## On the limits of fitting complex models of population history to genetic data

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#### 1 Abstract

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Our understanding of human population history in deep time has been assisted by fitting "admixture 3 4 graphs" to data: models that specify the ordering of population splits and mixtures which is the only 5 information needed to capture the patterns of allele frequency correlation among populations. Not 6 needing to specify population size changes, split times, or whether admixture events were sudden or 7 drawn out simplifies the space of models that need to be searched. However, the space of possible 8 admixture graphs relating populations is vast and cannot be sampled fully, and thus most published 9 studies have identified fitting admixture graphs through a manual process driven by prior 10 hypotheses, leaving the vast majority of alternative models unexplored. Here, we develop a method for systematically searching the space of all admixture graphs that can incorporate non-genetic 11 12 information in the form of topology constraints. We implement this findGraphs tool within a 13 software package, ADMIXTOOLS 2, which is a reimplementation of the ADMIXTOOLS software with 14 new features and large performance gains. We apply this methodology to identify alternative 15 models to admixture graphs that played key roles in eight published studies and find that graphs 16 modeling more than six populations and two or three admixture events are often not unique, with 17 many alternative models fitting nominally or significantly better than the published one. Our results 18 suggest that strong claims about population history from admixture graphs should only be made 19 when all well-fitting and temporally plausible models share common topological features. Our re-20 evaluation of published data also provides insight into the population histories of humans, dogs, and 21 horses, identifying features that are stable across the models we explored, as well as scenarios of 22 populations relationships that differ in important ways from models that have been highlighted in 23 the literature, that fit the allele frequency correlation data, and that are not obviously wrong.

#### 24 Introduction

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26 Admixture graph models provide a powerful intellectual framework for describing the relationships 27 among populations that allows not only branching of populations from a common ancestor but also 28 mixture events. Admixture graphs can precisely summarize important features of population history 29 without requiring specification of all parameters such as population sizes, split times, mixture times, 30 and distinguishing between sudden splits or drawn-out separations. All these parameters describe 31 important features of demographic history and are considered by several tools for fitting 32 demographic models (Gutenkunst et al. 2009, Gronau et al. 2011, Schiffels et al. 2016, Flegontov et 33 al. 2019, Kamm et al. 2019, Rogers 2019, Hubisz et al. 2020). However, the fact that it is possible to 34 first infer important aspects of the topology (admixture graphs), and then fit these additional 35 parameters, simplifies demographic inference (Patterson et al. 2012, Pickrell and Pritchard 2012, 36 Lipson et al. 2013, Leppälä et al. 2017, Lipson 2020, Molloy et al. 2021, Yan et al. 2021). Admixture 37 graphs thus serve both as conceptual frameworks that allow us to think about the relationships of 38 populations deep in time, and as mathematical models we can fit to genetic data. 39

40 A challenge for fitting admixture graph models is that they are often not uniquely constrained by the data, with many providing equally good fits to the  $f_2$ -,  $f_3$ -, and  $f_4$ -statistics used to constrain them 41 42 within the limits of statistical resolution. Previously published methods for finding fitting admixture 43 graphs were not well-equipped to handle the large range of equally well-fitting models for three 44 reasons: (1) They did not reliably provide information on whether there is a uniquely fitting 45 parsimonious model or alternatively whether there are many models that fit equally well to within 46 the limits of statistical resolution, (2) they did not provide formal goodness-of-fit tests, and related 47 to this, (3) they not provide tests for whether the difference between the fits of any two models is 48 statistically significant. As a consequence and as we demonstrate in what follows, many published 49 admixture graph models have been interpreted as providing more confidence than is merited about 50 the extent to which genetic data allows us to disentangle ancestral relationships.

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52 There are two main approaches to studying demographic history with admixture graphs.

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54 The first approach is to identify admixture graphs automatically, either without human intervention 55 or with guidance. It is possible in theory to exhaustively test all possible graphs for a given set of 56 populations and pre-specified number of admixture events, as implemented, for example, in the 57 admixturegraph R package (Leppälä et al. 2017). An exhaustive approach can provide a complete 58 and unbiased view on the kinds of models that are consistent with the data for a specified level of 59 parsimony (total number of admixture events allowed in the graph), but this approach is limited to 60 small graphs (typically up to 6 groups, 2 admixture events) due to the rapid increase in the number of possible admixture graphs as the number of populations and admixture events grows. In addition, 61 62 as we show in our discussion of case studies, the assumption of parsimony that needs to be made in 63 order to use an exhaustive approach can lead to misleading models of population history because 64 not including additional populations can blind users to additional mixture events that occurred (and 65 whose existence is revealed by examining data from additional populations). Specifically, models 66 with additional admixture events that are qualitatively profoundly different to the best fitting 67 parsimonious graph and that capture the true history, will sometimes be completely missed when 68 applying the parsimony assumption. Alternatively, the programs TreeMix (Pickrell and Pritchard 69 2012, Molloy et al. 2021), MixMapper (Lipson et al. 2013), and migoGraph (Yan et al. 2021) all 70 address the problem of how to rapidly explore the vast space of admixture graphs relating a set of 71 populations by applying algorithmic ideas or heuristics; all of these methods speed up model search 72 by orders of magnitude. The new method we introduce here, *findGraphs*, belongs to this class of 73 algorithms. Algorithmic innovations in *findGraphs* enable us to get around some of the limitations 74 associated with parsimony assumptions by more efficiently exploring a larger proportion of plausible 75 of admixture graph space, and by increasing the speed of evaluation of individual graphs. 76

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77 The second approach to fitting admixture graphs is to manually build them up by grafting additional 78 populations onto simpler smaller graphs that fit the data. This approach involves stepwise addition 79 of populations in an order that is chosen based on the best judgment of the user, and for each newly 80 added population involves adding admixture events or tweaks in the graph until a fit is obtained; the 81 user then moves on to adding the next population (see Reich et al. Nature 2009, Reich et al. AJHG 82 2011, Reich et al. Nature 2012, Lazaridis et al. 2014, Seguin-Orlando et al. 2014, Fu et al. 2016, 83 Skoglund et al. 2016, Yang et al. 2017, McColl et al. 2018, Moreno-Mayar et al. 2018, Tambets et al. 84 2018, van de Loosdrecht et al. 2018, Flegontov et al. 2019, Sikora et al. 2019, Wang et al. 2019, 85 Lipson et al. 2020, Shinde et al. 2019, Yang et al. 2020, Hajdinjak et al. 2021, Wang et al. 2021 for 86 examples). The program *qpGraph* in the *ADMIXTOOLS* package (Patterson et al. 2012) has been the 87 most common computational method used for testing fits of individual admixture graphs. Most 88 published admixture graphs have been constructed manually, often acknowledging the existence of 89 alternative models by presenting plausible models side-by-side, and this approach has been the 90 basis for many claims about population history (Lazaridis et al. 2014, Yang et al. 2017, Posth et al. 91 2018, Sikora et al. 2019, Shinde et al. 2019, Bergström et al. 2020, Lipson et al. 2020, Hajdinjak et al. 92 2021, Wang et al. 2021). A strength of this approach is that it takes advantage of human judgment 93 and outside knowledge about what graphs best fit the history of the human populations being 94 analyzed. This external information is powerful as it can incorporate non-genetic evidence such as 95 geographic plausibility and temporal ordering of populations or linguistic similarity, or other genetic 96 data such as estimates of population split times or shared Y chromosomes or rejection of proposed 97 scenarios based on joint analysis of much larger numbers of populations than can reasonably be 98 analyzed within a single admixture graph. Thus, while manual approaches explore many orders of 99 magnitude fewer topologies than automatic approaches often do, they still may provide inferences 100 about population history that are more useful than those provided by automatic approaches. These 101 methods' strength is also their weakness: by relying on intuition, following a manual approach has 102 the potential to validate the biases users have as to what types of histories are most plausible (these 103 may be the only types of histories that will be carefully explored). This can blind users to surprises: 104 to profoundly different topologies that may correspond more closely to the true history, and we 105 discuss examples of this in the Results section.

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The *findGraphs* method combines the advantages of automated and manual topology exploration by
 allowing users to encode various sources of information as constraints on the space of admixture
 graphs, which is then explored automatically.

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111 Regardless of the approach used to search through the space of possibly fitting admixture graphs, a 112 challenge in the effort to find a uniquely well-fitting admixture graph is that it is difficult to quantify 113 the absolute quality of the fit of a model, as well as the relative quality of the fits of multiple models. 114 Performance gains relative to the original implementation of *qpGraph* allow us to address this 115 problem by obtaining bootstrap confidence intervals and p-values for estimated parameters of 116 single models, as well as for the difference in fit quality of two models. Existing methods for 117 comparing admixture graph models (for example based on Akaike information criterion (AIC) or 118 Bayesian information criterion (BIC), see Flegontov et al. 2019, Shinde et al. 2019) do not take the 119 variability across SNPs into account appropriately since they do not rely on dataset resampling 120 approaches such as bootstrap, and thus they tend to overestimate our ability to differentiate 121 competing models. In combination with the previously described approach to automating the search 122 of admixture graphs, this leads to a situation where we are able to find and test a large number of 123 models, many of which fit equally well despite often having very different topological features. 124 125 The methods for automated graph topology inference and model comparison relying on bootstrap

resampling described above are implemented in *ADMIXTOOLS 2*, a comprehensive platform for learning about population history from *f*-statistics. It is built to provide a stand-alone workspace for research in this area and is implemented as an R package. For all computations, *ADMIXTOOLS 2* 

129 exhibits large speedups relative to previously published platforms for *f*-statistic analysis (e.g.

130 popstats and ADMIXTOOLS version 6.0 which we call "Classic ADMIXTOOLS" in what follows to

distinguish it from updated ADMIXTOOLS version 7.0.2 which implements some of the speed-up 131 132 ideas also implemented in ADMIXTOOLS 2). This is achieved by deploying a series of algorithmic 133 improvements, most notably storage of pre-computed f-statistics in random access memory, which 134 avoids having to rely on reading in extremely large genotype matrices to perform most 135 computations. In addition to the new algorithmic ideas allowing efficient searching through the 136 space of admixture graphs and comparing the fits of two admixture graphs, ADMIXTOOLS 2 also 137 provides a solution to the question of which parameters of an admixture graph are identifiable in 138 the limit of infinite data. The most important algorithmic improvements presented in ADMIXTOOLS 139 2 and its philosophical differences relative to classic ADMIXTOOLS are described in the next section, 140 while the full methodological details are presented in the Methods and Supplement. 141 142 143 <u>Results</u> 144 New ideas implemented in ADMIXTOOLS 2 145 146 147 We present a new implementation of the popular ADMIXTOOLS software (called "Classic 148 ADMIXTOOLS") (Patterson et al. 2012, Haak et al. 2015). Our implementation (ADMIXTOOLS 2, see 149 documentation at <a href="https://uqrmaie1.github.io/admixtools">https://uqrmaie1.github.io/admixtools</a>) enhances performance by greatly 150 reducing runtime and memory requirements across a wide range of different methods, relative to 151 Classic ADMIXTOOLS (Figure 1a). We note that some of these improvements have now been 152 implemented in version 7.0.2 of ADMIXTOOLS. The present study focuses not on the performance 153 differences between Classic ADMIXTOOLS and ADMIXTOOLS 2, but on the description of new ideas 154 implemented in one or both of these tools. 155 156 157 Computation and use of f-statistics 158 A key idea that facilitates the performance increases shared by ADMIXTOOLS 2 and ADMIXTOOLS v. 159 160 7.0.2 is that any f-statistic (which form the basis of almost all ADMIXTOOLS programs as well as 161 other toolkits for studying population history such as *popstats*) can be computed from a small 162 number of  $f_2$ -statistics. For most f-statistic-based analyses (for example qpWave, qpAdm, and 163 *qpGraph*; **Figure 1c**), the time required to process *f*-statistics is trivial compared to the time required 164 to compute f-statistics from genotype data. These  $f_2$ -statistics can be stored and re-used to 165 compute  $f_3$ - and  $f_4$ -statistics, thus reducing the size of the input data, runtime, and memory 166 requirements by orders of magnitude (Figure 1a, 1d). 167 168 Using precomputed  $f_2$ -statistics is not always the best solution. In data sets with large amounts of 169 missing data, computing  $f_3$ - and  $f_4$ -statistics from pre-computed  $f_2$ -statistics may introduce bias. In this case, it is necessary to compute  $f_3$ - and  $f_4$ -statistics directly, using different SNPs for each f-170 171 statistic (all available SNPs in each population triplet or quadruplet). However, even without the use 172 of pre-computed  $f_2$ -statistics, ADMIXTOOLS 2 often achieves large performance gains (Figure 1a). 173 174 The program *qpfstats* in Classic ADMIXTOOLS implements an idea which strikes a balance between 175 these two extremes. It increases the accuracy of estimation of f-statistics by using a regression 176 approach to jointly estimate the values of all  $f_2$ -,  $f_3$ - and  $f_4$ -statistics relating a set of populations. 177

177 taking advantage of the algebraic relationships of the expected values of these statistics. This 178 approach makes it possible to obtain more precise estimates of the values of these statistics than 179 can be obtained by inferring them only using SNPs that are covered in each of the populations being 180 compared. This feature is available in *ADMIXTOOLS 2* through the *qpfstats* option in the *extract\_f2* 181 function.

182

183 Another improvement introduced in *ADMIXTOOLS 2* relates to accurate evaluation of the match

184 between observed and expected  $f_3$ -statistics when fitting admixture graphs where at least one

185 population is represented by a single individual with genotypes derived from randomly selected 186 sequencing reads ("pseudo-diploid" or "pseudo-haploid" data). f-statistic computations need to be 187 modified when analyzing pseudo-diploid data, because heterozygosity cannot be computed using 188 comparisons of sequences within the same individual; however, computation of heterozygosity is 189 essential to calculate "admixture"  $f_3$ -statistics where negative values provide proof of the mixed 190 nature of the target population. When a population is represented by multiple individuals, unbiased 191 estimation of admixture  $f_3$ -statistics can be carried out even for pseudo-diploid data by analyzing 192 positions covered by sequences from at least two individuals and only computing variation rates 193 across individuals. This approach is implemented in Classic ADMIXTOOLS with the "inbreed: YES" 194 option. However, no admixture  $f_3$ -statistic can be computed when the "inbreed: YES" option is 195 turned on and the target population is represented by a single individual (as no variation across 196 individuals within a population can be detected in this case). Classic ADMIXTOOLS deals with this 197 case by failing to run if any population in an analysis is represented by a single individual and the 198 "inbreed: YES" option is turned on. Because the datasets from all the admixture graphs revisited 199 here included at least one population represented by a single individual (Table S1), the "inbreed: 200 YES" option could not be used in the original studies (the program failed with this option, by design). 201 Thus, admixture graph fitting in those studies relied on the incorrect algorithm for calculating  $f_3$ -202 statistics (except for Librado et al. 2021, who used TreeMix instead of qpGraph) and, as a result, 203 some  $f_3$ -statistics that are negative and could provide important constraints for admixture graph 204 fitting were evaluated as positive. These concerns are relevant for the Shinde et al. (2019), Lipson et 205 al. (2020), and Wang et al. (2021) studies we revisit below (see Table S1 for datasets where negative 206  $f_3$ -statistics were encountered). To be able to detect negative  $f_3$ -statistics and thus take advantage of 207 their power for constraining the space of possibly fitting historical models, in ADMIXTOOLS 2 we 208 introduced an option which makes it possible to compute negative  $f_3$ -statistics on pseudo-diploid 209 data, at a cost of removing sites with only one chromosome genotyped in any population that is 210 represented by at least two individuals (so that it is possible in theory to compute heterozygosity in 211 these populations). Admixture  $f_i$ -statistics continue to be incorrectly computed using ADMIXTOOLS 2 212 for targets that are singleton populations represented by pseudo-diploid data, as there is no 213 avoiding this particular problem. See Methods for a description of the new algorithm for calculating 214  $f_3$ -statistics.

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- 216
- 217 Figure 1
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N x M

#### 226

#### 227 Figure 1:

N x M

a Performance comparison of *f*-statistic computation and admixture graph fitting. Top: Memory usage and runtime for computing *f*-statistics using (1) the *qpDstat* program in *ADMIXTOOLS* v7.0.2 released in 06/2021
 (2) the function in *ADMIXTOOLS 2* without precomputing -statistics, and (3) the function in *ADMIXTOOLS 2* with precomputed -statistics. (1) and (2) give identical results, whereas (3) only gives identical results in the absence of missing data, which limits usefulness beyond a moderate number of populations. Bottom: Runtime comparison of *qpGraph* with and without precomputed *f*-statistics.

(<sup>k</sup><sub>4</sub>)×3

234 **b** Illustration of -and -statistics. measures the amount of drift separating any two populations, while 235 measures the amount of drift shared between two population pairs. Every  $f_4$ -statistic is a linear combination of 236 four  $f_2$ -statistics.

237 c Overview of the major ADMIXTOOLS programs, their primary use cases, and their associated f-statistics.

d Schematic representation of the computations behind the ADMIXTOOLS programs qpGraph, qpWave, and
 qpAdm. ADMIXTOOLS 2 separates the computation of -statistics from the later steps in the pipeline. Shown
 below are the number of data points for individuals, SNPs, and populations. The exact number of all
 possible non-redundant , , and -statistics for populations are , - , and - . A small number of
 -statistics can be used to obtain a much larger number of - and - statistics and requires much less space
 than the raw genotype data.

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247

#### 246 Admixture graph fitting, model comparison, and interpretation

248 There are several challenges that arise when modeling the ancestral relationships among

249 populations with admixture graphs, and *ADMIXTOOLS 2* implements solutions to several key

250 problems that were not adequately addressed with previous approaches: (1) Automated admixture

251 graph inference; (2) Estimating confidence intervals for admixture graph parameters; (3) Comparing

252 fits of different admixture graphs; (4) Determining identifiability of admixture graph parameters; and

- 253 (5) Drawing conclusions from a large number of fitting graphs.
- 254

255 Here, we describe these challenges, how we address them, and how our approaches compare to

256 other approaches, while the Methods section gives detailed descriptions.

257

#### 258 (1) Automated admixture graph inference

259 Constructing admixture graphs manually runs the risk of overlooking models that challenge 260 conventional hypotheses. On the other hand, current methods for inferring admixture graphs 261 automatically (Leppälä et al. 2017, Molloy et al. 2021, Pickrell and Pritchard 2012, Yan et al. 2021) do 262 not allow external information to be integrated into the analysis, and often result in models that 263 may fit the genetic data but can be rejected on other grounds. In addition, *TreeMix* (Pickrell and 264 Pritchard 2012), as well as OrientAGraph (Molloy et al. 2021), an improved version of TreeMix, can 265 miss admixture graph topologies that exist on parts of the non-convex likelihood surface that are 266 bypassed by these algorithms for exploring admixture graphs (for example, topology M7 in Figure 4) 267 of (Molloy et al. 2021)). MixMapper (Lipson et al. 2013) and miqoGraph (Yan et al. 2021) have a 268 different limitation: exploring topologies with more than one admixture event in the history of any 269 group is not possible. Due to these limitations, many published findings are based on manual 270 proposal of topologies and evaluation of fit, and the great majority of studies using this manual 271 approach (see, for example, Reich et al. 2011, Reich et al. 2012, Lazaridis et al. 2014, Fu et al. 2016, 272 Skoglund et al. 2016, Yang et al. 2017, McColl et al. 2018, Moreno-Mayar et al. 2018, Tambets et al. 273 2018, van de Loosdrecht et al. 2018, Flegontov et al. 2019, Sikora et al. 2019, Wang et al. 2019, 274 Lipson et al. 2020, Shinde et al. 2019, Yang et al. 2020, Hajdinjak et al. 2021, Wang et al. 2021) rely 275 on the software *qpGraph*. We introduce an approach for finding well-fitting admixture graphs 276 automatically that can integrate external information, and that recovers graph topologies more 277 accurately than TreeMix (Figure 2). External information can be integrated by specifying a set of 278 constraints that admixture graphs must satisfy. This not only ensures that resulting models are 279 temporally plausible, but also cleanly separates prior assumptions from the independent constraints 280 provided by genetic data. Our strategy implemented in the function "findGraphs", differs from 281 TreeMix/OrientAGraph in several deep ways, most notably in that it optimizes graphs directly, rather 282 than optimizing trees first and adding admixture events later. This makes it less prone to getting 283 stuck in local optima: our simulation results show that *findGraphs* is more accurate for random 284 graphs (Figure 2), and that it can recover specific topologies that pose problems for TreeMix and 285 OrientAGraph.

286

287 (2) Estimating confidence intervals for admixture graph parameters

Since our new implementation of *qpGraph* can evaluate models much more rapidly, it becomes feasible to evaluate the same model multiple times on different SNP sets. This allows us to derive bootstrap confidence intervals (Boos 2003) for all parameters estimated by *qpGraph*, including drift lengths, admixture weights, log-likelihood (LL) scores, and  $f_4$ -statistic residuals. Parameters with extremely wide confidence intervals can thus be immediately shown to be poorly determined. It should also be noted that the estimated confidence intervals do not take into account uncertainty about the graph topology.

- 295
- 296 (3) Comparing the fits of different admixture graphs

297 Using the bootstrap method for evaluating a graph multiple times on different SNP sets not only 298 allows us to obtain confidence intervals for single graphs, but also allows us to test whether the fit of 299 one graph is significantly better than the fit of another graph, by obtaining confidence intervals for 300 the log-likelihood score difference or worst-residual (WR) difference of two graphs. When we apply 301 this approach to a range of data sets, we find that models with modest log-likelihood differences are 302 often not distinguishable after accounting for the variability across SNPs, even if one might expect 303 them to be distinguishable based on the magnitude of the likelihood difference (Figure 3a,b). Thus, 304 previous methods relying on AIC or BIC (such as Shinde et al. 2019, Flegontov et al. 2019) that used 305 specified likelihood difference thresholds to reject some models over others, were over-aggressive. 306

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- 308 a



#### 312 Figure 2:

313 Comparison of accuracy of automated search for optimal topology in the findGraphs function of ADMIXTOOLS 314 2 and TreeMix using simulated graphs with 8, 10, 12, and 16 populations, and 0 to 10 admixture events. Error bars show standard errors calculated as  $SE^2 = p(1-p)/n$  where p is the fraction on the y-axis and n is the 315 316 number of simulations in each group (typically 20). In the case of ADMIXTOOLS 2, we applied findGraphs three 317 times on each simulated data set and picked a result with the best fit score. More details are provided in the 318 Methods section. a Fraction of simulations where the simulated graph is recovered exactly. b Fraction of 319 simulations where the simulated graph is either recovered exactly, or the score is at least as good as the score 320 of the simulated graph, when both graphs are evaluated by ADMIXTOOLS 2. More admixture edges greatly 321 increase the search space and make it more difficult to recover the simulated graph, but they do make it easier 322 to find alternative graphs with good fits.

323

324 A second challenge in comparing different admixture graph models arises when comparing models

- of different complexity (i.e., with a different number of admixture events). Established methods
- such as AIC and BIC are applicable and also can account for different model complexity if the

327 number of independent parameters in a model is known. However, the number of independent

- 328 parameters estimated in an admixture graph is not simply determined by the number of groups,
- 329 drift edges, or admixture events, as it also depends on the graph topology in a complex way. We
- 330 implement a method to compare admixture graph models of different complexity by using a new
- 331 scoring function, which uses different blocks of SNPs for deriving fitted and estimated *f*-statistics.
- 332 This ensures that our model comparison test does not favor more complex models by allowing them
- 333 to overfit the data. This cross-validation approach can also be used to rank alternative models of the
- 334 same complexity and deal with overfitting. We note that the calculation of cross-validated log-
- 335 likelihood scores is not turned on in *findGraphs* by default, and to make our results more
- 336 comparable to those of the published studies we revisited, we relied on standard log-likelihood scores below.
- 337 338

339 To test if our method is well calibrated, we simulated 100,000 SNPs under the same graph in 1000 340 replicates. We then created two new topologies by removing one out of two symmetric edges from 341 the first graph (Figure 3c). These new incorrect models are symmetrically related to the first graph 342 and can be used to test the null hypothesis that the true difference in log-likelihood scores of these 343 two graphs is zero. The uniform distribution of p-values confirms that our method is well calibrated 344 (Figure 3d). A caveat is that only one symmetric topology was explored in this way.

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352

#### 353 Figure 3: Calibrating the bootstrap model comparison approach

a Bootstrap sampling distributions of the log-likelihood scores for two admixture graphs (shown in Figure S1)
 of the same populations fitted using real data. Vertical lines show the log-likelihood scores computed on all
 SNP blocks.

b Distribution of differences of the bootstrap log-likelihood scores for both graphs (same data as in a). The
 purple area shows the proportion of resamplings in which the first graph has a higher score than the second
 graph. The two-sided *p*-value for the null hypothesis of no difference is equivalent to twice that area (or one
 over the number of bootstrap iterations if all values fall on one side of zero). In this case it is 0.078.

c The admixture graph which was used to evaluate our method for testing the significance of the difference of
 two graph fits on simulated data. We simulated under the full graph and fitted two graphs that result from
 deleting either the red admixture edge or the blue admixture edge. These two graphs have the same expected
 *qpGraph* fit score but can have different scores in any one simulation iteration.

365 d QQ plot of *p*-values testing for a score difference between the two graphs (on simulated data) under the null
 366 hypothesis of no difference, confirming that the method is well calibrated.

367 368

369 (4) Determining identifiability of admixture graph parameters

370 Fitting admixture graphs results in an estimate of the overall model fit, as well as in estimates of

371 branch lengths and admixture weights. However, even with infinite data some of these parameters

372 cannot be estimated, as they are not identifiable from the system of equations that corresponds to

373 the admixture graph. Issues like this have been well described for simple topological features of a

374 graph. For example, the lengths of the two branches connected to the root node cannot be

estimated without either fixing one of them or forcing the lengths to be evenly distributed.

Furthermore, in a graph with populations and admixture events, at least one parameter will not
 be identifiable unless the inequality –
 is satisfied (Lipson 2020). However, even in

378 graphs that meet this criterion, some parameters are not identifiable, and until *findGraphs*, there

379 was no method for testing whether any given parameter in an admixture graph is identifiable. We

introduce a method for testing which parameters in an admixture graph are identifiable, and which

381 are not, based on the Jacobi matrix of the graph's system of  $f_2$  equations. Like our method for 382 deriving confidence intervals for admixture graph parameters, this can improve the interpretability 383 of admixture graph analyses.

384

Our methods for automated topology inference, for bootstrapping log-likelihood scores or worst
 residuals for comparing model fits, for cross-validation of admixture graphs of different
 complexities, for estimating confidence intervals and determining identifiability of admixture graph
 parameters can greatly improve the interpretability of admixture graph analyses. We implement all
 these methods in *findGraphs* to assist the user in building a series of models that can explain
 admixture processes of ancient populations in a manner that surpasses all other programs in
 accuracy.

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393

394 (5) Drawing conclusions from a large number of fitted models

As discussed in the Results section, when we apply our methods for finding optimal graphs and

396 comparing admixture graphs to a number of previously published models, we find that there often

397 exists a much larger number of fitting models than has previously been appreciated. In these cases,

398 we are unable to prioritize a single model, or even a small number of models, based on the evidence

399 we have. However, we are still able to reject the vast majority of all tested models. This suggests 400 that insight can be gained by identifying common features among the well-fitting models. We 401 therefore introduce methods for summarizing collections of well-fitting admixture graphs to 402 determine which features they share. In practice, we find that these methods can aid the manual 403 inspection of *findGraphs* results, but the high diversity of well-fitting topologies we see in most case 404 studies and the importance of fitted parameters (especially admixture proportions) for historical 405 interpretation of topologies makes it difficult to reliably automatize the process of interpreting fitted 406 admixture graph models.

408 **Revisiting published admixture graphs** 

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409 410 We studied admixture graphs from eight publications (Lazaridis et al. 2014, Shinde et al. 2019, Sikora 411 et al. 2019, Bergström et al. 2020, Lipson et al. 2020, Hajdinjak et al. 2021, Librado et al. 2021, Wang 412 et al. 2021) with the goal of comparing published models to models identified by our algorithm for 413 automatically inferring optimal (best-fitting) admixture graphs (Table 1). In all but one study 414 *qpGraph* or its automated reimplementation (*admixturegraph*, Leppälä et al. 2017) was used for 415 fitting topologies to genetic data, while Librado et al. (2021) relied on the automated OrientAGraph 416 method. The main question we were interested in is whether we can find alternative models which 417 (1) fit as well as, or better than the published graph, (2) differ in important ways from the published graph, and (3) cannot immediately be rejected based on other evidence such as temporal 418 419 plausibility. The studies were selected according to the criterion that an admixture graph model 420 inferred in the study is used as primary evidence for at least one statement about population history 421 in the main text of the study. In other words, the admixture graph method was used in the original 422 studies to generate new insights into population history, and not simply to show that there is a 423 model that exists (even if it is not unique) that does not contradict results of other genetic analyses, 424 an approach that is a valid use of admixture graphs and has been taken in some studies (e.g., Seguin-425 Orlando et al. 2014, Narasimhan et al. 2019, Wang et al. 2019). There are many published studies 426 that could have been included in our re-evaluation exercise as they meet our key criterion (e.g., 427 Yang et al. 2017, McColl et al. 2018, Posth et al. 2018, Flegontov et al. 2019, Calhoff et al. 2021, 428 Lipson et al. 2022, Vallini et al. 2022). However, critical re-evaluation of each published graph was an 429 intensive process, and the sample of studies we revisited is diverse enough to identify some general 430 patterns and to support strong conclusions.



<sup>\*</sup> the population composition was modified, see Table S1 and the text \*\* certain gene flows were removed from the published model for simplicity, see Table S1 and the text

### 433 Table 1: Published graphs in the context of automatically found graphs.

<sup>431</sup> 432

- 434 We compared graphs from 8 publications to alternative graphs inferred on the same or very similar data (see
- 435 **Table S1** for details).
- 436 **Publication**: Last name of the first author and year of the relevant publication.
- 437 **Figure in the original publication**: Figure number in the original paper where the admixture graph is
- 438 presented.
- 439 **Groups (populations)**: The number of populations in each graph.
- 440 Admixture events: The number of admixture events in each graph.
- 441 **SNPs used**: The number of SNPs (with no missing data at the group level) used for fitting the admixture graphs.
- 442 For all case studies, we tested the original data (SNPs, population composition, and the published graph
- topology) and obtained model fits very similar to the published ones. However, for the purpose of efficient
- topology search we adjusted settings for  $f_3$ -statistic calculation, population composition, or graph complexity as noted in the footnotes, in **Table S1**, and discussed in the text.
- 446 **Publ. model: worst residual, SE**: The worst *f*-statistic residual of the published graph fitted to the SNP set
  447 shown in the "SNPs used" column, measured in standard errors (SE).
- 448 **Distinct alternative topologies found**: The number of distinct newly found topologies differing from the 449 published one.
- 450 Significantly better fitting topologies, %: The percentage of distinct alternative topologies that fit significantly
   451 better than the published graph according to the bootstrap model comparison test (two-tailed empirical *p*-
- 451 value <0.05). If the number of distinct topologies was very large, a representative sample of models (1/20 to
- 452 value <0.05). If the number of distinct topologies was very large, a representative sample of models (1/20 to</li>
   453 1/3 of models evenly distributed along the log-likelihood spectrum) was compared to the published one
- 454 instead, and the percentages in this and following columns were calculated on this sample.
- 455 **Non-significantly better fitting topologies**, %: The percentage of distinct topologies that fit non-significantly 456 (nominally) better than the published graph according to the bootstrap model comparison test (two-tailed 457 empirical p-value  $\ge 0.05$ ).
- 458 **Non-significantly worse fitting topologies**, %: The percentage of distinct topologies that fit non-significantly
- 459 (nominally) worse than the published graph according to the bootstrap model comparison test (two-tailed 460 empirical *p*-value  $\geq$ 0.05).
- 461 **Significantly worse fitting topologies**, %: The percentage of distinct topologies that fit significantly worse than 462 the published graph according to the bootstrap model comparison test (two-tailed empirical *p*-value <0.05).
- 463
- 464

Here we present a high-level summary of these analyses. Discussions of individual graphs, as well an
overview of the methodology, can be found in the next section.

467

468 For 19 out of 22 published graphs we examined we were able to find at least one, but usually many, 469 graphs of the same complexity (number of groups and admixture events) and with a log-likelihood 470 score that was nominally better than that of the published graph (see results for 11 selected graphs 471 in Table 1 and full results for all 22 in Table S1). The 22 graphs were drawn from the 8 publications 472 as there were multiple final graphs presented in some of the publications (Shinde et al. 2019, Sikora 473 et al. 2019, Librado et al. 2021), or we examined selected intermediates in the model construction 474 process (Bergström et al. 2020, Lazaridis et al. 2014, Lipson et al. 2020, Wang et al. 2021), or we 475 introduced an outgroup not used in the original study (Hajdinjak et al. 2021, Sikora et al. 2019), or 476 we tested additional graph complexity classes dropping "unnecessary" admixture events (Lipson et 477 al. 2020, Sikora et al. 2019).

478

479 Oftentimes these alternative graphs are not significantly better than the published one after taking 480 into account variability across SNPs via bootstrapping. In the following cases, at least one model that 481 fits significantly better than the published one according to our bootstrap model comparison 482 method was found: the Bergström et al. and Lazaridis et al. seven-population graphs; the Librado et 483 al. graph with 3 admixture events; the Hajdinjak et al. graphs with or without adding a chimpanzee 484 outgroup; the Lipson et al. intermediate graphs with 7 groups and 4 admixture events and with 10 485 groups and 8 admixture events; the Wang et al. 12-population graph; and the Sikora et al. graphs for 486 West Eurasians and for East Eurasians with 10 or 6 admixture events (Table S1). In nearly all cases 487 (except for the Lazaridis et al. six-population graph, Shinde et al. graph with 8 populations and 3 488 admixture events, and the Librado et al. graph with 4 admixture events), we also identified a large 489 number of graphs that fit the data not significantly worse than the published ones. In every example,

490 some of these graphs have topologies that are qualitatively different in important ways from those 491 of the published graphs. Features such as which populations are admixed or unadmixed, direction of 492 gene flow, or the order of split events, if not constrained *a priori*, are generally not the same 493 between alternative fitting models for the same populations. While some of these graphs can be 494 rejected since their topologies appear highly unlikely because of non-genetic or unrelated genetic 495 evidence, for all of the publications except one (Shinde et al. 2019), there are alternative equally-496 well-or-better-fitting graphs we identified and examined manually that differ in qualitatively 497 important ways with regard to the implications about history, are temporally plausible (for instance, 498 very ancient populations do not receive gene flows from sources closely related to much less ancient 499 groups), and not obviously wrong based on other lines of evidence. These findings suggest the 500 possibility that complex admixture graph models, even with a very good fit to the data, may differ in 501 important ways from true population histories.

502

The previous statements are valid if the original parsimony constraints are applied, i.e., if the graph complexity (the number of admixture events) is not altered. Below in selected case studies (Shinde *et al.*, Librado *et al.*) we explore the effect of relaxing the parsimony constraint.





507

#### 508 Figure 4:

509 Log-likelihood scores of published graphs (those shown in **Table 1**) and automatically inferred graphs. Each dot 510 represents the log likelihood score of a best-fitting graph from one *findGraphs* iteration (low values of the 511 score indicate a better fit); only topologically distinct graphs are shown. Log-likelihood scores for the published 512 models and best-fitting alternative models found are shown by blue and pink x's, respectively. Bootstrap 513 distributions of log-likelihood scores for these models (vertical lines, 90% CI) and their medians (solid dots) are 514 also shown. Lower scores of the fits obtained using all SNPs, relative to the bootstrap distribution, indicate 515 overfitting to the full data set. Green and red horizontal lines show the approximate locations where newly 516 found models consistently have fits significantly better or worse, respectively, than those of the published 517 model. In the case of the Bergström et al., Lazaridis et al. and Hajdinjak et al. studies, one or more worst-fitting 518 models were removed for improving the visualization. The setups shown here (population composition, 519 number of groups and admixture events, topology search constraints) match those shown in Table 1. 520

- 521
- 522 Detailed reconsideration of eight published admixture graphs

#### 523

524 We investigated admixture graphs from eight publications. We usually focused on one final graph 525 from each publication, and in some cases we also discuss simpler intermediates in the published 526 model-building process, apply various topological constraints to the model inference process, or 527 decrease/increase the number of admixture events to explore the influence of parsimony 528 constraints. Table 1 and Figure 4 summarize these results for one or a few graphs from each 529 publication, while Table S1 contains the full results for all studied graphs and setups. Table 2 530 summarizes our assessment of inferences in the original publications that were supported by the 531 published graphs.

532

533 To identify alternative models, we ran many iterations of *findGraphs* for each set of input 534 populations, constraints, and the number of admixture events we investigated, and we selected the 535 best-fitting graph in each iteration, that is, the graph with the lowest log-likelihood (LL) score. Each 536 algorithm iteration was initiated from a random graph, and the algorithm is non-deterministic so 537 that in each iteration it takes a different trajectory through graph space, possibly terminating in a 538 different final best graph. The number of admixture events in the initial random graphs and in the 539 output graphs was always kept equal to that of the published graph. For each example, we counted 540 how many distinct topologies were found with significantly or non-significantly better or worse LL 541 scores than that of the published graph (Table 1, Table S1). To obtain a formally correct comparison of model fit, the published graph and each alternative model were fitted to resampled replicates of 542 543 the dataset and the resulting LL score distributions were compared (see Methods). As shown in 544 Figure 4, for 4 of the 8 publications we re-analyzed, the LL score of the published graph run on the 545 full data is better than almost all the bootstrap replicates on the same data (it falls below the  $5^{tn}$ 546 percentile), which is a sign of overfitting, and underscores the importance of applying bootstrap to 547 assess the robustness of fitted models and conclusions drawn from them.

548

549 The fraction of graphs with scores better than the score of the published graph should not be 550 overinterpreted, as it is influenced by the *findGraphs* algorithm which does not guarantee ergodic sampling from the space of well-fitting admixture graphs. In particular, it is possible that despite 551 552 findGraph's strategies for efficiently identifying classes of well-fitting admixture graphs (see 553 Methods), it has a bias toward missing certain classes of graphs. However, even a single alternative 554 graph which is no worse-fitting than the published graph suggests that we are not able to identify a 555 single best-fitting model. Many of these alternatives, despite providing a good fit to the data, appear 556 unlikely, for example, because they suggest that Paleolithic era humans are mixed between different 557 lineages of present-day humans. We were mainly interested in alternative models which are also 558 plausible, and so we constrained the space of allowed topologies in *findGraphs* to those we 559 considered plausible a priori, in cases where this was necessary for reducing the search space size. 560 Constraints were either integrated into the topology search itself, or were applied to outcomes of 561 unconstrained searches, as detailed below.

562

#### 563 Figure 5:

564 Published graphs from six studies for which we explored alternative admixture graph fits. In all cases, we 565 selected a temporally plausible alternative model that fits nominally or significantly better than the published 566 model and has important qualitative differences compared to the published model with respect to the 567 interpretation about population relationships. In all but one case, the model has the same complexity as the published model shown on the left with respect to the number of admixture events; the exception is the 568 569 reanalysis of Librado et al. 2021 horse dataset since the published model with 3 admixture events is a poor fit 570 (worst Z-score comparing the observed and expected f-statistics has an absolute value of 23.9 even when 571 changing the composition of the population groups to increase their homogeneity and improve the fit relative 572 to the composition used in the published study). For this case, we show an alternative model with 8 admixture 573 events that fits well and has important qualitative differences from the point of view of population history 574 interpretation relative to the published model. The existence of well-fitting admixture graph models does not 575 mean that the alternative models are the correct models; however, their identification is important because 576 they prove that alternative reasonable scenarios exist to published models.

577



586	<b>b,</b> Librado <i>et al</i> . 2021 (modified	Significantly better fitting, temporally/geographically plausible.
587	population composition) published	In contrast to the published graph, in this graph with 8 mixture
588		events (the minimum necessary to obtain an acceptable statistical
589		fit to the data), a lineage maximized in horses associated with
590		Yamnava steppe pastoralists or their Sintashta descendants (C-
591		PONT TURG or DOM2) contributes a substantial amount of
592		ancestry to the horses from the Corded Ware archaeological
593		context (CWC) Thus in this model both CWC humans and horses
59/		are mixtures of Vampava and European farmer-associated
595		lineages. This is qualitatively different from the suggestion that
596		there was no Vampava associated contribution to CWC horses
597		which was a possibility raised in the paper. The eight admixture
537		graph also is different from the published model in that it shows a
500		fitting model where the Terpen herse does not have the history
599		ale image in the study (see an administration of the CMC and DOM2
600		claimed in the study (as an admixture of the CWC and DOW2
601		norses).
602		27.25
	score: 679.89 WR: 23.92	score: 37.36 WR: 3.38
	admix: 3	admix: 8
	survey	129
	181	

603

22%

604



![](_page_16_Figure_2.jpeg)

618

![](_page_17_Figure_1.jpeg)

![](_page_18_Figure_1.jpeg)

650

651 Bergström et al. 2020. The admixture graph for dogs in Figure 1e of Bergström et al. 2020 was 652 inferred by exhaustively evaluating all graphs with two admixture events and outgroup 'Andean fox' 653 for the six populations that remain in the graph after excluding an Early Neolithic dog from 654 Germany. The only six-population graph with a worst residual (WR) below 3 standard errors (SE) was 655 then chosen as a scaffold onto which the Early Neolithic dog genome from Germany was mapped, 656 allowing for one more admixture event, and a seven-population graph with the lowest LL score was 657 shown as the final model in the paper. Alternative six-population scaffolds were not explored in the 658 original publication, although two six-population graphs with fits very similar to the best one were 659 found. LL was not used as a ranking metric for alternative models in the original study; instead, the 660 number of  $f_4$ -statistics having residuals above 3 SE was considered. Since no  $f_3$ -statistics were 661 negative when all sites available for each population triplet were used (the "useallsnps: YES" option), 662 we did not use the upgraded algorithm for calculating  $f_3$ -statistics on pseudo-diploid data.

663

664 Our findGraphs results confirm the published six-population graph in that no graph with lower LL 665 score is identified, but 3 of 14 unique alternative graphs found fit not significantly worse than the 666 published graph (the published graph was also recovered by findGraphs) (Table 1, Table S1). When 667 we used *findGraphs* to infer seven-population graphs with three admixture events (again fixing 668 Andean fox as the outgroup), we identified 5 graphs with log-likelihood score nominally better than 669 that of the published graph and one with a score that is slightly lower than that of the published 670 graph but actually significantly better according to our model comparison methodology (this model 671 is very similar to the published graph, Figure S2). In the newly found seven-population graph with 672 the best log-likelihood score (Figure S2), the Siberian (Baikal), American, and Levantine dogs are 673 admixed, and the West European, East European (Karelia), and dogs of Southeast Asian origin (New 674 Guinea singing dog) are unadmixed, while the opposite pattern is found in the published graph 675 (Table 2). The best-fitting graph does not fit the data significantly better than the published graph 676 (two-tailed empirical p-value = 0.332), but it bears a closer resemblance to the human population 677 history (see the third-best graph found by findGraphs on human data from Bergström et al. 2020 in 678 Figure S3) than the published seven-population graph (Figure 5a, Figure S2).

679

680 In this new seven-population model (Figure 5a), both American and Siberian dog lineages represent 681 a mixture between groups related to the Asian and East European dog lineages, and robust genetic 682 results suggest that in the time horizon investigated in the original publication (after ca. 10,900 years 683 ago) nearly all Siberian (Jeong et al. 2019, Sikora et al. 2019) and all American (Raghavan et al. 2014, 684 Raghavan et al. 2015, Moreno-Mayar et al. 2018) human populations were admixed between groups 685 most closely related to Europeans and Asians. According to this model, Levantine dogs are modeled 686 as a mixture of a basal branch (splitting deeper than the divergence of the Asian and European dogs) 687 and West European dogs, again in agreement with current models of genetic history of Middle 688 Eastern human populations who are modeled as a mixture of "basal Eurasians" and West European 689 hunter-gatherers (Lazaridis et al. 2016, Lipson et al. 2017). Although greater congruence with human 690 history increases the plausibility of *findGraph*'s newly identified model relative to the published 691 model, to make unbiased comparisons between the history of the two species, model selection 692 should be done strictly independently for each species, and so the genetic data alone does not favor 693 one model more than another. Our results provide a specific alternative hypothesis that differs in 694 qualitatively important ways from the published model and can be tested against new genetic data 695 as it becomes available as well as other lines of genetic analysis of existing data.

696

697 To explain why the original paper on the population history of dogs missed the model that 698 findGraphs identified that is plausibly a closer match to the true history, we observe that the 699 Bergström et al. 2020 admixture graph search was exhaustive under the parsimony constraint (no 700 more than 2 admixture events for 6 populations), and thus missed the potentially true topology 701 including 3 admixture events for these 6 populations. This case study also illustrates that even in a 702 relatively low complexity context (7 groups and 3 admixture events) applying manual approaches for 703 finding optimal models is risky. When any new group such as an Early Neolithic dog from Germany is 704 added to the model, it may introduce crucial information into the system, and re-exploring the

705 whole graph space in an automated way is advisable. In contrast, mapping a newly added group on a 706 simple skeleton graph (even when that skeleton is a uniquely best-fitting model) may yield a 707 topology that is at odds with the true history. As the original Bergström *et al.* paper noted (Fig. 3C of 708 that study), no congruent six-population graph models were found for humans and dogs under the 709 parsimony assumption: the three most congruent graphs for dogs resulted in poorly fitting models 710 for the corresponding human populations (WR above 10 SE), and the three most congruent graphs 711 for humans resulted in poorly fitting models for the corresponding dog populations (WR between 5 712 and 10 SE). We added a West European hunter-gatherer group to the set of human groups from the 713 original publication and using *findGraphs* on the original set of 77K transversion SNPs we found that 714 the third best-fitting model for humans (Figure S3) (which is not significantly different in fit from the 715 first one) is topologically identical to the newly found dog graph.

716

717 Even though *findGraphs* identified an admixture graph topology that fits the data as well as the 718 seven-population graph in Bergström et al. and is qualitatively quite different with respect to which 719 populations were admixed, the new topology continues to support another of the key inferences of 720 that study: that many of the early divergences among domesticated dog lineages occurred prior to 721 the date of the Karelian dog (~10,900 ya). Thus, both graphs concur in providing strong evidence 722 that the radiation of domesticated dog lineages occurred by the early Holocene, prior to the 723 domestication of other animals. We further emphasize that the Bergstrom et al. 2020 graph is the 724 best-case scenario (along with Lazaridis et al. 2014 discussed below) for published admixture graphs. 725 Most published graphs are far less stable even than this.

726

# Table 2: Features of the published admixture graphs that support inferences in the original studies and the level of their support in our re-analysis.

The table lists key features whose support we assessed in sets of alternative well-fitting and temporally plausible models generated by *findGraphs*. Since this assessment had to be performed manually, only in two cases (marked by asterisks) all models fitting better and non-significantly worse than the published one were

732 scrutinized; in other cases only a subset of best-fitting models was examined (see the sections for details).

733

Study	Groups / admixture	Features of the published model supported by temporally	Features of the published model <u>not</u> supported by temporally plausible alternative
	events	generated by <i>findGraphs</i>	models generated by findGraphs
Bergström et al. 2020	7/3*	Early divergence of domesticated dog lineages (prior to the date of the Karelian dog, 10,900 ya).	Siberian (Baikal), American, and Levantine dog lineages are unadmixed, and the West European (Germany Early Neolithic), East European (Karelia), and dogs of Southeast Asian origin (New Guinea singing dog) are admixed.
Lazaridis et al. 2014	7 / 4	Present-day Europeans represent a mixture of three ancestral sources related to the following groups: Mal'ta (MA1), West European hunter-gatherers, and early European farmers.	N/A
Shinde et al. 2019	8/3*	<ul> <li>(1) Iranian farmer-related ancestry in the Indus Periphery group is not derived from the Hajji Firuz Neolithic or Tepe Hissar Chalcolithic groups.</li> <li>(2) There is Asian-related ancestry in the Indus Periphery group.</li> </ul>	N/A

	8/4	(2) There is Asian ancestry in the Indus Periphery group.	(1) Iranian farmer-related ancestry in the Indus Periphery group is not derived from the Hajji Firuz Neolithic or Tepe Hissar Chalcolithic groups.
Librado et al. 2021	10 / 8 or 9	(2) DOM2 and C-PONT are sister groups (they form a clade); (4) there was gene flow from a deep-branching ghost group to the NEO-ANA group.	<ul> <li>(1) NEO-ANA-related admixture is absent in the DOM2 group;</li> <li>(3) there is no gene flow connecting the CWC group and the cluster associated with Yamnaya horses and horses of the later Sintashta culture whose ancestry is maximized in the Western Steppe (DOM2, C- PONT, TURG);</li> <li>(5) Tarpan is a mixture of a CWC-related and a DOM2- related lineage.</li> </ul>
Hajdinjak et al. 2021	12 / 8	(3) the Vestonice16 lineage is a mixture of a Sunghir-related and a BK1653-related lineage.	<ul> <li>(1) there are gene flows from the lineage found in the ~45,000-43,000-years-old Bacho Kiro Initial Upper Paleolithic</li> <li>(IUP) associated lineage to the Ust'-lshim, Tianyuan, and GoyetQ116-1 lineages;</li> <li>(2) the ~35,000-years-old Bacho Kiro Cave individual BK1653 belonged to a population that was related, but not identical, to that of the GoyetQ116-1 individual.</li> </ul>
Lipson et al. 2020	12/11	N/A	<ul> <li>(1) A lineage maximized in present-day West African groups (Lemande, Mende, and Yoruba) also contributed some ancestry to the ancient Shum Laka individual and to present-day Biaka and Mbuti;</li> <li>(2) another ancestry component in Shum Laka is a deep- branching lineage maximized in the rainforest hunter-gatherers Biaka and Mbuti;</li> <li>(3) "super-archaic" ancestry (i.e., diverging at the modern human/Neanderthal split point or deeper) contributed to Biaka, Mbuti, Shum Laka, Lemande, Mende, and Yoruba;</li> <li>(4) a ghost modern human lineage (or lineages) contributed to Agaw, Mota, Biaka, Mbuti, Shum Laka, Lemande, Mende,</li> </ul>

			and Yoruba.
Wang et al. 2021	12 / 8	N/A	Admixture from a source related to Andamanese hunter- gatherers is almost universal in East Asians, occurring in the Jomon, Tibetan, Upper Yellow River Late Neolithic, West Liao River Late Neolithic, Taiwan Iron Age, and China Island Early Neolithic (Liangdao) groups.
Sikora et al. 2019 "West"	13 / 6	N/A	The Mal'ta (MA1_ANE) lineage received a gene flow from the Caucasus hunter-gatherer (CaucasusHG_LP or CHG) lineage.
Sikora et al. 2019 "East"	14/6	(2) European-related ancestry in the Kolyma, USR1, and Clovis lineages is closer to Mal'ta than to Yana.	<ul> <li>(1) the Mal'ta (MA1_ANE) and Yana (Yana_UP) lineages received gene flow from a common East Asian-associated source diverging before the ones contributing to the Devil's Cave (DevilsCave_N), Kolyma (Kolyma_M), USR1 (Alaska_LP), and Clovis (Clovis_LP) lineages;</li> <li>(3) the Devil's Cave lineage received no European-related gene flows, and Kolyma has less European-related ancestry than ancient Americans (USR1 and Clovis).</li> </ul>

734

735

736 Lazaridis et al. 2014. The graph in Figure 3 (Lazaridis et al. 2014) was inferred in the following 737 manner. First, a phylogenetic tree without admixture was constructed which was the best fit for all  $f_4$ -statistics among the populations "Mbuti", "WHG Loschbour" (Lazaridis et al. 2014), "LBK 738 Stuttgart" (Lazaridis et al. 2014), "Onge", and "Karitiana", with "Mbuti" fixed as an outgroup. Next, 739 740 all possible admixture graphs were considered that result from adding a single admixture edge to 741 this tree. After it was found that each of them had a WR > 3 SE, several graphs with two admixture events were considered, and some of them had WR < 3 SE. The "MA1" genome (Raghavan et al. 742 743 2014) was added to these graphs in several different ways, and only one of these configurations was 744 found to have WR < 3 SE. This was then used as a skeleton graph onto which a European population 745 (represented by different present-day groups) was added. No fitting graph was found in which 746 present-day Europeans could be modeled as a two-way mixture (adding one admixture event to the 747 graph). After inspecting the non-fitting f-statistics of one of these graphs, it was found that modeling 748 modern Europeans as a three-way mixture (adding two admixture events to the graph) is consistent 749 with all f-statistics. Six  $f_3$ -statistics were negative when all sites available for each population triplet 750 were used (the "useallsnps: YES" option, **Table S1**), but the upgraded algorithm for calculating  $f_{3}$ -751 statistics on pseudo-diploid data had no effect since the only pseudo-diploid group in the dataset 752 (MA1) was a singleton population, and the algorithm removes sites with only one chromosome 753 genotyped in any non-singleton population. Thus, below we show results generated using the 754 standard algorithm for calculating *f*-statistics. 755

First, we considered the published skeleton graph onto which a European population was later added (Lazaridis et al. 2014). As in the (Bergström et al. 2020) example, the best six-population graph with two admixture events found by *findGraphs* is identical to the published six-population

graph, which has a LL score of 3.0 (**Table S1**). The second-best graph found has a LL score of 31.8.

760 When computing the bootstrap *p*-value for the difference between these two graphs, we find that in

761 1.6% of all SNP resamplings the second-best graph has a better score than the published graph,

resulting in a two-tailed empirical *p*-value of 0.032 for a difference in fits between these two graphs.

All 14 alternative graphs found by our algorithm fit significantly worse than the published graph
 (Table S1).

765

When we add the European population (French) and consider seven-population graphs with four
admixture events, we find 40 out of 306 distinct graphs with a score better than that of the
published graph (10 of those graphs are shown in Figure S4). The best-fitting newly found model and
two other models fit the data significantly better than the published model (Table S1), but their
topology is qualitatively very similar to that of the published graph (Figure S4). In the best-fitting
newly found model, French and Karitiana share some drift to the exclusion of MA1, while in the
published model the source of MA1-related ancestry in French is closer to MA1 than to Karitiana.

773

774 It is important to point out that not all of the 40 alternative graphs that fit nominally or significantly 775 better than the published one are consistent with the conclusion that modern European populations 776 are admixed between three different ancestral populations (Figure S4). For example, the fifth 777 alternative graph in **Figure S4** that is fitting nominally better than the published model (p-value = 778 0.464) includes no basal Eurasian ancestry in early European farmers (LBK Stuttgart), and instead 779 models Onge as having ~50% West Eurasian-related ancestry and MA1 as having ~25% Asian 780 ancestry. According to that graph, the present-day European population was formed by admixture 781 of a MA1-related lineage and a European Neolithic-related lineage, with no West European hunter-782 gatherer (WHG) contribution. Of course, other lines of evidence make it clear that LBK Stuttgart is a 783 mixture of Anatolian farmer-related ancestry and WHG Loschbour-related ancestry (Lazaridis et al. 784 2016, Lipson et al. 2017), thus providing external information in favor of the Lazaridis et al. 2014 785 model, and the use of such ancillary information in concert with graph exploration is important in 786 order to obtain more confident inferences about population history taking advantage of admixture 787 graphs.

788

789 The second alternative graph in Figure S4 that fits just negligibly worse than the highest-ranking 790 model has another distinctive feature: LBK Stuttgart is modeled as a mixture of a WHG-related and a 791 basal Eurasian lineage, but modern Europeans receive a gene flow not from the LBK-related lineage, 792 but from its basal Eurasian source. Although temporally plausible, this model is much less plausible 793 from the archaeological point of view than the published model, and thus in this case too we can 794 reject it as unlikely based on non-genetic evidence. We note, however, that a large group of newly 795 found models (247 graphs) fits not significantly worse than the published one (Table S1), and those 796 are topologically diverse. Thus, strictly speaking, the admixture graph method on the given dataset 797 cannot be used to prove that the published model is the only one fitting the data.

798 799

800 Shinde et al. 2019. The skeleton admixture graph in the original study (Shinde et al. 2019) was 801 constructed manually on the basis of a SNP set derived from the 1240K enrichment panel, and 802 subsequently all possible branching orders (105) within the five-population Iranian farmer-related 803 clade were tested. The published model (Fig. 3 in that study) included 9 groups and 3 admixture 804 events, but one group (Belt Cave Mesolithic) had a very high missing data rate, and as a result model 805 fitting relied not just on the merged dataset which included 19,000 polymorphic sites without 806 missing data across groups, but also on a dataset with approximately 470,000 sites that excluded the 807 Belt Cave individual. The topological inferences were consistent for both analyses (Table S3 of that 808 study). Following the approach of the published paper, we repeated *findGraphs* analysis both with 809 and without the Belt Cave individual. Thus, we initially explored the following topology classes: 9 810 groups with 3 admixture events on ca. 19,000 polymorphic sites and 8 groups with 3 admixture 811 events on ca. 470,000 sites (Table S1). The sample composition of the groups and the SNP dataset

812 matched that in the original study. We summarize results across 4,000 independent iterations of the 813 *findGraphs* algorithm for each topology class.

814

815 For the nine-population graph we found 89 models with LL nominally better than that of the 816 published model (Table S1). For the eight-population graph, we found 61 nominally and 4 817 significantly better fitting models (Table S1), and their topological diversity was high (Figure S5). We 818 note that the following groups were admixed by default in the graph models compared in the 819 original study: Hajji Firuz Neolithic (labeled "Chalcolithic" in that study but the dates are Neolithic) 820 and Tepe Hissar Chalcolithic were considered as mixtures of an Anatolian farmer-related lineage and 821 an Iranian farmer-related lineage; Indus Periphery was considered as a mixture of an Andamanese-822 related lineage representing ancient South Indians (ASI) and an Iranian farmer-related lineage. However, calculation of negative "admixture"  $f_3$ -statistics for these target groups is impossible using 823 824 the original dataset and the original model fitting algorithm for several reasons. First, the Indus 825 Periphery group was represented by a single pseudo-diploid individual (18726) from the "Indus 826 Valley cline" for whom the best-quality data were available. But direct calculation of "admixture"  $f_{3}$ -827 statistics for such a group as a target is impossible since its heterozygosity cannot be estimated. 828 Second, as discussed above, in Classic ADMIXTOOLS it is impossible to apply a correction intended 829 for accurate calculation of f<sub>3</sub>-statistics on pseudo-diploid data (the "inbreed: YES" option) if there is 830 at least one population composed of one individual only (a singleton population). Third, the original 831 Hajji Firuz Neolithic group composed of five individuals included a family of three 2<sup>nd</sup> or 3<sup>rd</sup>-degree relatives, and that artificially inflated the drift on the Hajji Firuz branch and made detecting a 832 833 negative statistic  $f_3$ (Hajji Firuz Neolithic; X, Y), even if present, highly unlikely. Indeed, no  $f_3$ -statistic 834 turned out to be nominally negative for the groups on the eight-population graph when statistics 835 were calculated according to the original settings (we used settings equivalent to "useallsnps: NO" 836 and "inbreed: NO" in classic ADMIXTOOLS, 470,389 polymorphic sites were available). Considering this fact, it is not surprising that our automated topology space search is not well constrained. The 837 838 original study differed from ours since the constraints were introduced manually, but we wanted our 839 topology search to be automatic and to explore a wider range of parameter space.

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841 In order to provide power to detect negative  $f_3$ -statistics useful for constraining the model search, 842 we 1) removed two members of the family from the Hajji Firuz Neolithic group, 2) extended the 843 number of individuals and sites available for the Indus Periphery group by generating new shotgun-844 sequencing data for a previously published library (Narasimhan et al. 2019) derived from individual 18726 (see Table S2) and by adding published data for three other individuals from the Indus Valley 845 846 cline (from Gonur in Turkmenistan and Shahr-i-Sokhta in Iran; Narasimhan et al. 2019, Shinde et al. 2019), 3) removed from other groups two individuals based on  $2^{nd}-3^{rd}$  degree relatedness, and 4) 847 848 removed two individuals from other groups based on evidence of contamination with modern 849 human DNA. All the changes to the dataset are shown in Table S3. In addition to these dataset 850 adjustments, the new algorithm for calculating f-statistics makes it possible to compute negative  $f_{3}$ -851 statistics on pseudo-diploid data, but at a cost of removing sites with only one chromosome 852 genotyped in any non-singleton population (see Methods). We eventually detected significantly 853 negative "admixture"  $f_3$ -statistics  $f_3$ (Tepe Hissar Chalcolithic; Ganj Dareh Neolithic, Anatolia 854 Neolithic),  $f_3$  (Indus Periphery; Ganj Dareh Neolithic, Onge), and other similar statistics for the same 855 target groups. We also observed a nominally negative (Z-score = -0.6) statistic  $f_3$ (Hajji Firuz Neolithic; 856 Ganj Dareh Neolithic, Anatolia Neolithic), which is suggestive but does not by itself prove admixture 857 in Hajji Firuz Neolithic. For this updated analysis, 249,009 variable sites without missing data at the 858 group level were available for the eight populations.

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We repeated topology search with this set of *f*-statistics providing additional constraints, performing
4,000 runs of the *findGraphs* algorithm. The Mota ancient African individual was set as an outgroup
and 3 admixture events were allowed in the eight-population graph. Among 4,000 resulting graphs
(one from each *findGraphs* run), 144 were distinct topologically, and the published model was
recovered in 13 runs of 4,000 (**Table S1**). Only 4 distinct topologies fitting nominally better than the
published one were found, and those had LL scores almost identical to that of the published eight-

866 population model (16.97 and 17.66 vs. 17.85). These four alternative models (Figure S6b) shared all 867 topologically important features of the published model (Figure S6a). Five other topologies differed 868 in important ways from the published one and emerged as fitting the data worse, but not 869 significantly worse, than the published one (Figure S6c): two-tailed empirical p-values reported by 870 our bootstrap model comparison method ranged between 0.060 and 0.112. Three of these 871 topologies included a trifurcation of Iranian farmer-related lineages leading to the Indus Periphery, 872 Hajji Firuz Neolithic, and Ganj Dareh Neolithic groups. The other two topologies included Hajji Firuz 873 Neolithic as an unadmixed Anatolian-related lineage. In both cases, Indus Periphery was modeled as 874 receiving a gene flow from either the Onge lineage (a proxy for ASI) or a deep Asian lineage. 875 876 The finding that the predominant ancestry component of the Indus Periphery group was the most 877 basal branch in the Iranian farmer clade was a prominent claim of the original study (Shinde et al. 878 2019); for example, the abstract stated: "The Iranian-related ancestry in the IVC derives from a 879 lineage leading to early Iranian farmers, herders, and hunter gatherers before their ancestors 880 separated". Our finding that the Hajji Firuz Neolithic lineage may be as deep within the Iranian clade 881 as the Indus Periphery lineage or may even diverge from the Anatolian branch shows that this

- statement cannot be confidently made based on admixture graph analysis alone.
- 883

884 However, the findings we have described up to this point do not invalidate the broader conclusion 885 that the admixture graph modeling in Shinde et al. was used to support; namely (using the phrasing 886 from the abstract) that the genetic data "contradict... the hypothesis that the shared ancestry 887 between early Iranians and South Asians reflects a large-scale spread of western Iranian farmers 888 east." This finding if correct is important, since it implies that the Iranian-related ancestry in the IVC 889 (Indus Valley Civilization genetic grouping, which is the same group as IP), split from the Iranian-890 related ancestry in the first Iranian plateau farmers before the date of the Hajji Firuz farmers, who at 891  $\sim$ 8000 years ago are among the earliest people living on the Iranian plateau known to have grown 892 West Asian crops. The ancient DNA record combined with radiocarbon dating evidence suggests that 893 beginning around the time of the Hajji Firuz farmers, both West Asian domesticated plants such as 894 wheat and barley, and Anatolian farmer-related admixture, began spreading eastward across the 895 Iranian-plateau. If the Iranian-related ancestry in IP was spread eastward into the Indus Valley across 896 the Iranian-plateau as part of the same agriculturally-associated expansion—perhaps brought by 897 people speaking Indo-European languages as well as introducing West Asian crops—then we would 898 expect to see at least some of the Iranian-related ancestry in IP being a clade with that in Hajji Firuz 899 relative to Ganj Dareh. The fact that we do not find any models compatible with this scenario is thus 900 a potentially important finding. In summary, there are two reasons the genetic analyses we have 901 reported up to this point continue to support the finding that the Iranian-related ancestry in IP is not 902 a clade with the Iranian-related ancestry in Hajji Firuz (and Tepe Hissar) and thus is unlikely to reflect 903 the same eastward movement of agriculturalists. First, in findGraphs analysis, all models specifying 904 IP and Tepe Hissar and/or Hajji Firuz as a clade relative to Ganj Dareh were significantly worse-fitting 905 that the published one. Instead, either the Iranian-related ancestry in IP definitively splits off first 906 (the topology from Shinde et al.), or the branching order of IP, Ganj Dareh, and the Hajji Firuz / Tepe 907 Hissar lineages cannot be determined, or IP, Ganj Dareh, and Tepe Hissar are a clade relative to Hajji 908 Firuz. In all these fitting topologies, the ~10,000-year-old radiocarbon date of the Ganj Dareh 909 individuals sets a lower bound on the split time between IP and Hajji Firuz / Tepe Hissar, which is 910 pre-agriculturalist. This suggests that the Iranian-related ancestry in IP is not due to an eastward 911 agriculturalist expansion.

912

But in fact, the admixture graph analysis reported above is not an adequate exploration of the
problem. Although absolute fits of the best models found are good (WR = 2.5 SE), the parsimony
constraint allowing only 3 admixture events precluded correct modeling of basal Eurasian ancestry
shared by all Middle Eastern groups (Lazaridis et al. 2016) or of the Indus Periphery group itself, for
which a more complex 3-component admixture model was proposed (Narasimhan et al. 2019).
Concerned that this oversimplification could be causing our search to miss important classes of
models, we explored *qpAdm* models for the Indus Periphery group further, following the "distal"

920 protocol with "rotating" outgroups outlined by Narasimhan et al. (2019) and using the dataset and 921 outgroups ("right" populations) from that study. All sites available for analyses were used, following 922 Narasimhan et al. (the "useallsnps: YES" option). The combined Indus Periphery group we analyzed 923 included 7 individuals from Shahr-i-Sokhta and 3 individuals from Gonur (3 individuals were 924 removed from the Narasimhan et al. 2019 dataset due to potential contamination with modern 925 human DNA and low coverage). We removed one individual from the Ganj Dareh Neolithic group as potentially contaminated, and one 2<sup>nd</sup> or 3<sup>rd</sup> degree relative was removed from the Anatolia 926 927 Neolithic group, see the dataset composition in **Table S4**. We note that no "distal" *apAdm* models 928 were tested for the combined Indus Periphery group by Narasimhan et al. (2019), and individuals 929 from this group were modeled one by one (Table S82 from Narasimhan et al. 2019), which 930 potentially reduced the sensitivity of the method.

931

A model "Indus Periphery = Ganj Dareh Neolithic + Onge (ASI)" was strongly rejected for the Indus 932 Periphery group of 10 individuals with a *p*-value =  $2 \times 10^{15}$ , and a model that was shown to be fitting 933 934 for all Indus Periphery individuals modeled one by one by Narasimhan et al. (Ganj Dareh Neolithic + 935 Onge (ASI) + West Siberian hunter-gatherers (WSHG)) was rejected for the grouped individuals with 936 a p-value = 0.0044. In contrast, a model "Indus Periphery = Ganj Dareh Neolithic + Onge (ASI) + 937 WSHG + Anatolia Neolithic" was not rejected based on the p>0.01 threshold used in Narasimhan et 938 al. (p-value was marginal but passing at 0.03) and produced plausible admixture proportions for all 939 four sources that are confidently above zero: 53.2 ± 5.3%, 28.7 ± 2.1%, 10.5 ± 1.3%, 7.7 ± 2.9%, 940 respectively (Table S5). The same "distal" model albeit with Anatolian Neolithic always in higher 941 proportion was found as one of the simplest models (or the only simplest model) fitting the data for many other groups from Iran and Central Asia explored by Narasimhan et al. (2019): Aligrama2\_IA 942 943 (13% Anatolia Neolithic), Barikot\_H (21%), BMAC (26%), Bustan\_BA\_o2 (15%), Butkara\_H (24%), 944 Saidu Sharif H o (12%), Shahr I Sokhta BA1 (19%), SPGT (23%) (Table S5 gives a compendium of 945 distal modeling results by Narasimhan et al.). When we modeled Indus Periphery individuals 946 separately, as in Narasimhan et al. (2019), the simplest two-component model "Ganj Dareh Neolithic 947 + Onge (ASI)" was rejected for 5 of 10 individuals (at least ~315,000 sites were genotyped per 948 individual), including the individual I8726 used for the admixture graph analysis in Shinde et al. 949 (Table S6). The model "Ganj Dareh Neolithic + Onge (ASI)" was not rejected only for individuals with 950 fewer than 141,000 sites genotyped, suggesting that this result is attributed not to population 951 heterogeneity, but to lack of power.

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953 These *qpAdm* results show that the parsimony assumption that was made when constructing the 954 admixture graph analysis in Shinde et al. (2019) is contradicted by f-statistic evidence, and indeed 955 Narasimhan et al. themselves showed this when they presented a distal *qpAdm* model that was 956 more complex (Ganj Dareh Neolithic + Onge (ASI) + WSHG) than the one used for constraining the 957 admixture graph model comparison (Ganj Dareh Neolithic + Onge (ASI)). Another line of evidence 958 used to support the principal historical conclusion by Shinde et al. was a series of  $f_4$ -statistic cladality 959 tests following correction of allele frequencies using an admixture model "target group = Iranian 960 farmer + Anatolia Neolithic + Onge (ASI)", with a great majority of tests supporting the deepest 961 position of the Iranian farmer ancestry component in the Indus Periphery group within the Iranian 962 farmer clade (Shinde et al. 2019). However, the model used for allele frequency correction (Ganj 963 Dareh Neolithic + Onge (ASI) + Anatolia Neolithic) was simpler than the 4-component model and 964 different from the 3-component model for the Indus Periphery group suggested by Narasimhan et 965 al. (2019), which is a weakness of that analysis. A valuable direction for future work would be to 966 repeat this analysis with a 4-component allele frequency correction model (Ganj Dareh Neolithic + 967 Onge (ASI) + WSHG + Anatolia Neolithic), although that is beyond the scope of the present study, 968 which simply aims to re-visit the reported analyses and test if they fully support their inferences by 969 ruling out alternative explanations.

970

971 To explore how the parsimony constraint influences results, we allowed 4 admixture events in the

972 eight-population graph (Table S1). Among 4,000 resulting graphs (one from each *findGraphs* run),

973 443 were distinct topologically, and 270 had WRs between 2 and 3 SE, i.e., fitted the data well. We

974 explored 35 topologies with LL scores in a narrow range between 9.3 (the best value) and 13.3. In 975 Figure S7b we show four graphs with four admixture events that model the Indus Periphery group as 976 a mixture of three or four sources, with a significant fraction of its ancestry derived from the Hajji 977 Firuz Neolithic or Tepe Hissar Chalcolithic lineages including both Iranian and Anatolian ancestries. 978 The fits of these models are just slightly different (e.g., LL = 11.7 vs. 9.3, both WRs = 2.4 SE) from that 979 of the best-fitting model (Figure S7a), and similar to that of the published graph. Besides these four 980 illustrative graphs, dozens of topologies with very different models for the Indus Periphery group fit 981 the data approximately equally well, suggesting that there is no useful signal in this type of 982 admixture graph analysis when the parsimony constraint is relaxed (this finding is similar to that in 983 our reanalysis of the dog admixture graph in Bergström et al. 2020, where relaxation of the 984 parsimony constraint identified equally well fitting admixture graphs that were very different with 985 regard to their inferences about population history). These results show that at least with regard to 986 the admixture graph analysis, a key historical conclusion of the study (that the predominant genetic 987 component in the Indus Periphery lineage diverged from the Iranian clade prior to the date of the 988 Ganj Dareh Neolithic group at ca. 10 kya and thus prior to the arrival of West Asian crops and 989 Anatolian genetics in Iran) depends on the parsimony assumption, but the preference for three 990 admixture events instead of four is hard to justify based on archaeological or other arguments. 991

992 Why did the Shinde et al. 2019 admixture graph analysis find support for the IP Iranian-related 993 lineage being the first to split, while our *findGraphs* analysis did not? The Shinde et al. 2019 study 994 sought to carry out a systematic exploration of the admixture graph space in the same spirit as 995 findGraphs—one of only a few papers in the literature where there has been an attempt to do so— 996 and thus this qualitative difference in findings is notable. We hypothesize that the inconsistency 997 reflects the fact that the deeply-diverging WSHG-related ancestry (Narasimhan et al. 2019) present 998 in the IVC (Indus Valley Civilization genetic grouping, which is the same group as Indus Periphery) at 999 a level of ca. 10% was not taken into account explicitly neither in the admixture graph analysis nor in 1000 the admixture-corrected  $f_4$ -symmetry tests also reported in Shinde *et al.* (2019). The difference in 1001 qualitative conclusions may also reflect the fact that the Shinde et al. study was distinguishing 1002 between fitting models relying on a LL difference threshold of 4 units (based on the AIC). As 1003 discussed above, AIC is not applicable to admixture graphs where the number of independent model 1004 parameters is topology-dependent even if the numbers of groups and admixture events are fixed, 1005 and models compared with AIC should have the same number of parameters. Moreover, model 1006 comparisons with AIC do not account for the variability across SNPs, unlike the bootstrap model-1007 comparison method we use. Thus, the analysis by Shinde et al. was over-optimistic about being able 1008 to reject models that were in fact plausible using its admixture graph fitting setup.

1009

1010 The archaeological and linguistic implications of the Shinde et al. study are important, and there are 1011 several avenues available for further attempting to distinguish historical scenarios using f-statistics 1012 that are outside the scope of a methodological study like this one. Some of our observations that are 1013 most challenging for the conclusions of Shinde *et al.* are those related to the graphs with four 1014 admixture events in Figure S7b that fit the Iranian farmer-related ancestry in the Indus Periphery 1015 group as deriving partially from the Hajji Firuz Neolithic or Tepe Hissar Chalcolithic-related lineages. 1016 *qpAdm* is able to use information from distal outgroups (such as WSHG) not included in the 1017 admixture graph modeling exercise revisited here. Leveraging this information might be able to 1018 obtain constraints that would further test the key historical conclusions from Shinde et al. Non-f-1019 statistic-based methods could also be informative. Finally, we emphasize that the  $f_4$ -statistic cladality 1020 tests correcting for the Anatolian farmer-related and Onge-related admixture in the Indus Periphery 1021 grouping do continue to provide support for the historical conclusion of Shinde et al. (these analyses 1022 reject models where the Tepe Hissar or Hajji Firuz groups share genetic drift with the Indus Periphery individuals), with the caveat that they do not correct for the WSHG admixture. 1023 1024 1025

1026 **Librado et al. 2021.** In contrast to the other studies revisited in our work, the admixture graph 1027 published by Librado *et al.* (2021) was inferred automatically using *OrientAGraph*. Models with three

1028 (Fig. 3b in that study) and zero to five (Ext. Data Fig. 5a-d) admixture events were shown. The 1029 dataset included 10 populations (9 horse populations and donkey as an outgroup) and was based on 1030 7.4 million polymorphic transversion sites with no missing data at the group level. We observed that 1031 some groups used for the OrientAGraph and qpAdm analyses were very broad geographically and 1032 temporally (see Table S1 in the original study), and thus we tested two alternative group 1033 compositions: the original one and a streamlined one. In the latter case we included individuals from 1034 one archaeological site and one archaeological period per group: the Botai, C-PONT, DOM2, ELEN, 1035 and NEO-ANA groups were modified in this way, and the CWC, LP-SFR, Tarpan, and TURG groups 1036 were left with the composition used in the original paper (Table S7). In addition, 7 individuals with 1037 missing data proportion exceeding 80% were removed from the analysis, affecting the donkey 1038 outgroup, DOM2, and NEO-ANA groups (**Table S7**). Since among all possible  $f_3$ -statistics for the 10 1039 populations three were negative (using all sites available for each population triplet, "useallsnps: 1040 YES"), we applied the upgraded algorithm for calculating f-statistics, which removed sites with only 1041 one chromosome genotyped in any non-singleton population, resulting in the following site counts for the original and modified population compositions: 11,092 and 1,767,419 sites, respectively. The 1042 1043 very low number of sites available in the former case is due to the fact that all individuals are 1044 pseudohaploid, and that two groups (the donkey outgroup and NEO ANA) are composed of two 1045 individuals, a high-coverage one and a low-coverage one. Thus, just sites genotyped in both donkey 1046 individuals and in both NEO ANA individuals were kept. Considering this problem, we focused on 1047 the modified group composition only. We tested a range of model complexities (from 3 to 9 gene 1048 flows) and performed 1,000 findGraphs topology search iterations per model complexity class. 1049

1050 Unlike all the other admixture graphs we re-evaluate in this study whose fits to the data were 1051 evaluated in the published studies using *qpGraph*, the topologies published in Librado *et al.* 2021 1052 (with 3 to 5 admixture events) were not evaluated for statistical goodness-of-fit, and in fact fit the f-1053 statistic data so poorly that even simple statistics show they cannot be correct (Figure 5b, Figure 1054 S8a, c, e, Table S1). In this case, the approach of using *findGraphs* to identify alternative topologies 1055 with the same number of admixture events that fit the data better is meaningless, as both the 1056 published models and the alternative models do not have enough degrees of freedom to 1057 accommodate the complexity present in the real data; all models are guaranteed to be wrong. In 1058 particular, we found that WR of the published model with 3 admixture events is 23.9 SE (Figure S8a). 1059 In this complexity class *findGraphs* found 22 topologically diverse models that fit significantly better 1060 than the published one (Table 1, Table S1), but nevertheless have extremely poor absolute fits (from 1061 16.2 to 21.3 SE, see a temporally plausible example in Figure S8b). In the complexity class with 4 1062 admixture events, no model fitting better than the published one was found; however, five 1063 alternative models fitting not significantly worse than the published one had lower WR (10 or 12 SE 1064 vs. 14.1 SE, Figure S8d). The WR of the published model with 5 admixture events was 6.9 SE (Figure 1065 S8e); just two models fitting nominally better and 223 models fitting non-significantly worse than 1066 the published model and having similar or higher WR were found (**Table 1. Table S1**). These results 1067 suggest that while OrientAGraph was often (but not always) able to find the same tentative global 1068 likelihood optimum as findGraphs, neither 3 nor 5 admixture events are enough to explain the data 1069 since nearly all the groups are admixed.

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For this reason, we moved to topology searches in more complex model spaces incorporating 6 to 9 admixture events. Temporally plausible models with even a modest fit (WR between 3 and 4 SE) were encountered only among models with 8 and 9 admixture events (**Figure S8j-r**). In the complexity class with 8 admixture events, 5 such temporally plausible fitting models were found, with WRs ranging from 3.4 to 3.9 SE (all these models are shown in **Figure S8j-l**). In the complexity class with 9 admixture events, 11 such models were found, with WRs ranging from 3.4 to 4.0 SE (all these models are shown in **Figure S8m-r**).

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Librado *et al.* 2021 discussed the following inferences relying fully or partially on their published
 admixture graphs reported in that study (**Table 2**): 1) NEO-ANA-related admixture is absent in

1081 DOM2; 2) DOM2 and C-PONT are sister groups (they form a clade); 3) there is no gene flow

1082 connecting the CWC and the cluster associated with Yamnaya horses and horses of the later
1083 Sintashta culture whose ancestry is maximized in the Western Steppe (DOM2, C-PONT, TURG); 4)
1084 there was gene flow from a deep-branching ghost group to NEO-ANA; and 5) Tarpan is a mixture of a
1085 CWC-related and a DOM2-related lineage.

1086

The simplest temporally plausible and best-fitting (WR = 3.4 SE) model we found (modified group 1087 1088 composition, 8 admixture events, Figure 5b, Figure 58j, upper panels) supports inferences 2 and 4, 1089 and is incompatible with inferences 1, 3, and 5 (Table 2). This newly found model can be interpreted 1090 as follows. There is a trifurcation of three deep lineages: a lineage maximized in Western and Central 1091 Europe (up to 100% of ancestry in a Late Paleolithic group from France, LP\_SFR), a Western Steppe-1092 specific lineage (up to 55% in TURG), and a Tarpan-specific lineage (22% in Tarpan). Western and 1093 Central European horses, represented by LP-SFR, by the majority ancestry in horses found in the 1094 Corded Ware culture context (CWC), and by the majority ancestry in wild Neolithic Anatolian horses 1095 (NEO\_ANA), contributed about half of the ancestry in the Western Steppe groups TURG, C-PONT, 1096 and DOM2. The other half of ancestry in the Western Steppe groups is represented by the Western 1097 Steppe-specific lineage. That lineage also contributed about 50% of ancestry in wild horses from the 1098 Yana Upper Paleolithic site (ELEN), and the other half of ELEN's ancestry is derived from an even 1099 deeper lineage. The Botai group is modeled as a mixture of European horses (69%) and Siberian 1100 horses (31% ELEN-related ancestry). In contrast to Librado et al. (2021), Tarpan is modeled as a 1101 mixture of its specific lineage (22%) and a DOM2-related group (78%), and CWC also received 1102 ancestry (21%) from a DOM2-related group. All the populations included in the model except for 1103 LP SFR are admixed, and there is evidence of substantial genetic influence from a lineage that was 1104 eventually maximized in the Western Steppe (although it did not necessarily originate there) in the 1105 ELEN and Botai groups. We consider this model to be plausible from both temporal and geographical 1106 perspectives.

1107

1108 We are not arguing here that our 8-admixture-event model represents the true history; in fact, it is 1109 highly unlikely to be the truth, given how large the space of all possible admixture events is and how 1110 much admixture evidently occurred relating all these groups (which makes finding the unique truly 1111 fitting model extremely unlikely based on *f*-statistic fitting, see the results on simulated data in 1112 Figure 2b). We have also not attempted in any way to replicate the admixture graph exploration 1113 procedure performed in the Librado *et al.* study; the graph fitting procedure was quite different 1114 from ours, based on OrientAGraph optimization rather than findGraphs optimization, and a Block Jackknife procedure with a different genome block size for determining standard errors (4 Mbp in 1115 1116 our protocol and ca. 500 kbp in the Librado et al. study). Regardless of how the graph was obtained, 1117 it is valuable for providing readers with guidance about which topological features of the graphs are 1118 meaningful and stable, and which are less certain, especially—as in the case of the admixture graph 1119 presented in the paper—when some features of the presented model cannot be right, as evident by 1120 the WR of 6.9 in the published model for 5 admixture events. Our set of 16 temporally plausible and 1121 fitting (WR < 4 SE) models with 8 or 9 admixture events (Figure S8j-r) is consistent with some 1122 features of the published graph being stable: the features (2) that DOM2 and C-PONT are sister 1123 groups, and (4) that there was a gene flow from a deep-branching ghost group to NEO-ANA (Table 1124 2).

1125 1126 Equally important, however, is our finding that there are plausible models that are inconsistent with 1127 other inferences in Librado et al (Table 2). For example, 13 of these 16 models are inconsistent with 1128 the suggestion that there was no gene flow connecting the CWC and the cluster maximized in the 1129 Western steppe (DOM2, C-PONT, TURG) (Figure S8j-r). In the 8-admixture-event best-fitting 1130 plausible model (Figure 5b, Figure S8j, upper panels), CWC actually derives appreciable ancestry 1131 from the early domestic horse lineage DOM2 associated with the Sintashta culture to the exclusion 1132 of the more distant Yamnaya-associated TURG and C\_PONT horses. This scenario presents a parallel 1133 to the one observed in humans, with individuals associated with the CWC receiving admixture from 1134 Steppe pastoralists albeit in different proportions: ~75% for humans, versus ~20% in horses. These 1135 models specifying a substantial Steppe horse contribution to CWC horses would weaken support for

1136 the inference in Librado *et al.* that "Our results reject the commonly held association between

1137 horseback riding and the massive expansion of Yamnaya steppe pastoralists into Europe around

1138 3000 BC". We are not aware of other lines of evidence in the paper (apart from the fitted admixture

1139 graph) that support the claim of no Yamnaya horse impact on CWC horses.

1140

1141 Another example of a feature of the published graph that turned out to be unstable is the model for 1142 the Tarpan horse. Only 8 of 16 temporally plausible and fitting models (**Figure S8j-r**) support the 1143 conclusion by Librado *et al.* that the Tarpan is a mixture of a DOM2-related and a CWC-related 1144 lineage. The other 8 models suggest that Tarpan is a mixture of a deep lineage and a DOM2-related 1145 lineage (**Figure 5b, Figure S8j**, upper panels), echoing a hypothesis that Tarpan may be a hybrid with 1146 Przewalski's horses not represented in the admixture graph (Librado et al. 2021).

1147

1148Again, we are not arguing here that our fitting alternative model is right—indeed we are nearly1149certain it is wrong in important aspects—but we are merely pointing out that the complexity of the1150admixture graph space means that qualitatively quite different conclusions are compatible with the1151genetic data. Other aspects of the Librado *et al.* study, most notably the dramatic geographic1152expansion of the DOM2 modern domestic horse lineage after 4000 years ago in association with the1153Sintashta culture which is the most extraordinary finding of Librado *et al.*, are in no way challenged1154by our results.

1155 1156

1157 Hajdinjak et al. 2021. The admixture graph inferred by Hajdinjak et al. was constructed manually on 1158 the basis of a SNP set derived from in-solution enrichment of two SNP panels (1240K + a further 1159 million transversion polymorphisms discovered as polymorphic within one or two sub-Saharan 1160 African individuals or among archaic humans) and incorporated 11 groups and 8 admixture events 1161 (Figure 2d in the original study). The published graph has no clear outgroup since the deepest 1162 branch (Denisovan) is admixed. This property of the graph makes automated graph space 1163 exploration difficult. We explored two topology classes: 1) 11 groups with 8 admixture events, the 1164 original SNP set, Denisovan assigned as an outgroup only at the stage of generating random starting 1165 graphs (gene flows to/from the Denisovan branch were allowed at the topology optimization step); 1166 and 2) 12 groups with 8 admixture events, chimpanzee added and the original SNP set changed due 1167 to the zero missing rate condition, and chimpanzee assigned as an outgroup at both algorithm 1168 stages (Table S1). For both graph complexity classes, two topology search settings were tested: 1) either no additional constraints were applied beyond the outgroup constraints described above, or 1169 1170 2) the Vindija Neanderthal and Mbuti were allowed to have no admixture events in their history, and the Denisovan lineage was allowed to have up to one admixture event in its history (these 1171 1172 constraints were in line with the model in the original study and with literature on the genetic 1173 history of archaic humans, e.g., Prüfer et al. 2014). The composition of the groups matched that in 1174 the original study, as did the parameter settings for *qpGraph*, with the exception of "least squares 1175 mode", which was used in the original study, but not in our analysis. "Least squares mode" 1176 computes LL scores without taking into account the f-statistic covariance matrix, and we confirmed 1177 that changing this parameter does not qualitatively change our results. Since no  $f_3$ -statistics were 1178 negative when all sites available for each population triplet were used (the "useallsnps: YES" option), 1179 we did not use the upgraded algorithm for calculating  $f_3$ -statistics on pseudo-diploid data. We 1180 summarize results across 2,000 to 4,000 independent iterations of the *findGraphs* algorithm (Table 1181 **S1**). 1182

When chimpanzee was not included into the analysis and no topology constraints were applied,
nearly all newly found models turned out to be distinct (3,996 of 4,000), nearly all (96.8%) fit
nominally better and 15.9% fit significantly better than the published model (Table S1), and absolute
fits of 91.3% of novel models are good (WR < 3 SE). Similar results were obtained when the topology</li>
search algorithm was constrained: nearly all (89.5%) of 1,999 newly found models fit nominally
better and 26% fit significantly better than the published model (Table S1).

1189

1190 When chimpanzee was set as an outgroup and no topology constraints were applied, the picture 1191 remained similar. Nearly all newly found models turned out to be distinct (1,996 of 2,000), and a 1192 very large fraction of them (56.8%) fit significantly better than the published model (**Table S1**); 1193 16.4% of novel models demonstrated WR < 3 SE. Similar results were obtained when the topology 1194 search algorithm was constrained: most (71.4%) newly found models fit nominally better and 15.7% 1195 fit significantly better than the published model (Table 1, Table S1, Figure 4), which has a poor 1196 absolute fit on this set of sites and groups (WR = 4.8 SE, Figure 5c, Figure S9). The statistics described above and the fact that LL scores on all sites lie outside of the LL distribution on resampled 1197 1198 datasets (Figure 4) suggest that models in this complexity class are overfitted, but the published 1199 topology emerged as fitting relatively poorly.

1200

1201 Overfitting arises naturally during manual graph construction as performed in many studies (not 1202 only in Hajdinjak et al. (2021), but also in Fu et al. 2016, Skoglund et al. 2016, Yang et al. 2017, Posth 1203 et al. 2018, McColl et al. 2018, Moreno-Mayar et al. 2018, Tambets et al. 2018, van de Loosdrecht et 1204 al. 2018, Flegontov et al. 2019, Sikora et al. 2019, Wang et al. 2019, Lipson et al. 2020, Shinde et al. 1205 2019, Yang et al. 2020, and Wang et al. 2021). The graph grew one group at a time, and each newly 1206 added group was mapped on to the pre-existing skeleton graph as unadmixed or as a 2-way mixture. 1207 This imposed constraints on the model-building process. Another constraint imposed was the 1208 requirement that all intermediate graphs have good absolute fits (WR below 3 or 4 SE). When the 1209 model-building process is constrained in a particular path and fits of all intermediates are required 1210 to be good, unnecessary admixture events are often added along the way, and the resulting graph 1211 belongs to a complexity class in which models are overfitted and many alternative models fit equally 1212 well. There is no single obviously correct order of adding branches to a growing graph. For example, 1213 the Kostenki and Sunghir lineages were included into the initial graph (Fig. S6.1 in the original study) 1214 as unadmixed lineages, and their admixture status was not revisited at subsequent steps (unlike that 1215 of Tianyuan and Ust'-Ishim), except for adding the archaic gene flow common for non-Africans. For 1216 that reason, the published graph differs from many alternative better-fitting and temporally 1217 plausible graphs where the Kostenki and Sunghir lineages are modeled as more complex mixtures 1218 (Figure S9).

1219

1220 Hajdinjak et al. (2021)'s published graph had the following notable features that were interpreted by 1221 the authors and used to support some conclusions of the study (Table 2): 1) there are gene flows 1222 from the lineage found in the ~45,000-43,000-years-old Bacho Kiro Initial Upper Paleolithic (IUP) 1223 individuals to the Ust'-Ishim, Tianyuan, and GoyetQ116-1 lineages; 2) the ~35,000-years-old Bacho 1224 Kiro Cave individual BK1653 belonged to a population that was related, but not identical, to that of 1225 the GoyetQ116-1 individual; and 3) the Vestonice16 lineage is a mixture of a Sunghir-related and a 1226 BK1653-related lineage. To assess if these features are supported by our re-analysis, we focused on 1227 our most constrained *findGraphs* run: with chimpanzee set as an outgroup and with the topology 1228 constraints applied at the topology search step. We identified 1,421 topologies fitting nominally or 1229 significantly better than the published model and satisfying the constraints and moved on to inspect 1230 50 best-fitting topologies for temporal plausibility (all of them fitting significantly better than the 1231 published model). All non-African individuals included in the model are Upper Paleolithic and their 1232 dates are not drastically different in relative terms: from ca. 45 kya (thousand years before present) 1233 for some Bacho Kiro IUP individuals (Hajdinjak et al. 2021) to ca. 30 kya for the Vestonice16 1234 individual (Fu et al. 2016). Nevertheless, we considered most gene flows from later-attested to 1235 earlier-attested lineages as temporally implausible (for instance, GoyetQ116-1 (~35 kya)  $\rightarrow$  Ust'-1236 Ishim (~44 kya), GoyetQ116-1 (35 kya)  $\rightarrow$  Bacho Kiro IUP (45-43 kya), Kostenki14 (38 kya)  $\rightarrow$  Ust'-1237 Ishim (44 kya), Sunghir III (34.5 kya)  $\rightarrow$  Tianyuan (40 kya), Vestonice16 (30 kya)  $\rightarrow$  Tianyuan (40 kya)) 1238 since they imply great antiquity of the later-attested lineages, e.g., >40 kya for the Vestonice16 1239 lineage, and even greater antiquity for the related lineages such as Sunghir III and Kostenki14. 1240

1241 Of the 50 topologies inspected, 32 were considered temporally plausible. Of those topologies, none 1242 supported feature 1 of the published admixture graph (there is no replication of the finding of gene 1243 flows from the Bacho Kiro IUP lineage specifically into all three of the Ust'-Ishim, Tianyuan, and 1244 GoyetQ116-1 lineages). One topology supported features 2 and 3, and partially supported feature 1 1245 (there was Bacho Kiro ightarrow GoyetQ116-1 gene flow, but no Bacho Kiro ightarrow Tianyuan and Bacho Kiro ightarrow1246 Ust'-Ishim gene flows). A total of 17 topologies supported features 2 and 3 but were inconsistent with feature 1; and 14 topologies supported feature 3 only (Table 2). Best-fitting representatives of 1247 1248 each of these topology classes are shown along with the published model in Figure S9. Considering topological diversity among models that are temporally plausible, conform to current knowledge 1249 1250 about relationships between modern and archaic humans, and fit significantly better than the 1251 published model, we conclude that feature 3 is probably robust but other details of the fitted 1252 admixture graph in Hajdinjak et al. (Figure 2d of that study)—for example, gene flows to the Ust'-1253 Ishim, Tianyuan and Goyet Q116-1 lineages from sources sharing drift exclusively with the Upper 1254 Paleolithic Bacho Kiro lineage—should not be interpreted as providing meaningful inferences about 1255 population history of Upper Paleolithic modern humans. For example, the upper right-hand 1256 alternative model plotted in Figure S9c supports features 2 and 3 but includes no gene flows from the Bacho Kiro IUP lineage. 1257

1258

1259 A central finding of Hajdinjak et al. is that the Bacho Kiro IUP group shares more alleles with present-1260 day East Asians than with Upper Paleolithic Holocene Europeans despite coming from Europe. 1261 Specifically, the study documents significantly positive statistics of the form D(an Asian group, 1262 Kostenki14; Bacho Kiro IUP, Mbuti) (Fig. 2b and Extended Data Fig. 5 in the original study). For 1263 example, D(Tianyuan, Kostenki14; Bacho Kiro IUP, Mbuti) is significantly positive (D = 0.0032, SE = 1264 0.0010, Z = 3.2) on the dataset used for testing the twelve-population graphs (263,698 sites without 1265 missing data across all 12 groups). The same statistic is also significantly positive (D = 0.0029, SE = 1266 0.0006, Z = 4.4) when all 1,312,292 non-missing sites in the population quadruplet are analyzed. 1267 Hajdinjak et al.'s interpretation of this observation, using the language from the abstract, is that 1268 "there was at least some continuity between the earliest modern humans in Europe [Bacho Kiro IUP] 1269 and later people in Eurasia [East Asians]".

1270

1271 However, a significant D-statistic can have multiple explanations. The statistic  $f_4$  (Tianyuan, 1272 Kostenki14; Bacho Kiro IUP, Mbuti) is fitted equally well by the published twelve-population 1273 admixture graph (Z-score for the difference between the observed and fitted statistics = 0.64) and 1274 by, for example, the lower left-hand graph in Figure S9c (Z-score = 0.94) reproduced in Figure 5c. 1275 Under the latter model that fits the data significantly better than the published model (p-value = 1276 0.02), the Bacho Kiro IUP and Tianyuan branches are not connected by a gene flow and do not 1277 receive gene flows from a third common source, but the common ancestor of Ust'-Ishim and all 1278 European Paleolithic lineages receives an 8% gene flow from a divergent modern human lineage 1279 splitting deeper than Bacho Kiro IUP and Tianyuan (Figure 5c, Figure S9c). This scenario or some 1280 version of it seems archaeologically and geographically plausible and is not disproven by any other 1281 line of genetic or non-genetic evidence of which we are aware. It could correspond to a scenario 1282 where a primary modern human expansion out of the Near East contributed serially to the major 1283 lineages leading to Bacho Kiro, then later East Asians, then Ust'-Ishim, and finally the primary 1284 ancestry in later European hunter-gatherers. This has a very different interpretation from the 1285 scenario of distinctive shared ancestry between the earliest modern humans in Europe such as 1286 Bacho Kiro IUP and later people in East Asia—to the exclusion of later European hunter-gatherers— 1287 that is suggested by the Hajdinjak *et al.* published graph.

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We are not claiming that this specific alternative model is correct—indeed, it is almost certainly not the correct one given the topological complexity of the set of all admixture graphs consistent with the data—but the existence of it and many other models that fit the data makes it clear that we do not yet have a unique historical explanation for the excess sharing of alleles that has been documented between some Upper Paleolithic European groups (Bacho Kiro IUP, Hajdinjak et al. 2021, and GoyetQ116-1, Yang et al. 2017 and Hajdinjak et al. 2021) and all East Asians.

1295

Lipson et al. 2020. The admixture graph in the original study (Lipson et al. 2020) was constructed 1297 1298 manually based on a SNP set derived from the 1240K enrichment panel, and the final model was 1299 alternatively tested on the combined HumanOrigins sub-panels 4 and 5 (each ascertained on one 1300 African individual) or on sites ascertained as polymorphic in archaic humans. The final published 1301 model (Extended Data Fig. 4 in that study) is very complex (12 groups and 12 admixture events): it 1302 exists in a space of  $\sim 10^{44}$  topologies of this complexity. We note that one admixture event was 1303 added by Lipson et al. (2020) to account for potential modern DNA contamination in ancient Shum 1304 Laka individuals, and removing it caused a negligible difference in the fit of the published model 1305 (**Table S1**). Thus, to decrease the complexity of the graph search space, we considered graphs with 1306 12 groups and 11 admixture events. Twenty-two  $f_3$ -statistics for these 12 groups turned out to be 1307 negative (when the "useallsnps: YES" setting was used), and thus for exploring this graph complexity 1308 class we had to remove sites with only one chromosome genotyped in any non-singleton population 1309 (Table S1). The following constraints were applied during the topology search: chimpanzee was 1310 assigned as an outgroup at both stages of the process (while generating random starting graphs and 1311 while searching the topology space); Altai Neanderthal was required to be unadmixed; and non-1312 Africans (French) were required to have at least one admixture event in their history. The 1313 composition of the groups we analyzed matched that in the original study. We summarize results 1314 across 2,000 independent iterations of the *findGraphs* algorithm.

1315

1316 All newly found models turn out to be distinct (2,000), and 11.9% fit nominally (but not significantly) 1317 better than the published model (Table 1, Table S1, Figure 4). Absolute fits of 36.7% of novel models 1318 are good (WR < 3 SE). Fits of the highest-ranking model and the published model are not significantly 1319 different according to the bootstrap model comparison method (p-value = 0.176). These metrics, 1320 along with the fact that LL scores on all sites lie outside of the LL distribution on resampled datasets 1321 (Figure 4), suggest that models in this complexity class, including the published model, are 1322 overfitted. Of the admixture graphs we re-evaluate in this study, Lipson et al. 2020 shares with 1323 Hajdinjak et al. 2021, Sikora et al. 2019, and Wang et al. 2021, evidence of being overfitted (Figure 1324 4).

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1326 We also wanted to check if overfitting would be found in the graph complexity classes 1327 corresponding to two simpler intermediate graphs from the original study (Table S1): 7 groups and 4 1328 admixture events (Figure S3.24 in that study) and 10 groups and 8 admixture events (Figure S3.25 in 1329 that study). The population composition of the dataset we used for this analysis was slightly 1330 different from the dataset used by Lipson *et al.*: the ancient South African hunter-gatherer group 1331 was replaced by a related group (present-day Ju22 hoan North), and instead of the Shum Laka 1332 ancient group, only one high-coverage individual from the same group (I10871) was used. We 1333 summarize results across 2,000 or 10,000 independent iterations for each SNP set, for the small and 1334 large graphs, respectively. For 7 groups, we found 201 novel topologies fitting better than the 1335 published one, and for 10 groups we found nearly 9,000 such topologies (Table S1). In the latter case 1336 6.8% of newly found topologies fit significantly better than the published topology. For the more 1337 complex graph class with 10 groups and 8 admixture events we also found evidence of overfitting: 1338 the LL score of the published graph run on the full data is better than almost all the bootstrap replicates on the same data (it falls below the 5<sup>th</sup> percentile). 1339

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Below we discuss selected prominent features of the admixture graph published in the original study 1341 1342 (that were interpreted by the authors and used to support some conclusions of the study) and the 1343 extent to which these features consistently replicate across the large number of fitting 12-1344 population graphs with 11 admixture events (Table 2): 1) A lineage maximized in present-day West 1345 African groups (Lemande, Mende, and Yoruba) also contributed some ancestry to the ancient Shum 1346 Laka individual and to present-day Biaka and Mbuti; 2) another ancestry component in Shum Laka is 1347 a deep-branching lineage maximized in the rainforest hunter-gatherers Biaka and Mbuti; 3) "super-1348 archaic" ancestry (i.e., diverging at the modern human/Neanderthal split point or deeper) 1349 contributed to Biaka, Mbuti, Shum Laka, Lemande, Mende, and Yoruba; and 4) a ghost modern 1350 human lineage (or lineages) contributed to Agaw, Mota, Biaka, Mbuti, Shum Laka, Lemande, Mende,

and Yoruba. We identified 232 twelve-population topologies that fit nominally better than the
published one, 34 best-fitting topologies (of 232) were manually assessed for temporal plausibility,
and we focus on 30 topologies identified as temporally plausible and including a low-level
Neanderthal contribution (≤10%) in non-Africans (French). These 30 topologies are shown along with
the published model in Figure S10.

1356

1357 In this set of alternative models, high topological diversity is observed (see an example in Figure 5d 1358 and further topologies in Figure S10). We classified the topologies as follows. If an ancestral lineage 1359 defined above (for example, a deep-branching lineage maximized in rainforest hunter-gatherers 1360 Biaka and Mbuti) exists in the graph, we compared the sets of populations where it is found in the 1361 published model and in the model examined. If there was no more than one population where the 1362 ancestry is expected according to the published model but not present, or present but not expected, 1363 we considered this feature of the published graph supported by the alternative graph. If no ancestral 1364 lineage meets the definition above, the feature of the published graph was considered not 1365 supported. In all other cases partial support for the feature was declared (Figure S10). Considering 1366 extreme cases, two alternative graphs completely lacked support for three features of the published 1367 graph (Figure 5d, Figure S10c), and one graph supported all four features of the published graph 1368 fully (Figure S10q, bottom panels). There are some graphs where defining two distinct ancestral 1369 lineages maximized in West Africans and in Mbuti and Biaka (features 1 and 2) is essentially 1370 impossible since all or nearly all Africans are modeled as a mixture of at least two deep lineages (see 1371 graph no. 4, Figure S10d). In some graphs there is no single lineage specific to rainforest hunter-1372 gatherers (Biaka, Mbuti, and Shum Laka) since the primary ancestries in these groups form 1373 independent deep branches in the African graph (see graph no. 2 in Figure 5d and graph no. 16 in 1374 Figure S10j, bottom panels). The ghost modern and super-archaic gene flows to Africans also had no 1375 universal support in the set of alternative graphs we examined (see, for example, Figure 5d and 1376 Figure S10c).

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1378 Considering the high degree of topological diversity among models that are temporally plausible, 1379 conform to known findings about relationships between modern and archaic humans, and fit 1380 nominally better than the published model, we conclude that the features from the original study 1381 are not supported by our re-analysis (Table 2). As in the case study above, the published manually 1382 constructed model is a representative of a large class of models that are equally well fitting to the 1383 limits of our resolution. This situation may be attributed to 1) overfitting and/or to 2) the lack of 1384 information in the dataset (in the combination of groups and SNP sites) and/or to 3) inherent 1385 limitations of *f*-statistics, when distinct topologies predict identical *f*-statistics.

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1387 In reconsidering the findings of Lipson et al. 2020 it is important to keep in mind that analysis of 1388 allele frequency correlation statistics is not the only type of information that can be used to make 1389 inferences about population relationships in deep time. Other methodologies have provided 1390 important insights into deep African population history, and the model-building in Lipson et al. 2020 1391 was guided in an informal way by these other lines of evidence. For example, unknown archaic 1392 lineages admixing into some African populations were hypothesized through identification of deeply 1393 splitting haplotypes that are too long to have been freely mixing with other haplotypes in present-1394 day populations for all of their history (Hammer et al. 2011, Lachance et al. 2012, Speidel et al. 1395 2019). Similarly, analysis of haplotype divergence times of pairs of populations has been used to 1396 provide evidence of an early radiation of modern human lineages maximized today in southern 1397 African hunter-gatherers, Mbuti rainforest hunter-gatherers, and the great majority of other 1398 present-day populations; and a later split of lineages related to East African hunter-gatherers, West 1399 African agriculturalists, and non-Africans, which is a feature of the Lipson *et al.* model (Campbell and 1400 Tishkoff 2008, Mallick et al. 2016). Notably, some alternative models we found do not contradict the 1401 above-mentioned results and are profoundly different from the published model at the same time 1402 (see, for example, Figure 5d). These constraints are not enough, however, to provide evidence for all 1403 the topological details of the Lipson *et al.* 2020 admixture graph highlighted in this section, or for 1404 other features of Lipson et al. 2020 that were not invoked in the previous literature and newly

proposed in that study, such as the "ghost modern" lineage splitting around the same time as the
lineages leading to southern African hunter-gatherers and central African rainforest huntergatherers and mixing in highest proportion to Ethiopian hunter gatherers and to a lesser proportion
to West Africans, and the "basal West African" lineage that contributes uniquely to Shum Laka.
Many of the models that emerged as good fits in our admixture graph building exercise as the
published one did not share some of these features (Figure S10).

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1412 The high diversity of well-fitting admixture graph models that satisfy known constraints relating 1413 diverse African populations highlights the need for further research based on multiple lines of 1414 genetic analysis (in addition to allele frequency correlation patterns) to obtain further insights into 1415 deep African history. Our results particularly highlight the mystery around the highly distinctive 1416 genetic ancestry of the Shum Laka individuals themselves, who represent the newly reported data in 1417 the Lipson et al. 2020 study, and a highly important set of genetic datapoints that was not available 1418 prior to the study. The ancestral relationships of these four individuals to both rainforest hunter-1419 gatherers, and to the primary lineage in present-day West Africans, remains an open question, one 1420 whose resolution promises meaningful new insights into modern human population history.

1421 1422

1423 Wang et al. 2021. The admixture graph inferred by Wang et al. 2021 was constructed manually on 1424 the basis of a SNP set derived from the 1240K enrichment panel. We focused our analysis on the 1425 final graph (Extended Data Figure 6 in Wang et al. 2021, 12 groups and 8 admixture events) and on 1426 two simpler intermediates in the model building process (Figures SI3-9 and SI3-10a in Wang et al. 1427 2021). To simplify the latter two models further, we removed a low-level gene flow (1%) from a 1428 West European hunter-gatherer-related lineage (Loschbour) to the Mongolia Neolithic group, which 1429 resulted in negligible LL differences (0.5 and 2.4 log-units, respectively). Thus, using findGraphs we 1430 explored the following topology classes: 9 groups with 4 admixture events, 10 groups with 5 1431 admixture events, and 12 groups with 8 admixture events (Table S1). The composition of the groups 1432 matched that in the original study. We summarized results across 2,000 independent iterations of 1433 the *findGraphs* algorithm for each topology class. In the case of the most extensive population set 1434 (12 groups), three  $f_3$ -statistics turned out to be negative (when the "useallsnps: YES" setting was 1435 used), and thus for exploring this graph complexity class we had to remove sites with only one 1436 chromosome genotyped in any non-singleton population (**Table S1**). For this complexity class, we 1437 also applied several constraints on the graph space exploration process all of which were shared 1438 with the Wang et al. graphs: the Denisovan genome was assigned as an outgroup in the random 1439 starting graphs, but not at the topology search stage; up to one admixture event was allowed in the 1440 history of the Denisovan group; no admixture events were allowed in the history of Mbuti, 1441 Loschbour, and Onge; and the (Denisovan, (Mbuti, (Loschbour, Onge)))) branching order was 1442 required.

1443

1444 For each topology class we found hundreds to thousands of topologically unique graphs fitting 1445 nominally better than the published models (Table 1, Table S1). For both simple topology classes, no 1446 model fitting significantly better than the published one was found (Table S1). However, the final 1447 published model fits the data significantly worse than 12.6% of newly found models of the same 1448 complexity (Table 1, Table S1). The fact that many topologically diverse models had good absolute 1449 fits (65%, 55%, and 15% of distinct newly found graphs with 9, 10, and 12 groups, respectively, had 1450 WR < 3 SE) suggests that admixture graph models in these complexity classes are overfitted. Further 1451 evidence of overfitting comes from the poor fits of the published model on bootstrap-resampled 1452 datasets as compared to their fits on all sites (Figure 4).

1453

An important feature of the published graphs in Wang *et al.* 2021 that was remarked upon in the
study is admixture from a source related to Andamanese hunter-gatherers that is almost universal in
East Asians, occurring in the Jomon, Tibetan, Upper Yellow River Late Neolithic, West Liao River Late
Neolithic, Taiwan Iron Age, and China Island Early Neolithic (Liangdao) groups (**Table 2**). For
example, the abstract states "Hunter-gatherers from Japan, the Amur River Basin, and people of

Neolithic and Iron Age Taiwan and the Tibetan Plateau are linked by a deeply splitting lineage that 1459 1460 probably reflects a coastal migration during the Late Pleistocene epoch." We performed 2,000 1461 findGraphs iterations and obtained 1,778 distinct topologies satisfying all the constraints, nearly all 1462 of them (1,724) fitting nominally better than the published model, and 12.6% fitting significantly better (Table S1). The models were ranked by LL, and 56 highest-ranking topologies, all of them 1463 1464 fitting significantly better than the published one, were assessed for temporal plausibility (models 1465 with gene flows from a later group to Tianyuan dated to 40 kya were removed), and 20 topologies 1466 were considered temporally plausible (all of them are shown in **Figure S11**). According to these 1467 topologies, 0 to 2 East Asian groups had a fraction of their ancestry derived from a source 1468 specifically related to Onge, and 19 topologies included gene flows from the European (Loschbour)-1469 related branch to all 8 East Asian groups (Figure S11). The inferred topological relationships among 1470 East Asians are variable in this group of 20 models, and we decided to apply further constraints that 1471 guided model ranking and elimination by Wang et al., based on considerations from archaeological 1472 evidence, Y chromosome haplogroup divergence patterns, and population split time estimation. 1473

1474 The constraints that are not based on correlation of allele frequencies across populations that Wang 1475 et al. applied and that we applied in our re-examination are as follows. First, combined evidence 1476 from archaeology, linguistics, and genetics (a closely shared Y chromosome haplogroup) suggests 1477 that the present-day Tibetan Plateau population harbors a substantial proportion of ancestry from a 1478 large-scale migration from the Neolithic farming groups from the Upper and Middle Yellow River 1479 (Chen et al. 2015, Lu et al. 2016, Zhang et al. 2019). These arguments and radiocarbon dates favor 1480 the following branching order of predominant ancestry components: (Mongolia East N, (China Upper 1481 YR LN, Nepal Chokhopani)). Second, evidence from archaeology, linguistics and genetics suggests 1482 that the expansion of Austronesian speakers and the peopling of Taiwan was from southeast coastal 1483 China to Taiwan and Southeast Asia, but not from Taiwan to mainland China (Bellwood 2011, Gray 1484 and Jordan 2000, Ko et al. 2011). These arguments make a China Island EN  $\rightarrow$  Taiwan IA gene flow 1485 direction plausible and make the opposite direction of flow less likely. Third, in the original study 1486 MSMC cross-coalescence rates were computed for a few pairs of present-day proxies for the ancient 1487 groups, and it was argued that they impose constraints on the graph topology. The inferred 1488 coalescence date for the Tibetan and Ulchi groups was slightly younger than the Tibetan-Ami and 1489 Tibetan-Atayal dates (Fig. SI3-1 in the original study), suggesting that the Nepal Chokhopani and 1490 Mongolia East N group may share ancestral source populations more recently than these two groups 1491 and Taiwan IA. We note that it was not clear in the original paper if the difference in coalescence 1492 dates is statistically significant, the finding was clearer in MSMC than in MSMC2 analysis, and there 1493 was no attempt to calculate expected cross-coalescence profiles using these methods from models 1494 incorporating many gene flows. Nevertheless, we applied this constraint as well in an attempt to 1495 understand whether, if we used a constraint system similar to that in Wang et al., we would obtain 1496 results that agreed with respect to the finding of Onge-related admixture ubiquitous among East 1497 Asian groups.

1498

1499 Applying these three additional constraints, we identified two models (among the 56 ones subjected 1500 to manual inspection) that satisfied all of them. The highest-ranking of those models is shown in 1501 Figure 5e and Figure S11c (lower panels), and it includes a 13% (deeply) European-related gene flow 1502 to the common ancestor of all East Asians, and gene flows from the Onge-related branch to just two 1503 East Asian groups: Nepal Chokhopani and China WLR LN. This model fits the data significantly better 1504 than the published model (p-value = 0.028). We do not claim that this is the correct model (indeed 1505 we are almost certain that it is not given the high degeneracy of fitting models), but it is not 1506 obviously wrong and differs in qualitatively important ways from the published one.

1507

1508The Wang *et al.* 2021 admixture graph provides an illuminating example that helps us to understand1509the value added by admixture graph construction. The admixture graph construction process in1510Wang *et al.* followed a philosophy of not relying entirely on the allele frequency correlation data1511(not treating the genetic data as independent to explore how much new insight could come from1512genetic data alone). Instead, the study integrated other lines of genetic evidence as well as linguistic

and archaeological insights explicitly into the admixture graph construction process, with the goal of 1513 1514 identifying models consistent with multiple lines of evidence. The fact that after this procedure a 1515 fitting graph was obtained is not of great interest, as it is essentially always possible to obtain a fit to 1516 allele frequency correlation data when enough admixture events are added. The important question 1517 is whether any of the emergent features of the graph that were not applied as constraints in the 1518 construction process—for example the evidence of ubiquitous Andamanese-related gene flow 1519 throughout East Asia suggesting a coastal route expansion that admixed with an interior route 1520 expansion proxied by Tianyuan—were stably inferred. Our analysis does not come to this finding 1521 consistently among well-fitting and plausible admixture graphs. We conclude that an important 1522 feature of the published graph, i.e. variable levels of Andamanese-related ancestry found in all East 1523 Asians except for Siberians (Mongolia Neolithic) and the Upper Paleolithic Tianyuan (Fig. 2 in (Wang 1524 et al. 2021)), is not supported by f-statistic analysis alone (**Table 2**), and indeed we are not aware of 1525 a single feature of the Wang et al. 2021 admixture graph that is stably inferred beyond the 1526 constraints applied to build it.

1527

1528

1529 Sikora et al. 2019. Two admixture graphs inferred by Sikora et al. (2019) were constructed manually 1530 based on a SNP set derived from whole-genome shotgun data and incorporated 12 or 13 groups and 10 admixture events (Extended Data Figure 3f in the original study). One graph was focused on West 1531 1532 Eurasians, and the other one on East Eurasians, and both included a Neanderthal, a Denisovan, and 1533 an African group (Dinka). Although the chimpanzee outgroup was not included in the original graphs, 1534 we added it as it drastically constrains the topology search space. The following additional 1535 constraints were applied at the *findGraphs* model optimization stage: the Neanderthal and African 1536 groups were unadmixed and the Denisovan group had no more than one admixture event it its 1537 history. These three constraints match the features of the published graph. We also repeated 1538 topology searches without constraining the admixture status of the Neanderthal, Denisovan, and 1539 Dinka. Since no  $f_3$ -statistics were negative when all sites available for each population triplet were 1540 used (the "useallsnps: YES" option), we did not apply the algorithm that allows unbiased calculation 1541 of  $f_3$ -statistics on pseudo-diploid data at the expense of loss of analyzed SNPs.

1542

1543 In contrast to most other published graphs discussed above, gene flows in the graphs inferred by 1544 Sikora et al. do not have equal standing: four low-level gene flows (0-1%) connect the Neanderthal 1545 lineage to Upper Paleolithic lineages (Kostenki, Sunghir, Yana, Ust'-Ishim in the 'Western' graph and Sunghir, Yana, Mal'ta, Ust'-Ishim in the 'Eastern' graph). We repeated each topology search under 1546 1547 two alternative settings: either keeping the number of admixture events at 10 to match the 1548 published graphs, or at 6 to match simplified versions of the published graphs lacking these low-level 1549 Neanderthal gene flows. We performed that modification to simplify the search space and to 1550 alleviate the overfitting problem which becomes severe if 10 gene flows across the graph are 1551 allowed (Table S1). Here we compare LL and WR for the original published models and their 1552 simplified versions: the Western graph including chimpanzee (LL = 65.7, WR = 3.32 SE) vs. its simplified version (LL = 76.5, WR = 3.78 SE) and the Eastern graph including chimpanzee (LL = 85.3, 1553 1554 WR = 3.11 SE) vs. its simplified version (LL = 102.4, WR = 4.16 SE). In both cases we found no 1555 statistically significant differences in model fits (relying on the bootstrap model comparison 1556 method). In summary, topology search was repeated under 8 settings: for the Western or Eastern 1557 graphs, with no constrains on the admixture status or with the constraints specified above, and with 1558 10 or 6 gene flows (Table S1). Below we focus on results for constrained models with 6 admixture 1559 events. In contrast, Figure 4 and Table 1 show results for constrained Western graphs with 10 1560 admixture events.

1561

1562 In the case of the constrained Western graphs with 6 admixture events, 1,000 topology search
1563 iterations were performed, 894 distinct topologies were found, 4 models fit significantly better, and
1564 151 models fit nominally better than the published one (Table 1, Table S1). We inspected those 155
1565 topologies and identified 29 topologies (Figure S12) that are temporally plausible and include no
1566 non-canonical gene flows from archaic groups such as Denisovan or gene flows ghost archaic →

non-Africans. Sikora et al. came to the following striking conclusion relying on the Western 1567 1568 admixture graph (Table 2): the Mal'ta (MA1 ANE) lineage received a gene flow from the Caucasus 1569 hunter-gatherer (CaucasusHG LP or CHG) lineage. However, in our *findGraphs* exploration this 1570 direction of gene flow (CHG  $\rightarrow$  Mal'ta) was supported by two of the 29 topologies, and the opposite gene flow direction (from the Mal'ta and East European hunter-gatherer clade to CHG) was 1571 1572 supported by the remaining 27 plausible topologies (Figure S12). The highest-ranking plausible topology (Figure 5f) has a fit that is not significantly different from that of the simplified published 1573 1574 model (*p*-value = 0.392). We note that the gene flow direction contradicting the graph by Sikora *et* 1575 al. was supported by a published *qpAdm* analyses (Lazaridis et al. 2016, Narasimhan et al. 2019), and 1576 *qpAdm* is not affected by the same model degeneracy issues that are the focus of this study. 1577 Considering the topological diversity among models that are temporally plausible, conform to robust 1578 findings about relationships between modern and archaic humans, and fit nominally better than the 1579 published model, we conclude that the direction of the Mal'ta-CHG gene flow cannot be resolved by 1580 admixture graph analysis (Table 2).

1581

Some important conclusions based on the Eastern graph also do not replicate across all plausible 1582 1583 admixture graphs (Table 2). In the case of the constrained Eastern graphs with 6 admixture events, 1584 4,446 topology search iterations were performed, and 2,785 distinct topologies were found. Only 3 1585 topologies fit significantly and 13 nominally better than the published one (p-value for the highest-1586 ranking newly found model vs. the simplified published model = 0.112), and 9.8% of topologies fit 1587 not significantly worse than the published one (Table 1, Table S1). Of the topologies belonging to 1588 these groups, we inspected 116 best-fitting ones and identified 97 topologies that are temporally 1589 plausible and include no gene flows from archaic groups such as Denisovan or ghost archaic  $\rightarrow$  non-1590 Africans that are qualitatively different from the gene flows that are currently widely accepted. The 1591 Sikora et al. Eastern admixture graph had the following distinctive features that were used to 1592 support some conclusions of the study (Table 2): 1) the Mal'ta (MA1 ANE) and Yana (Yana UP) 1593 lineages receive a gene flow from a common East Asian-associated source diverging before the ones 1594 contributing to the Devil's Cave (DevilsCave N), Kolyma (Kolyma M), USR1 (Alaska LP), and Clovis 1595 (Clovis LP) lineages; 2) European-related ancestry in the Kolyma, USR1, and Clovis lineages is closer 1596 to Mal'ta than to Yana; 3) the Devil's Cave lineage received no European-related gene flows, and 1597 Kolyma has less European-related ancestry than ancient Americans (USR1 and Clovis). Only feature 2 1598 was universally supported by all the 97 plausible alternative models fitting significantly better, 1599 nominally better, or not significantly worse than the simplified published model, while feature 3 was 1600 supported by 83 of 97 plausible models, and feature 1 was supported by 28 of 97 plausible models 1601 (Table 2). We plotted 14 plausible graphs as examples of topologies supporting all three features, 1602 two features, or one feature of the published graph (Figure S13). We note that all the Eastern graphs 1603 discussed here, both the published and alternative ones, have relatively poor absolute fits with WR 1604 above 4 or 5 SE. Increasing the number of gene flows to 10 allowed us to reach much better 1605 absolute fits (with WR as low as 2.42 SE), but that resulted in high topological diversity (on a par with 1606 some other case studies discussed above). In the case of the constrained Eastern graphs with 10 1607 admixture events, 1,000 topology search iterations were performed, and 1,000 distinct topologies 1608 were found. Of these topologies, 13.2% fit significantly better, 30% nominally better, and 17.6% 1609 non-significantly worse than the published model (p-value for the highest-ranking newly found 1610 model vs. the published model < 0.002) (Table S1).

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- 1613 1614

#### A Proposed Protocol for Using Admixture Graph Fitting in Genetic Studies

1615Admixture graphs represent a conceptually powerful framework for thinking about demographic1616history, but the practice of manually constructing a small number of complex models without1617exploring admixture graph space in an automated way can lead to overconfidence in the validity of1618these models. An ideal outcome of an admixture graph model exploration exercise would be the1619identification of a model or a group of topologically very similar models which fits the data well and1620significantly better than all alternative models with the same number of admixture events; however,

this is almost never achieved for graphs with more than eight populations and three admixture 1621 1622 events in our experience, and even this approach can lead to potentially unstable results as relaxing 1623 the assumption of parsimony (that fewer admixture events is more likely) can lead to qualitatively 1624 quite different equally well fitting topologies as in our reanalysis of the Bergström et al. and Shinde 1625 et al. datasets. Most of the examples of admixture graphs in eight recently published studies we 1626 revisited do not fit this ideal pattern, as we were able to identify many topologically different 1627 alternative models that could not easily be rejected based on temporal plausibility or other 1628 constraints (Figures S5-S13). In particular, for all studies except Shinde et al. 2019 (under a strict 1629 parsimony assumption however), we identified admixture graphs that were not significantly worse 1630 fitting than the published ones, and with topological features that were different in qualitatively 1631 important ways. There were also some more encouraging findings of the exercise we performed to 1632 re-evaluate published models. For example, at least one of the key inferences about population 1633 history relying on the admixture graph modeling were stable for all analyzed models for the Lazaridis 1634 et al., Librado et al., Hajdinjak et al., Shinde et al. 2019 (under the parsimony assumption), and 1635 Sikora et al. (simplified Eastern graph) studies. The existence of some stable features in these graphs 1636 helps to point the way toward a protocol that we believe should be applied in all future studies that 1637 use admixture graph fitting exercises to support claims about population history.

1638

1639 We propose the following tentative protocol to identify features of fitting admixture graphs that are 1640 stable enough to be used to make inferences about population history.

1641

1642 1. For a given combination of populations, carry out an initial scan using *findGraphs* to identify 1643 reasonable parameter values for the number of allowed admixture events (the graph complexity 1644 class). For example, run *findGraphs* allowing between zero and eight admixture events (100 1645 algorithm iterations per graph complexity class), each iteration saving one or a few best-fitting 1646 outcomes. The smallest number of admixture events that yields models where the (negative) LL 1647 score or the worst *f*-statistic residual is lower than some threshold can then be explored more 1648 deeply by running more iterations of *findGraphs*.

1649

1650 2. Run findGraphs on the determined complexity class, where some of the resulting graphs should 1651 be inspected manually to determine whether they could in principle be historically plausible models. 1652 Implausible models (for example, models where a very ancient population appears to be admixed 1653 between two modern populations) can be filtered out by imposing topological constraints. If no or 1654 only a few graphs remain, *findGraphs* can be run again under these constraints. This can be repeated 1655 until one or more graphs with an acceptable LL score or worst residual has been identified. At this 1656 stage we apply the bootstrap method to determine whether the best-fitting graph is significantly 1657 better than the next best-fitting graph. If it is not, we identify a set of graphs which are not clearly 1658 worse than the best-fitting graph by performing the bootstrap model comparison for many model 1659 pairs.

1660

16613. Researchers should compare the resulting graphs to each other with the goal of identifying1662common features. Although ADMIXTOOLS 2 includes automated tools for cataloguing common1663topological features (Suppl. Methods), we found a manual approach to be valuable, as the fitted1664parameters (especially admixture proportions) are as important for this task as graph topology.

1665

1666 4. Once a set of fitting graphs and stable topological features shared between them is identified, 1667 researchers should carry out a *findGraphs* exploration of the space of graphs with one additional 1668 admixture event. If inferences are stable even when fitting graphs with one more level of complexity 1669 than the graphs with the minimal number of admixture events needed to fit the data, this increases 1670 confidence in the inferences. Furthermore, the addition of a new population may introduce crucial 1671 information to an existing set of populations, which can change the space of fitting topologies in a 1672 profound way, as in our reanalysis of the data from Bergström et al. 2020 (Figure 5a, Figure S2). 1673 Thus, it is advisable that the topology optimization procedure is repeated on several alternative

population sets, in addition to considering models that allow an additional admixture event beyond
 the minimum required for parsimony, to explore if inferences about topology change qualitatively.

1677 5. Admixture graphs fitted with f-statistics do not distinguish between time and population size as 1678 the two sources of genetic drift, and many different complex genetic histories for a set of 1679 populations can result in the exact same expected *f*-statistics. This provides an important 1680 opportunity to further constrain a model fitting procedure. Methods that take advantage of 1681 information from the site frequency spectra (momi2, fastsimcoal, Kamm et al. 2019, Excoffier et al. 1682 2013) or derived site patterns, a special case of site frequency spectra (Legofit, Rogers 2019), can 1683 supply alternative information not captured by *f*-statistics (further information can come from 1684 methods that fit haplotype divergence patterns such as MSMC (Schiffels and Durbin 2014) and 1685 SMC++ (Terhorst et al. 2017), or inferences based on fitted gene trees such as RELATE (Speidel et al. 1686 2019), and ARGweaver (Hubisz et al. 2020, Hubisz and Siepel 2020)). These tools are too 1687 computationally intensive to explore a large number of models, but the advantages of the different 1688 approaches can be combined by first identifying a set of candidate models using *findGraphs*, and 1689 then testing these candidate models with other methods. This approach is also expected to deal 1690 with overfitting since different data types almost always include different variable site sets.

1691
1692 We believe that researchers should only begin to make strong claims about population history with
admixture graphs once a protocol such as we propose is applied.

1694

1695 We see the guidelines above as analogous in spirit to the protocols that were introduced in medical 1696 genetics at a time when a reproducibility crisis was found in the field of candidate gene association 1697 studies. Many studies looking for risk factors for common complex diseases resulted in publications 1698 with marginally significant p-values without correcting for the multiple hypothesis testing that was implicitly performed due to many candidate genes being tested and only those with significant 1699 1700 findings being published. Unsurprisingly, most of these claims failed to replicate in follow-up studies 1701 in independent sets of samples (Ioannidis 2005, Border et al. 2019, Collins et al. 2012, Duncan et al. 1702 2019). The human medical genetic community addressed this challenge by coming together to 1703 support a rigorous set of commonly accepted standards for declaring genome-wide statistical 1704 significance, such as the requirement that p-values be corrected for the effective number of 1705 independent common variants in the genome and requiring correcting for the known confounders 1706 of population structure and undocumented relatedness among individuals (Hirschhorn and Daly 1707 2005).

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1709

#### 1710 <u>Conclusion</u>

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1712 Sampling admixture graph space is a useful method for modeling population histories, but finding 1713 robust and accurate models can be challenging. As we demonstrated by revisiting a handful of 1714 published admixture graphs and re-analyzing the same datasets used to fit them, f-statistic and, 1715 more generally, allele frequency data alone are usually insufficient for building accurate graph 1716 models, making it necessary to incorporate other sources of evidence. This provides a challenge to 1717 previous approaches for automated model building. We investigated several published admixture 1718 graph models and, in nearly all cases, found many alternative models, some of which are historically 1719 and geographically plausible but contradict conclusions that were derived from the published 1720 models. To conduct these analyses, we developed a method for automated admixture graph 1721 topology optimization which can incorporate external sources of information as topological 1722 constraints. This method is developed in a framework called ADMIXTOOLS 2, which aside from 1723 admixture graph modeling, implements many other methods for population history inference based 1724 on *f*-statistics. In the process of revisiting published admixture graphs we found a well-fitting model 1725 for the history of dogs that is substantially different from the published model (Bergström et al. 1726 2020) and is strikingly congruent with the known history of relevant human lineages. We also found 1727 a novel admixture graph for domestic and wild ancient horses (Librado et al. 2021) that is

1728 substantially different from the published model, fits the data significantly better, and is 1729 geographically and historically plausible. These alternative graphs, however, have some of the same 1730 challenges as the published ones: they are almost certainly oversimplified relative to the true 1731 histories, and they exist in a large space of admixture graphs with meaningful topological differences 1732 that fit the allele frequency correlation data equally well. An important topic for future work should 1733 be to test these new alternative models as well as the previously published models as hypotheses 1734 with newly reported ancient samples and additional lines of genetic, archaeological, and other forms 1735 of analysis to obtain further clarity about population history.

1736

1737 It is important to recognize that the key concern we have highlighted in this study—the fact that 1738 there can often be multiple different topologies that are equally good fits to the allele frequency 1739 correlation patterns relating a set of populations—does not invalidate the use of allele frequency 1740 correlation testing in many other contexts in which it has been applied to make inferences about 1741 population history. For example, negative  $f_{\beta}$ -statistics ("admixture"  $f_{\beta}$ -statistics) continue to provide unambiguous evidence for a history of mixture in tested populations, and  $f_4$  and D symmetry 1742 1743 statistics remain a powerful way of evaluating whether a tested pair of populations is consistent 1744 with descending from a common ancestral population since separation from the ancestors of two 1745 groups used for comparison. The *qpWave* methodology remains a fully valid generalization of f<sub>4</sub>-1746 statistics, making it possible to test whether a set of populations is consistent with descending from 1747 a specified number of ancestral populations (which separated at earlier times from a comparison set 1748 of populations). In addition, the *qpAdm* extension of *qpWave*—which allows for estimating 1749 proportions of mixtures for the tested population under the assumption that we have data from the 1750 source populations for the mixture—remains a valid approach, unaffected by the concerns identified 1751 here. Instead of relying on a specific model of deep population relationships, *qpAdm* relies on an 1752 empirically measured covariance matrix of  $f_4$ -statistics for the analyzed populations, which is highly 1753 constraining with respect to estimation of mixture proportions but can be consistent with a wide 1754 range of deep history models. All these methods are implemented in ADMIXTOOLS 2.

1755

Finally, approaches that use admixture graphs to adjust for the covariance structure relating a set of 1756 1757 populations without insisting that the particular admixture graph model that is proposed is true with 1758 can be useful, for example for the purpose of analyzing shared genetic drift patterns of a group of 1759 populations that derive from similar mixtures. One example was a study that attempted to test for 1760 different source populations for Neolithic migrations into the Balkans after controlling for different 1761 proportions of hunter-gatherer admixture (Mathieson et al. 2018). Another example was a study 1762 that attempted to study shared ancestry between different East African forager populations after 1763 controlling for different proportions of deeply divergent source populations (Lipson et al. 2022). 1764 However, with respect to the inferences about deep history produced by admixture graphs 1765 themselves, our results highlight the importance of caution in proposing specific models of 1766 population history that relate a set of groups.

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#### 1769 <u>Methods</u>

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### 1771 Technical presentation of *ADMIXTOOLS 2* in the context of *f*-statistic modeling methods

1773 Much of the content that follows recapitulates theory presented in previous work, notably Reich *et* 1774 *al.* 2009, Green *et al.* 2010 and Patterson *et al.* 2012, but we summarize it here for coherence.

- 1775
- 1776 *f-statistics*
- 1777

1778 All *ADMIXTOOLS* programs are based on the statistics  $f_2$ ,  $f_3$ , and  $f_4$ , for population pairs, triplets, 1779 and quadruples, respectively.

1780

 $f_2$  quantifies the genetic drift separating two populations A and B. For a single SNP, it is given by 1781  $f_2(A, B) = \frac{1}{M} \sum_j (a_j - b_j)^2$ , where  $a_j$  and  $b_j$  are the allele frequencies for SNP j in populations A 1782 and B. When allele frequencies are estimated using a small number of samples, this estimator of  $f_2$ 1783 will be biased upwards. An unbiased estimator of  $f_2$  is given by  $f_2 = \frac{1}{M} \sum_j (a_j - b_j)^2 - \frac{a_j(1-a_j)}{n_{A,j-1}} - \frac{b_j(1-b_j)}{n_{B,j-1}}$ , where  $n_{A,j}$  and  $n_{B,j}$  are the observed allele counts in 1784 1785 populations A and B. 1786

1787

 $f_3(A; B, C) = \frac{1}{M} \sum_j (a_j - b_j)(a_j - c_j)$  is the covariance of the allele frequency differences between 1788 populations A and B, and the allele frequency differences between populations A and C (assuming 1789 1790 that alleles are coded randomly, so that a - b and a - c are both 0 in expectation). Significantly negative values of  $f_3(A; B, C)$  suggest that A is a mixture of sources related to B and C (although the 1791 1792 converse does not hold: A might be admixed between B and C even if  $f_3$  is positive).

1793

 $f_4(A, B; C, D) = \frac{1}{M} \sum_j (a_j - b_j)(c_j - d_j)$  is the covariance of the allele frequency differences 1794 between A and B, and the allele frequency differences between C and D. Significantly positive 1795 1796 values of  $f_4(A, B; C, D)$  (or equivalently significantly negative values of  $f_4(A, B; D, C)$ ) reveal that A 1797 and B do not form a clade with respect to C and D, and that some of the drift separating A from C is 1798 shared with the drift separating B from D. 1799

1800  $f_3$  and  $f_4$  can be written as linear combinations of  $f_2$  statistics:

1801 
$$f_3(A; B, C) = \frac{1}{2}(f_2(A, B) + f_2(A, C) - f_2(B, C))$$

1802

 $f_4(A, B; C, D) = \frac{1}{2}(f_2(A, D) + f_2(B, C) - f_2(A, C) - f_2(B, D)) \quad (Eq. 2)$ This implies that all  $f_3$ - and  $f_4$ -statistics can be computed from  $f_2$ -statistics as long as they are 1803 defined on the same SNPs. 1804

(Eq. 1)

1805

For revisiting published studies, we used the "extract  $f^{2}$ " function with the "maxmiss" argument set 1806 1807 at 0, which corresponds to the "useallsnps: NO" setting in classic ADMIXTOOLS. It means that no 1808 missing data are allowed (at the level of populations) in the specified set of populations for which pairwise f<sub>2</sub>-statistics are calculated. For the values of the "blgsize", "adjust\_pseudohaploid", and 1809 1810 "minac2" arguments we use in our analyses, see Table S1. The "blgsize" argument sets the SNP 1811 block size in Morgans, and we used either the default value of 0.05 (5 cM), or 4,000,000 bp when a genetic map was not available. Genotypes of pseudo-haploid samples are usually coded as 0 or 2 1812 1813 (i.e., they are, strictly speaking, pseudo-diploid), even though only one allele is observed. The 1814 "adjust pseudohaploid" argument ensures that the observed allele count increases only by 1 for each pseudo-haploid sample. If "TRUE" (default), samples that do not have any genotypes coded as 1815 1816 1 among the first 1,000 SNPs are automatically identified as pseudo-haploid. This leads to slightly 1817 more accurate estimates of f-statistics. Setting this parameter to "FALSE" treats all samples as 1818 diploid.

1819

Another important argument ("minac2=2") of the "extract\_f2" function removes sites with only one 1820 1821 chromosome genotyped in any non-singleton population and is needed for unbiased estimation of 1822 negative  $f_3$ -statistics in non-singleton pseudo-haploid populations. In the absence of negative  $f_3$ -1823 statistics or pseudo-haploid populations, this argument has no influence on admixture graph log-1824 likelihood scores. This algorithm for calculation of *f*-statistics triggered by the "*minac2=2*" argument 1825 is described below.

1826

1827 For  $f_3(a; b, c)$ , we compute the uncorrected numerator for each SNP,  $(a - b) \times (a - c)$ . We then subtract a bias correction factor at each SNP, p(1-p)/(ac-1), which we only need for population a 1828 1829 (because the other factors cancel out); p is the allele frequency, and ac is the observed allele count. 1830 In pseudo-haploid samples, (ac - 1) would be zero and produce an error in any sites with only one 1831 observed allele. With the "inbreed: NO" setting in Classic ADMIXTOOLS, the smallest non-zero value

for ac is 2, so the division by 0 problem is avoided, but the correction factor is slightly smaller than it 1832 1833 should be. ADMIXTOOLS2 adds only an allele count of 1 for each site in a pseudo-haploid sample 1834 (with the default option "adjust pseudohaploid = TRUE"), so there can be cases where ac = 1. To imitate what the setting "inbreed: NO" in Classic ADMIXTOOLS is doing, ac is set to 2 at those sites 1835 1836 (or the denominator is set to 1). There is still a small difference between Classic ADMIXTOOLS and ADMIXTOOLS2 at other sites because each observed site adds 2 alleles in ADMIXTOOLS with the 1837 1838 default setting "inbreed: NO", but only 1 allele in ADMIXTOOLS2 with the default setting 1839 "adjust pseudohaploid = TRUE", but for admixture graph fitting that does not matter. One solution to avoid biased correction factors is to only consider sites with ac of at least two, which is what the 1840 1841 "inbreed: YES" setting in Classic ADMIXTOOLS does. The problem with this is that we cannot use 1842 populations with a single pseudo-haploid sample, which is often useful, and would only give misleading results if that population is admixed. The new option "minac2=2" in ADMIXTOOLS2 is 1843 1844 different from the "inbreed: YES" setting in Classic ADMIXTOOLS since it makes an exception for 1845 populations consisting of a single pseudo-haploid sample in that it sets ac to 2 at each site 1846 (denominator is set to 1) when computing the correction factor of those populations.

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#### 1849 Fitting admixture graphs

1851 An admixture graph is a directed acyclic graph specifying the topology of the ancestral relationships 1852 among a set of populations. Each node in this graph represents a (present-day or ancient) 1853 population. Terminal nodes (also called leaf nodes) represent observed populations, while internal nodes represent unobserved ancestral populations. Modeling all observed populations as leaf nodes 1854 1855 confers some robustness to drift specific to single populations and to genotyping errors. The edges 1856 connecting the populations are weighted and correspond either to the magnitude of genetic drift 1857 that has occurred along that branch (drift edges), or to the admixture proportions (admixture edges, 1858 where two edges point to the same node).

1859

1860 The goal of *qpGraph* is to test how well a given graph topology fits the observed *f*-statistics. This is 1861 achieved by varying the edge weights until the maximum likelihood fit is obtained. The following 1862 section describes the graph fitting in more detail.

1863

First, for k populations, all  $\frac{k(k+1)}{2} f_3$ -statistics of the form  $f_3(0; X_1, X_2)$  are computed, where 0 is 1864 1865 one of the k populations (typically an outgroup), and  $X_1$  and  $X_2$  are all pairs formed from the other populations (including pairs where  $X_1 = X_2$ ). These  $f_3$ -statistics can then be used to fit the graph 1866 and to compute the likelihood. The likelihood score of a graph is the dot product of the differences 1867 1868 between the expected and observed  $f_3$ -statistics, weighted by the inverse covariance matrix of  $f_3$ -1869 statistics:

1870

1871 
$$L(g) = -\frac{1}{2} (f_{3,obs} - f_{3,fit})' Q^{-1} (f_{3,obs} - f_{3,fit})$$
 (Eq. 3)  
1872

Here,  $f_{3,obs}$  are the observed  $f_3$ -statistics and  $f_{3,fit}$  are the fitted  $f_3$ -statistics. Both are vectors of 1873 length  $q = \frac{k(k+1)}{2}$  for k populations excluding the outgroup. Q is the  $q \times q$  covariance matrix of  $f_3$ -1874 1875 statistics, where the diagonal entries are the  $f_3$ -statistic variances, and the off-diagonal entries are 1876 the covariances for all pairs of  $f_3$ -statistics. Just like the variances (the squared standard errors), the covariances are estimated from the jackknife leave-one-block-out  $f_3$ -statistics. 1877

1878

1879 Finding the edge weights which maximize the likelihood score involves two nested optimization 1880 steps. The inner optimization finds the drift weights which maximize the likelihood score while fixing 1881 the admixture weights. The outer optimization finds the admixture weights which maximize the 1882 likelihood score, while optimizing the drift weights for each set of admixture weights. The inner

1883 optimization uses a quadratic programming solver to find the optimal drift weights, while the outer

1884 optimization uses a general purpose optimization algorithm to find the optimal admixture weights.

1885 While the gradient function in the outer optimization adjusts the admixture weights, the objective 1886 function iterates over the following steps:

#### 1887 1888

1889

1890 1891

- 1. Optimization of drift weights conditional on admixture weights
- 2. Estimation of fitted  $f_3$ -statistics
- 3. Calculation of the graph likelihood using observed and fitted  $f_3$ -statistics

1892 These steps are repeated until convergence is reached and the likelihood score can no longer be 1893 improved by adjusting the admixture weights.

1894

1895Step 1 optimizes the drift edge weights, while holding the admixture weights constant. All drift edge1896weights are required to be non-negative, which makes this a constrained quadratic programming1897problem (hence qpGraph). Additional upper and lower bounds can be specified for individual graph1898edges.

1899

1900 Step 2 turns the edge weights into fitted  $f_3$ -statistics. To see how edge weights in an admixture 1901 graph translate to  $f_3$ -statistics, it helps to first consider how they translate into  $f_2$ -statistics for a pair 1902 of populations. Without any admixture events, there is exactly one path p connecting any two 1903 populations. The fitted  $f_2$ -statistic ( $f_{2,fit}$ ) is the sum of edge weights  $w_e$  along this path p connecting 1904 two populations. The fitted  $f_2$ -statistic is the sum of edge weights  $w_e$  along this path:

$$f_{2,fit} = \sum_{e \in p} w_e$$

1905 In the presence of admixture events, two populations may be connected via multiple paths. Each 1906 admixture node that lies between the two populations increases the number of possible paths. The 1907 fitted  $f_2$ -statistic for the two populations now becomes the weighted sum of all these paths, where 1908 the weight of each path is given by the product of all estimated admixture proportions  $w_a$  along this 1909 path ( $\prod_{a \in p} w_a$ ):

1910  $f_{2,fit} = \sum_{p \in P} \prod_{a \in p} w_a \sum_{e \in p} w_e$  (Eq. 4)

1911 The fitted  $f_2$ -statistics are then used to obtain fitted  $f_3$ -statistics using Eq. 1.

1912

1913 Step 3 uses the fitted and observed  $f_3$ -statistics to estimate the likelihood score using Eq. 3.

1914 1915 Prior to these three steps, initial admixture weights are drawn randomly. To ensure that the end 1916 results do not depend on the random initialization, the whole optimization process is repeated 1917 multiple times with different random initial values. The original *ADMIXTOOLS* implementation 1918 retains only the results from the initial values resulting in the lowest absolute likelihood score. The 1919 new *ADMIXTOOLS* implementation provides an option to retrieve the results for all random 1920 initializations. This can be useful, as large fluctuations between different random initializations can 1921 be an indicator of an overparameterized or otherwise poorly fitting model.

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#### 1924 Automated admixture graph inference

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To find graph topologies that could conceivably have given rise to the observed *f*-statistics, we start with a randomly generated graph with a fixed number of admixture events, apply a number of modifications to this graph, and evaluate each of the resulting graphs. We then pick the best-fitting graph and repeat this procedure until graph modifications no longer lead to improved scores. We use a number of random graph modifications, as well as targeted modifications which are informed by parameters obtained during the fitting of the current graph.

For the targeted modifications we change the optimization of a single graph from a constrained optimization problem, in which drift edges are constrained to be positive and admixture weights are constrained to be between zero and one, to an unconstrained optimization problem in which both types of parameters can take any real values. Rearranging the nodes adjacent to edges which were estimated to be negative results in an improved fit at a much higher rate than random graph adjustments.

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1940 The random modifications include (1) pruning and randomly re-grafting leaf nodes, (2) pruning and 1941 randomly re-grafting a set of connected nodes in the graph, (3) swapping the orientation of 1942 admixture edges, (4) shifting admixture edges, (5) re-rooting the graph, (6) combinations of two or 1943 more of any of these modifications.

1944

1945 The number of admixture events is not affected by the graph modifications described so far. A significant score improvement can often be achieved by adding a single admixture edge to several 1946 1947 random positions in a graph. This is unsurprising since it increases the degrees of freedom of the 1948 original graph. However, picking the best fitting graph with one admixture edge added, and testing 1949 all graphs that result from removing a single admixture edge from that graph often results in a graph 1950 with the same number of admixture events and a better fit than the original graph. We employ this 1951 strategy whenever the regular graph modifications described above do not lead to any further 1952 improvements.

1953

We keep track of the search tree of all previously evaluated graphs and their scores in order to not evaluate any graph more than once, and so that backtracking in the search space is possible in cases where no more local improvements can be identified. Nevertheless, multiple iterations with different random starting graphs are usually necessary to find graphs with good fits. The number of iterations needed to approach a global optimum depends on the size of the search space, but the optimal number of iterations is hard to estimate in practice.

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For revisiting published studies, we used the following settings of the *findGraphs* algorithm:

- mutfuns = namedList(spr\_leaves, spr\_all, swap\_leaves, move\_admixedge\_once, flipadmix\_random, place\_root\_random, mutate\_n), a list of functions used to modify graphs.
- *numgraphs* = 10, number of alternative graphs produced by randomly applying the mutation functions at the start of each generation.
- *stop\_gen = 10000*, total number of generations after which to stop.
- *stop\_gen2 = 30*, number of generations without LL score improvement after which to stop.
- *plusminus\_generations = 10.* If the best score does not improve after *plusminus\_generations* generations, another approach to improving the score is attempted: A number of graphs with an additional admixture edge is generated and evaluated. The resulting graph with the best score is picked, and new graphs are created by removing any one admixture edge (bringing the number back to what it was originally). The graph with the lowest score is then selected. This approach often makes it possible to break out of local optima.
- 1974 $opt\_worst\_residual = FALSE.$  Optimize for lowest worst residual instead of best score.1975"FALSE" by default, because the LL score is generally a better indicator of the quality of the1976model fit, and because optimizing for the lowest worst residual is much slower since  $f_{4^-}$ 1977statistics need to be computed.
- 1978 $reject_f4z = 0.$  If this is a number greater than zero, all  $f_4$ -statistics with |Z-score| >  $reject_f4z$ 1979will be used to constrain the search space of admixture graphs: Any graphs in which  $f_4$ -1980statistics greater than  $reject_f4z$  are expected to be zero will not be evaluated.
- *diag = 1e-04*. This argument is passed to the *qpgraph* function and determines the regularization term added to the diagonal elements of the covariance matrix of fitted branch lengths (after scaling by the matrix trace). Default is 0.0001.

- *numstart = 10.* This argument is passed to the *qpgraph* function and determines the number
   of random initializations of starting weights (defaults to 10). Increasing this number will
   make the optimization slower but reduce the risk of not finding the optimal weights.
- 1987 1988

Isqmode = FALSE. This argument is passed to the qpgraph function. If set to "FALSE", the inverse f<sub>3</sub>-statistic covariance matrix is not discarded by the algorithm.

The arguments "admix\_constraints" (constraints on the number of admixture events in the history of a given population), "event\_constraints" (constraints on the branching order of specified lineages), and "outpop" (the population assigned as an outgroup) were set according to **Table S1**. Each *findGraphs* run was initiated by a random graph with a specified number of admixture events. Usually, the same topology constraints were applied at the stage of random graph generation and the topology search stage, for exceptions see **Table S1**.

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1997 Evaluating automated admixture graph inference through simulations

1999 We evaluated the performance of *findGraphs* by simulating genetic data under a large number of 2000 different admixture graph models, applying findGraphs to each simulated data set in three 2001 independent iterations, and comparing the resulting best graph across three iterations to the 2002 simulated graph. We applied TreeMix to the same simulated data for comparison. We simulated 2003 between 8 and 16 populations per graph, and between 0 and 10 admixture events. For each 2004 parameter combination, we simulated 20 different admixture graphs generated by the 2005 random\_admixturegraph function. We counted both the fraction of random simulated graphs where 2006 the best inferred graph was identical to the simulated graph, as well as the fraction of random 2007 simulated graphs where the best inferred graph was either identical to the simulated graph or had a 2008 better score than the simulated graph. For models with a large number of admixture events the 2009 number of possible models is so large that it becomes increasingly likely that there will be some 2010 alternative models which fit the data better than the model under which the data were simulated.

2011

2012 We used *msprime* and the *msprime sim* wrapper function in *ADMIXTOOLS* 2 to simulate data for 2013 100,000 unlinked SNPs and 100 diploid samples per population for each admixture graph. The 2014 simulation parameters we chose were aimed at facilitating fast simulations of large numbers of 2015 informative SNPs rather than at being as realistic as possible. We therefore expect that the 2016 simulation results allow us to make comparisons across groups, but not that they are informative 2017 about the rate at which "true" models can be recovered in empirical data. We simulated under a 2018 constant mutation rate of 0.001 per site per generation, a constant haploid effective population size 2019 of 1000, with neighboring nodes in the graph separated by 1000 or more generations, and all 2020 admixture events occurring in discrete pulses of 50/50 proportions.

To allow for a fair comparison between *findGraphs* and *TreeMix*, we made sure that small differences in the way admixture graphs are modeled in *findGraphs* and in *TreeMix* were accounted for before testing graphs for identical topology. For example, *TreeMix* admixture graphs can have lineages terminating at an admixture node, whereas in *findGraphs* lineages always end at a 'leaf' node with a single ancestor.

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#### 2029 Comparing the fits of different admixture graphs

2031 We are interested in determining whether one admixture graph fits the data significantly better 2032 than another admixture graph, or whether an observed score difference  $\Delta = S_1 - S_2$  can be 2033 attributed to variability across independent SNPs.

2035 We first consider two admixture graphs with the same number of admixture events, where we can 2036 ignore the problem of comparing two models with different complexity. As in other bootstrap 2037 standard error calculations, we divide the genome into n blocks indexed by i, and we draw b sets of 2038 n blocks with replacement, indexed by j. We fit both graphs b times - once for each bootstrap set of 2039 SNP blocks. This results in a set of b score differences  $\Delta_i$ . The bootstrap confidence interval for the 2040 difference in scores is given by the quantiles of the distribution of  $\Delta_j$ . We also compute an empirical bootstrap p-value, testing the null hypothesis that two different graphs fit the data equally well. It is 2041 computed as  $p = max(\frac{1}{\mu}, 2\delta)$  (Boos 2003), where  $\delta$  is either the fraction of  $\Delta_i > 0$ , or the 2042 fraction of  $\Delta_i < 0$ , whichever is smaller. The reason for applying bootstrap resampling, as opposed 2043 2044 to jackknife resampling in this case, is that the distribution of score differences tends to have a high 2045 kurtosis, which can make jackknife estimates inaccurate. Simulating data under the null hypothesis is 2046 not straightforward in this case, because it involves finding two non-identical graphs which in 2047 expectation fit the data equally well. We decided to simulate under one graph and compare two 2048 graphs which are symmetrically related to the simulated graph (Figure 3). This confirmed that the p-2049 value follows a uniform distribution under the null hypothesis.

2050

Next, we consider comparisons of two graphs of different complexity. The problem here is that more
 complex graphs have more degrees of freedom which allow them to overfit the data better, without
 necessarily being any closer to the truth. To solve this problem, we introduce an out-of-sample
 likelihood score. The regular likelihood score is given by:

2055  $L(g) = -\frac{1}{2} (f_{3,obs} - f_{3,fit})' Q^{-1} (f_{3,obs} - f_{3,fit})$ , with  $f_{3,obs}$  and  $f_{3,fit}$  defined on the same set of 2056 SNPs. The out-of-sample likelihood score is defined in the same way, except that  $f_{3,obs}$  and  $f_{3,fit}$  are 2057 defined on mutually exclusive sets of SNP blocks, thereby preventing any overfitting. The covariance 2058 matrix Q is defined on the same set of SNP blocks as  $f_{3,fit}$ . As described earlier, we use block-2059 bootstrap to fit both graphs multiple times on different SNP blocks. In each bootstrap iteration, we 2060 use all SNP blocks which are not used in fitting the graph for estimating  $f_{3,obs}$ .

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#### 2063 Admixture graph identifiability

2064 2065 An edge in an admixture graph is unidentifiable, if small changes to the weight of this edge 2066 (admixture proportions in the case of an admixture edge, drift length in the case of a drift edge) do 2067 not necessarily lead to changes in expected *f*-statistics. This is the case if the small change in weight 2068 can be offset by small changes in other graph edges, leading to a situation where observed f-2069 statistics can be explained by more than one weight estimate for that edge. To find unidentifiable 2070 edges, we derive the Jacobi matrix of the graph's system of  $f_2$  equations (Eq. 4 applied to each 2071 population pair). In principle, whether or not a parameter is identifiable can depend on the values of 2072 all other parameters. However, in practice this is rarely the case, and so we draw values for all 2073 parameters from a uniform distribution, which gives us a Jacobi matrix with numeric values. We 2074 then determine the rank of the Jacobi matrix, along with the rank of all matrices that result from 2075 dropping a single column (a parameter corresponding to a graph edge). For identifiable edges, the 2076 rank of the full matrix will be greater than the rank of the reduced matrix, and for unidentifiable 2077 edges, the ranks will be the same.

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#### 80 Drawing conclusions from a large number of fitting models

2082 We developed several methods that aim to summarize a collection of graphs which all fit the data 2083 similarly well. By highlighting features which are observed repeatedly across graphs, it becomes 2084 possible to extract interpretable conclusions from an otherwise hard to interpret collection of 2085 possible models. These graph summaries identify features in each graph that can be compared to 2086 different graphs describing the same populations. We summarize each graph in several ways:

- 2087 (1) Admixture status of each population
- 2088For each population, we count the total number of admixture events that is encountered2089along all paths to the root.

2090	(2) Order of population split events
2091	For each pair of population pairs, we determine if the most recent split of the first pair has
2092	occurred before or after the most recent split of the second pair, or whether the graph does
2093	not specify the order in which those splits occurred.
2094	(3) Proxy populations
2095	For each admixed population in a graph, we attempt to identify proxy sources: populations
2096	closest to the admixing populations. In contrast to the other approaches to summarizing
2097	graphs which are based only on the topology of each graph, this can also rely on information
2098	about the estimated graph parameters.
2099	(4) Cladality
2100	For each group of four populations, we test whether the graph implies that any $f_4$ -statistic
2101	describing the relationship between the four populations is expected to be zero.
2102	(5) Node descendants
2103	Each internal node in an admixture graph is an ancestor to a specific set of leaf populations.
2104	An admixture graph can be characterized by the sets of leaf populations formed by the
2105	internal nodes. Multiple admixture graphs may be compared by counting the number of
2106	overlapping sets. This also makes it possible to quantify for each internal node in a single
2107	graph, how often a matching internal node can be found across a collection of alternative
2108	graphs, which is conceptually similar to bootstrap support values in phylogenetic trees.
2109	While these methods provide some help in comparing features across many graphs, they are not
2110	able to reliably answer the question whether the fitting graphs are relatively similar or dissimilar
2111	from each other, and whether they are similar to any particular graph. This is in part due to the fact
2112	that small topological changes involving populations of interest may be more relevant than similar
2113	topological changes involving only populations that are not the focus of the study.
2114	
2115	
2116	<u>Acknowledgements</u>
2117	
2118	We thank Anders Bergström, Esther Brielle, Mateja Hajdinjak, Iosif Lazaridis, Pablo Librado, Mark
2119	Lipson, Vagheesh Narasimhan, Ludovic Orlando, Nick Patterson, Mary Prendergast, Jakob Sedig,
2120	Kendra Sirak, Pontus Skoglund, and Chuanchao Wang, for suggestions for how to improve specific
2121	analyses, and for conversations and critical comments. We thank Matthew Mah, Shop Mallick, Adam
2122	Micco, Nadin Rohland, Ron Pinhasi for help in generating additional data from an ancient DNA
2123	library from individual 18726 for which 1.24 million SNP capture data was generated and published in
2124	Narasimhan et al. 2019 and for which we report 2.6-fold shotgun data here (Table S2). P.F., P.C., and
2125	O.F. were supported by the Czech Ministry of Education, Youth and Sports (program ERC CZ, project
2126	no. LL2103) and by the Czech Science Foundation (project no. 21-27624S). P.F. and P.C. were also
2127	supported by the Czech Ministry of Education, Youth and Sports (Large Infrastructures for Research,
2128	Experimental Development and Innovations project "IT4Innovations National Supercomputing
2129	Center – LM2015070"; Inter-Excellence program, project no. LTAUSA18153). R.M. and D.R. were
2130	supported by grants from the National Institutes of Health (GM100233 and HG012287), the John
2131	Templeton Foundation (grant 61220), and the Allen Discovery Center program, a Paul G. Allen
2132	Frontiers Group advised program of the Paul G. Allen Family Foundation. D.R. was also supported by
2133	a private gift from JF. Clin and is an Investigator of the Howard Hughes Medical Institute.
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2527 **Supplementary Tables** 

2521	Juppi
2528	

Publication	tiste inth	e orieind	Populitation of the strategy o	ST SIONE USE	ABPIOR PO	Julions actalians events el.nobelity	elite ino	A LIN BOOST	Settings for calculating f2-statistics	To pology search constraints and population modifications	Herei	ors used	sions cont	nrine subi net sterre	the dash	seesound	S topologic	Etheopolities	Die wood	Berlitine	opologies percentri significantly significantly	Pre Dopole	Les ar in the l
Bergst röm et al. 202	0 1e	6 7	4 5	0 2 0 3	4./ 8.4	4.9	2.1	312,282	max miss=0, big_size=4000000, adjust_pseudohaploid=T	And ean Fox OG	100	38 337	14 221	0	0 5	3 37	11 178	0.0	0.0	21.4	78.6 80.5	0.060*	yes
Lazaridis et al. 2014	3	6 7	1	62 64	3.0 8.7	4.2 11.1	2.2	624,583 642,247	maxmiss=0, blg_size=0.05, adjust_pseudohaploid=T, minac2=2	MbutiOG	100 1,000	32 0	14 306	0 3	0 37	0 247	14 19	0.0	0.0 12.1	0.0 80.7	100.0 6.2	0.032* 0.032	yes
China and 2010	2	9 8	3 2	6 3 2 3	22.1 29	40.9 40.8	2.8 3.2	19,017 470,389	max miss=0, blg_size=0.05, adjust_pseud oha ploid=T	Mota OG  4	4,000 4,000	0	398 216	0 4	89 61	266 79	43 72	0.0	22.4 28.2	66.8 36.6	10.8 33.3	0.136 0.072	
Shindeetal.2019	3	8 8	1	10 3 4	17.9 N/A	) 28.2 N/A	2.6 N/A	249,009	max miss=0, blg_size=0.05, adjust_pseudo haploid=T, minac 2=2	<ul> <li>Mota OG; Indus_Peripherygroup expanded to 4 ind., relatives and contaminated ind. removed</li> </ul>	4,000 4,000	13 N/A	143 443	0 N/A	4 N/A	5 N/A	134 N/A	0.0 N/A	2.8 N/A	3.5 N/A	93.7 N/A	0.088* N/A	yes
	3 b Ext5d			3 4	907. 363.	3942 3386.2	28.3 15.7				1,000 1,000	25 10	335 531	nottes	ted							0.002	
	Ext Se	10	3	3 6	267.	7 286.5	10.4	6,343,116	max miss=0, blg_size=4000000,		1,000	0	747 894									0.002	
Librado et al. 2021	N/A			7 8 9	N/A	N/A	N/A		ad just_pseudonapioid=i	Donkey OG; group composition was altered : for all populations	1,000 1,000 1,000	N/A	986 999 1,000	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	
	3 b Ext5d Ext5e			3 4 5	679. 202. 121	9 711.1 8 224.4 139.6	23.9 14.1 6.9			— individuals from o hiy o ne archaeo logicals ite and temporal horizon were included	1,000 1,000 1,000	1 30 3	324 535 784	22 0 0	51 0 2	78 24 223	173 511 559	6.8 0.0 0.0	15.7 0.0 0.3	24.1 4.5 28.4	53.4 95.5 71.3	0.040 0.788* 0.720	yes yes
	N/A	10	3	3 6 7 8	3 6 7 N/A N/A N/A	1,767,419 N/A	67,419 ad just_pseudo hap loid=⊤, minac 2=2		1,000 1,000 1,000	N/A	954 995 1,000	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A			
	3 b Ext 5d Ext 5e N/A	10	1	9 3 4 5 1 6 7 8 9	1133 477. 380. N/A	3 1163 1 500.5 5 395 A N/A	33.8 16.3 13.2 N/A	7, 403,037	, maxmiss=0, blg_size=4000000, adjust_pseudohaploid=T	Don key OG	1,000 1,000 1,000 1,000 1,000 1,000 1,000 1,000	0 3 0 N/A	1,000 363 567 729 906 981 999 998	nottes N/A	ted N/A	N/A	N/A	N/A	N/A	N/A	N/A	0.002 0.002 0.002 N/A	
		11	5	0 8	35.8	65.1	2.8	283,890		Denisovan OG [4] Vindija Mhuti unadmixed, Denisovan max, 1 admix, event	4,000	0	3,996					15.9 26.0	80.8	3.2	0.0	0.008	
Hajdinjak et al. 202:	12d -	12	6	0 8	71.6	5 112.4	4.8	263,698	– max miss=0, blg_size=0.05, adjust_pseud ohaploid=T	Chimp OG Vindia Mbuti unad mixed. Denisovan max. 1 admix. event. Chimp. OG	2,000	0	1,995 1,988	– a fracti	onofmo	odels te	ested	56.8 15.7	41.4 55.7	1.8	0.0	0.002	yes
	\$3.24 \$3.25	7 10	1	0 4 15 8	5.4 33.6	8.7 5 51.2	1.9 2.7	801,362 838,910	max miss=0, blg_size=4000000, adjust_pseudo haploid=E_mipac 2=2	Chimp OG  4	2,000	0 0	778 9,927	2 afracti	199 on of mo	489 odelste	89 ested	0.3	25.6 83.5	62.9 9.1	11.4 0.6	0.260 0.032	
Lipson et al. 2020	Ext A	12	2	22 12	2 29.5	61.6	2.2	363,131	max miss=0, blg_size=0.05, adjust_pseudo haploid=T	Chimp OG	2,000	0	2,000	nottes	ted			0.0	28.5	69.5	3.0	0.064	
	\$13-9  2	4	2	11	1 21.8	52.5 29.6	2.3	211,738	minacter minacter maxmissen ble sizes 0.5 adjust nseudobabloidet	Chimp OG, Altai unadmixed, French min. 1 admix. event	2,000	0	2,000	a fract i	onofmo	odels te	ested	0.0	28.5 11.9 38.8	77.1	10.4	0.176	yes
Wang et al. 2021	SI3-10a  3	, 10	3	0 5	23.3	41.8	2.5	544,068	maxmiss=0, blg_size=0.05, adjust_pseudohaploid=F maxmiss=0, blg_size=0.05, adjust_pseudohaploid=F	Den isovan OG  4	2,000	0	1,524	a fracti	onofmo	odels te	ested	0.0	44.0	56.0	0.4	0.288	
wang et al. 2021	Ext6	12	3	38	62.1	99.2	3.1	496,233	max miss=0, blg_size=0.05, adjust_pseudohaploid=T same as above + minar 2=2	Denisovani OG [4] Denisovan,  Mbuti,  Loschbour, Onge     branching order, Mbuti,	2,000	0	1,993	- nottes	ted	odalsta	stad	12.6	8/1 3	3.1	0.0	0.004	VAS
				10	) 65.7	1164	3.2	203,733	2011-03-000 VC + 11118-2-2	<u>Loschpour, Unge unad mixed. Denisovan max 1 ad mix: Denisovan OG 14</u> Altai, Dinka unadmixed, Denisovan max. 1 ad mix. event, Chimp OG	1,000	0	1,000	anacti	onormu	oueiste	.a.cu	33.2	13.6	50.0	3.2	0.002	yes.
	3f  left	13	4	0	76 5	132	3.8	344,903		Chimp OG Altai, Dinka unadmixed, Denisovan max. 1 admix. event, Chimp OG	1,000	0	996 894	-				28.5 0.3	68.7 17.1	2.8 34.6	0.0 48.0	0.002	yes
Siko ra et al. 2019				10	) 85.3	147.8	3.1		– max miss=0, blg_size=0.05, adjust_pseud ohaploid=T	Chimp OG Altal, Dinka unadmixed, Denisovan max. 1 ad mix. event, Chimp OG Chimp OG	1,000	0	966 1,000	– a fracti	onofmo	odels te	ested	0.1	37.0 30.0 44.4	52.3 17.6	10.6 39.2	0.024	
	3f right	14	2	0 6	102.	4 171.9	4.2	613,509		Altai, Dinka unadmixed, Denisovan max. 1 admix. event, C himp OG Chima OG	4,446	0	2,785					0.1	0.9	9.8 40.3	89.2 52.8	0.112	yes

Counting at most one topology per iteration, not double counting topologies found in multipler iterations
 a 1% gene flow from Loschbour to the Mongolia Neolithic group was dropped from the published model to decrease the complexity of the baseline model. That increased model LL slightly (by 0.5 log-units).
 a 1% gene flow from Loschbour to the Mongolia Neolithic group was dropped from the published model to decrease the complexity of the baseline model. That increased model LL slightly (by 0.4 log-units).
 a 0% gene flow from Loschbour to the Mongolia Neolithic group was dropped from the published model to decrease the complexity of the baseline model. That increased model LL slightly (by 2.4 log-units).
 a 0% gene flow from Loschbour to the Mongolia Neolithic group was dropped from the published model to decrease the complexity of the baseline model. That increased model LL slightly (by 2.4 log-units).
 a 0% gene flow from Loschbour to the "find\_graphs" algorithm itself

\* here the direction of comparison is reversed since the published model is the highest-ranking among all models found

**Table S1: Published graphs in the context of automatically found graphs.** We compared 22 different graphs from 8 publications to alternative graphs inferred on the same or very similar data; these *findGraphs* runs are highlighted in blue in the "Iterations" column. In total, 51 *findGraphs* runs are summarized here since in some cases models more

complex or less complex than the published one were explored and/or different population compositions were tested (see the "Topology search constraints and population modifications" column and footnotes for details). The columns with names in blue show various information on the published graphs or their modified versions and some

properties of the published population sets. The columns with names in magenta show settings used for calculating *f*-statistics and for exploring the admixture graph space, and

the number of SNPs used that depends on them. The columns with names in black summarize the outcomes of *findGraphs* runs, i.e., the properties of alternative model sets

2537 found.

2538 **Publication**: Last name of the first author and year of the relevant publication.

2539 **Figure in the original publication**: Figure number in the original paper where the admixture graph is presented.

**Groups (populations)**: The number of populations in each graph.

2541 Singleton pseudo-diploid populations: The number of populations in the graph composed of a single pseudo-diploid individual. Calculation of negative "admixture"  $f_3$ -statistics is

impossible for such populations since their heterozygosity cannot be estimated (see the text for details).

2543 No. of negative f3-stats (allsnps: YES): The number of negative  $f_3$ -statistics among all possible  $f_3$ -statistics for a given set of populations when all available sites are used for each

statistic. If no negative f<sub>3</sub>-statistics exist for a set of populations, admixture graph fits are not affected by the "minac2=2" setting intended for accurate calculation of f-statistics for

2545 non-singleton pseudo-diploid groups.

- 2546 Admixture events: The number of admixture events in each graph.
- 2547 Publ. model: log-likelihood (LL): Log-likelihood score of the published graph fitted to the SNP set shown in the "SNPs used" column.
- 2548 Publ. model: LL, median of bootstrap distr.: Median of the log-likelihood scores of 100 or 500 fits of the published graph using bootstrap resampled SNPs.
- Publ. model: worst residual (WR), SE: The worst *f*-statistic residual of the published graph fitted to the SNP set shown in the "SNPs used" column, measured in standard errors
   (SE).
- 2551 SNPs used: The number of SNPs (with no missing data at the group level) used for fitting the admixture graph. For all case studies, we tested the original data (SNPs, population
- 2552 composition, and the published graph topology) and obtained model fits very similar to the published ones. However, for the purpose of efficient topology search we adjusted 2553 settings for  $f_3$ -statistic calculation, population composition, or graph complexity as shown here and discussed in the text.
- 2554 Settings for calculating f2-statistics: Arguments of the *extract\_f2* function used for calculating all possible f2-statistics for a set of groups, which were then used by *findGraphs* for
- 2555 calculating  $f_3$ -statistics needed for fitting admixture graph models. See Methods for descriptions of each argument.
- 2556 **Topology search constraints and population modifications**: Constraints applied when generating random starting graphs and/or when searching the topology space.
- 2557 Modifications of the original population composition are also described in this column, where applicable.
- **Iterations**: The number of *findGraphs* iterations, each started from a random graph of a certain complexity. For each case study, *findGraphs* setups that were considered optimal are highlighted in blue in this column.
- 2560 Iterations confirming published graph: The number of iterations in which the resulting graph was topologically identical to the published graph. In the cases when the published
- model was irrelevant since more complex graphs were explored, "N/A" appears in this and subsequent columns. If less complex models were explored, the published model was still relevant since its version without selected admixture edges was tested.
- 2563 Distinct alternative topologies found: The number of distinct newly found topologies. If graph complexity was equal to (or less than) that of the published graph, the published
- topology (or its simplified version) is not counted here. If graph complexity exceeded that of the published graph, all newly found topologies are counted. If the published
- 2565 topology was recovered by *findGraphs*, the numbers in this column are shown in bold.

2566 Significantly better fitting topologies: The number of distinct topologies that fit significantly better than the published graph according to the bootstrap model comparison test

- (two-tailed empirical p-value <0.05). If the number of distinct topologies was very large, a representative sample of models (1/20 to 1/3 of models evenly distributed along the
- log-likelihood spectrum) was compared to the published one instead. These cases are marked as "a fraction of models tested" in this column. If model complexity was higher than
- that of the published model, model comparison was irrelevant and was not performed.
- 2570 Non-significantly better fitting topologies: The number of distinct topologies that fit non-significantly (nominally) better than the published graph according to the bootstrap
   2571 model comparison test (two-tailed empirical *p*-value ≥0.05).
- 2572 Non-significantly worse fitting topologies: The number of distinct topologies that fit non-significantly (nominally) worse than the published graph according to the bootstrap
   2573 model comparison test (two-tailed empirical *p*-value ≥0.05).
- 2574 Significantly worse fitting topologies: The number of distinct topologies that fit significantly worse than the published graph according to the bootstrap model comparison test
   2575 (two-tailed empirical *p*-value ≥0.05).
- 2576 Significantly better fitting topologies, %: The percentage of distinct topologies that fit significantly better than the published graph according to the bootstrap model comparison
- test (two-tailed empirical p-value <0.05). If the number of distinct topologies was very large, a representative sample of models (1/20 to 1/3 of models evenly distributed along
- the log-likelihood spectrum) was compared to the published one instead, and the percentages in this and following columns were calculated on this sample.
- Non-significantly better fitting topologies, %: The percentage of distinct topologies that fit non-significantly (nominally) better than the published graph according to the
- 2580 bootstrap model comparison test (two-tailed empirical *p*-value  $\ge 0.05$ ).
- **Non-significantly worse fitting topologies,** %: The percentage of distinct topologies that fit non-significantly (nominally) worse than the published graph according to the bootstrap model comparison test (two-tailed empirical p-value  $\geq 0.05$ ).
- 2583 Significantly worse fitting topologies, %: The percentage of distinct topologies that fit significantly worse than the published graph according to the bootstrap model comparison test (two-tailed empirical *p*-value ≥0.05).
- 2585 **P-value best alternative vs. publ.**: An empirical two-tailed *p*-value of a test comparing log-likelihood distributions across bootstrap replicates for two topologies, the highest-
- ranking newly found topology and the published topology. In some cases, the highest ranking newly found topology (according to LL) has a fit that is not significantly better than
- that of the published model, but other newly found models fit significantly better despite having higher LL. *P*-values below 0.05 are highlighted in green.
- 2588 Used in Table 1: Here the *findGraphs* runs featured in Table 1 are marked.

#### Table S2: Statistics for shotgun sequencing of individual I8726

18726
SHAR_201 (Grave 201)
Petrous bone
Seistan, Shahr-i-Sokhta, Iran
3100-3000 BCE
30.649857
61.400311
Narasimhan, Patterson et al. Science 2019
S8726.E1.L1
3
Male
2.60306

- 2592 Supplementary Items Available as Separate Files
- 2594 5 supplementary tables are available as separate files (Tables S3-S7)
- 2595

2593

2596 **13** supplementary figures are available as separate files (Figures S1-S13)