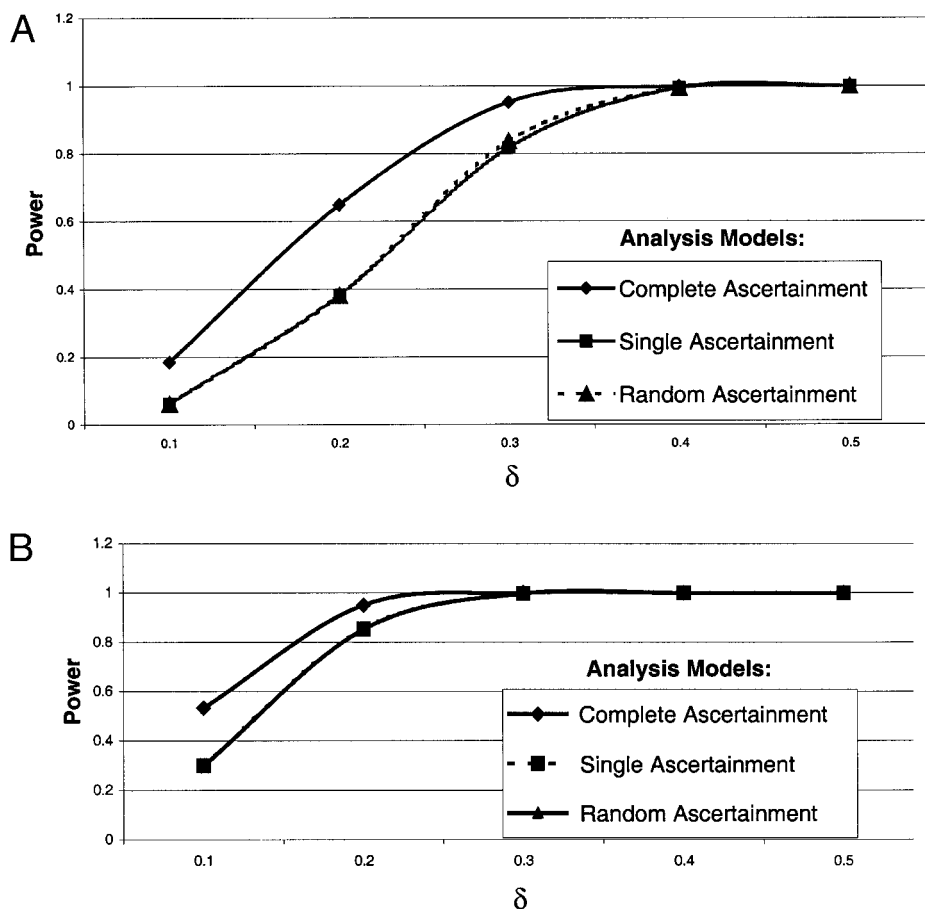


Figure 2 Observed power values from simulations of situation 2, for data sets generated under complete ascertainment (A) and data sets generated under single ascertainment (B). Mean f is 0.5. Note that the curves for single and random ascertainment are superimposed.

Table 7
Observed $\hat{\delta}$ Values from Simulations of Situation 3, Generated under Complete Ascertainment

AM	f	δ				
		.1	.2	.3	.4	.5
Complete ascertainment	.9	.060
	.7	.096	.190	.258
	.5	.093	.193	.288	.391	.499
Random ascertainment	.9	.056
	.7	.063	.130	.194
	.5	.050	.107	.172	.263	.407

certainment). These correspond to special cases of more-general ascertainment models (Weinberg 1928; Morton 1959; Ewens and Shute 1986; Greenberg 1986). However, our finding that ascertainment does matter for analysis of parent-of-origin effect would presumably also hold for other cases of those ascertainment models.

We specifically considered differential ascertainment probabilities for males and females, because one can readily imagine a scenario in which females, for example, are more readily ascertained than males. This situation could arise, for example, if women are more likely to seek treatment for illnesses—in particular, psychiatric illnesses—than men are.

Our simulations (tables 3 and 5) reveal that, if data sets are generated under single ascertainment but are analyzed as if they had been randomly ascertained, then the estimates of δ appear to be unbiased. If one writes out the likelihood, $L(\delta)$, under single ascertainment, one can see why this is so: when the mean disease penetrance f is known or specified and only δ is being estimated, the correct $L(\delta)$ for single ascertainment is the same as the “wrong” likelihood that one would use if the families had been randomly ascertained. Hence, the two likelihoods yield identical $\hat{\delta}$ values (for details, see Appendix A). This result demonstrates yet another way in which single ascertainment is “special” (Hodge and Vieland 1996). However, note that the result would not hold if the user were estimating both f and δ ; in that situation, one would have to model the single-ascertainment scheme explicitly, just as with all other ascertainment schemes.

Future Extensions

We outline four possible future extensions of this likelihood model: (1) parameterizing the likelihood in terms of two parental-transmission parameters, rather than one difference parameter δ ; (2) modeling differential male and female disease penetrances; (3) including multigenerational pedigrees; and (4) extending the likelihood to more-complex models, including linkage analysis.

Two parental-transmission parameters.—In this study, we have expressed parent-of-origin effect in terms of a

single parameter, δ , defined as the deviation, from the mean disease penetrance f , of the maternal- and paternal-transmission penetrances. In this way, if the overall average f is already known, then the model has only one unknown parameter, δ . Note that the chosen value of f used in the likelihood calculation influences the estimation of δ . For example, when we simulated data by use of $f = 0.7$ and $\delta = 0.2$ and then analyzed the data assuming the wrong f values (e.g., $f = 0.8$ and $f = 0.6$), we observed the following: When $f = 0.8$ was used, the $\hat{\delta}$ value was lower than expected ($\hat{\delta} = 0.161$). In this case, the allowed range of δ is ± 0.2 , as opposed to ± 0.3 , which includes the “true” value of $f = 0.7$. Since this value (i.e., $f = 0.8$) did not cover the entire true range of δ , the likelihood maximized at a lower value of δ than expected. However, when $f = 0.6$ was used, $\hat{\delta}$ was unbiased (i.e., $\hat{\delta} = 0.2$), because the allowed range of $\delta = \pm 0.4$ spanned the true range. These results illustrate the dependency that the estimated parental effect has on the value of the disease penetrance, which is an important consideration in the study of complex diseases for which the disease penetrance cannot be reliably determined. To avoid such a dependency, the likelihood could be reparameterized with respect to two parameters, corresponding to maternal- and paternal-transmission penetrances directly (i.e., $f_m =$ maternal, and $f_p =$ paternal), while the computational complexity is increased because of addition of a second parameter. Given this parameterization, to test only for parent-of-origin effect one would examine the difference between δ , expressed as $(1/2)(f_m - f_p)$.

Differential male and female disease penetrances.—The likelihood can be extended to model differential male and female disease penetrances. The likelihood is parameterized with respect to the combinations of disease penetrance, given the sex of the individual and the parental origin of the disease gene(s) (i.e., $f_{g,m} =$ disease penetrance, given that the individual is a girl and that the disease is maternally transmitted; $f_{b,m} =$ disease penetrance, given that the individual is a boy and that the disease is maternally transmitted; and $f_{g,p}$ and $f_{b,p}$, similarly defined). Given this parameterization, the null hypotheses are either $H_0: f_{g,m} = f_{b,m}$ or $H_0: f_{g,p} = f_{b,p}$, for testing for sex-based disease liability and either maternal or paternal effect, respectively.

This approach would be useful for study of sex-based threshold models that are well known in classical genetics. Threshold models in general have an underlying liability distribution for the disease; that is, when the liability threshold is crossed, the disease is expressed. For a sex-based model, the liability threshold differs, depending on the sex of the individual. One such disease is pyloric stenosis, a disorder that is manifested shortly after birth and that is characterized by a narrowing or obstruction of the pylorus. The prevalence of the disease

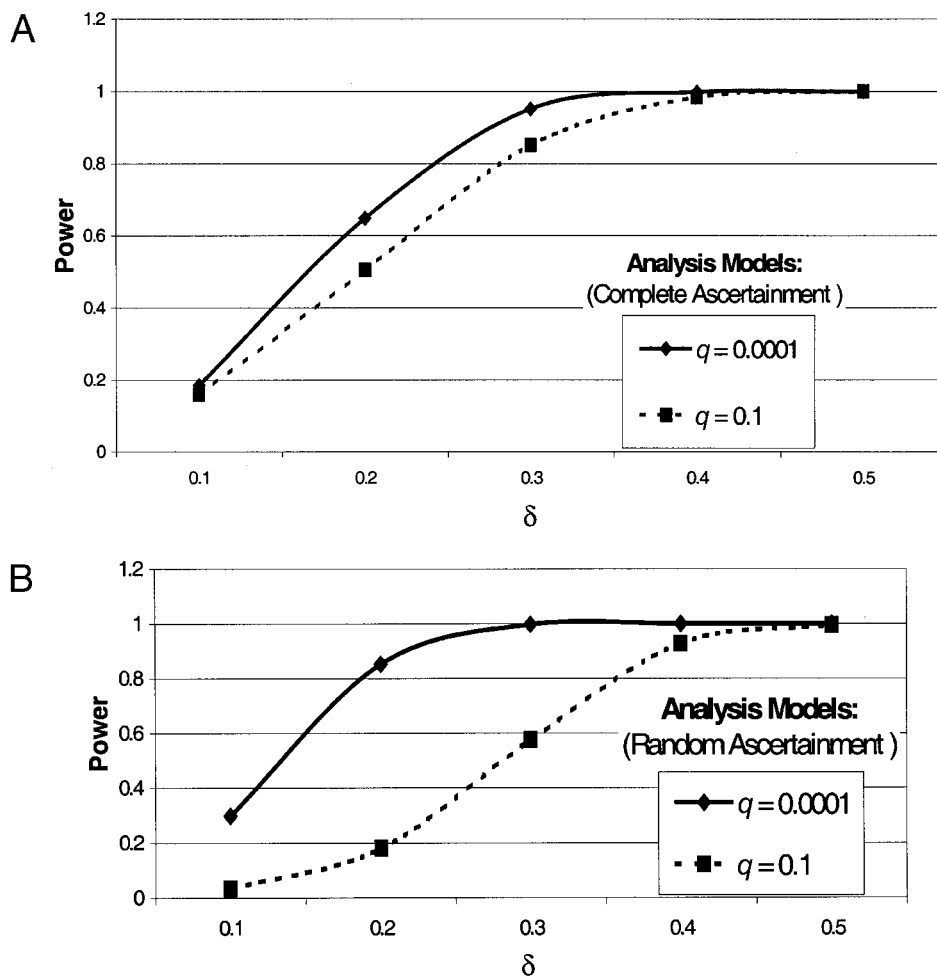


Figure 3 Observed power values from simulations of situation 3, for data sets generated under complete ascertainment and analyzed assuming complete ascertainment (A) and data sets generated under complete ascertainment and analyzed assuming random ascertainment (B). Mean f is 0.5.

is much higher in males than in females, affecting 1/200 males and 1/1,000 females in a sample of individuals of European descent (Jorde et al. 2000). This implies that females have a higher liability threshold and that, to exhibit the disease, must therefore be exposed to more “disease-causing” factors. Another disease for which this approach could be useful is panic disorder, in which there is an established sex-based disease liability and a potential parental effect. Panic disorder is a psychiatric disease characterized by spontaneous and repeated panic attacks often accompanied by agoraphobia. The risk of developing panic disorder in females has been estimated to be twice that in males, and this 2:1 (female:male) sex ratio has been observed cross-culturally (Bland et al. 1988; Robins et al. 1988; Keyl and Eaton 1990; Eaton et al. 1994). Also, there is suggestive evidence for a maternal effect in panic disorder, in which a nominally significant difference (greater than the expected 2:1) in the

sex ratio has been observed, when the disease is transmitted maternally (Haghighi et al. 1999). The parent-of-origin analysis conducted by Haghighi et al. (1999), was based on a simple counting approach. Application of the full likelihood model developed here to the same collection of families with panic disorder would make it possible to test for both sex-based disease penetrance and parent-of-origin effect.

Multigenerational pedigrees.—The existing likelihood models for parent-of-origin effects are limited to nuclear families or small- to moderate-size multigenerational pedigrees (Strauch et al. 2000). However, since we have derived the exact likelihoods for all possible parental MTs (table 1), these likelihood calculations can be extended straightforwardly to general pedigree structures, by application of the clipping algorithm (Elston and Stewart 1971; Ott 1974). Here, all the information concerning the pattern of disease transmission across suc-

cessive generations is captured for testing for parent-of-origin effect. This approach would be more robust than the existing methods, since it would handle pedigrees of arbitrary size and structure. The likelihood model with ascertainment cannot in general be extended to general pedigree structures, because of the inherent intractability of incorporating the general ascertainment model in the likelihood for extended pedigrees (Vieland and Hodge 1995), except in the special case of single ascertainment (Hodge and Vieland 1996).

More-complex models.—Although we have adopted a single-locus dominant genetic model with reduced penetrance, the likelihood model can be extended to allow for more-complex models that approximate the genetic etiology of complex diseases. For example, these models may allow for phenocopies, genetic heterogeneity (i.e., locus or allelic heterogeneity), epistasis (i.e., interaction among multiple genes), and/or a variety of environmental factors.

Of particular interest will be the attempt to extend the likelihood formulation to two (or more) loci, with recombination fraction, so as to be able to perform linkage studies. Investigators have attempted to model parent-of-origin effect in linkage analysis by maximizing the LOD score over separate male and female recombination fractions. The difference in the estimates of the male and female recombination fractions was used as an indicator of potential parental effect. For modeling of genomic imprinting, the male and female recombination fractions were maximized separately, when depending on the sex of the assumed imprinted parent, the corresponding recombination fraction was fixed at 1/2 (Smalley 1993; Strauch et al. 2000). This means that, in the likelihood calculation, the nonpenetrant children of an imprinting parent are considered to be recombinants. However, this approach does not account for the situation in which, in successive generations, these nonpenetrant children can be disease carriers and may have affected children (Strauch et al. 2000). Another approach at modeling of

parental effect could involve assigning of liability classes for maternal- and paternal-transmission penetrances separately, depending on the observed parental mating phenotypes. However, the liability classes cannot be accurately assigned when the MT is ambiguous with respect to maternal or paternal transmission (i.e., both parents either are affected or are unaffected). An advantage of the likelihood formulation derived here is that it allows for all those combinations and possibilities accurately, weighting their probabilities appropriately.

Strauch et al. (2000) have modeled parent-of-origin effect (specifically genomic imprinting) in parametric linkage analysis, by replacing the single-heterozygous penetrance parameter with two penetrance parameters corresponding to maternal- and paternal-transmission penetrances. Although this approach has been parameterized in the context of genomic imprinting, it is in principle equivalent to our likelihood formulation of the parent-of-origin-effect model without ascertainment (model I). This method has been incorporated as an extension to the GENEHUNTER program (Kruglyak et al. 1995, 1996), referred to as “GENEHUNTER-IMPRINTING.” Also note that that program was tested on a single real data set in which the underlying genetic model was unknown. One of the strengths of our study is that we tested our program with extensive simulations, in which the underlying genetic model is known.

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Appendix A

Demonstration That, When δ Is the Only Parameter of Interest, δ Estimated under Single Ascertainment Is Identical to δ Estimated under Random Ascertainment

We consider nuclear families with s children, and to begin we consider the special case of a *rare* dominant disease, so that there are only three phenotypic MTs: affected mother \times unaffected father (MT = 1); unaffected mother \times affected father (MT = 2); and both parents unaffected (MT = 3). Let r denote the number of affected children, $r = 1, \dots, s$. As in the text, f denotes the mean penetrance (and, thus, the penetrance that applies to the parents, since the sexes of *their* respective transmitting parents are not known). Let f_1 and f_2 denote maternal- and paternal-transmission penetrances, respectively, and $\delta \equiv (1/2)(f_1 - f_2)$, as in the text. For convenience, we also define $p_i \equiv (1/2)f_i$ as the segregation ratio for the mother or father, respectively.

Probabilities under Single Ascertainment

We define $u_{i,r}$ as the probability that a family of $MT = i$, with r affected children, is in the data set—that is,

$$u_{i,r} = P(MT = i, r \text{ aff} | \text{family is ascertained}) .$$

Under single ascertainment, these probabilities are as follows:

$$\begin{aligned} u_{i,r} &= \left[\frac{1}{2} f \binom{s}{r} p_i^r (1 - p_i)^{s-r} \times r\pi \right] / D , \quad \text{for } i = 1, 2 , \\ u_{3,r} &= \left\{ \frac{1}{2} (1 - f) \left[\binom{s}{r} p_1^r (1 - p_1)^{s-r} + \binom{s}{r} p_2^r (1 - p_2)^{s-r} \right] \times r\pi \right\} / D , \end{aligned} \quad (A1)$$

where π represents the probability ($\pi \rightarrow 0$ under single ascertainment) that an affected child becomes a proband, and where D , the denominator, represents the sum of the numerators of all the $u_{i,r}$, summed over $i = 1, 2, 3$ and over $r = 1, \dots, s$. (The “1/2” is an indication that, a priori, the transmitting parent is equally likely to be the mother or the father in this model.) Algebraic manipulation reveals that $D = (1/2)\pi s(p_1 + p_2)$. On the basis of the above definitions, $p_1 = (1/2)(f + \delta)$ and $p_2 = (1/2)(f - \delta)$, so the denominator becomes $D = (1/2)\pi s f$. Thus, in the probabilities $u_{i,r}$, the parameter of interest, δ , appears only in the p_i terms in the numerators. We now rewrite the probabilities in equation (A1), to explicitly show the role played by δ :

$$\begin{aligned} u_{1,r} &= A_1 \left\{ (f + \delta)^r \left[1 - \frac{1}{2}(f + \delta) \right]^{s-r} \right\} , \\ u_{2,r} &= A_2 \left\{ (f - \delta)^r \left[1 - \frac{1}{2}(f - \delta) \right]^{s-r} \right\} , \\ u_{3,r} &= A_3 \left\{ (f + \delta)^r \left[1 - \frac{1}{2}(f + \delta) \right]^{s-r} + (f - \delta)^r \left[1 - \frac{1}{2}(f - \delta) \right]^{s-r} \right\} , \end{aligned} \quad (A2)$$

where A_1 , A_2 , and A_3 represent “constant” terms that do not contain δ .

Probabilities under “Wrong” Ascertainment Model

Let $t_{i,r}$ represent the “wrong” probability that a family of $MT = i$, with r affected children, is in the data set—that is,

$$t_{i,r} \equiv P(MT = i, r \text{ aff} | \text{family is randomly ascertained}) .$$

We derive these probabilities:

$$\begin{aligned} t_{i,r} &= \frac{1}{2} f \binom{s}{r} p_i^r (1 - p_i)^{s-r} , \quad \text{for } i = 1, 2 , \\ t_{3,r} &= \frac{1}{2} (1 - f) \left[\binom{s}{r} p_1^r (1 - p_1)^{s-r} + \binom{s}{r} p_2^r (1 - p_2)^{s-r} \right] . \end{aligned}$$

Again, we rewrite these probabilities as functions of δ :

$$\begin{aligned} t_{1,r} &= B_1 \left\{ (f + \delta)^r \left[1 - \frac{1}{2}(f + \delta) \right]^{s-r} \right\} , \\ t_{2,r} &= B_2 \left\{ (f - \delta)^r \left[1 - \frac{1}{2}(f - \delta) \right]^{s-r} \right\} , \\ t_{3,r} &= B_3 \left\{ (f + \delta)^r \left[1 - \frac{1}{2}(f + \delta) \right]^{s-r} + (f - \delta)^r \left[1 - \frac{1}{2}(f - \delta) \right]^{s-r} \right\} , \end{aligned} \quad (A3)$$

where B_1 , B_2 , and B_3 represent constant terms that do not contain δ .

Equivalence of the Two Likelihoods

Let $n_{i,r}$ represent the number of families in the data set of $MT = i$, with r affected children. The log-likelihood for the data set under single ascertainment is given by

$$\log L_{\text{correct}}(\delta) = \sum_i \sum_{r=1}^s n_{i,r} \log u_{i,r} , \quad (A4)$$

whereas under the wrong ascertainment model it is

$$\log L_{\text{wrong}}(\delta) = \sum_i \sum_{r=0}^s n_{i,r} \log t_{i,r} . \quad (A5)$$

There are two differences between equations (A4) and (A5), but neither difference affects $\hat{\delta}$. The first difference is that equation (A4) uses $u_{i,r}$, whereas equation (A5) uses $t_{i,r}$. However, equations (A2) and (A3) reveal that, although the probabilities $u_{i,r}$ and $t_{i,r}$ are not respectively equal, they are *proportional* in δ . Hence, equations (A4) and (A5) will both maximize at the same value of δ . The other difference is that the sum in equation (A5) includes the case $r = 0$, whereas the sum in equation (A4) does not. However, $n_{i,0} = 0$ for all i , so that difference is immaterial.

More General Result

Above, we have derived the explicit probabilities for nuclear families with only three possible MTs, so as to show the algebra. However, the more general reasoning is that, under single ascertainment, the “denominator” in equation (A1) is always proportional to the prevalence of a proband (Hodge and Vieland 1996), and this prevalence is not a function of δ . Hence, as long as we are maximizing only with respect to δ , the “wrong” likelihood based on random ascertainment will yield the same $\hat{\delta}$ as the correct likelihood based on single ascertainment.

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