## **Evidence for abrupt speciation in a classic case of gradual evolution**

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**In contrast with speciation in terrestrial organisms, marine plankton frequently display gradual morphological change without lineage division (e.g., phyletic gradualism or gradual evolution), which has raised the possibility that a different mode of evolution dominates within pelagic environments. Here, we reexamine a classic case of putative gradual evolution within the** *Globorotalia plesiotumida–G. tumida* **lineage of planktonic foraminifera, and find both compelling evidence for the existence of a third cryptic species during the speciation event and the abrupt evolution of the descendant** *G. tumida***. The third morphotype, not recognized in previous analyses, differs in shape and coiling direction from its ancestor,** *G. plesiotumida***. This species dominates the globorotaliid population for 414,000 years just before the appearance of** *G. tumida***. The first population of the descendant,** *G. tumida***, evolves abruptly within a 44,000-year interval. A combination of morphological data and biostratigraphic evidence suggests that** *G. tumida* **evolved by cladogenesis. Our findings provide an unexpected twist on one of the best-documented cases of within-lineage phyletic gradualism and, in doing so, revisit the limitations and promise of the study of speciation in the fossil record.**

cladogenesis | evolutionary dynamics | foraminifera | fossil record | plankton

**T**he fossil record in marine plankton is characterized by gradual morphological change both with and without apparent cladogenesis (1–10). Phyletic gradualism has been attributed to a lack of barriers to gene flow in species that are both cosmopolitan and phenotypically plastic (11–13). However, a growing number of phylogenetic studies have revealed the presence of multiple cryptic species within named marine morphospecies (14–16). In some cases, within-species morphological clines have subsequently been found to consist of numerous genetically, biogeographically, and ecologically distinct species (17, 18). The presence of cryptic species complexes in the modern ocean suggests a fossil record laden with hidden cladogenetic events (19) potentially affecting the perception and interpretation of evolutionary patterns.

The existence of cryptic species complexes, and the consequent discrepancy between morphological and genetic species, is of general concern because open ocean microfossils provide one of the best records (temporally and spatially) of the last 130 million years of life (20–22). For instance, planktonic foraminifera have been used in global studies of the determinants of species richness (23), body size (24, 25), and speciation (26). For all the utility of open ocean microfossils, there have been relatively few coordinated studies of both the morphologic and genetic similarity of individuals in the modern ocean (although see refs. 17, 18, 27, and 28). There is also evidence that open ocean microfossils may not actually conform to the morphological species concept. A high-resolution study of the *Globorotalia* lineage of planktonic foraminifera in the Early to Middle Miocene failed to find evidence of discrete, nonoverlapping morphological clouds as is expected in the typical morphological species concept (29).

The morphological similarity of foraminiferal species has implications for the detection of cladogenesis. Most past studies have either assumed a priori that cladogenesis occurred (7–9, 30) or, if not, have interpreted evolutionary trends as cases of within-lineage evolution after considering trait distributions (1, 5). Trait variation within a given species is typically normally distributed and, in theory, deviations from normality should occur when two or more species coexist. In practice, this normality test for cladogenesis has little statistical power when two morphologically similar species with high trait variability coexist (as described in ref. 29) and are sampled at the sample sizes typical of past studies (31). Sample variance can also be used to detect the presence of multiple taxa with some of the same statistical limitations (4). The lack of clear correspondences between named morphospecies and discrete morphological clusters increases the difficulty of detecting cladogenesis in fossil planktonic foraminifera. In one instance, reproductive isolation and, putatively, speciation was found to be uncoupled from morphological evolution in the *Fohsella* lineage of globorotaliid foraminifera (32).

Here we test for speciation in the fossil record within a lineage of planktonic foraminifera. In this study, we use the term *speciation* to refer to cladogenetic events (e.g., phyletic splitting), and not within-lineage evolution. We reexamine a classic case of putative gradual evolution in which the ancestor *Globorotalia plesiotumida* is thought to have evolved over  $\approx$  500,000 years into the descendant *G. tumida* (1, 33). Both before and after the morphological transition there are several million years of morphological stasis suggesting that this case represents a hybrid of phyletic gradualism and morphological stasis dubbed ''punctuated gradualism'' (33). The *G. tumida* lineage has often been reexamined in studies of evolutionary mode (34–40), due to the compelling results and data availability of Malmgren et al.'s original study (33). In readdressing this widely cited case of within-lineage gradual evolution, we consider the effect of methodology on our perception of evolutionary trends, test for the possibility of phyletic splitting and within-lineage change, and reconsider the morphological species concept in planktonic foraminifera.

## **Results and Discussion**

We analyze morphological change in the *Globorotalia plesiotumida–G. tumida* evolutionary series in a deep-sea sediment core record from the western tropical Pacific [Ocean Drilling Program (ODP) site 806B, Ontong Java Plateau, 0°19.11'N, 159°21.69′E, 2,520 m water depth]. Using eigenshape analysis (a morphometric technique for comparing outlines) (41, 42) and an updated time scale (supporting information (SI) [Fig. S1\)](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=SF1), we

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**Fig. 1.** Morphological trends over time in the *Globorotalia plesiotumida–G. tumida* lineage in the western equatorial Pacific. Mean morphology as a function of time expressed as (*A*) an eigenshape amplitude (EA 3) from eigenshape analysis (all individuals), (*B*) a relative warp score (RW 2) from semilandmark TPS analysis (all individuals analyzed, sinistrally coiled individuals plotted), (*C*) centroid size (all individuals), and (*D*) a relative warp score (RW 1) from semilandmark TPS analysis (all individuals). Error bars in (*A–D*) are parametric 95% CI and mean values are fit with a loess curve. (*E*) Percent sinistral coiling individuals and (*F*) the standard deviation of RW 1 as a function of time. Boxed values in (*F*) indicate samples with 20% or greater overlap in sinistral and dextral coiled individuals. Gray points in (*B*) indicate samples containing three or fewer individuals.

obtained a similar pattern of morphological evolution in the western tropical Pacific as was found in an earlier study in the central Indian Ocean (Fig. 1*A* and [Fig. S2\)](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=SF2). The similarity of both results, despite using materials from widely separated sites, suggests that the evolutionary transition is synchronous across a large stretch of the ocean. Indeed, *G. tumida* displays a nearsimultaneous first appearance throughout the tropical Indo-Pacific with a geographic range overlapping that of *G. plesiotumida* at the Miocene/Pliocene boundary (refs. 43–45; see also *[SI](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=STXT) [Materials and Methods](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=STXT)* and [Fig. S3\)](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=SF3). Our study design was therefore predicated on the hypothesis that the morphological evolution of *G. tumida* in the western tropical Pacific occurred in situ.

Our analysis departs from previous analyses by controlling for the effects of shell size on morphology, and by using a different morphometric method. In Malmgren et al.'s study (33), a 3-fold increase in mean size accompanies the morphological transition in the *G. tumida–G. plesiotumida* lineage and is strongly correlated with mean morphology along eigenaxis 2 (EA 2, Pearson's  $r = 0.89$  and  $P < 0.001$ ). Here we explicitly control for the effect of size on the perceived evolutionary trend by sampling individuals in a narrower size range (250– 500  $\mu$ m in contrast to the ~150-500  $\mu$ m originally used). Given this greater control for size, the correlation between size and EA 3 is weak (Pearson's  $r = 0.23$  and  $P < 0.001$ ). However, we still note a  $\approx$  1.4-fold increase in mean centroid size in the equatorial Pacific (Fig. 1*C*).

In an additional departure, we use a second morphometric technique for analyzing outlines, semilandmark thin-plate spline analysis (semilandmark TPS) (46). Methodological aspects of semilandmark TPS techniques suggest advantages of this approach over eigenshape analysis (*[SI Materials and](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=STXT) [Methods](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=STXT)*). The second relative warp (RW 2, a morphological eigenaxis capturing 20% of morphological variance) is correlated to EA 3 from eigenshape analysis (Pearson's  $r = 0.70$  and  $P < 0.001$  for sinistrally coiled individuals; see also [Fig. S4\)](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=SF4) and reveals a comparable shift in mean morphology (Fig. 1*B*). However, our findings also show critical differences between the two methods.

Surprisingly, the first relative warp (RW 1) of the semilandmark TPS analysis indicates that some individuals present during the transition are morphologically more distinct than *Globorotalia plesiotumida* and *G. tumida* are from each other (Fig. 1*D*; RW 1 captures 53% of morphological variance). These divergent individuals are distinctly flattened (see Fig. 3*C Middle* and [Movie S1\)](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=SV1) and are easily differentiated by eye. Furthermore, the flattened morphotype is almost exclusively dextrally coiled (clockwise chamber addition from the spiral perspective) and rarely coexists with sinistral *G. plesiotumida* (Figs. 2 and 3*A*). Grouping individuals by coiling direction reveals a significant difference in the RW 1 scores of sinistral and dextral coiled individuals (Fig. 2;  $t$  test,  $P < 0.001$ ). A few dextral individuals occur in our oldest sample at 6.403 Ma, but these are more *G. plesiotumida*-like in their outline morphology than any subsequent dextral population. Dextral morphotypes display the typical elongated final chamber often used as a defining characteristic of *G. plesiotumida*. Where dextral and sinistral individuals do co-occur (20% or greater overlap), the populations exhibit higher morphological variance along RW 1 (Fig. 1*F*) than is the case for populations dominated by one coiling morphology. Additionally, we find significant differences in RW 1 scores between coiling groups in each of the seven time periods containing at least three individuals per coiling direction ( $t$  test,  $P < 0.001$ ). A maximum likelihood analysis of mixture models provides statistical support for the interpretation of two coexisting morphotypes rather than a single morphotype in six time periods (Fig. 3*A*; stars indicate significance at  $0.05$  and squares at  $0.06$  significance level; details in *Materials and Methods* and ref. 47).



**Fig. 2.** Distribution of relative warp 1 scores for dextrally and sinistrally coiled individuals. Each histogram is scaled to a unit area. Sinistral and dextral distributions are significantly different (*t* test,  $P < 0.001$ ).

The correspondence between coiling direction and morphology suggests that the flattened, dextral morphotype represents a species fully differentiated from the sinistral *G. plesiotumida.* Our evidence for a cryptic species in the *G. plesiotumida* complex is consistent with evidence in other groups of foraminifera that coiling direction is a heritable trait (48, 49). In two modern planktonic foraminiferal species, *Neogloboquadrina pachyderma* and *Globorotalia truncatulinoides*, coiling direction was found to be indicative of cryptic species (17, 28).



**Fig. 3.** Morphological change in the *Globorotalia plesiotumida–G. tumida* lineage. (*A*) Morphology viewed as histograms along relative warp 1, with sinistrally coiled individuals in black and dextrally coiled individuals in red, and (*B*) along relative warp 1 as calculated for sinistrally coiled individuals. Stars indicate two overlapping populations supported at a 0.05 significance level and squares at a 0.06 significance level. (*C*) Scanning electron microscopy (umbilical view) and digitized outlines (edge view) of sinistrally coiled *Globorotalia plesiotumida* and *G. tumida* (top two and bottom two panels) and the dextral morphospecies (*Middle*). Individuals panels of morphology with time along RW 1 and RW 2 are shown in [Movie](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=SV2) [S2.](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=SV2)

The dextral, flattened morphotype becomes abruptly dominant in our record at 6.225 Ma, and persists for 414,000 years until 5.819 Ma. During this period, which overlaps the start of the gradual evolution of *G. tumida* in eigenshape analyses, the dextrally coiled morphogroup oscillates in dominance with the sinistrally coiled morphogroup. In many oscillations, the abundance of the rare morphotype (sinistral or dextral) is zero. These recorded absences may coincide with time periods during which a given morphotype is globally rare and therefore unsampled. Alternatively, the absences may indicate periods of changing biogeographic distributions, with rare morphotypes found in abundance in other locations. Based on our sampling scheme, we cannot determine which of these two alternatives is more likely.

After the period of oscillating dominance, rare dextrally coiled individuals are present for an additional 444,000 years with a second peak in dextral abundance at 5.503 Ma. Together, these data suggest that the flattened, dextral taxon evolved by cladogenesis and coexisted with its ancestor for at least 850,000 years. If the global first appearance of the dextral species is recorded in our sample set at 6.256 Ma, then its evolution was very rapid, occurring within a 26,000-year window. However, we caution that it is possible that the first appearance of the flattened dextral taxon could reflect immigration or have occurred at or before 6.403 Ma in the equatorial Pacific. Dextral individuals are recorded in the Indian Ocean before 6.4 Ma (33), but whether these have the flattened morphology of the cryptic taxon described here is yet to be determined.

A second cladogenetic event may accompany the first appearance of fully differentiated *Globorotalia tumida* between 5.865 and 5.819 Ma. In this case, both the ancestor, *G*. *plesiotumida*, and the descendant, *G. tumida*, are sinistrally coiled and therefore differentiated entirely upon a change in outline morphology (Fig. 3*B*). Cladogenesis is indicated by several factors, including (*i*) the abrupt shift in mean morphology toward *G. tumida* within a 44,000-year span (mean shift  $>1.5$ ) SD), (*ii*) the co-occurrence of pre- and postshift morphologies at 5.819 Ma, a period of elevated population variance (one of the three highest observed), (*iii*) the observation of several reversals in the population morphology toward the *G. plesiotumida* type, and (*iv*) periods of elevated variance between 5.819 and  $\approx$  5.5 Ma. Finally, a short interval of co-occurrence (several 100 kya) between *G. plesiotumida* and *G. tumida* is indicated in Indian Ocean records (50). In the Atlantic, *G. plesiotumida* persists well into the middle Pliocene (51, 52).

Throughout the entire time series, maximum likelihood analysis of mixture models provides support for only one sinistral population per time interval (Fig.  $3B$ ; RW  $1_{\text{sinistral}}$ ) results shown, approximately equivalent to RW 2<sub>all</sub> as explained in *Materials and Methods* and see [Fig. S5\)](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=SF5), in possible agreement with Malmgren et al.'s (33) interpretation of gradual, within-lineage evolution. However, the high variance in morphology within some samples (particularly between 5.5 and 6.0 Ma) suggests that there could be two coexisting species at several points in the time series. Unfortunately, our maximum likelihood analysis has very low power (0.01–0.48) to detect overlapping populations if they exist, due in part to our small sample size. We conclude that we cannot unambiguously determine whether *G. plesiotumida* co-occurs in the same samples with *G. tumida* during the transition period from approximately 6 –5.5 Ma. Both larger sample sizes and more informative traits could help resolve this ambiguity in future studies.

Our conclusions differ from those of Malmgren et al. (1, 33) in two critical ways. First, our methods clearly identify the ecological dominance of a flattened, dextral cryptic species just before the appearance of *G. tumida*, which is inferred to have arisen cladogenetically from a *G. plesiotumida* ancestor. Malmgren et al. (33) noted the presence of dextral, biconvex forms during the transitional period like the compressed, dextrally coiled individuals we see in the western equatorial Pacific. However, they did not find morphometric evidence that would distinguish these dextral forms as species in their own right. Second, previous work has not detected as rapid an appearance of fully formed *G. tumida* as we observe here at 5.81 Ma. Malmgren et al. (33) did observe a step in both shell size and morphology about this time, but they attributed these morphological shifts to oscillations within a longer evolutionary trend.

Our findings show that morphometric techniques and measurement strategy are key to the interpretation of evolution in the fossil record. It is disturbing that eigenshape analysis (used by previous authors) fails to detect differences in compression of individuals in the equatorial Pacific, where both semilandmark TPS analysis and visual inspection confirm the distinct morphology of dextrally coiled individuals. We hypothesize that eigenshape analysis cannot detect differences in shell compression that are not accompanied by major changes in the angles between many points (see *[SI Materials and Methods](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=STXT)*). Malmgren et al. (33) also ruled out the possibility of cladogenesis due to the apparent lack of bimodality within populations. This test is less powerful than the mixture models used here, because distributions never appear bimodal along RW 1 even when multiple morphospecies coexist (Fig. 3*A*). Finally, both our outline data and those collected by Malmgren et al. (33) fail to capture essential aspects of shell morphology, such as the shape of the final chamber, that are used by taxonomists to differentiate *Globorotalia plesiotumida* and *G. tumida*. Without examining a larger subset of informative characters, it is difficult to assess whether the perceived mode of evolution and the applicability of the morphological species concept is an artifact of the morphological traits under consideration. Consequently, this and other morphometric studies may lack adequate information to clearly differentiate speciation from within lineage evolution.

A synthesis of past studies of cladogenesis has documented a common pattern of sympatric speciation in open ocean taxa (6), with the gradual divergence of the mean daughter morphotype from a stable ancestral morphotype for  $\approx$  500,000 years following speciation in foraminifera (8, 9), radiolarians (30, 53), and diatoms (7). In contrast to instances of sympatric speciation, morphological change associated with allopatric speciation in marine microfossils can be rapid (10 kya) (54).

Our findings add to these observations by showing that, at least in this classic case, the evolution of *G. tumida* from *G. plesiotumida* was not a simple affair. Evolution involved both the cryptic evolution of a dextral, compressed morphotype and the rapid evolution of *G. tumida*, in both cases likely by cladogenesis. The dramatic decrease in abundance of the flattened dextral forms coincides with the abrupt appearance of *G. tumida*. It is clear that the first *G. tumida* appeared very rapidly, in less than 45,000 years, rather than the hundreds of thousands of years inferred from previous work. Furthermore, stratigraphic analysis from other sites has shown that *G. plesiotumida* persists after the evolution of *G. tumida*. Changes in the dominance of coexisting ancestral and descendant morphotypes occur several times and may be due to an oscillation in the environment. Our work underscores the inherent difficulty of inferring evolutionary mechanisms from fossils. At the same time, our study highlights observations unique to the fossil record, including the measurement of evolutionary patterns, species coexistence, and changing population variance through time.

## **Materials and Methods**

**Sample Preparation.** Within each time interval, we sampled the first 30 individuals encountered from the *Globorotalia plesiotumida–tumida* lineage in the  $>$ 250- $\mu$ m size fraction from ODP site 806B on the Ontong Java Plateau (1,140 individuals from 38 depth intervals; [Dataset S1\)](http://www.pnas.org/content/vol0/issue2009/images/data/0902887106/DCSupplemental/SD1.txt). An age model was calculated for the Ontong Java Plateau using 10 biostragraphic markers (55) and assuming constant sedimentation rates between markers [\(Fig. S1;](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=SF1) age model for Atlantic site 959C also shown). The two exceptions to the sampling protocol of 30 individuals occurred in samples at 163.27 and 194.77-meter composite depth (mcd) where 31 and 29 individuals were analyzed, respectively. Individuals were cleaned, taped to glass coverslips, and mounted on a universal stage for the digitization of the edge view using a video capture system. A digitized 2D outline of each individual (100 coordinate points, approximately evenly spaced around the foraminiferal edge view) was initialized at the proloculous for morphometric analyses (41, 42) [\(Dataset S1\)](http://www.pnas.org/content/vol0/issue2009/images/data/0902887106/DCSupplemental/SD1.txt).

**Morphometric Methods.** Semilandmark thin-plate spline analysis is a landmark morphometric technique adapted to assessing similarity among outlines by ignoring differences that arise from the location of coordinates along an outline (46). After an initial consensus form (or mean shape) is calculated, points are allowed to slide along individual outlines to minimize the difference between individual shapes and the consensus form (56, 57). If the semilandmark analysis is recursive, then the postsliding location of points for each outline is used in the next iteration. If the semilandmark analysis is not recursive, then the consensus form is updated for each iteration, but individual outline points always start in same initial location. Eigenshape analysis (41, 42) was also used to compare results between the Pacific and Indian Ocean (see *[SI Materials and Methods](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=STXT)* for details on this method). Note, the signs of eigenaxes and relative warps can vary between analyses. Signs were reversed along eigenaxes and relative warps when necessary to conform to the orientation of Malmgren et al. (33). For instance, signs were reversed for RW 1<sub>all</sub> and RW 2<sub>all</sub> in Figs. 1-3 and related statistics.

Although theoretically preferable, recursive semilandmark TPS was computationally prohibitive given the large number of outlines (1,140) and sliding points (100 per outline) in this study. We assessed the effect of using recursive versus nonrecursive TPS by analyzing a representative subset of individuals with both techniques (304 total outlines, including eight randomly chosen individuals per time interval). Recursive and nonrecursive semilandmark results were highly correlated along the first three relative warps (Pearson's  $r > 0.91$ ,  $P < 0.001$ ), accounting altogether for 73% and 82% of morphological variance, respectively [\(Table S1\)](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=ST1). Judging from this subset of outlines, and specifically comparing the relative assignments of dextral and sinistral individuals, the effect of using nonrecursive semilandmark analysis on our interpretation of the large-scale RW 1 displacements is minimal. This was potentially not the case for the subtle trends in sinistrally coiled individuals along relative warp 2. Therefore, sinistral individuals (719 total) were separately analyzed using a recursive semilandmark analysis (three iterations). The first relative warp from sinistral semilandmark TPS analysis was strongly correlated with the second relative warp from the full analysis ( $r^2$  = 0.94,  $P$  < 0.001), and was used in Fig. 3 and in all statistical considerations of the evolution of *G. tumida* from *G. plesiotumida*.

As a preprocessing step to semilandmark TPS analysis, we first performed a Generalized Procrustes alignment (involving translation, rotation, and scaling) to minimize the sum-squared distance between outline points and a consensus shape using the function procGPA in the package ''shapes'' (version 1.0 – 8) in R (version 2.2.1) (58). This particular function allows shapes to be mirrored, an option that we used to minimize the apparent shape differences between left- and right-coiled individuals (shape outlines and mirroring results visualized in [Movie S1\)](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=SV1).

Here we use relative warps (RWs) to described main patterns of shape change in the *G. plesiotumida–G. tumida* lineage. We used the program tpsRelw (version 1.46, created by F. James Rohlf) for all semilandmark analyses, including the calculation of relative warps. In tpsRelw, we used orthogonal projections and scaling by the cos (rho) for Procrustes alignments; results were unaffected by scaling by a unit centroid size  $(r^2 >$ 0.9999 for the first 10 relative warps in TPS analysis). Uniform weighting of all partial warps ( $\alpha = 0$ ) was used to include uniform components in the partial warp scores matrix. Uniform components of shape change were estimated as the complement to the nonuniform shape variation (59). Finally, due to computational limitations and the similarity of TPS and semilandmark TPS results [\(Table S1\)](http://www.pnas.org/cgi/data/0902887106/DCSupplemental/Supplemental_PDF#nameddest=ST1), semilandmark TPS analysis was run for a maximum of three sliding iterations.

**Maximum Likelihood Analysis of Mixture Models.** A maximum likelihood analysis of mixture models was used to assess the number of distinct populations along RW 1<sub>all</sub> and RW 1<sub>sinstral</sub>. The maximum likelihood framework allowed us to test the relative support for one or more overlapping populations within a single time period and morphological distribution (specifically, the histograms in Fig. 3). All analyses were performed using the program Mixture Model Analysis (version 1.32, created by G. Hunt) and a previously described approach (47).

In brief, we calculated the likelihood of one to two populations for each time interval along RW 1<sub>all</sub> and RW 1<sub>sinistral</sub> (roughly equivalent to RW 2<sub>all</sub>) using 200 random initiations, and assuming equal population variance and a normal distribution. A bootstrap approach was used to determine the relative support for one or more distributions, as increasing parameters generally improves model fit (e.g., the log-likelihood ratio will favor the model with more populations). To compare the relative support for one versus two overlapping populations, we generated 1,000 sample distributions based on the mean and variance calculated for a single population and compared these maximum likelihood estimates with that determined

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empirically for two overlapping populations. If the two-population loglikelihood ratio was greater than 95% of those generated from a single population ( $\alpha$  = 0.05), then we considered two overlapping populations more likely than a single population in a given time period. In interpreting negative results (e.g., the failure to reject a single population), we assessed the power of the maximum likelihood test using a second bootstrapped approach (log-likelihood ratio obtained from alpha in the first bootstrap test and assumed population parameters for two populations to generate distributions; see details in ref. 47).

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