Early hominid biogeography

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ABSTRACT We examined the biogeographic patterns implied by early hominid phylogenies and compared them to the known dispersal patterns of Plio-Pleistocene African mammals. All recent published phylogenies require between four and seven hominid dispersal events between southern Africa, eastern Africa, and the Malawi Rift, a greater number of dispersals than has previously been supposed. Most hominid species dispersed at the same time and in the same direction as other African mammals. However, depending on the ages of critical hominid specimens, many phylogenies identify at least one hominid species that dispersed in the direction opposite that of contemporaneous mammals. This suggests that those hominids may have possessed adaptations that allowed them to depart from continental patterns of mammalian dispersal.

A critical biogeographic question in early human evolution concerns the relationships between the eastern and southern African hominids. Given that the mammalian fossil record preserves evidence of faunal interchange between these two regions (1, 2), it is very unlikely that hominids in one region evolved entirely independently from those in the other. Yet, despite the strong probability that early hominids dispersed between eastern and southern Africa, few studies (1–4) have examined the paleobiogeography of early hominid species. These studies agree that either two, or three, dispersal events must have taken place and that, in general, the early hominids followed the continental dispersal patterns of other large-bodied mammals (Fig. 1a). Some of these studies are predicated by the hypothesis that hominid speciation and dispersal events are driven by environmental change, and in particular by the hypothesis that mammals shift their ranges to match changes in the distributions of vegetational zones (2, 5, 6).

Prior studies have addressed the impact of ecology on hominid biogeography, but the biogeographic implications of phylogeny have yet to be fully explored. Obtaining a reliable phylogeny is a crucial, initial step in the formulation of any biogeographic hypothesis because, if a species in one region is descended from an ancestor in another region, then a vicariance (splitting of ranges) or dispersal event must have taken place and that, in general, the early hominids followed the continental dispersal patterns of other large-bodied mammals (Fig. 1a). Some of these studies are predicated by the hypothesis that hominid speciation and dispersal events are driven by environmental change, and in particular by the hypothesis that mammals shift their ranges to match changes in the distributions of vegetational zones (2, 5, 6).

Patterns implied by cladograms and phyletic trees (Fig. 2) were documented by treating geography as an ordered cladistic character in which regions correspond to character states. Two main regions were recognized: eastern and southern Africa. A third region, “the Corridor,” represented by the Malawi section of the Rift Valley, also was defined because at least one hominid specimen has been reported from there (16) and because the Corridor is likely to have served as a faunal conduit between the other two regions (2). Although mammal
dispersal event. The geographic distribution of hominid speciation events also was recorded to determine whether those speciations tended to be focused in any particular region.

Note that the formal methods of cladistic biogeography, in which biogeographic patterns are tested by examining area cladograms (8), cannot be applied here because data are available from too few regions. Cladistic biogeography requires faunal information from at least three areas, and although three regions are defined here, only eastern and southern Africa have been well sampled. Faunal representation from the Corridor is limited (21).

Taxa examined here comprise the Plio-Pleistocene hominid species known from between 3.5 and 1.5 Myr that have been included in phylogenetic reconstructions (Fig. 2). *Ardipithecus ramidus* (22) and *Australopithecus anamensis* (23) are likely to be near the ancestry of later hominids and were used as an outgroup to determine the polarity of the “geography” cladistic character. In other words, it was assumed that eastern Africa was the region occupied by the last common ancestor of the hominids considered here. Note that this does not preclude the possibility that the last common ancestor of chimpanzees and humans arose outside of eastern Africa. Certainly, the hominid fossils from Chad (20) indicate that there must have been at least one hominid dispersal between eastern and central Africa.

Some of the phylogenetic analyses (14, 24–26) did not investigate phylogenetic relationships within the genus *Homo*. For those trees, all possible relationships within the genus were examined, with the exception that *H. ergaster* was never assumed to be directly ancestral to *H. rudolfensis* or *H. habilis*. Moreover, in the case of biogeographic patterns that are equally parsimonious, it was assumed for the sake of simplicity that species originated in either eastern or southern Africa. This is consistent with previous investigations of early hominid biogeography, and, although there are no strong grounds on which one can justify the assumption, its utilitarian value is that it greatly reduces the number of possible biogeographic patterns implied by some of the phylogenies.

Biogeographic patterns implied by phyletic trees were complicated by the fact that two hominid species are known from more than one region [i.e., *H. habilis* (eastern and southern Africa) and *H. rudolfensis* (eastern Africa and the Corridor)]. The regions in which these species first occur are unclear because of ambiguities concerning the age and taxonomic affinities of critical specimens (16, 17, 19, 27, 28). Accordingly, each phyletic tree was examined four times, taking into account all possible combinations of the centers of origin of those species.

**RESULTS**

Biogeographic patterns implied by early hominid phylogenies are summarized in Table 1. It is evident that published phylogenies do not all agree on a single biogeographic pattern. However, as researchers make decisions concerning fundamental questions about phylogeny, taxonomy, and chronology, then biogeographic patterns become clearer. For example, consider the phylogenetic hypotheses based on our own work (Fig. 2b, c, and e). Of these, Wood (29) does not consider the earliest putative *Homo* specimens from southern Africa as belonging to *H. habilis*. Consequently, only one biogeographic pattern is consistent with a strict reading of his phylogenetic hypothesis (Fig. 1b). In contrast, Strait et al. (30) included the southern African specimens within *H. habilis*, meaning that the biogeographic implications of those phylogenies must accommodate uncertainty as to the center of origin of that species. If *H. habilis* did not arrive in southern Africa until 1.8 Myr, then the resulting biogeographic patterns are very similar or identical to those implied by Wood’s (29) phylogeny (Fig. 1b).

If, on the other hand, *H. habilis* originated in southern Africa,
then more complicated biogeographic patterns are implied (Table 1).

Ultimately, the choice of biogeographic pattern depends on the cladogram or phyletic tree that is preferred. However, despite the fact that early hominid phylogenies differ markedly in the details of their branching patterns, certain generalizations about hominid biogeography emerge from this study (Table 1). All of the cladograms and phyletic trees agree that *Australopithecus africanus*, or its ancestor [a category that might include recently discovered fossils from Sterkfontein’s Member 2 (31)], dispersed from eastern to southern Africa between 3.5 and 3.0 Myr. All of the cladograms imply four episodes of southward dispersal. The phyletic trees all indicate that *H. rudolfensis* dispersed between eastern Africa and the Malawi Rift around 2.4 Myr, but the direction of the dispersal is unclear. Moreover, *H. habilis* dispersed either northward at ~2.3 Myr or southward at ~1.8 Myr. In addition, all of the cladograms and many of the phyletic trees indicate that *P.*
**DISCUSSION**

In contrast to the two, or three, dispersals identified by previous investigations, the trees examined in the present study require between four and seven. This suggests that patterns of early hominid biogeography were more complicated than those indicated by earlier studies. Moreover, these patterns allow one to test whether hominid dispersals conformed to the mammalian patterns identified by Turner and Wood (1). Such a test depends critically on the taxonomy and chronology of those indicated by earlier studies. Moreover, these patterns require between four and seven. This suggests that patterns of mammalian dispersal, then it is probable that one or both of them possessed behavioral or anatomical adaptations that allowed them to do so. This possibility needs to be tested against the paleontological and archaeological records. A departure from mammalian trends would further suggest that biogeographic models based on ecological hypotheses, such as Vrba’s (5) Habitat Theory, have important and interesting exceptions.

All cladograms and phyloetic trees identify eastern Africa as the region responsible for producing most (if not all) hominid species, either through speciation events within the region or because some eastern African populations dispersed to southern Africa and differentiated into regional species. Although taphonomic bias may contribute to this pattern, it nonetheless begs the question of what were the evolutionary mechanisms responsible for the frequent speciation of hominids in eastern Africa. Although several reconstructions of early hominid habitats have attempted to explain hominid speciation patterns (2–4, 32, 33), these studies either have not considered fully the implications of early hominid phylogeny or have made assumptions about biogeography that are inconsistent with the patterns found here.

**CONCLUSION**

Phylogeny has significant implications for interpretations of early hominid biogeography. In particular, hominid dispersals between eastern and southern Africa appear to have been

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Table 1.  Dispersal and speciation events implied by early hominid phylogenies

<table>
<thead>
<tr>
<th>Tree type</th>
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<th>Dispersal with speciation</th>
<th>Dispersal without speciation</th>
<th>Speciation without dispersal</th>
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<td>1, 4</td>
<td>12 or 14, 13</td>
<td>A, C, E, F, G, H</td>
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<td>A, C, E, F, G, H</td>
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<td>Phyletic tree*</td>
<td>Fig. 2d</td>
<td>1, 3, 4, 6</td>
<td>14, 16 or 13</td>
<td>A, C, E, F, G, H</td>
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<td></td>
<td>Fig. 2e</td>
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<td>14, 16 or 13</td>
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<td>14, 16 or 13</td>
<td>A, C, E, F, G, H</td>
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<td>A, C, E, F, G, H</td>
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<td>Fig. 2h</td>
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<td>Fig. 2i</td>
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<td>14, 16 or 13</td>
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</table>

Italicized numbers represent southward dispersals. Bold-faced numbers represent northward dispersals. Underlined numbers represent dispersals that occur only if *H. rudolfensis* appeared first in the Malawi Rift. Letters represent speciation events that are not associated with dispersals. “East” and “South” refer to regions in Africa.

* Assumes that *H. habilis* appears first in eastern Africa.
† Assumes that *H. habilis* appears first in southern Africa.
‡ Dispersals: 1. *Africanus* or its ancestor disperses from East to South between 3.5 and 3.0 Myr; 2. *H. habilis* or its ancestor, East to South, ~2.5 Myr; 3. *H. rudolfensis* or its ancestor, East to the Corridor, ~2.4 Myr; 4. *P. robustus* or its ancestor, East to South between 2.5 and 1.7 Myr; 5. *H. habilis* or its ancestor, the Corridor to South, ~2.5 Myr; 6. the common ancestor of *Homo* and *Paranthropus*, South to East, ~2.7 Myr; 7. the common ancestor of *Paranthropus*, South to East, ~2.6 Myr; 8. *P. aethiopicus* or its ancestor, South to East, ~2.5 Myr; 9. *H. rudolfensis* or its ancestor, South to the Corridor, ~2.4 Myr, 10. *P. boisei* or its ancestor, South to East, ~2.2 Myr; 11. *H. ergaster* or its ancestor, the Corridor to East, ~1.9 Myr; 12. *H. habilis* East to South, ~2.5 Myr; 13. *H. rudolfensis* East to the Corridor, ~2.4 Myr; 14. *H. habilis*, East to South, ~1.8 Myr; 15. *H. habilis*, South to East, ~2.3 Myr; 16. *H. rudolfensis*, the Corridor to East, ~2.4 Myr; 17. the common ancestor of *H. rudolfensis* and *H. ergaster*, South to East, ~2.5 Myr.

roxotus, or its ancestor, dispersed from eastern to southern Africa between 2.5 and 1.7 Myr [depending on the ages of the fossils at Drimolen and Kromdraai (6, 13, 15, 18)].

The same could be said for *P. robustus*, or its ancestor, if Drimolen is older than Swartkrans I or if Kromdraai B dates to ~2.0 Myr (13). If these species departed from continental patterns of mammalian dispersal, then it is probable that one or both of them possessed behavioral or anatomical adaptations that allowed them to do so. This possibility needs to be tested against the paleontological and archaeological records. A departure from mammalian trends would further suggest that biogeographic models based on ecological hypotheses, such as Vrba’s (5) Habitat Theory, have important and interesting exceptions.

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more frequent that has previously been thought, and some of those dispersals may have opposed prevailing mammalian trends. If the latter is true, then attention should be paid to determining whether such dispersals were facilitated by the evolution of anatomical or behavioral adaptations in those hominid species. Most phylogenies also indicate that eastern Africa hosted a majority of hominid speciation events. An integration of ecological, phylogenetic, and archaeological data, as well as any relevant evidence about local habitats within regions, should lead to a more complete understanding of hominid dispersal and speciation patterns.

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