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# ARTÍCULO DE INVESTIGACIÓN/RESEARCH ARTICLE

# PLANKTIC FORAMINIFERAL DIVERSITY: LOGISTIC GROWTH OVERPRINTED BY A VARYING ENVIRONMENT

# Diversidad en foraminíferos planctónicos: crecimiento logístico sobrepuesto por un ambiente variable

# Andrés L. CÁRDENAS-ROZO<sup>1</sup>; Peter J. HARRIES<sup>2</sup>.

<sup>1</sup> Ciencias de la Tierra, Escuela de Ciencias, Universidad EAFIT. Cra. 49 nº. 7 Sur-50. Medellín, Colombia.

<sup>2</sup> Marine, Earth, and Atmospheric Sciences Faculty, Collegue of Sciences, North Carolina State University. 2800 Faucette Drive, 1125 Jordan Hall, Campus Box 8208. Raleigh, North Carolina, USA.

For correspondence. acarde17@eafit.edu.co

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# ABSTRACT

This study statistically assesses the relationship between the planktic foraminiferal long-term diversity pattern (~170 Ma to Recent) and four major paleobiological diversification models: (i) the 'Red Queen' (Van Valen, 1973; Raup *et al.*, 1973), (ii) the turnover-pulse (Vrba, 1985; Brett and Baird, 1995), (iii) the diversity-equilibrium (Sepkoski, 1978; Rosenzweig, 1995), and (iv) the 'complicated logistic growth' (Alroy, 2010a). Our results suggest that the long-term standing diversity pattern and the interplay between origination and extinction rates displayed by this group do not correspond to the first three models, but can be more readily explained by the fourth scenario. Consequently, these patterns are likely controlled by a combination of planktic foraminiferal interspecific competition as well as various environmental changes such as marine global temperatures that could impacted the niches within the upper mixed layer within the oceans. Moreover, as other global long-term patterns have been interpreted as reflecting 'complicated logistic growth', this study further suggests that the interplay between abiotic and biotic factors are fundamental elements influencing the evolutionary processes over the extensive history of the biota.

**Keywords:** abiotic and biotic controls, complicated logistic growth, diversity dynamics, macroevolution, planktic foraminifera, paleobiology.

# RESUMEN

Este estudio evalúa estadísticamente la relación entre el patrón de diversidad global de los foraminíferos planctónicos en el largo plazo (~170 Ma al Reciente) y los cuatro modelos de diversificación propuestos desde la rama de la paleobiología: (i) "Reina Roja" (Van Valen, 1973; Raup *et al.*, 1973), (ii) remplazo pausado (Vrba, 1985; Brett y Baird, 1995), (iii) diversidad en equilibrio (Sepkoski, 1978; Rosenzweig, 1995), y (iv) el "crecimiento logístico complicado" (Alroy, 2010a). Nuestros resultados sugieren que la forma de este patrón global de diversidad y la inter-relación entre las tasas de extinción y originación de este grupo no corresponden con los primeros tres modelos anteriormente citados. Sin embargo, estos pueden ser explicados bajo el cuarto escenario. Consecuentemente, las dinámicas de diversidad (i.e. patrón de diversidad y tasas de extinción y originación) de este grupo posiblemente son controladas por la combinación de la competencia interespecífica de los foraminíferos planctónicos y varios cambios ambientales tales como temperaturas globales marinas que pudieron impactar el número de nichos dentro de la capa superior de los océanos. Además, otros patrones globales de diversidad en el largo plazo han sido interpretados como el reflejo del modelo de crecimiento logístico complicado, lo que sugiere que la relación entre factores abióticos y bióticos tiene un carácter fundamental en los procesos evolutivos que han sucedido a lo largo de la historia de la vida.

**Palabras clave**: crecimiento logístico complicado, controles abióticos y bióticos, dinámicas de diversidad, foraminíferos planctónicos, macroevolución, paleobiología.



#### INTRODUCTION

Planktic foraminifers, a group of extant sexually reproducing protists, have been an extremely useful group for testing possible controls on long-term diversity patterns (e.g., Cifelli,1969; Frerichs, 1971; Lipps, 1979; Wei and Kennett, 1986; Schmidt et al., 2004; Ezard et al., 2011; Peters et al., 2013). In part, this is a reflection of their extremely wide biogeographic distribution and the fact that virtually continuous microfossil successions can be accessed through the study of outcrops and especially deep-sea cores dating back to the Bajocian (~170 Ma). Despite the interest in this group, only a few studies have examined possible controls on their long-term diversity dynamics using quantitative approaches (e.g., Wei and Kennett, 1986; Schmidt et al., 2004; Ezard et al., 2011), and these studies have been concentrated on specific time intervals rather than on the group's entire fossil history (e.g., Neogene [~23 Ma to Recent; Wei and Kennett, 1986] and Cenozoic [~65.5 Ma to Recent; Schmidt et al., 2004 and Ezard et al., 2011]). Moreover, even though comprehensive models have been developed to explain underlying long-term diversity dynamics, these likely need to combine 'abiotic' and 'biotic' elements rather to contrast them (Alroy, 2010a); to date only Ezard's et al., (2011) work has employed that approach in examining the role of changing oceanic temperatures and species' ecologies as controls on the Cenozoic diversity pattern.

Consequently, this study aims to augment and extend this approach by examining planktic foraminifer diversity dynamics across the group's entire fossil history (~170 to 0 Ma) and to analytically describe their long-term diversity pattern through the use of an integrated abiotic-biotic model. This approach involves the comparison of quantitative turnover predictions of each one of the four major paleobiological diversification models: (i) the 'Red Queen' scenario (Van Valen 1973; Raup et al., 1973); (ii) the turnover-pulse model (Vrba, 1985; Brett and Baird, 1995); (iii) the diversity-equilibrium model (Sepkoski, 1978; Rosenzweig, 1995); and (iv) the 'complicated logistic growth' scenario, where diversity limits are imposed by variable extrinsic factors (Alroy, 2010a). The fundamental basis of this approach is to employ time-series analyses of changes in global planktic foraminiferal diversity dynamics (i.e., the relationships between extinction and origination rates as well as between those rates and diversity), and, mirroring previous studies, testing between changes in both global long-term planktic foraminiferal diversity and mean global marine temperature (i.e., changes in the oceanic upper mixed layer as influenced by temperature variation, represents an important control on global long-term diversity standing pattern for the planktic foraminifers over the longterm; Peters et al., 2013).

#### MATERIALS AND METHODS

This study is based on a compilation of 699 planktonic foraminifera species' stratigraphic ranges from published

sources spanning the Middle Bajocian (~ 170 Ma) to Recent, calibrated to Gradstein *et al.*, (2012) geologic timescale, and binned into 1 Ma intervals (see raw data in Appendix; Cárdenas, 2012). These stratigraphic ranges were used to determine standing diversity, extinction and origination rates. Furthermore, these data were then analyzed using both the boundary-crossing (BC; Bambach, 1999) and per-capita (Foote, 2000) methodologies, respectively (Fig. 1). The BC



**Figure 1.** Time series used in the analyses. **a)** Per-interval middle Jurassic to Recent (~170 to 0 Ma) values of standing diversity for planktic foraminifers species. **b)** Per capita extinction rate of planktic foraminifers. For each curve, data were computed at 1 Ma intervals. Mixed-house (~170 to ~100 Ma) and icehouse (~37.5 to 0 Ma) modes are representing by a grey background, whereas the greenhouse interval (~100 to ~37.5 Ma) by a white one. Abbreviations: OAE 1a, Oceanic Anoxic Event 1a; OAE 1b, Oceanic Anoxic Event 1b; OAE 1c, Oceanic Anoxic Event 1c; OAE 1d, Oceanic Anoxic Event 3; K/Pg, Cretaceous/Paleogene extinction event; PETM, Palaeocene-Eocene Thermal Maximum; EECO, Early Eocene Climate Optimum; Oi-1, early Oligocene oxygen isotope event 1; MMCO, Mid-Miocene Climate Optimum; MPT, Mid-Pleistocene Transition.

approach was preferred in the computation of planktic foraminifers diversity curve as opposed to subsampling procedures (i.e., sampling in bin and shareholder quorum; Alroy, 2010b), primarily because these other methodologies require information related to individual taxonomic occurrences (i.e., abundances), and, even though these kind of data are available for planktic foraminifers in numerous biostratigraphic reports from the Deep Sea Drilling Project/Ocean Drilling Project, they are primarily from the Late Cretaceous to Recent interval. Therefore, an application of this procedure would result in the loss of 60 % of the planktic foraminfers' total temporal longevity (i.e., ~170 to ~70 Ma would be excluded). In the same way, given that the raw data for the computation of extinction and origination rates was based on first and last appearances of the taxa, the per-capita-rates technique for calculating extinction and origination rates (Foote, 2000) was selected over newer techniques, such as the 'three-timers' approach, as the latter necessitates a fossil-occurrence database for the calculations (Alroy, 2010b); such a database does not currently exist for the Foraminifera. Additionally, because not all taxa present during a given 1 Ma bin ranged throughout the entire interval, the total diversity of an interval likely overestimates the number of species at extinction risk at any given time. Per-capita rates were used as they address those sorts of biases by drawing a census of taxa at precise moments in time (i.e., the beginning and the end of the interval) (Foote, 2000; Foote and Miller, 2007). Finally, to avoid spurious correlations resulting from serial auto-correlation and 'noisy' secular trends (McKinney and Oyen, 1989), the datasets were analyzed using first-generalized differences before the use of the statistical correlation to compare turnover rates (i.e., extinction versus origination) both to each other as well as with the standing diversity pattern. Owing to the nonnormal character of those variables throughout the planktic foraminifers' fossil history (Appendix), Spearman's rankorder coefficient was used to test for correlation of changes in both the extinction and origination per-capita rates, as well as changes in species richness and in turnover rates (i.e., extinction and origination). Furthermore, we tested whether cross-correlations among planktic foraminiferal diversity dynamics time series (i.e., diversity pattern and extinction-origination per-capita rates) varied when binned into different global climatic mode intervals, comprising a 'mixed house' interval characterized by evidence for both warm (e.g., Littler et al., 2011) and cool temperatures (e.g., Frakes et al., 1992; Veizer et al., 2000) (~170 to ~100 Ma; Bajocian-Albian), a greenhouse phase (~100 to ~37.5 Ma; Cenomanian-late early Eocene; Miller et al., 2005), and an icehouse phase (~37.5 to 0 Ma; latest Eocene-Recent; Miller et al., 2005) (Fig. 1)). Finally, a Kolmogorov-Smirnov test for equality of distribution was employed to perform a statistical comparison of the complete distribution frequency shapes

of both 'turnover-pulse' model (Appendix) and planktic foraminiferal first and last occurrence data (Appendix).

## RESULTS

The computed global planktic foraminiferal diversity pattern showed that species richness remained very low initially and then increased ~30 Ma after the initial appearance of the clade (Fig. 1). Furthermore, the data also show that apart from major turnover events forced by significant global environmental events (i.e., at the Aptian-Albian, Cretaceous-Paleogene, and Eocene-Oligocene boundaries), it has 'lesser' troughs during the mid-Cretaceous, mid-Paleogene, and mid-Neogene. Moreover, the planktic foraminiferal diversity's general trend has not been neither constantly increasing nor reaching a saturation level (Fig. 1). However, extinction and origination per-capita rates through the clade's history have their highest peaks during the group's earliest evolutionary phase (~170 Ma to 145 Ma) and surrounding the Cretaceous-Paleogene boundary (Fig. 1). Furthermore, the extinction and origination per-capita rates had values of zero only during the first 30 Ma of the group's evolutionary history (Fig. 1, Appendix).

Spearman's rank-order correlation test applied to the first-generalized differences in both planktic foraminiferal extinction and origination per-capita rates throughout the fossil history of the group (170-0 Ma), reveal a significant positive association ( $\rho = 0.473$ ,  $p = 1.07*10^{-10}$ ) (Fig. 2). Furthermore, using an identical statistical approach over the same interval, the correlation between planktic foraminiferal standing diversity display a significant relationship when compared to both extinction and origination per-capita rates ( $\rho = -0.245$ ,  $p = 1.42 \times 10^{-3}$  and  $\rho = 0.384$ ,  $p = 2.96 \times 10^{-3}$ <sup>7</sup>, respectively). Additionally, when these variables were binned relative to the dominant global climatic modes, significant negative and positive associations between diversity-extinction and diversity-origination regressions, respectively, were also found (Table 1). Moreover, when Spearman's rank-order test was used in evaluating variation in both extinction and origination per-capita rates with future changes in diversity and changes in extinction percapita rates with future changes in origination per-capita rates, both tested by offsetting richness and evolutionary rates by one bin (= 1 Ma), no significant relationships remain (Table 2). Finally, the Kolmogorov-Smirnov test of the equality of frequency distribution displayed a significant difference among the overall shapes of planktic foraminifers' first appearance and last appearance distributions (D = $0.1001 p = 1.80 \times 10^{-3}$ ; Fig. 3).

# DISCUSSION

During the first ~30 Ma of planktic foraminiferal evolutionary history, the computed standing diversity has a relatively invariant character in terms of overall species richness with an average value of 2.61 species (CI: 2.25 to 2.96, computed



**Figure 2.** Relationship obtained between changes in both generalized differences of planktic foraminifers extinction per-capita rates and generalized differences of planktic foraminifers origination per-capita rates throughout the entire fossil history of the group (~170 Ma to Recent). The correlation test was carried out by Spearman's method, and this middle Jurassic to Recent time series consisted of 167 points ordered in time. Alpha level was established at 0.05. Outliers from the Aptian-Albian (A-A), Cretaceous-Paleogene (K-Pg), and Eocene-Oligocene (E-O) extinction events are shaded in grey. Abbreviations: OAE 1a, Oceanic Anoxic Event 1a; OAE 1b, Oceanic Anoxic Event 1b; OAE 1c, Oceanic Anoxic Event 1c; OAE 1d, Oceanic Anoxic Event 1d; OAE 2, Oceanic Anoxic Event 2; OAE 3, Oceanic Anoxic Event 3; K/Pg, Cretaceous/Paleogene extinction event; PETM, Palaeocene-Eocene Thermal Maximum; EECO, Early Eocene Climate Optimum; Oi-1, early Oligocene oxygen isotope event 1; MMCO, Mid-Miocene Climate Optimum; MPT, Mid-Pleistocene Transition.

**Table 1.** Results obtained from the Spearman correlation tests between changes in planktic foraminiferal standing diversity and changes in planktic foraminiferal turnover rates, and changes between per-capita extinction and origination rates throughout mixed-house, greenhouse, and icehouse intervals. For all of the correlation tests carried out by the Spearman's method, the mixed-house consists of 68 points, the greenhouse consists of 63 points, and the icehouse consisted of 37 points. Alpha level was established at 0.05.

Climatic mode	Correlation	ρ	Þ
Mixed-house	$\Delta$ diversity- $\Delta$ per-capita extinction rates	-0.2536	0.03
Mixed-house	∆ diversity-∆ per-capita origination rates	0.4794	4.06*10-5
Mixed-house	${\tt \Delta}$ per-capita extinction rates- ${\tt \Delta}$ per-capita origination rates	0.2643	0.03
Greenhouse	$\Delta$ diversity- $\Delta$ per-capita extinction rates	-0.3485	0.005
Greenhouse	∆ diversity-∆ per-capita origination rates	0.2563	0.04
Greenhouse	${\tt \Delta}$ per-capita extinction rates- ${\tt \Delta}$ per-capita origination rates	0.6659	4.85*10 <sup>-9</sup>
lcehouse	$\Delta$ diversity- $\Delta$ per-capita extinction rates	-0.5703	2.28*10-4
lcehouse	∆ diversity-∆ per-capita origination rates	0.3713	0.02
Icehouse	${\tt \Delta}$ per-capita extinction rates- ${\tt \Delta}$ per-capita origination rates	0.4248	0.009

<b>Table 2.</b> Results obtained from the Spearman correlation tests between ( <i>i</i> ) changes in planktic foraminifers standing diversity offset by one bir
(1 Ma) and changes in planktic foraminifers per-capita extinction rates, (ii) changes in planktic foraminifers standing diversity offset by one bin (1
Ma) and changes in planktic foraminifers per-capita origination rates, (iii) changes in planktic foraminifers per-capita extinction rates offset by one
bin (1Ma) and changes in per-capita origination rates, and (iv) changes in planktic foraminifers per-capita origination rates offset by one bin (1Ma
and changes in per-capita extinction rates, throughout the planktic foraminifers history (~170 Ma to recent). Alpha level was established at 0.05.

Correlation	ρ	þ
$\Delta$ Diversity (lag 1 Ma)- $\Delta$ per-capita extinction rates	-0.0395	0.6
Δ Diversity (lag 1 Ma)-Δ per-capita origination rates	0.0167	0.8
Δ per-capita extinction rates (lag 1 Ma)- $Δ$ per-capita origination rates	0.0508	0.5
$\Delta$ per-capita origination rates (lag 1 Ma)- $\Delta$ per-capita extinction rates	0.0721	0.3

after the 95 % confidence intervals bootstrapped 1000 times), whereas both extinction and origination per-capita rates display a very volatile pattern, characterized by several peaks of varying magnitude but also troughs in which both per-capita rates are zero (Fig. 1, Appendix). In part, this volatility is a function of the very low species richness during this interval such that even the origination or extinction of a single species can have a disproportionate impact as compared to later in the group's history.

In addition, this may also potentially be a result of lower planktic foraminiferal interspecific competition levels (Table 1; the value of extinction-origination cross-correlation is lower during the mixed-house interval as compared to the other two stages) associated with the initial invasion of the upper water column due to the evolutionary innovation of a 'free-floating' life habit from a benthic progenitor (Boudagher *et al.*, 1997). This adaptive radiation allowed the expansion of the foraminifers' habitat into a previously unoccupied ecological domain where the lack of constraints likely allowed evolutionary experimentation.

Following that phase, the standing-diversity curve dramatically increases, but rather than increasing exponentially or eventually reaching a plateau, the curve displays several peaks and troughs, which not only coincide with global events that disrupted the biota (i.e, Aptian-Albian bioevent as well as the Cretaceous-Paleogene and Eocene-Oligocene mass extinctions; Fig. 1), suggesting that changes in global diversity for this group, could also be related to a different set of environmental/paleobiological alterations (e.g. changes in niche availability, changes in types of competition). On the other hand, the turnover rates display a diminished intensity in their peaks from the Late Mesozoic through the Cenozoic (with exception of the values associated with the Cretaceous-Paleogene boundary), and even though there are various lower values, as reflected in the troughs, they are less prominent than during the initial ~30 Ma of the group during which the minimum values for both per-capita extinction and origination rates exceed zero (Fig. 1, Appendix, see online). Moreover, through time there

is an increase in the per-capita extinction-origination crosscorrelation values (Table 1). Lower values of extinctionorigination cross-correlation occurred during the mixedhouse interval, increased during the greenhouse stage, and are even higher through the icehouse period. In part, this continuous strengthening of the correlation between these two components likely suggests an intensification of planktic foraminiferal interspecific competition given that species have acquired a higher fitness in the pelagic environment and/or the enhancement of niche partitioning into a more densely inhabited ecological space.

The 'stationary' models of diversification (i.e., 'turnover pulse' [Vrba 1985] and 'coordinated stasis' [Brett and Baird, 1995]) necessitate that extinction and origination rates be cross-correlated to balance or offset any changes (Alroy, 1996; Alroy, 2008). Furthermore, these two models also predict no significant difference between the frequency distribution of the first and last appearance data given that long intervals (i.e., millions of years) of evolutionary stasis, resulting from a relatively consistent set of extrinsic environmental conditions (Vrba, 1985) and/ or strong stabilizing selection as postulated by Brett and Baird (1995), must be separated by virtually instantaneous (from a geological perspective of time) evolutionary pulses induced by major environmental shifts, allowing an increase in environmental fragmentation (Vrba, 1985) and/or accompanying biotic reorganizations (Brett and Baird 1995). Results obtained here, despite the Spearman-rank-order tests displaying a positive correlation between origination and extinction per-capita rates throughout the entire planktic foraminifers fossil history, do not support these two diversification hypotheses, as there is no correlation between origination and extinction per-capita rates when offsetting is performed (Table 2) and a significant difference in the equality of distribution between the curves of first and last appearances throughout the groups fossil history (~170 Ma to Recent; Fig. 3) as tested using the Kolmogorov-Smirnov test. Consequently, this lack of coordination between these two variables does not support the notion that long-term



**Figure 3.** Histograms of planktic foraminifer long-term diversity. **a**) Empirical diversity plotted as time-frequency histogram. **b**) Simulated exponential growth diversity plotted as time-frequency histogram. The exponential values were obtained from the equation  $N_t = N_a(\lambda)^t$ , where *t* is the number of time units,  $N_a$  is the size of the population at the beginning,  $\lambda$  is the per-capita rate of increase and  $N_t$  is the predicted size of the population after *t* time units. In the case of the planktic foraminifera *t* is the interval between 170 to 0 Ma,  $N_a$  is four species, and  $\lambda$  was equal to 0.2 (mean origination rate of planktic foraminifera).

intervals of morphological/evolutionary inertia within the planktic foraminifera shifted only when disrupted by environmental events.

Another oft-cited diversification hypothesis is the Red Queen model which postulates that the diversification pattern through time has been shaped by various competitive interactions between taxa (e.g. competition for resources, predator-previnteractions) leading to an unending escalation of adaptations (Van Valen, 1973). A major prediction based on this hypothesis is that diversity should grow exponentially through time, given that the constant turnover rates would be driven by biotic interactions and be, therefore, immune to environmental forcing (Van Valen, 1973; Benton, 2009; Alroy, 2010a). According to this criterion, the analysis of planktic foraminiferal diversity dynamics applied in this study do not support the Red Queen's predictions as the diversity pattern does not follow an exponential pattern (Kolmogorov-Smirnov test for equality of distribution between planktic foraminifera empirical diversity (Fig. 3a) and a simulated planktic foraminifera exponential diversity growth (Fig. 3b) showed a significant difference in distribution between the two curves:  $p = 2.2 \times 10^{-16}$ ), and both extinction and origination per-capita rates (Fig. 1) show a significant negative trend through time (Spearman-rankorder results after the regression of extinction per-capita rates against time:  $\rho = -0.269$ ,  $p = 4.25 \times 10^{-4}$ , Spearmanrank-order results after the regression of origination percapita rates against time:  $\rho = -0.209$ ,  $p = 6.61 \times 10^{-4}$ ).

An alternative explanation of long-term diversification is the so-called 'diversity equilibrium' model (Sepkoski, 1978; Rosenzweig, 1995; Alroy, 2008). Based on this, long-term diversification processes should obtain an equilibrium level at some point as a result of the following turnoverrelationship scenarios: (i) a secular increase and decline in extinction and in origination rates, respectively (Gilinsky and Bambach, 1987), (ii) the existence of a positive crosscorrelation between extinction rates with diversity levels and/or a negative cross-correlation among origination rates and standing diversity (i.e., density dependence of rates; Sepkoski, 1978; Rosenzweig, 1995), and (iii) a positive cross-correlation between extinction and origination rates given that origination pulses will respond to bursts of extinction and as well as vice versa (Webb, 1969; Mark and Flessa, 1977).

The current results do not support the first two components of this model given the significant negative trends through time for both extinction and origination per-capita rates ( $\rho$  = -0.269, *p* = 4.25\*10<sup>-4</sup> and  $\rho$  = -0.209,  $p = 6.60 \times 10^{-3}$ ), and the significant, although opposite to the postulated relationship (i.e., positive relationship between origination and standing diversity and negative correlation between extinction and species richness), cross-correlations between changes in both extinction and origination percapita rates with changes in diversity throughout time. Nevertheless, the presence of a significant positive crosscorrelation between origination and extinction per-capita rates suggests that the long-term planktic foraminiferal diversity pattern could be described by a logistic growth pattern with no extrinsic controls and limited only by intrinsic ecological factors (i.e., planktic foraminiferal

interspecific competition) as proposed by Webb (1969) and Mark and Flessa (1977). However, the overall logistic growth pattern predicted by this model is not displayed by the planktic foraminifers standing diversity (Fig. 1). Rather than gradually approaching an equilibrium level, the curve is instead characterized by several peaks and troughs, in addition to those reflecting elevated turnover forced by major biotic disruptions (i.e., Aptian-Albian, Cretaceous-Paleogene, Eocene-Oligocene). Given that the data do not match the predicted response, this hypothesis is also rejected for planktic foraminifers.

A final diversification scenario to be considered is the 'complicated logistic growth', which predicts that diversity limits are imposed by changing extrinsic factor(s) (Alroy, 2010a). As predicted by this model, diversity should reach a carrying capacity that in turn is controlled by varying external components (e.g., niche availability, resource abundance, richness and abundance of an interacting group, specifically either prey or predators). As a result, the diversity pattern predicted by this model does not mirror a logistic curve, instead it should display peaks and troughs suggesting a dynamic record being driven by medium- to long-term shifts in the carrying capacity forced by various external factors (Sepkoski, 1984; Alroy, 2010a). Moreover, competition should be sufficiently strong to maintain a level of diversity close to its saturation point in response to any environmental shifts, and the extrinsic factor(s) should change through time; if the extrinsic factor(s) does/do not change, the diversity pattern should 'plateau' (Alroy, 2010a). Consequently, the combination of the lack of a logistic standing diversity pattern (Fig. 1), the positive cross-correlation between changes in extinction and origination per-capita rates indicating significant foraminiferal levels of competition through all the fossil history of the group, and the significant association between changes in planktic foraminifers diversity pattern with various extrinsic environmental factor (e.g., changes in marine global temperature, as suggested by Schmidt et al., 2004; Peters et al., 2013; Frass et al., 2015), all indicate that the planktic foraminifers diversity pattern approaches logistic growth over relatively short intervals, due to fluctuating diversity limits imposed by changing extrinsic factor(s) that overprint that pattern. Furthermore, the 'non-equilibrium' state of global planktic foraminifers diversity through time is likely a result of continual changes in environmental conditions which influenced niche availability (e.g., Rutherford et al., 1999). This produces a diversity trend which, rather than equilibrating at a given level and producing a plateau, is instead highly dynamic and variable with diversity increasing and decreasing as upper mixed-layer niches expand and contract, respectively. This is supported by the results obtained here which display both significant cross-correlations between extinction per-capita rates and standing diversity (negative) as well as between origination per-capita rates with standing diversity (positive).

#### CONCLUSIONS

The quantitative examination of intrinsic planktic for a minifers' diversity dynamics throughout their entire fossil history (~170 Ma to Recent) supports the interpretation that this group's long-term diversity pattern follows the 'complicated' logistic growth' model, which has also been documented for the global marine Phanerozoic record of macrofossils (Alroy, 2010a). As proposed by Rosenzweig (1995), this indicate that diversity is a (i) self-regulating property of natural systems, with a dynamic eroding and restoring of the number of taxa, and (ii) does not increase infinitely through vast amounts of time given the evolutionary limitations imposed by various extrinsic components. For planktic foraminifers, diversity changes are likely controlled by niche availability, which is a function of both diversity as well as various biologic and environmental factors. Consequently, this long-term diversity pattern likely has changed in lockstep with the vertical range of the pelagic upper-mixed layer. These changes possibly influenced available niche space and potentially the overall number of niches. Furthermore, given that planktic foraminifer interspecific competition, as inferred by the significant cross-correlation displayed by changes in both extinction and origination per-capita rates, has been sufficiently strong to track the carrying capacity of the system (forced by the global oceanic temperature changes) through time, the 'complicated logistic growth' long-term diversification scenario is the most parsimonious model explaining the shape of the global long-term planktic foraminifers diversity pattern. Finally, our study strongly suggests that a full understanding of long-term diversity patterns must be rooted in the analysis of both intrinsic and extrinsic biotic and abiotic controls.

Appendix data of this manuscript are available at Acta Biológica Colombiana Online version.

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