Coordinated shifts to non-planktotrophic development in spatangoid echinoids during the Late Cretaceous

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Despite widespread interest in the interplay between evolutionary and developmental processes, we still know relatively little about the evolutionary history of larval development. Many clades exhibit multiple shifts from planktotrophic (feeding) to non-planktotrophic (non-feeding) larval development. An important question is whether these switches are scattered randomly through geological history or are concentrated in particular intervals of time. This issue is addressed using the Cretaceous spatangoid sea urchins, which are unusual in that larval strategy can be determined unambiguously from abundantly fossilized adult tests. Using a genus-level phylogeny, we identify five clades of non-planktotrophic taxa, each of which first appears in the fossil record in the Campanian or Maastrichtian (the final two Cretaceous stages). No examples of non-planktotrophy have been identified in any of the earlier stages of the Cretaceous echinoid clades, she found that the first known occurrence of non-planktotrophy in five different orders lay in the final two stages of the Cretaceous (the Campanian and Maastrichtian), with no examples of non-planktotrophic echinoids known from earlier than this interval. Despite this study, little is known about the finer scale patterns within orders and at lower taxonomic levels. The present study addresses this by focusing in detail on a single echinoid order, the Spatangoida.

2. MATERIAL AND METHODS

(a) Choice of model organism

The Cretaceous spatangoid echinoids were chosen as the model organism for this study as they are ideal for a number of reasons: (i) they are unusual in that larval strategy can be inferred from the fossilized adult test using either morphological criteria or crystallographic analysis of the apical system (discussed subsequently); (ii) they are abundantly preserved, usually with the apical system present, providing a large volume of suitable material for analysis; (iii) their complex morphology makes it relatively straightforward to establish their phylogenetic relationships; and (iv) the group contains both planktotrophic and non-planktotrophic taxa and spans the time when non-planktotrophy is thought to have first evolved within the group.

(b) Inferring modes of larval development

Larval modes were determined for 87 fossil spatangoid specimens from museum collections that span the early history of the group (see electronic supplementary material for details). Three criteria were used for inferring the mode of larval development from adult fossil sea urchin specimens.

(i) Identification of brood pouches (marsupia). Some non-planktotrophic taxa brood their larvae and a subset of these do so in specialized brood pouches on the test of the female; the identification of such marsupia is indicative of non-planktotrophic development (e.g. Kier 1969).

(ii) Extreme sexual dimorphism of gonopore size. Because non-planktotrophic taxa have much larger eggs (Emlet et al. 1987; Emlet 1989), the females frequently have enlarged development to non-planktotrophic development. This has been inferred by mapping larval modes onto phylogenetic trees of various taxa (e.g. echinoids (Wray 1992; Smith et al. 1995); temnopleurid echinoids (Jeffery & Emlet 2003; Jeffery et al. 2003); asteroid starfish (Hart et al. 1997); Conus gastropods (Duda & Palumbi 1999); litorinid gastropods (Reid 1989); turritellid gastropods (Lieberman et al. 1993)). We now know a great deal about the developmental and genetic changes involved in these switches owing, in particular, to the large body of work on the non-planktotrophic echinoid Helicodiscus erythrogramma and the closely related Helicodiscus tuberculata, which has planktotrophic development (reviewed by Raff & Byrne 2006). However, an important, and yet much overlooked, issue is when these switches to non-planktotrophy occurred: were shifts to non-planktotrophy scattered randomly through time, or were they instead concentrated in particular intervals of geological history? The question is important as it may help to determine the factors that drive shifts to non-planktotrophy. If the switches were concentrated in particular time intervals, then this would strongly imply that extrinsic factors operating at these times are responsible for driving the switches to non-planktotrophy.

The tacit assumption among biologists appears to be that the switches were scattered randomly throughout geological history (Jeffery 1997). However, when Jeffery (1997) carried out a broad survey of all major echinoid clades, she found that the first known occurrence of non-planktotrophy in five different orders lay in the final two stages of the Cretaceous (the Campanian and Maastrichtian), with no examples of non-planktotrophic echinoids known from earlier than this interval. Despite this study, little is known about the finer scale patterns within orders and at lower taxonomic levels. The present study addresses this by focusing in detail on a single echinoid order, the Spatangoida.
The early Campanian, and planktotrophy is the only non-planktotrophy extend no further back in time than or Maastrichtian. The known ranges of taxa exhibiting appear in the fossil record in either the Campanian spatangoids. All five non-planktotrophic clades first arose independently five times within the Cretaceous results reveal that non-planktotrophic development

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Figure 1. Strict consensus tree of Cretaceous spatangoid genera plotted against the geological time scale. Planktotrophic taxa are shown in red, non-planktotrophic taxa in blue and unknown development in grey. Coloured circles represent specimens for which development has been inferred. The green band highlights the Campanian to Maastrichtian interval. Analysed specimens of Holanthus, Cyclaster and Orthaster are Danian in age (see electronic supplementary material for details of phylogenetic analysis and specimens studied).

gonopores through which to extrude these eggs. Extreme sexual dimorphism can indicate non-planktotrophic development (Emlet 1989), although gonopores can also be enlarged by dissolution (S. K. Donovan 2008, personal communication).

(iii) Crystallographic orientation of the apical plates. Larval mode can be inferred from the orientation of the crystallographic axes of the plates in the apical system (Emlet 1985, 1989). This is possible because planktotrophic larvae have skeletal calcite rods, the main function of which is to support the larval arms that are used in feeding; during metamorphosis some of the apical plates grow from the proximal ends of the rods, imparting characteristic c-axis orientations to the plates. In non-planktotrophic larvae, these rods are absent and the apical plates all form de novo at metamorphosis, producing a distinct pattern of c-axis orientations.

c Determining number and timing of switches

The inferred larval modes were mapped onto a novel phylogeny of the group (see electronic supplementary material for details of analysis). This allowed determination of the number and direction of switches in developmental mode within the group. This information was combined with the stratigraphic ranges of the taxa to elucidate the timing of these shifts in developmental mode.

3. RESULTS

The data on larval modes inferred from morphological and crystallographic criteria were mapped onto the phylogeny of Cretaceous spatangoids, which is plotted against the geological time scale in figure 1. These results reveal that non-planktotrophic development arose independently five times within the Cretaceous spatangoids. All five non-planktotrophic clades first appear in the fossil record in either the Campanian or Maastrichtian. The known ranges of taxa exhibiting non-planktotrophy extend no further back in time than the early Campanian, and planktotrophy is the only strategy observed prior to this. The Campanian to Maastrichtian is also the interval during which the first known instance of non-planktotrophy in four other echinoid orders has been recorded; there are no reported occurrences of non-planktotrophy in echinoids from any time earlier than this (Jeffery 1997).

4. DISCUSSION

While this study identifies five shifts to non-planktotrophic development during the Late Cretaceous, Peterson (2005) recognized that switches occurred in the opposite direction at least four, and perhaps as many as eight, times between the Late Cambrian and Middle Ordovician. Taken together, these studies show that switches in developmental mode are, at least in some circumstances, concentrated within particular intervals of geological history. This pattern of coordinated shifts in independent clades strongly implies that extrinsic factors operating at the times of the shifts were responsible for driving these switches in developmental mode. The two most likely external driving mechanisms are predation (Peterson 2005) and environmental change (e.g. Jeffery 1997). While the data to quantitatively test which factor is most likely to be responsible are not yet available, it is possible to elucidate this issue qualitatively by assessing whether major changes in either proposed factor occurred at the time of the developmental switches.

Having argued that the radiation of epifaunal suspension feeders in the Late Cambrian to Middle Ordovician was most probably responsible for the coordinated switches to planktotrophy that occurred at this time, Peterson (2005) argued that benthic
predation was also likely to have driven later switches to non-planktotrophy. The basis for this argument is that adaptations associated with non-planktotrophic development (e.g. positive buoyancy) also serve to protect the larva from predation. In addition, it has been shown that reducing time to metamorphosis is advantageous in environments with high levels of predation (Wray 1995).

While predation may well account for some of the selection pressure to evolve non-planktotrophy, it is unlikely to account for the coordinated switches to non-planktotrophic development that we have identified in sea urchins at the end of the Cretaceous. If benthic predation were responsible, then we would expect to see a major radiation of benthic predators coinciding with the shifts. However, while there is little literature on change in predation rate in the Late Cretaceous, it seems that any increases in predation occurred earlier in the Cretaceous than the interval in question (Kosnik 2005).

The other factor that has been suggested as a driving mechanism for switches to non-planktotrophy is environmental change (Jeffery 1997). In the Cenomanian and Maastrichtian, the climate was changing dramatically. Sea surface temperature was falling rapidly (e.g. Gale 2000) and, while total primary productivity was increasing (Faul et al. 2003), there is evidence that this was associated with increasing climatic seasonality (Steuber 1996; Francis & Poole 2002; Steuber et al. 2005; Dutton et al. 2007) and vigorous seasonal upwelling (Handoh et al. 2003). These changes would have led to the nutrient supply available to planktotrophic larvae becoming more abundant yet less reliable. Modern invertebrates living in areas with intermittent nutrient supply, such as strongly seasonal regions, tend to have evolved either so that their reproductive cycle coincides with the nutrient blooms, or else so that they develop with non-planktotrophic larvae and thus become independent of the nutrient supply (e.g. Picken 1980). The fact that the coordinated shifts in developmental mode that we have identified coincide with these major environmental changes (but not with a marked change in predation—the other proposed driver) identifies environmental factors as the most plausible driving mechanism for these switches to non-planktotrophic development. Thus, it seems likely that there are intrinsic links between major developmental change and major environmental change.

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