Interrogating multiple aspects of variation in a full resequencing data set to infer human population size changes

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We present an expanded data set of 50 unlinked autosomal noncoding regions, resequenced in samples of Hausa from Cameroon, Italians, and Chinese. We use these data to make inferences about human demographic history by using a technique that combines multiple aspects of genetic data, including levels of polymorphism, the allele frequency spectrum, and linkage disequilibrium. We explore an extensive range of demographic parameters and demonstrate that our method of combining multiple aspects of the data results in a significant reduction of the compatible parameter space. In agreement with previous reports, we find that the Hausa data are compatible with demographic equilibrium as well as a set of recent population expansion models. In contrast to the Hausa, when multiple aspects of the data are considered jointly, the non-Africans depart from an equilibrium model of constant population size and are compatible with a range of simple bottleneck models, including a 50–90% reduction in effective population size occurring some time after the appearance of modern humans in Africa 160,000–120,000 years ago.

bottlenecks | combining P values | human demographic inference | population growth

Eucidating how and when populations change in size is an important element in reconstructing evolutionary history because these changes often reflect crucial events in the history of a species, such as range expansions, environmental changes, and mixture between groups (1). In addition, making inferences based on population variation data typically requires the specification of a demographic model. Such applications include detecting the signature of natural selection or estimating recombination rates from patterns of linkage disequilibrium (LD) (2–5). Finally, better knowledge of demographic histories in human populations is particularly important for whole-genome, LD-based association studies (6, 7).

Motivated by the excess of rare variants observed in mitochondrial DNA data, attention initially focused on models of ancient population growth and on the idea that population expansions may have accompanied the dispersal out of Africa or the emergence of modern humans in Africa 160,000–120,000 years ago. However, the investigation of formal bottleneck models has typically used a single aspect of genetic variation data, either the allele frequency spectrum (15, 20) or patterns of LD (18, 21), raising the question of whether such models were indeed consistent with the data when multiple aspects of genetic variation were considered simultaneously (22–24). Specifically, it is not known whether simple bottleneck models can generate the marked differences in LD levels seen between Africans and non-Africans with only a limited reduction in polymorphism levels outside Africa. Although previous work suggested that variation in recombination rate may explain the decay in LD observed in a multiethnic sample (7), it is not obvious that it could also explain the differences between Africans and non-Africans.

Ideally, making inferences about population history should be based on data from a large number of unlinked and neutrally evolving loci and on statistical methodology that makes efficient use of all or most of the information in the data. Full resequencing studies, in which the sequence of the surveyed segments is determined for every individual in the sample, represent one scheme for generating data sets in which multiple aspects of sequence variation are characterized. With regard to data analysis, full likelihood methods have been successfully applied to nonrecombining data (Y chromosome or mitochondrial DNA) to reconstruct population histories (25–27). However, for regions with recombination, the currently available methods are computationally infeasible. As a result, a variety of statistics, each summarizing different aspects of genetic variation data, may be used (13, 15, 28), with the subsequent reduction in information content traded for computational tractability. It is still desirable to combine the results of tests based on individual statistics because the joint distributions of multiple summaries of the data should contain more information than the marginal distributions of multiple single summaries considered separately.

We previously developed a full resequencing scheme in which pairs of tightly but not completely linked segments, referred to as “locus pairs,” are surveyed (19). This study design aims to maximize the information content for a given amount of sequencing effort because, by skipping the intervening segment, many more independent loci can be surveyed. Using this scheme, we previously surveyed 10 noncoding regions in three human population samples: Hausa of Cameroon, Italians, and Chinese. Here, we survey an additional 40 locus pairs in the same samples. This data set allows the simultaneous characterization of polymorphism levels, allele frequency spectrum, and LD in each sample; in addition, it obviates the need to correct for ascertainment bias with its associated uncertainties and possible loss of information (29–31). In choosing only noncoding regions distant from genes, we limit the possibility that our analysis of demographic history will be confounded by the

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effects of natural selection. To analyze these data, we implement an approach to determine \( P \) values associated with several observed summaries of genetic data considered jointly over a grid of demographic parameter values. These summaries include the average Tajima’s \( D \) (\( D \)) and the variance of Tajima’s \( D \) across loci \( \text{Var}(D) \) (32), the average number of segregating sites across loci (\( \bar{S} \)), and an overall composite likelihood estimator of the population cross-over rate parameter (\( \rho \)) (33). By combining \( P \) values obtained from these individual statistics into a single statistical test, we greatly improve the power to reject demographic scenarios incompatible with the data. Although it is well established that other demographic features apply to these populations (e.g., population subdivision and gene flow) (34, 35), we chose to focus solely on population size changes to reduce modeling complexity. We explore an extensive grid of the demographic parameter space that revealed a confidence set of relatively simple bottleneck models that explain the patterns of variation in the non-African samples. Our results combine aspects of genetic variation from allele frequency spectrum, LD, and polymorphism levels within noncoding autosomal regions to infer the history of human populations. Because our data set was collected without ascertainment, it may be useful for validating the results of SNP genotyping surveys.

**Materials and Methods**

**DNA Samples.** Sequence variation was surveyed in DNA samples from the same three human populations investigated in Friesen et al. (2001): 15 Hausa samples from Yaounde, Cameroon; 15 individuals from central Italy; and 15 Han Chinese from Taiwan. In addition, one common chimpanzee DNA sample was also sequenced at each region. This study was approved by the Institutional Review Board of the University of Chicago.

**Resequencing Data Collection.** We selected 40 unlinked genomic regions for resequencing using the locus pair approach (19): For each unlinked region, we sequenced two segments of \( \sim 1 \) kb separated by \( \sim 8 \) kb. The selection of genomic targets was aimed at regions that did not contain nor were tightly linked to known or strongly predicted coding regions. Most surveyed segments also did not contain and were not tightly linked to noncoding regions strongly conserved between human and mouse (as determined by inspection of the VISTA genome browser). These regions were selected as described in ref. 19 except that here we deliberately included regions with a broader range of cross-over rates and \( \%G+C \) content. The local eM:Mb (Mb-megabase) ratio was obtained based on the interval defined by the two closest flanking markers on the DeCode Genetics (Reykjavik, Iceland) genetic map (36). The average and variance of the eM:Mb ratio across the 50 segments (i.e., 40 locus pairs from this study and the 10 given in ref. 19) are 1.31 and 0.83, respectively. The average and variance of \( \%G+C \) across the 50 locus pairs are 38.3 and 46.6, respectively. Detailed information on each surveyed segment is provided in Table 2, which is published as supporting information on the PNAS web site. PCR and sequencing was performed as described in refs. 19 and 37. All sequencing reactions were run on automated capillary sequencers (ABI3100 and ABI3700). Sequence reads were scored by using POLYPHRED (38); all putative polymorphisms and software-derived genotype calls were visually inspected and individually confirmed.

**Testing Demographic Models.** For each demographic model of interest, we performed a separate test for each summary statistic of genetic variation. In addition, for some of the models (equilibrium and bottleneck), we also calculated a test statistic, \( C \), which combines the \( P \) values of multiple summary statistics as follows:

\[
C = -2 \sum_{i=1}^{k} \ln(p_i),
\]

where \( P_i \) is the estimated \( P \) value of the \( i \)th summary statistic of \( k \) summary statistics.

For models defined by more than one demographic parameter (i.e., simple growth and bottleneck models), these tests were performed over a grid of parameter values. The combinations of parameter values that are compatible with the observed values of the test statistic(s) constitute the accepted portion of the parameter space for each model. For simple growth models, the test was based on Fu and Li’s \( D^* \) (39), whereas for bottleneck models, the test was based on combining \( P \) values from multiple summary statistics, as discussed below. The \( P \) values, \( P_i \), for each individual summary statistic were estimated from Monte Carlo simulations using a modification of the program ms (40), as follows. We used coalescent simulations to generate 50,000 replicates, each consisting of 50 independent locus pairs, for each combination of parameter values; mutation and recombination rates were allowed to vary across locus pairs as described below. Samples of sequences 10 kb in length were generated in which the intervening 8 kb were ignored to mimic the locus pair data. The probability, \( P \), of observing a value greater than that found in the data were estimated by simulations and converted to a two-tailed \( P \) value by applying the formula \( 1 - 2 \times |0.5 - P| \).

The \( P \) values for the combined test statistic \( C \) were estimated by using the empirical distribution of the statistic from simulations. For each combination of parameter values, we recorded the values of each summary statistic in each replicate and generate the distribution of these simulated values. For each replicate, we treated the value of each summary statistic as the “observed” value and determined its \( P \) value relative to the empirical distribution from the remaining 49,999 replicates. For each replicate, we combined these \( P \) values to calculate a value of \( C \). By following this procedure for each of the 50,000 replicates (for a single demographic scenario of interest), we obtained a distribution of the combined statistic. This distribution can be used to estimate a one-tailed \( P \) value for the observed value of \( C \).

**Mutation Rate Model.** We assumed an infinite sites model, where we modeled the variation in mutation rate across locus pairs by using a gamma(12.46, 2.11 \times 10^{-5}) distribution. The mean and variance for this distribution matched the observed mean and variance for the mutation rates estimated based on human–chimpanzee sequence divergence in our locus pair data (assuming 6 million years since divergence and a generation time of 25 years). The 90% central interval of this distribution is (1.54 \times 10^{-8}, 3.96 \times 10^{-8}) with \( E\mu = 2.63 \times 10^{-8} \).

**Recombination Rate Model.** We modeled the variation in the crossing-over rate, \( c \), across locus pairs using a lognormal \([-18.148, (0.5802)^2]\) distribution; cross-over rate was assumed to be homogeneous within each locus pair. The 90% central interval of this distribution is (0.51 \times 10^{-8}, 3.41 \times 10^{-8}). The median of this distribution matched the overall recombination rate for the Hausa data (1.31 \times 10^{-8}) based on the composite likelihood estimator, \( \hat{\rho} \), of Hudson (33). Because we cannot accurately estimate the variance in recombination rate across surveyed segments as short as 10 kb, we matched the variance of the lognormal distribution to the variance of eM:Mb values estimated from the Marshfield genetic map for the interval containing each locus pair (41). We acknowledge that this model may capture some but not all of the recombination rate variation estimated across the human genome (42).

**Summary Statistics.** We summarize the locus pair data in terms of the average Tajima’s \( D \) (\( D \)), the variance of Tajima’s \( D \) \( \text{Var}(D) \)), the average Fu and Li’s \( D^* \) (\( D^* \)), the average number of segregating sites (\( \bar{S} \)), and the average nucleotide diversity across the 50 locus pairs (\( \bar{\pi} \)), as well as \( \hat{\rho} \), an overall estimate of the population crossing-over parameter (4Nc) as obtained by composite likelihood (33). Because there is not enough information in our data to accurately estimate \( \hat{\rho} \) and the gene conversion parameters (43), we
assumed a model of gene conversion with rate \( f \) twice that of cross-over and tract lengths exponentially distributed with mean \( L \) 500 bp and estimate \( \hat{\rho} \). Alternative models of gene conversion \( (f = 10, L = 55 \text{ bp}) \) based on sperm-typing data (44) yielded qualitatively similar results (data not shown).

**Results**

**Summary of Sequence Variation and Tests of the Equilibrium Model.**

We resequenced 40 unlinked locus pairs in 15 individuals from each of three population samples: Hausa, Italians, and Chinese. The results of this survey are analyzed together with data for an additional 10 unlinked locus pairs previously resequenced in the same population samples (19), for a total of 50 unlinked locus pairs. The average surveyed length per locus pair was 2,365 bp (for a total of 118,259 bp surveyed in each individual), and the average unsurveyed intervening segment was 7,921 bp long.

The values of summary statistics used for demographic testing are shown in Table 1, with a synopsis of the summary statistics for the 40 new locus pairs presented in Table 3, which is published as supporting information on the PNAS web site. The allele frequency spectrum was summarized by the average and variance of Tajima’s \( D \) and Fu and Li’s \( D^* \), polymorphism levels are summarized by the average number of polymorphic sites \( \bar{S} \) across loci, and LD decay was summarized in terms of an overall composite likelihood estimator of the population cross-over rate parameter \( \hat{\rho} \) (33). The results of this expanded data set are in qualitative agreement with those from our previous survey (19) and with other similar data sets (2, 5, 15, 16). With regard to the allele frequency spectrum, the Hausa show a skew toward rare variants and a low variance across loci, whereas both non-African samples have an excess of intermediate frequency variants and high variance across loci. In addition, polymorphism levels and LD decay are higher in the Hausa compared with both non-African samples, but this difference is greater for LD decay (1.9- to 3.2-fold) than polymorphism levels (1.6-fold).

To determine whether the levels of LD decay and the frequency spectrum were consistent with a model of constant population size, we conducted coalescent simulations under equilibrium to determine the \( P \) values of the observed summary statistics. We obtained the effective population size, denoted \( N_A \), for each population by using an estimator of the population mutation rate parameter \( (4N_A \mu) \) based on the number of polymorphic sites and sample size (45), and an estimate of \( \mu \) based on sequence divergence between human and chimpanzee for the 50 locus pairs. Each summary statistic for the Hausa data are consistent with the equilibrium model (Table 1). However, for the non-African populations, the skew toward intermediate frequency variants, and the elevated LD are incompatible with a simple equilibrium model: a combined statistic based on \( D \), \( \text{Var}[D] \), and \( \pi \), obtained by using Eq. 1, is significant for the Italian \( (P \approx 0.0148) \) and Chinese data \( (P \approx 0.0052) \).

**Table 1. Observed summary statistics**

<table>
<thead>
<tr>
<th>Population</th>
<th>( D )</th>
<th>( \text{Var}[D] )</th>
<th>( D^* )</th>
<th>( \bar{S} )</th>
<th>( \bar{\pi} % )</th>
<th>( \pi_0 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hausa</td>
<td>-0.20</td>
<td>0.55</td>
<td>-0.17</td>
<td>11.1</td>
<td>0.110</td>
<td>0.0006</td>
</tr>
<tr>
<td>Italian</td>
<td>0.28*</td>
<td>1.19**</td>
<td>0.18</td>
<td>7.1</td>
<td>0.085</td>
<td>0.0003</td>
</tr>
<tr>
<td>Chinese</td>
<td>0.18</td>
<td>1.08*</td>
<td>0.05</td>
<td>6.9</td>
<td>0.079</td>
<td>0.0002*</td>
</tr>
</tbody>
</table>

Observed summary statistics of polymorphism data for 50 locus pairs. *, \( P < 0.05 \); **, \( P < 0.01 \); under an equilibrium model.

We conducted coalescent simulations under equilibrium to determine the \( t_{\text{onset}} \) and \( N_A \) obtained by ML based on the Hausa data and the simple growth model \( \left( N_A = 10,659 \right) \). This assumption has important implications for our subsequent inferences about compatible bottleneck scenarios. We then used coalescent simulations to estimate the \( P \) values for each summary statistic, point on a grid of bottleneck severities \( (b) \), bottleneck duration \( (t_{\text{end}}) \), and time since the beginning of the bottleneck \( (t_{\text{start}}) \). This procedure allows defining the portion of the multidimensional parameter space that is compatible with the data.

By combining \( P \) values of different summaries as described by Eq. \( D^* \) in the Hausa may simply reflect limited power and suggest that some expansion models are appropriate for this population. By following the approach in ref. 28, we considered a model in which an ancestral population at equilibrium size \( N_A \) grows exponentially beginning \( t_{\text{onset}} \) generations in the past at rate \( \alpha \), such that the present population size is \( N_A e^{\alpha t} \) (8). To test this model, we fixed the ancestral population size for each combination of demographic parameter values, such that the expected number of segregating sites matched the average number observed in the Hausa sample (28).

Unlike in ref. 28, we estimated the best-fit growth parameters for the Hausa data, \( \alpha \) and \( t_{\text{onset}} \), along with the associated point estimate of \( N_A \), via approximate maximum likelihood (ML) based on the summary statistic, Fu and Li’s \( D^* \). We focused on the average \( D^* \) across the 50 locus pairs, denoted \( D^*_e \), because it was previously shown to be the most informative for discriminating between equilibrium and growth models (28). For each demographic growth model, we obtained distributions of \( D^* \) by simulation and estimated the probability that \( |D^* - D^*_e| < 0.001 \) and then chose the model for which this probability was highest. This procedure returns the approximate ML estimate of the growth parameters, \( \alpha \) and \( t_{\text{onset}} \), compatible with the Hausa data based on \( D^*_e \). Note that we refer to this as approximate ML on a summary statistic because we do not use the full data and because we approximate rather than obtain the probabilities exactly. We found that the model with the highest overall probability was at an \( \alpha \) of \( 0.75 \times 10^{-3} \) and \( t_{\text{onset}} \) of 1,000 generations, which corresponds to a model with \( \approx 2 \)-fold growth starting 25,000 years ago, assuming a generation time of 25 years, from an ancestral population size of 10,659. We present confidence sets of \( \alpha \) and \( t_{\text{onset}} \) for which \( D^*_e \) are consistent with the observed Hausa data in Fig. 3, which is published as supporting information on the PNAS web site. The span of acceptable models is consistent with previous reports (28), with a slight reduction in confidence set due to the inclusion of additional data.

To assess the uncertainty in \( N_A \), we obtain a range of \( N_A \) consistent with the ML estimate of \( \hat{\alpha} = 0.75 \times 10^{-3} \) and \( t_{\text{onset}} = 1,000 \) as follows. We performed additional coalescent simulations as described earlier, where we used the ML parameters for the demographic history and gradually lowered or raised the value of \( N_A \). It was incompatible with the observed data at the 5% level. We found these high and low values of \( N_A \) to be 9,450 and 12,300, respectively. Later, we will use this information to assess the effect of our choice of \( N_A \) in testing bottleneck models.

**Testing Bottleneck Models in the Non-African Data.**

The positive \( D \) values and large \( \text{Var}[D] \) along with the low polymorphism and high LD levels observed in the non-African populations (Table 1) suggest that models including a reduction in population size may be compatible with the data. We considered one family of bottleneck models for these data, where a population of constant size \( N_A \) instantaneously shrinks in size to \( b \cdot N_A \) at time \( t_{\text{start}} \) before the present. The population remains at that size for \( 2- \) fold growth starting 25,000 years ago, assuming a generation time of 25 years, from an ancestral population size of 10,659. We present confidence sets of \( \alpha \) and \( t_{\text{onset}} \), for which \( D^*_e \) are consistent with the observed Hausa data in Fig. 3, which is published as supporting information on the PNAS web site. The span of acceptable models is consistent with previous reports (28), with a slight reduction in confidence set due to the inclusion of additional data.

Under the assumption that non-African populations originated from an ancestral population in sub-Saharan Africa, we set the ancestral population size in the bottleneck simulations to the values of \( N_A \) obtained by ML based on the Hausa data and the simple growth model \( (N_A = 10,659) \). This assumption has important implications for our subsequent inferences about compatible bottleneck scenarios. We then used coalescent simulations to estimate the \( P \) values for each summary statistic, point on a grid of bottleneck severities \( (b) \), bottleneck duration \( (t_{\text{end}}) \), and time since the beginning of the bottleneck \( (t_{\text{start}}) \). This procedure allows defining the portion of the multidimensional parameter space that is compatible with the data.
and longest bottleneck occurs where and duration have inversely related effects on patterns of variation. 

lie on the diagonal of the plots, indicating that bottleneck severity varies with various bottleneck severities (duration of 20,000 years for a total demographic epoch of 40,000 years for Voight et al., 2005). For the Chinese data, if published as supporting information on the PNAS web site). For the start is 40,000 years, the compatible parameter space for a bottleneck of 20,000 and 120,000 years (Figs. 6 and 7, which are published as supporting information on the PNAS web site). In all cases, the lower values showed a confidence set that was shifted toward scenarios of longer and more severe bottlenecks. Conversely, at higher values, more severe bottlenecks were rejected in favor of milder bottleneck scenarios.

To assess the effect of the uncertainty associated with the estimates of , we repeated the above analyses by using different values of that were obtained from estimating the uncertainty around from the Hausa ML growth models described above. As shown in Fig. 2, the effect of on the accepted parameter space is substantial. As expected, for the larger value of , the accepted portion of the parameter space is reduced such that only relatively severe and long bottlenecks are compatible with the data, whereas a larger range of less severe bottlenecks are compatible with the smaller value of .

Discussion

By resequencing unlinked noncoding regions, we assessed patterns of polymorphism levels, frequency spectrum and LD for the same set of genomic segments and population samples. To achieve greater resolution of different demographic scenarios, we use an analytical approach that combines information from individual summary statistics of sequence variation; computer simulations showed that combinations of summaries allow for more powerful tests of each demographic scenario. Rather than focusing on a single best-fitting demographic model, we construct an acceptance region of the parameter space that is compatible with the demographic model of interest (in this case population growth or bottleneck), thus providing an inclusive picture of the uncertainty in inferences of human demography. A major conclusion of this analysis is that the non-African population samples are compatible with simple bottleneck models even when multiple aspects of sequence variation are considered simultaneously. Consistent with our previous analysis (28), the Hausa sample from sub-Saharan Africa is compatible with the equilibrium model and with relatively recent population growth.

Modeling human population history is central to a variety of questions in human biology, but most recently the search for signatures of natural selection has given new importance to this line of inquiry (2, 5, 46, 47). The impact of natural selection on the human genome can be detected by contrasting patterns of neutral variation, i.e., those shaped solely by demography, to those observed at test loci that may be shaped by natural selection in addition to demography. Traditionally, this contrast used the theoretical predictions of the standard neutral model in which the population was assumed to be constant in size and randomly mating. However, studies of human variation have shown genome-wide departures from this model, suggesting that human demography is complex (7, 15, 21, 48). Thus, the development of a more realistic null model of evolutionary neutrality is necessary for improving inferences about natural selection (2, 5).

Several conditions must be satisfied to achieve these goals. One is the availability of sequence variation data for many unlinked and neutrally evolving regions. Although several whole-genome variation data sets are available, they consist mainly of genotyping data for ascertained polymorphisms (49, 50). Resequencing data are also available, but they tend to focus on gene regions that may have been targets of selection and, hence, are less suitable for demographic inference (2, 3, 5). An additional challenge derives from the complexity of human demography and the fact that realistic models are defined by multiple unknown demographic parameters, which implies that, for any given value of one parameter (e.g., bottleneck severity), the accepted parameter space is reduced such that only relatively severe and long bottlenecks are compatible with the data, whereas a larger range of less severe bottlenecks are compatible with the smaller value of .
several aspects of genetic variation by combining the genetic variation data in each population. The inclusion of multiple aspects of the multidimensional parameter space that is consistent with uncertainty around the best-fitting model by identifying the portion of human demography. First, it provides a full characterization of the content of the data so that multiple aspects of genetic variation are considered simultaneously. Several previous studies of human sequence variation had modeled specific bottleneck scenarios on the basis of either frequency spectrum information (2, 5, 15, 48), LD decay (18), or polymorphism levels (21). Wall and Przeworski (15) analyzed full resequencing data and proposed that a bottleneck and selective sweeps at some loci could explain the frequency spectrum observed in non-Africans but did not provide information regarding the likely parameter values. The frequency spectrum was used also by Marth et al. (20) to estimate a best-fit bottleneck model for Europeans and East Asians. We used our simulation scheme to estimate the probability of the Italian and Chinese data for the corresponding best-fit models of Marth et al. (20). In our parameterization, the best-fit model for the Asian

An important limitation of our analysis is that we considered only models of randomly mating populations. Although this is a common assumption in modeling studies of population size change, it is unlikely to be satisfied by human populations, even if geographically defined (34, 51). In fact, it is possible that population structure alone could account for the observed patterns of human variation (2, 5, 15, 35). Interestingly, the addition of $\text{Var}[D]$ into the bottleneck analysis results in a further reduction of the accepted parameter space (Figs. 8–11, which are published as supporting information on the PNAS web site). Although combining this statistic with $\hat{D}$, $\hat{S}$, and $\hat{\rho}$ reduces the power to reject the constant size model (Fig. 1). This observation suggests that additional features, such as population structure, are required to produce $\text{Var}[D]$ values that are more consistent with our data. Although it is desirable and certainly more realistic to include elements of population structure in models of human demography (52), there is insufficient data to indicate the most plausible family of such models. For these reasons, testing simple growth and bottleneck models is a reasonable first step toward developing more complex and realistic models. Obviously, if changes in population size and population structure were considered jointly rather than separately, the accepted range of values for the growth and bottleneck parameters is likely to be different.

A main conclusion of this study is that simple bottleneck models can explain the non-African data even when multiple aspects of genetic variation are considered simultaneously. Several previous studies of human sequence variation had modeled specific bottleneck scenarios on the basis of either frequency spectrum information (2, 5, 15, 48), LD decay (18), or polymorphism levels (21). Wall and Przeworski (15) analyzed full resequencing data and proposed that a bottleneck and selective sweeps at some loci could explain the frequency spectrum observed in non-Africans but did not provide information regarding the likely parameter values. The frequency spectrum was used also by Marth et al. (20) to estimate a best-fit bottleneck model for Europeans and East Asians. We used our simulation scheme to estimate the probability of the Italian and Chinese data for the corresponding best-fit models of Marth et al. (20). In our parameterization, the best fit model for the Asian

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**Fig. 2.** Confidence sets for a bottleneck with $t_{\text{bottleneck}}$ of 40,000 years. Results are shown for the Italian (a–c) and Chinese (d–f) data sets for $N_A$ values of 9,450 (a and d), 10,659 (b and e), and 12,300 (c and f). The combined statistics are $\hat{D} - \hat{S}$. The contours represent the confidence region of parameter space with $P$ values of 0.1, 0.05, 0.02, and 0.01 from innermost to outermost, with darker shading indicating lower $P$ values.
sample in Marth et al. corresponds to an $N_A$ of 10,000, $b$ of 0.3, $\tau_{dur}$ of 400 generations, a $t_{dur}$ of 90,000 years; note that this model includes growth after the bottleneck to a size of 25,000. The best-fit model for the European sample in Marth et al. corresponds to an $N_A$ of 10,000, $b$ of 0.2, $\tau_{dur}$ of 500 generations, and a $t_{dur}$ of 87,500 years, with growth after the bottleneck to a size of 0.2. Using our simulation scheme, our data turned out to be incompatible with these models ($P < 0.0001$). It should be noted, however, that Marth et al. (20) analyzed a data set of ascertained SNPs and attempted to correct for the resulting bias. Hence, the discrepancy between the two studies may be due to incomplete ascertainment correction and highlights the value of resequencing data.

Based on the frequency spectrum observed in a large resequencing study of genes involved in inflammation, Akey et al. (2) concluded that the European data were consistent with a bottleneck starting 40,000 years and a bottleneck intensity, as measured by the inbreeding coefficient ($F$) of 0.175. This best-fit model can be translated to a range of models in our notation by using

$$F = 1 - \left(1 - \frac{1}{2bN_A}\right)^{\tau_{dur}}.$$  \[2\]

This bottleneck model corresponds to a number of points that are well within the accepted portion of the parameter space for our non-African data (for example, $b = 0.2$ and $t_{dur} = 820$ generations assuming our best-fit $N_A$ of 10,659). Because only the best-fit model is reported by Akey et al. (2), the overall agreement between these two data sets cannot be assessed.

Similar conclusions were obtained through an analysis of pairwise LD data of ascertained SNPs in a European population sample (18); however, a narrow portion of the parameter space was investigated. We determined that there are points in our accepted parameter space that correspond to the estimated time of onset and $F$ reported by Reich et al. (18), indicating agreement between the two methods and data sets. Finally, a recent analysis of resequencing data from a pool of ethnically diverse samples detected evidence for very recent population growth (3). Although this model is compatible with our Hausa data, it does not provide a good explanation for the Italian and Chinese data, hence, pointing to the need for population-specific demographic inferences.

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