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Simulating Human Origins and Evolution

The development of populations over time and, on longer time scales, the evolution of species are both influenced by a complex of interacting, underlying processes. Computer simulation provides a means of experimenting within an idealised framework to allow aspects of these processes and their interactions to be isolated, controlled and understood.

In this book, computer simulation is used to model migration, extinction, fossilisation, interbreeding, selection and non-hereditary effects in the context of human populations and the observed distribution of fossil and current hominoid species. The simulations described enable the visualisation and study of lineages, genetic diversity in populations, character diversity across species and the accuracy of reconstructions, allowing new insights into human evolution and the origins of humankind for graduate students and researchers in the fields of physical anthropology, human evolution and human genetics.

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*Each change of many-coloured life he drew
Exhausted worlds, and then imagined new.*

Samuel Johnson 1709–84

For Cindy, Jessamine and Xanthe

Contents

<i>Preface</i>	<i>page</i> xiii
1 Introduction	1
1.1 Phylogenetics and human origins	1
1.2 Origin of modern humans	7
1.3 Computer methods in phylogenetics	11
Part I Simulating species	15
2 Overview	17
2.1 Hominoids	20
2.2 Hominids	22
3 Simulation design	26
3.1 Phylogenetic reconstruction	29
3.2 Example simulation and reconstruction	33
3.3 Analysis and evaluation	38
4 Running the simulation	42
4.1 A simple example	42
4.2 Migration	46
4.3 Advanced features	52
5 Simulating diversity	56
5.1 Recent reduction in diversity profiles	57
5.2 Recent maximum of diversity profiles	69
5.3 Studying parameter sensitivity	74
6 Simulating migration	84
6.1 Species migration with an amphora profile	84
6.2 Simulating hominoid migrations	91

x	<i>Contents</i>	
	6.3 Restricted migrations and interbreeding	95
	6.4 Unrestricted migration with advantage	113
7	Discussion	118
	7.1 Single-continent summary	118
	7.2 Migration summary	122
	7.3 Implications	126
	7.4 Future work	128
	Part II Simulating genealogies	131
8	Overview	133
	8.1 Coalescent theory	134
	8.2 The historical human population	139
	8.3 Human mating patterns and fertility	141
	8.4 Coalescence and biological ancestry	143
9	Simulation design	151
	9.1 Parameters	152
	9.2 Simulating and analysing a genealogy	153
	9.3 Output data and visualisation	155
10	Simulating a single population	162
	10.1 Constant demographics	162
	10.2 Varying demographics	174
11	Simulating multiple populations	186
	11.1 Sample simulation with regular migrations	186
	11.2 Simulations with restricted migrations	191
12	Adding genetics to the genealogy	201
	12.1 Modelling genetics with coalescent theory	201
	12.2 Genetics models in the simulation	209
	12.3 Sex-specific migrations and selection	211
13	Discussion	220
	13.1 Single-population summary	220
	13.2 Migration summary	223

Cambridge University Press
0521843995 - Simulating Human Origins and Evolution
K. P. Wessen
Frontmatter
[More information](#)

<i>Contents</i>	xi
13.3 Genetics summary	225
13.4 Implications for modern human origins	225
13.5 Future work	228
<i>References</i>	231
<i>Index</i>	239

Preface

Recent times have seen a great deal of activity and progress in human origins research, from the advent of molecular methods in the 1960s to the many important fossil hominid discoveries of the past few years. Nevertheless, the debate over whether particular fossil species are direct human descendants or not, and whether the fossil record and molecular results support a recent African origin or multiregional continuity, continues to rage. There is clearly a substantial need for fundamental work studying the methods employed in the interpretation of these data. The primary aim of the research presented in this volume is to begin to address this need by means of direct computer modelling and simulation of the many underlying and interacting processes.

Specifically, this volume describes the development and application of two related, but distinct, simulations, each designed to model important aspects of evolution in general, and the origin and evolution of humans in particular, as well as to provide substantial analysis and a wide variety of visualisations of the results.

The first simulation, *Specialist*, models the evolution of species and subspecies over millions of years, by starting with a single ancestral species with a particular suite of morphological ‘characters’ and allowing it to evolve in discrete steps. The characters are either hereditary or non-hereditary, and at each step a small number of these characters may change, either owing to random mutation or as a result of a change in the nature of the home environment of each species. Random extinction, fossilisation, interbreeding of subspecies, migration between four continents, and selective advantage are included in the model.

The main focus is on using the resulting species data to construct a phylogeny and migration history, which is then compared against the known true phylogeny. Two techniques of reconstruction are employed. The first technique involves matching existing species and fossils to the most closely (morphologically) related earlier fossil, whereas the second involves a reconstruction based on differences between the characters of the existing species only.

The second simulation, *Genie*, models several generations of individuals in up to three independent populations, thus allowing study of the effects of different mating patterns, fertility, adult sex ratio, migrations of various types, limiting population size, selective advantage and the impact of external,

natural disasters on common ancestry and the mixing of lineages generally. Once a complete genealogy is generated, common ancestry, lineage mixing and migrations are determined and analysed for the purely paternal and maternal genealogies (corresponding to Y chromosome and mitochondrial DNA inheritance), as well as the *biological* genealogy, or pedigree, where lineages are traced back through both parents simultaneously. This analysis is carried out on both a small sample of individuals and the full population, and individuals in the population carry both sex-specific and autosomal genes that are subject to mutation and recombination in controlled ways.

The simulations presented are essentially simulations of evolutionary change, and as such may be applied across a very large range of problems. As is apparent from the title of this volume, I have chosen to focus on problems relating to human and hominoid evolution, but extensions to many other areas are relatively straightforward, especially for the species/subspecies simulation.

Both simulations can do either single runs, with various visualisations and interactions, or multiple runs, with more limited visualisation but with basic statistical analysis of the results and all the required information for more advanced analysis provided in a simple text report. In particular, the simulated demographic, genetic and genotype data from *Genie* may be easily exported into other programs to provide more detailed or custom analysis. This removes the need for *Genie* to try to cover the myriad of possible analyses.

The programs that implement the simulations may be freely downloaded by following the links from <http://school.anhb.uwa.edu.au/personalpages/kwessen>.

In the interests of quality control, some minor limitations have been placed on the downloadable versions of the software, but these limitations can be removed via a simple registration process that will also allow me to provide updates and maintain some degree of dialogue with users. It is my hope that making the software available will lead to much further and diverse development of the simulations in collaboration with other researchers.

In addition, many of the figures in this book are black and white, or otherwise adjusted, versions of colour visualisations produced by the simulations. For this reason, the majority of simulations presented in the text are available for download along with the software, enabling them to be viewed in colour, and also enabling the many interactions provided by the simulation program to be explored in the context of these particular simulations. In order to gain a full appreciation of the results presented in this book, readers are urged to download the associated software and familiarise themselves in a hands-on way with the models and visualisations employed.

I take this opportunity to heartily thank Professor Charles Oxnard, without whose ongoing encouragement and highly infectious enthusiasm this project would never have begun, let alone finished. I also thank Professor Paul O'Higgins for his comments on an earlier version of this work; those comments were particularly instrumental towards providing the necessary impetus for me to undertake the publication of this work in book form. Professor Colin Groves also had several useful comments on an earlier manuscript, and various suggestions from Algis Kuliukas have led to valuable enhancements to the species simulation. Thanks are also due to Mat Abdy for his help in preparing this book's associated website.

And, of course, my most sincere and personal thanks go to my beautiful wife Cindy and lovely daughters Jessamine and Xanthe, each of whom will, I'm sure, very much share my relief at seeing this book complete!