

Effect of gross morphology on modern planktonic foraminiferal test strength under compression

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Abstract: Planktonic foraminifera are a source of important geochemical, palaeoceanographic, and palaeontological data. However, many aspects of their ecology remain poorly understood, including whether or not gross morphology has an ecological function. Here, we measure the force needed to crush multiple planktonic foraminiferal morphotypes from modern core top and tow samples. We find significant differences in the resistance of different morphotypes to compressional force. Three species, *Globorotalia tumida* (biconvex, keeled), *Menardella menardii* (discoidal, keeled), *Truncorotalia truncatulinoides* (conical, keeled), require on average 59% more force (1.07 v. 0.47 N) to crush than the least resistant species (*Orbulina universa* and *Trilobatus sacculifer*) in core-top samples. Towed samples of pre-gametogenic individuals also show significant differences of the same magnitude (0.693 v. 0.53 N) between the conical (*T. truncatulinoides*) and globular/spherical morphologies (*Globoconella inflata* and *O. universa*). We hypothesize that the greater compressional strength of certain shapes confers a fitness advantage against predators and could contribute to the repeated, convergent evolution of keeled, conical and bi-convex forms in planktonic foraminifer lineages.

Keywords: planktonic foraminifera, morphology, predation, pelagic ecology, iterative evolution

Supplementary material: Raw data for all crushing experiments, wall thickness measurements, and results for all pair-wise Kolmogorov-Smirnov Tests are available at <https://doi.org/10.6084/m9.figshare.c.3725236.v1>

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Throughout their fossil record, planktonic foraminifera have repeatedly converged on a limited number of gross morphologies (i.e. morphotypes). Characteristic morphological features, such as reinforced ‘keeled’ margins and elongated ‘digitate’ chambers, have evolved, gone extinct and re-evolved in different lineages in a process known as iterative evolution (Cifelli 1969; Norris 1991; Coxall *et al.* 2007). The characteristic evolutionary sequence leads from a relatively simple, globular (globigerine) form to a more derived morphotype like those including keeled margins and biconvex or conical forms (Cifelli 1969; Norris 1991).

This pattern of iterative evolution suggests that there may be adaptive advantages to different morphologies, but what these advantages might be is still unclear. In a 1979 review, Lipps presents five hypotheses to account for the ubiquity of convergent evolution in planktonic foraminifera: (1) there is a limited number of possible forms (i.e. convergence is inevitable); (2) physical forces – particularly surface tension – shape foraminiferal tests; (3) test shape relates to buoyancy control; (4) test shape regulates orientation in the water to reduce settling speed; and (5) some morphologies provide greater defence against predation (Lipps 1979). The first two hypotheses were immediately rejected by Lipps as having already been disproven. In the first case, Lipps referenced Loeblich & Tappan’s (1964) illustration of the morphological diversity of planktonic foraminifera to argue against limits on the number of possible foraminiferal forms. More recent modelling studies suggest this argument may be premature, as they show that much of the theoretical foraminiferal morphospace has been exploited repeatedly (Tyszkla & Topa 2005; Tyszkla 2006). In short, morphological constraint can no longer be so completely discounted. For the second case, put forth by Thompson (1961), Lipps argues that surface tension alone is an oversimplification of the factors that affect cell dimensions and is insufficient to explain

the range of morphologies. Foraminiferal gross morphology is often attributed to hydrodynamic controls, including buoyancy, settling and preferred orientation (Marsalek *et al.* 1969; Lipps 1979; Caromel *et al.* 2014), the third and fourth hypotheses in Lipps (1979). Multiple factors affect buoyancy, including the composition of cytoplasm, the volume of the test occupied by the living cytoplasm, and the test density and weight (Marsalek *et al.* 1969; Lipps 1979). Together these factors can determine whether an individual sinks, floats or maintains its depth habitat. Likewise, individuals can also reduce their sinking speed by adopting and maintaining specific body orientations (Lipps 1979). This was extensively explored in a 2014 study by Caromel and colleagues, where they used hydrodynamic models to conclude that buoyancy control and orientation can require very minor morphological changes in test dimensions and density relative to those observed (Caromel *et al.* 2014). In other words, neither explanation can account for the range of morphologies observed in fossil and living taxa.

The third viable hypothesis of Lipps – that different morphologies serve to prevent or limit selective predation – has remained largely unexplored, even though zooplankton predators exert heavy selective pressure in the pelagic realm in general (Lipps 1979; Smetacek 2001). Diatoms – planktonic photoautotrophs with siliceous tests called ‘frustules’ – also have a wide array of frustule shapes and structures. It has been hypothesized that these shapes deter mesoplankton grazers, such as pelagic copepods and euphausiids (Smetacek 2001; Hamm *et al.* 2003). The resistance of diatoms to crushing has been tested experimentally and it has been shown that the frustules can resist surprisingly high crushing forces of up to 7 N mm⁻² (Hamm *et al.* 2003). Moreover, the force needed to crush diatom frustules varies among species of different shapes, potentially

offering different levels of defence (Hamm *et al.* 2003). The same defensive benefits and variation among morphologies may apply to planktonic foraminifera (Lipps 1979; Hemleben *et al.* 1989) but have yet to be tested.

There are reasons to suspect that selective predation may play a role in shaping foraminiferal morphology. Some features of planktonic foraminifera, such as the pyramidal shape of *Truncorotalia truncatulinoides* or the long, bulky spikes of the extinct *Hantkenina mexicana*, may be unwieldy or hard to bite and crush for a zooplankton predator, causing the predator to preferentially feed on unarmoured individuals. Even small differences in the amount of time a predator spends trying to capture and ingest its prey (i.e. handling time) can affect optimal foraging behaviour of selective zooplankton predators and the rate at which they can cull populations of their prey species (Holling 1959; Chang & Hanazato 2005). Arthropod predators, such as copepods and crabs, have been observed to feed preferentially on easier to crush or open-shelled prey when the option is available (Boulding 1984; Katz 1985; Chang & Hanazato 2005). Handling time can shape population size over extended time-scales through differential mortality rates by morphotype (Holling 1959; Werner & Hall 1979; Jeschke *et al.* 2002; Chang & Hanazato 2005). Even if the bulk of predation pressure on planktonic foraminifera were by indiscriminant feeders (like filter-feeding fish), a small proportion of selective feeding is enough to cause differential survivorship due to morphology. Along these lines, there have been anecdotal accounts of crushed planktonic foraminifera in the guts of arthropods (Bé & Hutson 1977; Lipps 1979; Hemleben *et al.* 1989).

A few published studies have explored the resistance of foraminifera to compressional forces and found differences

among morphotypes (Whetmore 1987; Whetmore & Plotnick 1992; Pearson *et al.* 2015). In an experiment on benthic foraminifera, crushing resistance was positively correlated with test diameter and wall thickness in several of the species tested, and species with biconvex morphologies (*Elphidiella hannai*, *Islandiella limabata* and *Oolina borealis*) were generally the most resistant to crushing (Whetmore 1987). Another study related crushing resistance to wave-stress in the benthic *Archaias angulatus*, *Amphistegina gibbosa* and *Laevipeneroplus proteus* (Whetmore & Plotnick 1992) and found that the strongest species occurred in the most dynamic benthic environments. This relationship also held true within populations of the same species, *Archaias angulatus*. More recently, in the only crushing study to date on planktonic foraminifera, Pearson *et al.* (2015) explored the compressional forces needed to crack planktonic foraminiferal tests. The aim was to determine whether diagenetically recrystallized fossil tests of two different species, *Cribohantkenina inflata* and *Turborotalia cerroazulensis*, were less resistant to compression than unaltered fossil tests. They found that unaltered tests were stronger than altered tests and that the more conical species, *T. cerroazulensis*, was stronger than the more globular species.

Inspired by the report by Pearson *et al.* (2015), we herein test whether morphologically distinct, well-preserved modern planktonic foraminiferal tests vary in their resistance to compressional forces (i.e. 'crushing resistance'). The eight species used in this analysis, *Trilobatus sacculifer*, *Orbulina universa*, *Globigerinoides conglobatus*, *Globoconella inflata*, *Neogloboquadrina dutertrei*, *Globorotalia tumida*, *Menardella menardii* and *Truncorotalia truncatulinoides*, span much of the morphological range of modern planktonic foraminifera and include forms with varying test-wall thickness (Fig. 1). Relative crushing resistance amongst



Fig. 1. Edge views of the eight species used in this study: (a) *Globigerinoides conglobatus*, (b) *Globoconella inflata*, (c) *Trilobatus sacculifer*, (d) *Truncorotalia truncatulinoides*, (e) *Neogloboquadrina dutertrei*, (f) *Orbulina universa*, (g) *Globorotalia tumida* and (h) *Menardella menardii*.

taxa is then considered with regards to predation on planktonic foraminifera and preservation potential in the fossil record.

Materials and methods

Three samples (two core top and one tow) were picked for approximately 15 individual empty foraminiferal tests per target species for crushing experiments. A different subset of the eight target species was obtained from each sample locality, given the availability of species within that locality. The eight species targeted were chosen based on abundance, morphology and wall thickness, and included the globigerinoid foraminifer *Trilobatus sacculifer* (formerly *Globigerinoides sacculifer*, see Spezzaferri *et al.* (2015)); the spherical *Orbulina universa*; the calcite-encrusted globigerinoid *Globigerinoides conglobatus*; the rounded, cuboidal *Neoglobobulimina dutertrei*; the inflated, semi-conical chambered turborotalid *Globoconella inflata*; the keeled, biconvex globorotalid *Globorotalia tumida*; the keeled, compressed, thin-walled globorotalid *Menardella menardii*; and the conical globorotalid *Truncorotalia truncatulinoides* (Fig. 1). Species were identified based on the species concepts of Kennet & Srinivasan (1983), with naming after Aze *et al.* (2011) (with the exception of *T. sacculifer*; Spezzaferri *et al.* (2015)).

Two types of samples were used: core-top samples and net-tow samples. The first was used to sample a range of abundant taxa and the second to test how gametogenic-calcite (added immediately prior to death) might influence our inferences. In the KC78 core-top sample, *T. sacculifer*, *O. universa*, *G. tumida*, *N. dutertrei*, *M. menardii* and *G. conglobatus* specimens were picked from the

425–600 and 600–710 µm sieve size fractions. The KC78 core-top was sampled at 5.267°N by 44.133° W in the South Atlantic Ocean at a water depth of 3273 m by Woods Hole Oceanographic Institution (Sun *et al.* 2006), and provided by Bruce Corliss (University of Rhode Island, Graduate School of Oceanography). KC78 is a diverse assemblage dominated by the mixed-layer dwelling species *T. sacculifer*, with abundant *O. universa*, *M. menardii* and *G. tumida* in the size fractions picked. Two other species, *N. dutertrei* and *G. conglobatus*, although relatively rare, were also included in the study because of their distinct morphologies. The AII60-10 core top sample was provided by Richard Norris (Scripps Institution of Oceanography) and obtained from 29.660° S and 34.667° W at 1840 m water depth by the Woods Hole Oceanographic Institution's *Atlantis II* research vessel. Specimens of *O. universa*, *T. truncatulinoides* and *G. tumida* were picked from AII60-10. Three species, *G. inflata*, *O. universa* and *T. truncatulinoides*, were picked from two net tows and combined for this study: TAN1106/40 and TAN1106/50. The TAN1106/40 and TAN1106/50 localities were sampled at 48.796° S by 165.497° W from <150 m water depth in 2011 by the National Institute of Water and Atmospheric Research's Solander Trough Cruise with a Multiple Opening/Closing Net and Environmental Sensing System (MOCNESS, Wiebe *et al.* 1976). *Globoconella inflata*, *O. universa* and *T. truncatulinoides* were very abundant in the 300–600 µm size fraction and provide a 'living assemblage' test of our inferences otherwise made from post-gametogenic core-top individuals. Yale Peabody Museum catalogue numbers (YPM#) for all samples are listed in Table 1. All specimens from AII60-10 and TAN1106, and a number of

Table 1. Yale Peabody Museum accession number, species name, sample locality, size range, type and size, mean force at failure, coefficients of variation for failure force and wall thickness, mean wall thickness and relative crushing resistance for each sample based on force at failure

YPM#	Species	Sample locality (size range)	Sample type	Number of individuals (n)	Force at failure (N)	Coeff. of variation	Mean wall thickness (µm)	Coeff. of variation	Relative crushing resistance
IP.307890	<i>Globorotalia tumida</i>	AII60-10 (425–710 µm)	Core top	12	1.053	0.304	24.87	0.20	Higher
IP.307891	<i>Orbulina universa</i>	AII60-10 (425–710 µm)	Core top	12	0.419	0.419	29.43	0.44	Normal
IP.307892	<i>Truncorotalia truncatulinoides</i>	AII60-10 (425–710 µm)	Core top	12	1.26	0.284	18.82	0.18	Higher
IP.307893	<i>Globigerinoides conglobatus</i>	KC78 (425– 600 µm)	Core top	16	0.585	0.249	52.80	0.13	Normal
IP.307894	<i>Globorotalia tumida</i>	KC78 (425– 600 µm)	Core top	15	1.32	0.24	34.22	0.13	Higher
IP.307895	<i>Menardella menardii</i>	KC78 (425– 600 µm)	Core top	15	0.944	0.357	27.02	0.21	Higher
IP.307896	<i>Neoglobobulimina dutertrei</i>	KC78 (425– 600 µm)	Core top	17	0.618	0.182	39.18	0.31	Normal
IP.307897	<i>Orbulina universa</i>	KC78 (425– 600 µm)	Core top	16	0.499	0.374	20.17	0.32	Normal
IP.307898	<i>Trilobatus sacculifer</i>	KC78 (425– 600 µm)	Core top	16	0.475	0.29	29.97	0.20	Normal
IP.307899	<i>Menardella menardii</i>	KC78 (600– 710 µm)	Core top	15	0.783	0.293	24.38	0.27	Higher
IP.307900	<i>Orbulina universa</i>	KC78 (600– 710 µm)	Core top	14	0.575	0.317	38.70	0.36	Normal
IP.307901	<i>Trilobatus sacculifer</i>	KC78 (600– 710 µm)	Core top	12	0.463	0.256	33.31	0.22	Normal
IP.307902	<i>Globoconella inflata</i>	TAN1106 (300–600 µm)	Tow	15	0.421	0.277	11.64	0.29	Normal
IP.307903	<i>Orbulina universa</i>	TAN1106 (300–600 µm)	Tow	14	0.379	0.491	26.30	0.51	Lower
IP.307904	<i>Truncorotalia truncatulinoides</i>	TAN1106 (300–600 µm)	Tow	15	0.639	0.256	11.95	0.20	Higher

Images of specimens used are available (searchable by IP# on the Invertebrate Paleontology page) at <http://collections.peabody.yale.edu/search>

individuals of *G. tumida* and *M. menardii* from KC78, were photographed from the umbilical view and measured along their major and minor axes prior to crushing using a stage micrometer and ImageJ software (Abramoff *et al.* 2004). The area of individually measured foraminifera was estimated by multiplying the major and minor axes. Wall thickness was measured by breaking off the final chamber (excluding kummerform chambers) of individual foraminifera from each species, imaging them so that a cross-section of the chamber wall was visible, and measuring the wall thickness using ImageJ.

To precisely measure crushing resistance of fossil foraminifera under compression, Pearson *et al.* (2015) used a Losenhausen servohydraulic testing machine with a 5 N Interface S-Beam load cell. For our study, we designed a relatively simple crushing rig to

mount and crush foraminifera via incremental weight addition. Specimens were mounted on either their umbilical or spiral side, depending on which one provided the most stable (i.e. flat) surface. The mass needed to crush a test was determined by subsequent weighing of the weights after test failure, and the compressional force was calculated in Newtons as mass \times acceleration (i.e. 9.807 m s^{-2}). Many design iterations were explored, but the most effective apparatus was a tube constructed from 20 ml plastic syringes (Fig. 2). The plunger of one syringe acts as a stage (Fig. 2f), to which a foraminifer is fixed, orientated dorso-ventrally, as pictured in Figure 1, with double-sided tape. Over this stage, a sleeve made from two syringes (Fig. 2d), guides a second plunger (Fig. 2b and c). The metal peg rests directly on the target foraminifer and applies the overlying weight to the

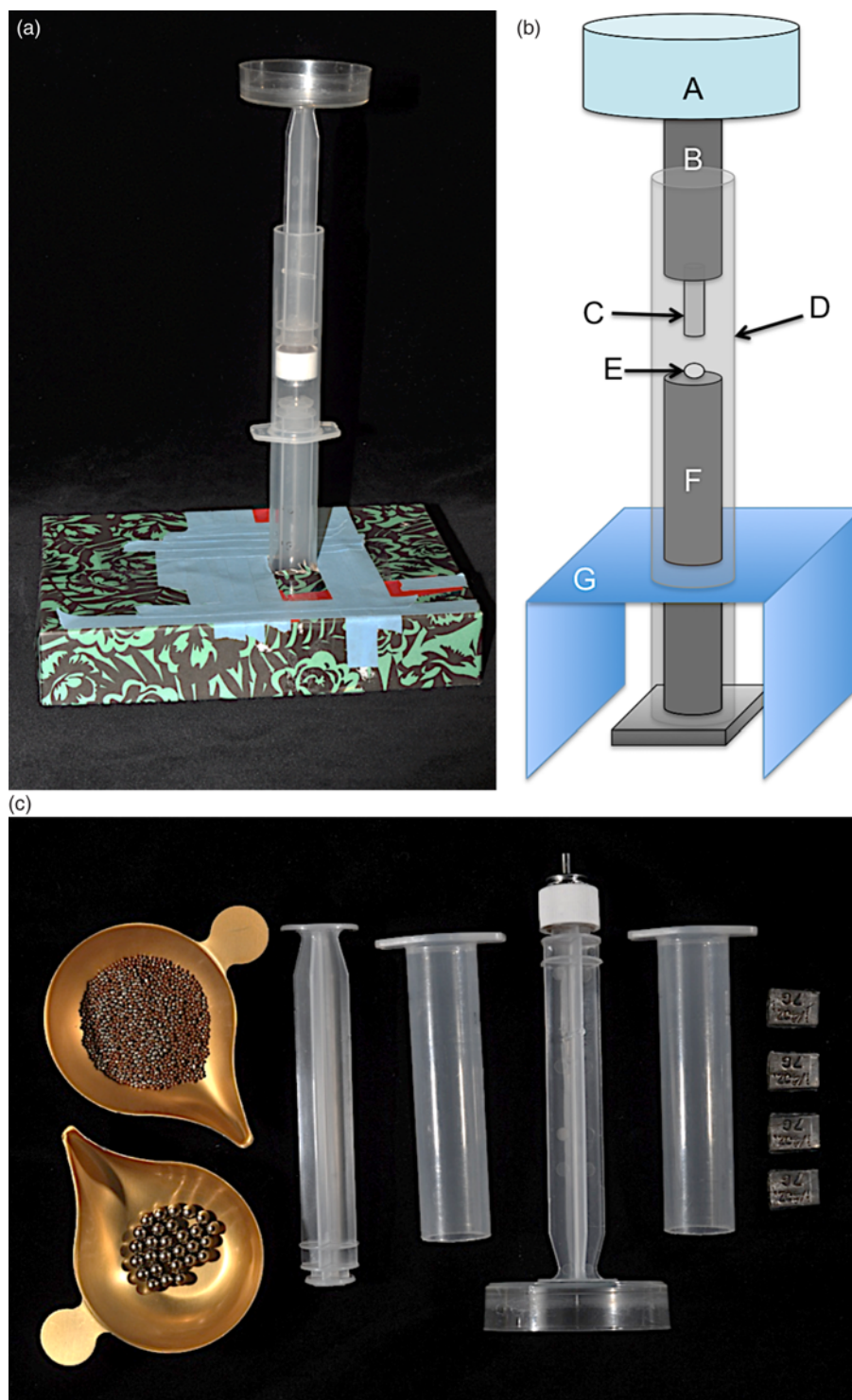


Fig. 2. Crushing rig: (a) photo and (b) sketch (not to scale). Crushing rig components in (b) labelled as follows: (A) weight collection dish, (B) upper plunger, (C) metal peg, (D) syringe sleeve, (E) specimen, (F) lower plunger/stage and (G) cardboard base (for stability). (c) Additional and disassembled crushing rig components from left to right: brass scoops with 1 and 4 mm beads, syringe plunger (used as a stage, F in b), cropped syringe sheath (D in b), syringe plunger with metal peg attached to one end and plastic dish attached to the other (A–C in b), second cropped syringe sheath, 7 g lead weights. Photographs courtesy of the Yale Peabody Museum division of Invertebrate Zoology.

specimen (Fig. 2). A small receptacle for weights rests on top of the second plunger (Fig. 2a). In each crushing experiment, small metal weights were carefully added to the dish until the specimen broke, causing the upper part of the rig to visibly drop. The combined mass of the weights at breakage and mass of the peg/plunger/dish apparatus were measured and converted to Newtons.

All statistical analyses were performed using built-in functions in RStudio version 0.97.55. To determine whether parametric or non-parametric statistics were appropriate, we tested the distribution of compressional forces for each species (by sample) for normality using the Shapiro–Wilk test. The data from two sample sets failed the tests of normality (see below for details), leading us to use non-parametric statistics. After deciding to use non-parametric statistics, we tested for significant differences among species and samples using a one-way Kruskal–Wallis ANOVA. Kolmogorov–Smirnov tests were then used to identify those pairs of taxa with significantly different compressional force distributions.

Results

Compressional force at crushing was measured for each species and case (core top or tow and size class) in 12–17 individuals. All species showed variability in the distribution of crushing resistance

values (Fig. 3; Table 1). For instance, the compression force at crushing for *Trilobatus sacculifer* individuals ranged from 0.249 to 0.726 N. In most cases, the force needed to crush individuals of a given species were normally distributed (i.e. they failed to be rejected by a Shapiro–Wilk test for normality). The two exceptions were the towed *Orbulina universa* from TAN1106 and the core-top *Trilobatus sacculifer* from the 600–710 μm sieve size fraction of KC78. Because these two cases failed the Shapiro–Wilk test for normality, non-parametric statistics were used.

Within species groups and localities, the relationship between test area and compressional force at failure was weak or non-existent. Across all individuals, compressional force at failure was significantly correlated with test area (P -value = 0.0001), but the variance explained was low ($r^2 = 0.121$). When analysed by species and locality, no locality or species had a significant relationship between test area and failure force.

Similarly, a linear regression detected no significant relationship between mean test thickness and the mean force at failure for each species from each locality ($r^2 = 0.003$, $P = 0.852$, Fig. 4). Within samples, *O. universa* typically had among the highest test wall-thicknesses measured (Table 1, Supplementary Table 3). *Orbulina universa* and *G. tumida* were significantly thicker than *T. truncatulinoides* in the AII60-10 core top sample. *Orbulina*

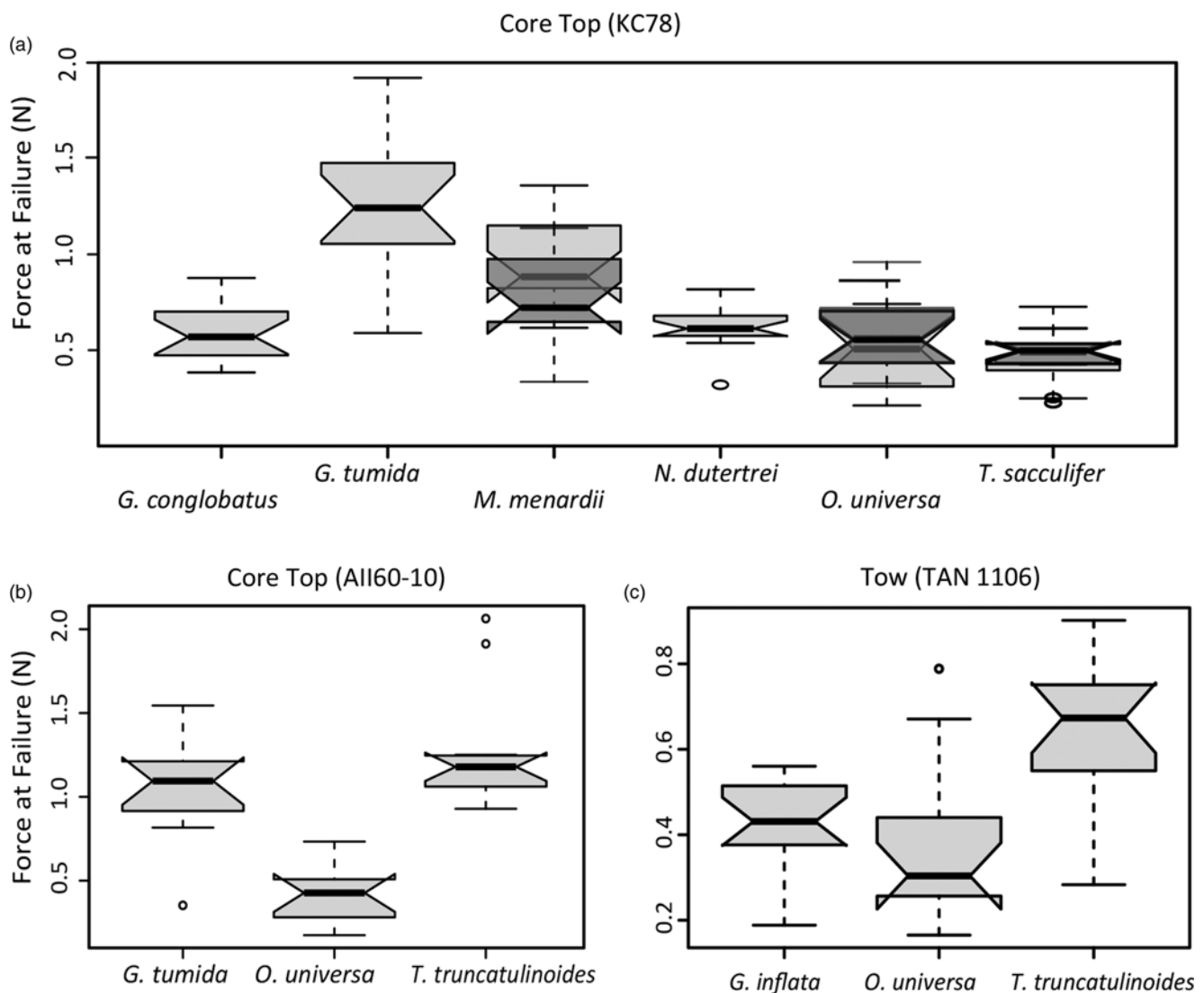


Fig. 3. Boxplots of failure weight distributions for all species groups by sample group. Boxplot notches indicate a 95% confidence interval on the median. (a) Core-top sample KC78, (b) core-top sample AII 60-10 and (c) MOCNESS tow sample TAN1106. In (a) the two size fractions of KC78 are overlain, with the smaller size fraction (425–600 μm) in light grey and the larger size fraction (600–710 μm) in dark grey.

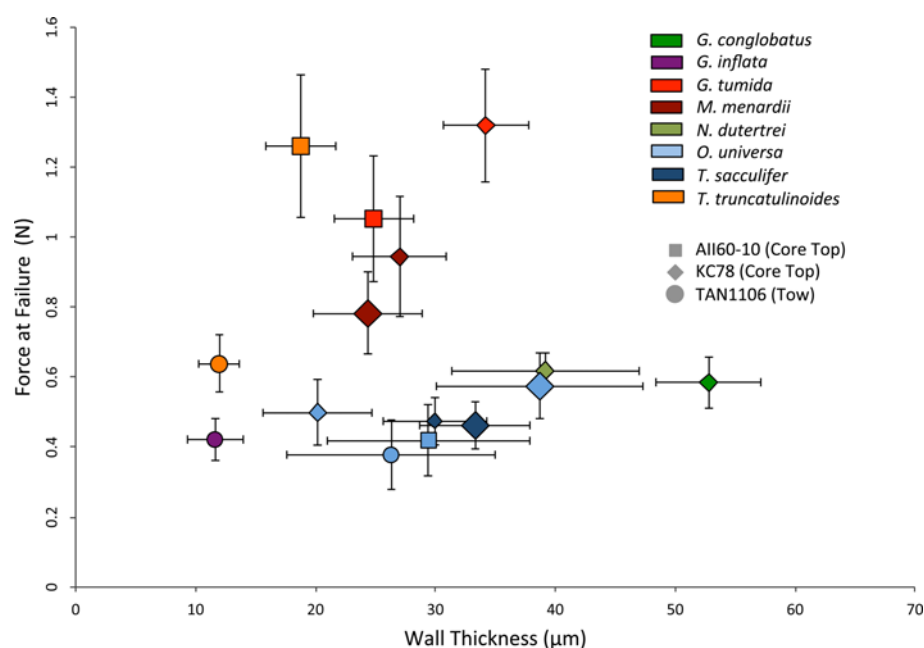


Fig. 4. Average values for wall thickness and force at failure for each species group from each locality, with 95% confidence intervals denoted by error bars. There is no significant relationship between the thickness of the final chamber and resistance to compressive force. For KC78, the larger diamond icons indicate averages from the 600–710 μm size fraction and the smaller diamond icons indicate averages from the 425–600 μm size fraction.

universa was also significantly thicker than the other species in the towed sample, TAN1106. In the larger size fraction of KC78, *O. universa* was thicker on average than the other species in the larger size fraction of KC78, but not significantly so. Finally, *G. conglobatus* was significantly thicker than all other species in the smaller size fraction of KC78. Variation in wall thickness was high, with coefficients of variation as high as 0.51 for the towed specimens of *O. universa* from TAN1106. *Globigerinoides conglobatus* and *G. tumida* from KC78 (425–600 μm size fraction) had the lowest coefficients of variability (0.13).

Significant differences in the distribution of the crushing resistance of species were tested with a one-way Kruskal–Wallis ANOVA for each case (*P*-values in Table 2). Kolmogorov–Smirnov tests (Supplementary Table 2) were then used to determine which pairwise comparisons were statistically different. The conical and biconvex species were significantly more resistant to crushing under compression as compared to the other species in all samples tested. In the core-top KC78, *G. tumida* ($\mu = 1.32$ N) and *M. menardii* ($\mu = 0.944$ N) were significantly more resistant to compressional force than all other species (*P*-value < 0.001) by 48% on average, but did not differ significantly from one another (*P*-value = 0.076). A separate Kruskal–Wallis test of the four relatively weak species from KC78, *O. universa* ($\mu = 0.499$ N), *G. conglobatus* ($\mu = 0.585$ N), *N. dutertrei* ($\mu = 0.618$ N) and *T. sacculifer* ($\mu = 0.475$ N), also found significant differences amongst species (*P*-value = 0.017; see Supplementary Table 2 for pairwise). In the core top All60-10, *G. tumida* and *T. truncatulinoides* were significantly more resistant (36% on average) to crushing force than *O. universa* (*P*-value < 0.001), with mean crushing forces of 1.053, 1.26, and 0.419 N, respectively. Similarly, in the TAN1106 towed sample, *T. truncatulinoides* was significantly more resistant (63% on average) to crushing force than the *O. universa* and *G. inflata*

(*P*-value < 0.001), with mean crushing forces of 0.639, 0.421, and 0.379 N respectively. In the same sample, *O. universa* crushed under significantly less force than *G. inflata*. Across samples, *G. tumida*, *M. menardii* and *T. truncatulinoides* were significantly more resistant to crushing force than the other species tested (Supplementary Table 2).

Orbulina universa was present in all four cases examined (core-top KC78 425–600 μm , core-top KC78 600–710 μm , core-top All60-10 and tow TAN1106) and consistently showed the greatest variability in compressional force, as indicated by the coefficient of variation (0.374 v. 0.182–0.357 for core-top KC78 425–600 μm ; 0.317 v. 0.256–0.293 for core-top KC78 600–710 μm ; 0.419 v. 0.284–0.304 for core-top All60-10; and 0.491 v. 0.256–0.277 for tow TAN1106). There was no significant difference in the force needed to crush *O. universa* in the three core-top samples (Fig. 5). Average crushing force for *O. universa* was 0.379 N (median = 0.304 N) in the tow sample TAN1106, 0.499 N (median = 0.506 N) in the 425–600 μm size fraction of core-top KC78, and 0.419 N (median = 0.427 N) in core-top All60-10. The towed *O. universa* from TAN1106 were significantly weaker ($\mu = 0.575$ N, median = 0.561 N) than the *O. universa* individuals in the largest size fraction of KC78 (*P*-value = 0.037).

Discussion

Modern planktonic foraminifera differ in their resistance to crushing, as measured by mean compressional force at test failure. Of the eight species tested, the most resistant species were *Menardella menardii*, *Globorotalia tumida* and *Truncorotalia truncatulinoides*, which have biconvex keeled (*M. menardii* and *G. tumida*) and conical keeled (*T. truncatulinoides*) morphologies. For instance, in the core-top sample KC78, *M. menardii* and *G. tumida* were on average 1.5 and 2 times more resistant to crushing than the next most resistant species, *Neogloboquadrina dutertrei*, in the 425–600 μm size fraction. More generally, biconvex and conical species had between 1.4 and 3 times the compressional strength of the more globular species (Tables 1, 2 and Results) and this compressional strength was unrelated to variation in wall thickness across species (Table 1; Fig. 4).

The aim of this study was to understand how mean compressional force at test failure varies amongst species and how this might affect (and reflect) their ecology. For this, the comparison between core-

Table 2. Results of Kruskal–Wallis analysis of variance for each sample group

Sample locality	Size range (μm)	Sample type	Chi squared	<i>P</i> value
All 60-10	425–710	Core top	19.320	<0.001
KC78 425–600	425–600	Core top	56.380	<0.001
KC78 600–710	600–710	Core top	24.618	<0.001
TAN1106	300–600	Tow	15.357	<0.001

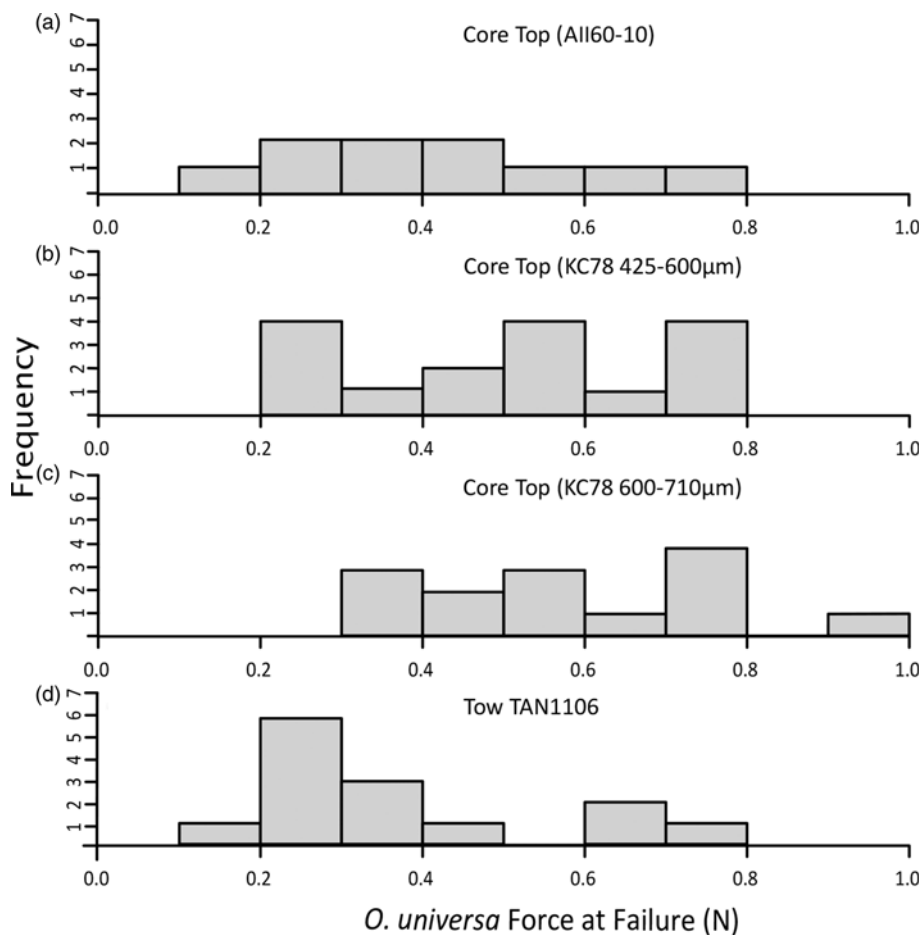


Fig. 5. Distribution of compressional force at failure for *Orbulina universa* in all four sample groups: (a) core-top AII 60-10; (b) core-top KC78 425 – 600 µm; (c) core-top KC78 600 – 710 µm; (d) MOCNESS tow TAN1106. Failure force distributions of *O. universa* samples are not significantly different from one another.

top and tow samples is important. Individuals from tow samples were pre-gametogenic. That is, they had yet to reproduce and add the extra layer of calcite that thickens planktonic foraminiferal tests immediately prior to death. We expected that the effect of gametogenesis would be to generally strengthen the foraminiferal test. Following on these expectations, *T. truncatulinoides* from the tow sample were significantly less resistant to crushing than those from the AII60-10 core top (P -value < 0.001; Fig. 3). In addition, failure weights for the towed, pre-gametogenic individuals of *O. universa* were noticeably skewed toward lower compressional forces (Fig. 5), although pairwise Kolmogorov–Smirnov tests failed to distinguish amongst core-top and towed samples of *O. universa*. Counter to our expectation regarding gametogenic calcification, compressional forces were relatively low in the second core-top sample for all species (AII60-10) and comparable to that of the tow sample (see Figs 3b, c and 5). This pattern remains unexplained and could indicate a number of things, including a relatively minor role of post-gametogenic wall thickening in the species tested at AII60-10, a relatively greater importance of intraspecific variation in crushing resistance, or relatively poor preservation at AII60-10 as compared to KC78 (note: visual inspection suggested excellent ‘glassy’ preservation at both core-top sites).

Importantly, we found significant differences in crushing resistance among pre-gametogenic species in the tow sample. Specifically, *T. truncatulinoides* was significantly more resistant, at least 1.5-fold stronger, than *O. universa* and *G. inflata*. As with the core-top sample, this significant difference in the mechanical strength existed between the conical *T. truncatulinoides* and globular/spherical morphologies of *O. universa* and *G. inflata*. Although the rest of our comparisons were in core-top individuals, and likely on post-gametogenic individuals, the tow data suggest that the relative relationship amongst species is maintained. In other

words, relatively resistant core-top morphologies were also likely relatively hard to crush when living.

In all species measured, we observed a wide variation in the crushing resistance of individuals. Pearson *et al.* (2015) also observed wide variation in the force at first cracking of tests (forces ranged from 0.2 to 1.7 N) and they hypothesized that this may have been due to undetected wall-defects (cracks) in some individuals. A second possible source of variation is measurement error introduced by the relatively simple crushing rig used here. Weights were added sequentially, with the heaviest weights (7 g lead tabs) being utilized only on the most rugged specimens of *G. tumida*, *M. menardii* and post-gametogenic *T. truncatulinoides*. At times, additional weight of up to 7 g may have been added after test failure. This source of error is relatively small as compared to the standard deviation of failure weights (standard deviations ranged from 11 to 48 g). Additionally, mounting position is critical, but the most difficult to mount specimens (biconvex taxa *G. tumida* and *M. menardii*) do not have the highest coefficients of variation, suggesting that this is not a large source of variation.

We suspect that intraspecific variation in test construction and wall thickness may account for some of the within-species variation in crushing resistance observed. For instance, *O. universa* had the greatest variation in mean weight at failure of all species tested (Table 1; Figs 3–5). This broad variation may relate to the known occurrence of ‘thin-’ and ‘thick-’walled morphotypes of *O. universa*, perhaps coinciding with cryptic species (Hamilton *et al.* 2008; Morard *et al.* 2009; Marshall *et al.* 2015). In our four samples, we found wall thicknesses of *O. universa* to range from 14.4 to 49.3 µm (Fig. 4; Table 1, Supplementary Table 3). However, even though high variability of resistance to crushing and high variability in wall thickness were associated in *O. universa*, wall thickness overall was a poor predictor of crush resistance across species

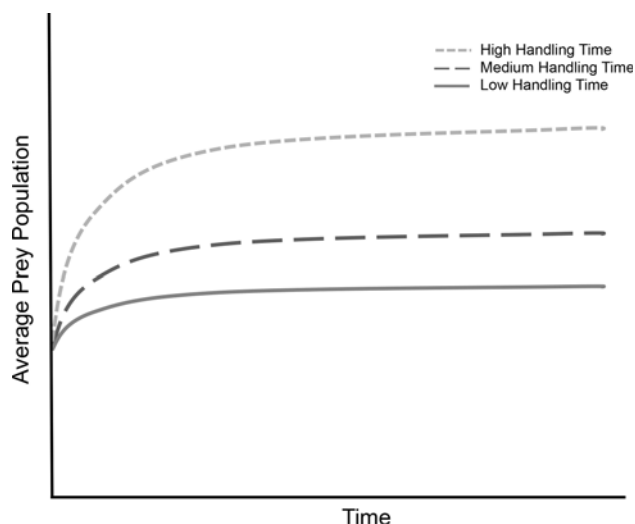


Fig. 6. A theoretical sketch of the effect of crushing resistance on average prey population size. Predation pressure is illustrated for three populations with different predator handling times based on Lotka–Volterra models described in Stevens (2009) and handling time disparities after the range observed in Chang & Hanazato (2005) for water fleas and copepods.

(Fig. 4). In many cases, the most crush-resistant species in a sample was, in fact, one of the thinnest walled. For example in the AII60-10 core-top sample, *T. truncatulinoidea* has the thinnest walls and the highest compressional force at failure. Similarly, in the towed TAN1106 sample, *T. truncatulinoidea* requires the highest compressional force to crush and has among the thinnest average test-walls. This suggests that gross morphology, or the shape and the arrangement of the preceding chambers, is key to crush resistance, and not wall thickness.

If higher crushing resistance in planktonic foraminifera leads to higher handling time during attempted predation, and higher handling time results in reduced mortality as it does in other groups (Boulding 1984; Chang & Hanazato 2005), then a fitness advantage may be conferred on relatively hard to crush species. In Figure 6, the potential of handling time to affect survivorship of different planktonic foraminiferal species is imagined in a sketch of idealized prey populations with differing crushing resistance under predation. This figure was constructed with Lotka–Volterra type coupled equations, as in Stevens (2009), which model the interdependent fluctuations of two populations linked by trophic interactions. In general, higher handling times should reduce prey consumption per unit time and lead to higher average population sizes and survivorship over time (Fig. 6). Populations of planktonic foraminifera might be expected to show this type of behaviour even if most of their mortality was due to non-selective predators. This is

because predation by large filter feeders would exert a constant, non-selective pressure on all morphotypes, with just the selective feeding driving the difference in mortality between species. Future evaluation of this predation hypothesis is needed to directly parameterize the relationship between shape and handling time, feeding anatomy of potential predators and the relative importance of other traits, such as spines and palatability. Additionally, we measured compressional strength on the spiral–umbilical axis, but reinforced features such as the keel suggest that the greatest forces experienced in nature may occur in other orientations. Further crushing and feeding experiments could provide more specific insight into the repeated evolution of keels and other features.

Mean compression strength can also have taphonomic effects, influencing the porosity of sediments and the composition of fossil foraminiferal assemblages (i.e. Pearson *et al.* 2015). Berger & Piper (1972) explored the differences in dissolution susceptibility as well as sinking and settling speed in relation to preservation potential. Here we can add crushing resistance to the taphonomic data of Berger & Piper (Table 3, note: compressional strength is considered here independently of subsequent dissolution effects), highlighting the particularly low preservation potential of some taxa (*T. sacculifer* and *O. universa*) as compared to others (i.e. *G. tumida*). Berger & Piper noted that poorly preserved sediments had higher counts of *G. tumida* than unaltered sediments. Crushing resistance, in addition to solubility and sinking speed, might contribute to this trend, as many of the species that were identified by Berger & Piper as having low preservation potential are also the species that displayed low crushing resistance in this study.

Conclusion

We have shown that modern planktonic foraminifera vary in their resistance to crushing, as measured by the mean compressional force at mechanical failure. Biconvex, keeled and conical forms, morphotypes that convergently evolved multiple times in the evolutionary history of planktonic foraminifera, are significantly more resistant to crushing than the other morphologies tested. Such resistance could increase the viability of living populations under selective predation, thereby providing a driving force behind the repeated evolution of certain morphologies.

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Table 3. Taphonomic potential

	Sinking speed (1 = slowest)	Solubility (1 = most soluble)	Crushing resistance	Relative preservation potential
<i>Globigerinoides conglobatus</i>	5	2	Medium	Medium
<i>Globoconella inflata</i>	7	5	Medium	Medium
<i>Globorotalia tumida</i>	8	8	High	High
<i>Menardella menardii</i>	5	6	High	High
<i>Neogloboquadrina dutertrei</i>	6	7	Medium	High
<i>Orbulina universa</i>	1	3	Low	Low
<i>Trilobatus sacculifer</i>	2	1	Low	Low
<i>Truncorotalia truncatulinoidea</i>	3	4	High	Medium

Sinking speed and solution rankings from Berger & Piper (1972), crushing resistance from experimental results in Table 1 (least crushable = high). Relative preservation potential was calculated by combining all three preceding factors into a single score and placing each species into one of three categories.

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References

- Abramoff, M.D., Magalhaes, P.J. & Ram, S.J. 2004. Image processing with ImageJ. *Biophotonics International*, **11**, 36–42.
- Aze, T., Ezard, T.H.G., Purvis, A., Coxall, H.K., Stewart, D.R.M., Wade, B.S. & Pearson, P.N. 2011. A phylogeny of Cenozoic macroperforate planktonic foraminifera from fossil data. *Biological Reviews*, **86**, 900–927.
- Bé, A.W.H. & Hutson, W. 1977. Ecology of planktonic foraminifera and biogeographic patterns of life and fossil assemblages in the Indian Ocean. *Micropaleontology*, **23**, 369–414.
- Berger, W. & Piper, D. 1972. Planktonic foraminifera: differential settling, dissolution, and redeposition. *Limnology and Oceanography*, **17**, 275–287.
- Boulding, E.G. 1984. Crab-resistant features of shells of burrowing bivalves: Decreasing vulnerability by increasing handling time. *Journal of Experimental Marine Biology and Ecology*, **76**, 201–223.
- Caromel, A.G.M., Schmidt, D.N., Phillips, J.C. & Rayfield, E.J. 2014. Hydrodynamic constraints on the evolution and ecology of planktic foraminifera. *Marine Micropaleontology*, **106**, 69–78.
- Chang, K.W. & Hanazato, T. 2005. Prey handling time and ingestion probability for *Mesocyclops* sp. predation on small cladoceran species *Bosmina longirostris*, *Bosminopsis deitersi*, and *Scapholeberis mucronata*. *Limnology*, **6**, 39–44.
- Cifelli, R. 1969. Radiation of Cenozoic planktonic foraminifera. *Systematic Zoology*, **18**, 154–168.
- Coxall, H.K., Wilson, P.A., Pearson, P.N. & Sexton, P.F. 2007. Iterative evolution of digitate planktonic foraminifera. *Paleobiology*, **33**, 495–516.
- Hamilton, C.P., Spero, H.J., Bijma, J. & Lea, D.W. 2008. Geochemical investigation of gametogenic calcite addition in the planktonic foraminifera *Orbulina universa*. *Marine Micropaleontology*, **68**, 256–267.
- Hamm, C.E., Merkel, R., Springer, O., Jurkojc, P., Maier, C., Prechtel, K. & Smatcek, V. 2003. Architecture and material properties of diatom shells provide effective mechanical protection. *Nature*, **421**, 841–843.
- Hemleben, C., Spindler, M. & Anderson, O.R. 1989. *Modern planktonic foraminifera*. Springer, Berlin.
- Holling, C.S. 1959. Some characteristics of simple types of predation and parasitism. *Canadian Entomologist*, **91**, 385–398.
- Jeschke, J.M., Kopp, M. & Tollrian, R. 2002. Predator functional responses: Discriminating between handling and digesting prey. *Ecological Monographs*, **72**, 95–112.
- Katz, C.H. 1985. A nonequilibrium marine predator–prey interaction. *Ecology*, **66**, 1426–1438.
- Kennett, J.P. & Srinivasan, M.S. 1983. *Neogene Planktonic Foraminifera: a Phylogenetic Atlas*. Hutchison Ross, Stroudsburg.
- Lipps, J.H. 1979. Ecology and paleoecology of planktic foraminifera. *Foraminiferal Ecology and Paleocology*. The Society of Economic Paleontologists and Mineralogists: Short Course Notes, **6**, 62–104.
- Loeblich, A.R. & Tappan, H. 1964. *Treatise on Invertebrate Paleontology, Part C: Protista 2 – Sarcodina, Chiefly 'Thecamoebians' and Foraminiferida*. Moores, R.C. (ed.) Geological Society of America and University of Kansas Press.
- Marsalek, D.S., Wright, R.C. & Hay, W.W. 1969. The function of the test in foraminifera. *Transactions Gulf Coast Association of Geological Societies*, **19**, 342–352.
- Marshall, B.J., Thunnell, R.C., Spero, H.J., Henahan, M.J., Lorenzoni, L. & Astor, Y. 2015. Morphometric and stable isotopic differentiation of *Orbulina universa* morphotypes from the Cariaco Basin, Venezuela. *Marine Micropaleontology*, **120**, 46–64.
- Morard, R., Quillevère, F., Escarguel, G., Ujiie, Y., de Garidel-Thoron, T., Norris, R.D. & de Vargas, C. 2009. Morphological recognition of cryptic species in the planktonic foraminifer *Orbulina universa*. *Marine Micropaleontology*, **71**, 148–165.
- Norris, R. 1991. Biased extinction and evolutionary trends. *Paleobiology*, **17**, 388–399.
- Pearson, P.N., Evans, S.L. & Evans, J. 2015. Effect of diagenetic recrystallization on the strength of planktonic foraminifer tests under compression. *Journal of Micropaleontology*, **34**, 59–64, <http://doi.org/10.1144/jmpaleo2013-032>
- Smetacek, V. 2001. A watery arms race. *Nature*, **411**, 745.
- Spezzaferri, S., Kucera, M. *et al.* 2015. Fossil and genetic evidence for the polyphyletic nature of the planktonic foraminifera 'Globigerinoides', and description of the new genus *Trilobatus*. *PLoS One*, **10**, e0128108.
- Stevens, M.H.H. 2009. *A Primer of Ecology with R*. Springer, New York.
- Sun, X., Corliss, B.H., Brown, C.W. & Showers, W.J. 2006. The effect of primary productivity and seasonality on the distribution of deep-sea benthic foraminifera in the North Atlantic. *Deep Sea Research I*, **53**, 28–47.
- Thompson, D.W. 1961. *On Growth and Form*. Cambridge University Press, Cambridge.
- Tyszk, J. 2006. Morphospace of foraminiferal shells: Results from the moving reference model. *Lethaia*, **39**, 1–12.
- Tyszk, J. & Topa, P. 2005. A new approach to modeling foraminiferal shells. *Paleobiology*, **31**, 522–537.
- Werner, E.E. & Hall, D.J. 1979. Foraging efficiency and habitat switching in competing sunfishes. *Ecology*, **60**, 256–264.
- Whetmore, K.L. 1987. Correlations between test strength, morphology and habitat in some benthic foraminifera from the coast of Washington. *Journal of Foraminiferal Research*, **17**, 1–13.
- Whetmore, K.L. & Plotnick, R.E. 1992. Correlations between test morphology, crushing strength, and habitat in *Amphistegina gibbosa*, *Archaias angulatus*, and *Laevipeneroplis proteus* from Bermuda. *Journal of Foraminiferal Research*, **22**, 1–12.
- Wiebe, P.H., Burt, K.H., Boyd, S.H. & Morton, A.W. 1976. A multiple opening/closing net and environmental sensing system for sampling zooplankton. *Journal of Marine Research*, **34**, 313–326.