

Testing character-evolution models in phylogenetic paleobiology: a case study with Cambrian echinoderms

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1 Abstract

Macroevolutionary inference has historically been treated as a two-step process, involving the inference of a phylogenetic tree, and then inference of a macroevolutionary model using that tree. Newer models, such as the fossilized birth-death model, blend the two steps. These methods make more complete use of fossils than the previous generation of Bayesian phylogenetic models. They also involve many more parameters than prior models, including parameters about which empiricists may have little intuition. In this paper, we set forth a framework for fitting complex, hierarchical models. We ultimately fit and use a joint tree and diversification model to estimate a dated phylogeny of the Cincta (Echinodermata), a morphologically distinct group of Cambrian echinoderms that lack the five-fold radial symmetry characteristic of extant members of the phylum. Although the phylogeny of cinctans remains poorly supported in places, we show how models of character change and diversification contribute to understanding patterns of phylogenetic relatedness and testing macroevolutionary hypotheses. Finally our new analysis raises interesting questions about how incorporating age information is expected to affect a phylogeny, and provides a framework for future systematic and macroevolutionary studies of cinctan echinoderms.

2 Introduction

Historically, drawing macroevolutionary inferences from phylogenetic trees has been a two-step process (Harvey and Pagel, 1991). First, a researcher would estimate a phylogenetic tree from a matrix of phylogenetic characters. Then, they would use that tree (or a set of trees, such as a posterior sample) to fit a macroevolutionary model. Over the past decade, models that blend macroevolutionary inference with phylogenetic inference have become increasingly common. For example, the fossilized birth-death process is used to estimate dated phylogenetic trees (Stadler, 2011; Heath et al., 2014). This process is usually implemented as a Bayesian hierarchical model, in which one model describes the process of character change for phylogenetic characters, one describes the distribution of evolutionary rates over the tree, and one describes the process of speciation, extinction, and sampling that lead to the observed tree (see Wright and Warnock in this issue for a more complete discussion of this). In this manuscript, we describe an approach to fitting complex hierarchical models using a focal dataset of cinctan echinoderms.

We can divide macroevolutionary hypotheses into two non-mutually exclusive groups: those making predictions about origination and extinction dynamics, and those making predictions about rates and modes of trait evolution. The latter group includes hypotheses about shifts in rates of anatomical change and hypotheses about driven trends in which particular character states become more (or less) common over time. Hypotheses predicting such patterns come both from macroecological theory and from evolutionary-developmental theory, and thus span a range of basic issues including developmental, ecological, and physical constraints, and selection (Valentine et al., 1969; Valentine, 1980). Research programs dedicated to assessing shifts in rates and modes of anatomical evolution have been a staple of quantitative paleobiology since the early 1990's. Accordingly, anatomical character evolution models that describe the predictions of these different macroevolutionary hypotheses have important theoretical implications for these endeavors.

Phylogeneticists have long been interested in the same sorts of character evolution models, albeit for very different reasons. Hypotheses of phylogenetic relationships make exact predictions about character state evolution among taxa given observed data and models of character change (e.g., Kimura (1980); Felsenstein (1981); Hasegawa et al. (1985); Tavaré (1986)). The most common phylogenetic model for morphology makes the assumption of time-invariant models with no biases in the rate of character acquisition and loss (Lewis, 2001). The expectations of character evolution, of associations of characters with one another, and disparities between taxa are very different when rates of acquisition and loss vary among characters and with time. This is particularly true when we include divergence times as part of phylogenetic hypotheses (Huelsenbeck et al., 2000; Sanderson, 2002; Drummond et al., 2006): but it is still true if we worry only about general cladistic relationships (i.e., which taxa are most closely related to each other; see Felsenstein (1981); Nylander et al. (2004); Wright et al. (2016)). In other words, many of the conceptual mice that paleo-

biologists seek to capture are the conceptual mouse-traps that systematists seek to use to capture phylogenies.

Many readers' first instincts will be that this presents paleobiological phylogeneticists with a quandary: which comes first, the character evolution models or the phylogenetic inference? Part of the dilemma here stems from a historical view that we should treat phylogenetic analysis and macroevolutionary analysis as two separate endeavors (e.g., Harvey and Pagel (1991)). When we estimate a phylogenetic tree, we typically need to make simplifying assumptions for tractability of the analysis. For our comparative methods, we are often exploring more complex models, possibly even seeking to falsify those same simplifying assumptions. Here, we advocate a very different approach that stems from hierarchical Bayesian phylogenetic approaches. That is, we should not view phylogenetic analysis and macroevolutionary analysis as two independent projects, but instead as two parts of the same endeavor of unravelling the evolutionary history of fossil taxa. These evolutionary histories include when clades and lineages diverged, the consistency of character change rates, biases in state acquisition, the process of diversification that lead to the observed tree, and (of course) exactly how taxa were related to each other. Along the same lines, we have to accept and even embrace the fact that there will always be some degree of uncertainty in all of these things. These uncertainties are not reason to abandon the endeavor as hopeless; on the contrary, it will mean that those conclusions that we can reach do not assume that specific historical details are true.

In this work, we will provide an example of the approach that we are advocating using a series of analyses of the *Cincta*, an extinct clade of 'carpoid' echinoderms from the middle Cambrian. We will detail how paleobiologists can adapt different clock models and character state evolution models initially devised to accommodate uncertainties in molecular evolution to represent and model macroevolutionary hypotheses. In doing so, we will also outline protocol that paleontologists can replicate to conduct analogous analyses on other clades. We will emphasize how the combination of Markov Chain Monte Carlo analyses and stepping-stone tests allow us to marginalize specific details of character evolution models and phylogenetic relationships in order to generate the best joint summary of a clade's evolutionary history. Because there are innumerable possible models that one might consider, we will draw attention to existing methods with which paleontologists might already be familiar that should be useful for suggesting particular models as worthy of consideration. Finally, we will briefly outline other theoretical and methodological areas that remain for paleobiologists and systematists to resolve and unite in the future.

3 Taxonomic Background and Data

3.1 *Cincta*: an enigmatic clade of Cambrian echinoderms

Echinoderms are a diverse phylum of marine animals represented today by more than 7,000 living species (Brusca and Brusca, 2003) distributed among five extant classes, including sea stars, brittle stars, echinoids, sea cucumbers, and crinoids. However, the spectacular diversity of extant echinoderms, measured by both species richness and anatomical variety, represents a paltry fraction of their prodigious evolutionary history recorded in the fossil record. Because echinoderms possess a mineralized endoskeleton made of high-magnesium calcite (calcium carbonate) and occur in virtually all habitats across the spectrum of marine depositional environments, the echinoderm fossil record is spectacularly complete and reveals approximately 30 clades distributed among 21 taxonomic classes spanning the entire Phanerozoic Eon (Sprinkle and Kier, 1987; Sumrall, 1997; Sumrall and Waters, 2012; Zamora and Rahman, 2014; Wright et al., 2017; Sheffield and Sumrall, 2019). Unlike familiar echinoderms inhabiting modern oceans, such as starfish and sea urchins, Cambrian lineages comprise an unfamiliar, taxonomically and morphologically diverse assemblage of predominately stem-group taxa exhibiting a diversity of body plans, life modes, and ecological traits unseen in extant lineages (Sprinkle, 1973; Zamora et al., 2013a; Zamora and Rahman, 2014).

Perhaps no group of early echinoderms has received greater attention and controversy than the ‘carpoids’ (Rahman, 2009). Sometimes called homalozoans or calcichordates in the literature, carpoids comprise a heterogeneous assemblage of extinct echinoderms including ctenocystoids, cinctans (*Homostelea*), solutes (*Homoiostelea*), and stylophorans. Although carpoids possess unique skeletal features that unambiguously identify their echinoderm affinities (David et al., 2000; Bottjer et al., 2006; Rahman, 2009; Zamora et al., 2020), they lack other traits considered synapomorphies of crown-group echinoderms. For example, all extant echinoderms exhibit pentaradial symmetry in adults and possess a water vascular system, unique to the phylum, used for locomotion, respiration, and excretion (Nichols, 1972). In contrast, ‘carpoid’ taxa exhibit bilateral to asymmetrical forms, and it’s debated whether some carpoids possessed a water vascular system (Smith, 2005; Lefebvre et al., 2019). Although the phylogenetic position of carpoids have long been contentiously debated (see Rahman, 2009 and Rahman et al., 2009, and articles cited therein), only recently have computer-based phylogenetic analyses played a major role in evaluating alternative hypotheses (Sumrall, 1997; Smith and Zamora, 2013; Zamora and Rahman, 2014), and only one previous study tested phylogenetic hypotheses using stratigraphic data (Rahman et al., 2009). Crucially, the character matrices constructed for these analyses have greatly benefited from recent improvements to identifying homologous characters across morphologically disparate early echinoderm lineages, often arising from new fossil discoveries (e.g., Zamora et al., 2012; Smith and Zamora, 2013). Taxonomic controversy remains (David et al., 2000), though both recent computational phylogenetic analyses and stratigraphic congruence

metrics support the hypothesis that carroids comprise a paraphyletic assemblage of stem-group echinoderms (Rahman et al., 2009; Smith and Zamora, 2013; Zamora and Rahman, 2014). If this view is correct, then carroids help document the radical transition in echinoderm evolution from an ancestral, bilateral body plan to the pentaradial symmetry characteristic of crown-group forms that have dominated marine ecosystems since the close of the Cambrian. Regardless of their specific branching relationships in the echinoderm tree of life, it is nevertheless clear that understanding the distribution of character combinations and patterns of trait evolution in these enigmatic, pre-radial echinoderm lineages are critical to deciphering the early evolution of the phylum.

Cinctans are a significant group of non-radiate, carroid echinoderms temporally restricted to the middle Cambrian (Miaolingian, 509–497 Ma) and paleogeographically restricted to western Gondwana, Avalonia, and Siberia. Cinctans are generally small (i.e., several 10^{-3} to 10^{-2} m in length), flattened, symmetrical to irregularly shaped fossils resembling a tennis racquet or badly formed pancake, generally interpreted as an adaptation to an epibenthic, suspension feeding lifestyle (Rahman and Zamora, 2009; Rahman et al., 2015). Like all echinoderms, cinctans have a complex, multi-element, calcitic endoskeleton, which makes them particularly amenable for coding discrete, phylogenetic characters in fossil taxa (Smith and Zamora, 2009). The main body, called the theca, is surrounded by a series of rigid, stout, marginal plates (called the cinctus), which surrounds a central body of smaller, tessellated integument plates on both dorsal and ventral sides. The mouth is a circular opening located at the end of a narrow food groove (or pair of grooves) on the right anterior side of the theca. A posterior appendage, called the stele, forms a rigid structure extending from the cinctus, commonly subequal in length to the theca.

Despite their diminutive size, geological antiquity, and narrow paleogeographic and stratigraphic ranges, the significance of cinctans to understanding early echinoderm evolution, as well as their evolutionary implications surrounding ancestral character states in ancient deuterostomes (Smith and Swalla, 2009), has led to a substantial amount of interest to decipher their paleobiology. Recent advances in cinctan paleobiology include efforts to better understand patterns of taxonomic diversity (Zamora and Álvaro, 2010), ontogeny and development (Smith, 2005; Zamora et al., 2013b), life mode and feeding ecology (Rahman et al., 2009, 2015; Zamora and Rahman, 2015), convergence and adaptive evolution (Zamora and Smith, 2008) and phylogenetic relationships (Friedrich, 1993; Sdzuy, 1993; Smith and Zamora, 2009; Zamora et al., 2013b). In this study, we combine morphological data with fossil age information to re-evaluate phylogenetic relationships and evolutionary dynamics among cinctan lineages using hierarchical Bayesian phylogenetic models and provide a phylogenetic template for future systematic and macroevolutionary studies.

3.2 Character Data

We use the character data initially published by (Smith and Zamora, 2009) and subsequently augmented by (Zamora et al., 2013b). The analyzed matrix

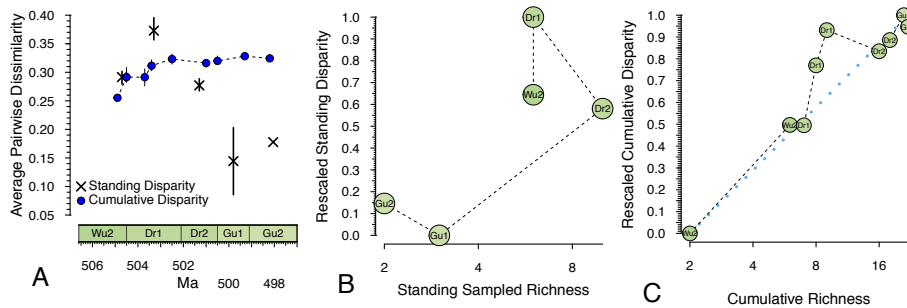


Figure 1: Disparity patterns for cinctans. A. Disparity over time based on average pairwise dissimilarities among taxa (see Foote (1992)). Vertical bars represent 90-percentile error bars from bootstrapping pairwise comparisons (Foote, 1993). X’s give disparity among standing richness only. Circles give the cumulative disparity among all members of the clade at least as old as that point in time. B) Rescaled standing disparity vs. log-taxonomic richness. Because cinctans begin with relatively high richness and then decline over time, the curve begins near the middle of the plot rather than near the bottom as is typical (Jablonski, 2020). C) Rescaled cumulative disparity given cumulative richness. The dashed line gives the expectation given continuous change. Cumulative disparity is generally higher than expected given initial disparity and cumulative richness through the first half of clade history.

includes 22 cinctan species plus one outgroup taxon (*Ctenocystis*, represented by *Ctenocystis utahensis*). An additional four cinctan species are excluded due to inadequate material for coding. We refer the readers to the papers cited above for additional information concerning the character data.

Both disparity analyses of these data conducted and arguments pertinent to early echinoderm evolution in the literature (e.g., Smith et al., 2013) suggests that cinctans may exhibit “Early Burst”-type dynamics (Figure 1A), in which disparity is higher than expected given either standing disparity (Hughes et al., 2013) or cumulative disparity (Wagner and Estabrook, 2015). Standing disparity vs. log-standing richness patterns deviate strongly from expectations given constant rates of change (Jablonski (2020); see also Wright (2017a)), but this reflects in part the clade decreasing in richness through its later history. The same relationship with cumulative disparity (i.e., disparity among all species known by some date) shows a weaker trend towards higher disparity than expected during the first half of clade evolution (Fig. 1C). This in turn suggests that rates of change might have been higher early in clade history (Foote, 1996b).

3.3 Chronostratigraphic Information

Our chronostratigraphic data come from 221 occurrences of Cambrian echinoderm species from 143 localities last downloaded from the Paleobiology Database on 2020-01-01. The locality and occurrence data came from 81 references with

the seven biggest sources including Nardin et al. (2017), Zamora (2009), Chlupac et al. (1998), Sprinkle and Collins (2006), Termier and Termier (1973) and Sprinkle (1973). We entered 108 of those occurrences and 72 of those localities, and updated 51 of the remaining localities for the purpose of this study. After accounting for synonymies and variant spellings, the localities represent 55 different rock units (i.e., formations and formations+members). The accepted names of the species occurrences reflect 361 taxonomic opinions, 155 of which we entered for the purposes of this study.

The Paleobiology Database returns ages based only on the entered interval, which usually is a stage. Here, nearly every cinctan-bearing locality is assigned to the Middle Cambrian and thus receives a possible age of 513-501 Ma. However, PBDB provides information allowing much more exact ages. For example, PBDB collection 67775 is one of four including *Trochocystites bohemicus*. This collection is assigned to the Middle Cambrian, and thus is dated by the PBDB as 513 - 501 Ma. However, it is the collection represents the Skryje Shale, which is known to span four trilobite zones that restrict the age to 505.2 - 500.7 Ma. Thus, if we had no further information, then that would be the oldest and youngest possible ages for this collection. However, PBDB collection 67775 also is assigned to the *Eccaparadoxides pusillus* trilobite zone, which further restricts the age to 505.2 - 504.5 Ma. We use a database of Paleozoic rock units and faunal zones compiled by one of us (P JW) as a thesaurus to provide more exact earliest and latest possible ages for each cinctan-bearing collection. The typical locality now can be restricted to a 0.7 million year window. Prominent sources for the information relevant to our study and for interregional correlations of rock units and trilobite zones include Alvaro et al. (2001); Liñán et al. (2004); Geyer and Landing (2006); Geyer and Shergold (2000); Alvaro et al. (2007); Geyer (2019). The overall timescale is that of Gradstein et al. (2012).

We use the refined dates to put lower and upper bounds on the possible first-appearance dates of the cinctan species (Table 1). For species known from only single intervals or trilobite zones, the lower and upper bounds for both first and last appearances are necessarily identical. This is not the case for species spanning multiple intervals. For example, *Gyrocystis platessa* occurs in rocks as old as the *Badulesia granieri* trilobite zone (504.9-504.5 Ma given Geyer (2019) and Gradstein et al.'s timescale) and also occurs in rocks as young as the *Solenopleuropsis marginata* trilobite zone (501.0-499.3 Ma). Here, the latest first possible appearance is 504.5 Ma. We choose the widest possible uncertainty. For example, *Gyrocystis erecta* occurs in rocks belonging to the *Solenopleuropsis* zone (503.1-501.0 Ma) but also in rocks dated more specifically to the *Solenopleuropsis thoralis* subzone (501.6-501.0 Ma). Because the former set of occurrences might be as old as 503.1 Ma (given existing information), we set the possible FA for *G. erecta* at 503.1 – 501.0 Ma.

Finally, we also use these data to derive initial estimates of origination, extinction and sampling for Cambrian echinoderms. We use a modified version of the Three-Timer method (Alroy, 2015) that uses lognormal distributions for sampling rates per stage slice rather than a single value (Wagner and Marcot, 2013). Note that these are used only parameterize the prior distribution of new

proposals for diversification and sampling in MCMC generations and not as fixed values.

4 Methods

The fossilized birth-death is a hierarchical model, meaning that different model subcomponents explain the evolution of the phylogenetic characters (the morphological evolution model), the distribution of evolutionary rates across the tree (the clock model), and the model that describes the speciation, extinction and sampling events leading to the tree (the tree model). Below, we describe a hierarchical approach to model-fitting, in which we fit a model to each subcomponent. The model subcomponents are then assembled into a total fossilized birth-death process.

For each model subcomponent, we first ran an MCMC in RevBayes to assess how long it takes for the analysis to reach convergence. Then, using this value, we ran 20 stepping-stone replicates to calculate a marginal likelihood for the data. Stepping-stone model fitting samples iteratively in the space between the prior and the posterior. The aim in doing this is to estimate the probability of the data summed over all possible values for parameters (Xie et al., 2011). This enables the calculation of an unbiased marginal likelihood, in contrast to MCMC, which will be biased towards regions of treespace that contain good solutions.

The result of each stepping-stone analysis is a marginal likelihood. Because phylogenetic likelihoods tend to be quite small, they are typically reported as log-transformed values. This means that for model comparisons, we used the log Bayes Factor (Kass and Raftery, 1995), which is represented by the character K , and given via the formula:

$$K = \ln[BF(M0, M1)] = \ln[P(X|M0)] - \ln[P(X|M1)],$$

In the above equation, the Bayes Factor for model comparison between Model 0 and model 1 is equal to the probability of model 0 minus model 1. The final Bayes Factor is calculated by exponentiating K :

$$BF(M0, M1) = e^K$$

The final Bayes Factor is a single value for which a value greater than one constitutes support for model one and a value less than negative one is support for model zero.

Within each model subcomponent, Bayes Factors were used to compare different candidate models. The winning candidate model for each subcomponent was then used to estimate the subsequent FBD trees.

Table 1: Chronostratigraphic information for analyzed taxa based on occurrences in the Paleobiology Database. FA and LA denote first and last appearance dates, with LB and UB giving the oldest and youngest possible FA and LA given known finds. “NS” gives number of collections (= collections or localities) that a species occupies. “NR” gives number of rock units (formations or members) that a species occupies. For *Ctenocystis*, FA and sampling data are for the oldest known species (*C. utahensis*) only, whereas the LA data are for the entire genus.

Taxon	FALB	FAUB	LALB	LAUB	NS	NR
<i>Ctenocystis</i>	506.6	506.5	501.0	500.5	4	3
<i>Gyrocystis platessa</i>	504.9	504.5	501.0	499.3	13	4
<i>Gyrocystis testudiformis</i>	503.1	502.5	503.1	502.5	4	1
<i>Gyrocystis cruzae</i>	503.1	501.0	503.1	501.0	1	1
<i>Gyrocystis badulesiensis</i>	503.1	501.0	503.1	501.0	2	1
<i>Gyrocystis erecta</i>	503.1	501.0	501.6	501.0	2	1
<i>Progyrocystis disjuncta</i>	503.1	501.0	503.1	501.0	1	1
<i>Protocinctus mansillaensis</i>	506.6	505.4	506.6	505.4	1	1
<i>Elliptocinctus barrandei</i>	501.6	501.0	501.6	499.3	6	3
<i>Elliptocinctus vizcainoi</i>	504.5	503.4	504.5	503.4	1	1
<i>Sucocystis theronensis</i>	501.6	501.0	501.6	501.0	2	2
<i>Sucocystis bretoni</i>	501.0	500.5	501.0	500.5	1	1
<i>Lignanicystis barriosensis</i>	501.6	501.0	501.6	501.0	3	1
<i>Undatacinctus undata</i>	501.0	499.3	501.0	499.3	1	1
<i>Sucocystis acrofera</i>	499.3	498.2	499.3	498.2	2	1
<i>Undatacinctus quadricornuta</i>	501.0	499.3	501.0	499.3	1	1
<i>Undatacinctus melendezi</i>	501.0	499.3	498.2	497.0	9	2
<i>Asturicystis jaekeli</i>	504.9	504.5	504.9	504.5	1	1
<i>Sotocinctus ubaghisi</i>	505.4	504.9	505.4	504.5	2	2
<i>Trochocystites bohemicus</i>	505.2	504.5	503.0	502.2	4	3
<i>Trochocystoides parvus</i>	504.5	503.7	504.5	503.7	1	1
<i>Ludwigicinctus truncatus</i>	501.6	500.5	501.6	500.5	1	1
<i>Graciacystis ambigua</i>	504.9	504.5	503.7	503.1	3	1

4.1 Morphological Evolution Models

We first fit a morphological character model, as no tree can be estimated without one. We compared three models for morphological character evolution. All three were based on the basic Mk model (Lewis, 2001). In this model, it is assumed that any character has an equal probability change and reversal between any two states. The data matrix was partitioned according to the number of character states, so that size of the transition matrix of the model was correctly specified. In the first model, we did not allow rate heterogeneity. In effect, this means we assume all characters to have the same rate of evolution. In the second, used Gamma-distributed rate heterogeneity to allow different characters in the matrix to have different evolutionary rates. A common concern is that characters directly involved with basic organismal ecology and function such as those involved in feeding might evolve under different rules than do other characters (Foote, 1994, 1996a; Wagner, 1995; Sánchez-Villagra and Williams, 1998; Ciampaglio, 2002; Hopkins and Smith, 2015; Wright, 2017a). Because Smith and Zamora (2009) explicitly identify characters related to the “food groove” anatomy of cinctans, our third model partitions the data into feeding and non-feeding characters. Then the above model was applied in each partition. In effect, the third model has two times as many parameters as the first, one set for each set of characters.

Our MCMC analyses reached convergence after about 80,000 generations, as checked in the software Tracer (Rambaut et al., 2018). Stepping stones should generally be run to approximate convergence per stone. Therefore, each stepping stone was run for 100,000 generations to account for any late-converging stones.

4.2 Clock Models

A phylogeny cannot be estimated without a model of character evolution. Hence, the morphology model was fit first. Next, we fit a clock model. Although “clock” might conjure images of a constant rates model, that is only one type of clock (i.e., a “Strict Clock”): clock models include a range of models that either directly predict or at least constrain plausible rates of change. In order to do this, we used our best-fit morphology model and a simple, time-homogeneous FBD model to compete different clock models against one another. The four candidate clock models were as follows.

- A strict clock: In this model, the rate of evolution along each branch is assumed to be equal. The rate of evolutionary change is sampled from an exponential distribution. Note that the strict clock model is most simple clock model, and also (an) equivalent to null models used in paleontological rate studies. It assumes that all branches follow a single, constant rate of morphological evolution. Although simplistic, some studies have found surprising degree of concordance with fossil data fitting a strict morphological clock even when models incorporating rate variation provide a better statistical fit (Drummond and Stadler, 2016; Wright, 2017b) The strict clock model has one advantage in its simplicity: it adds only one

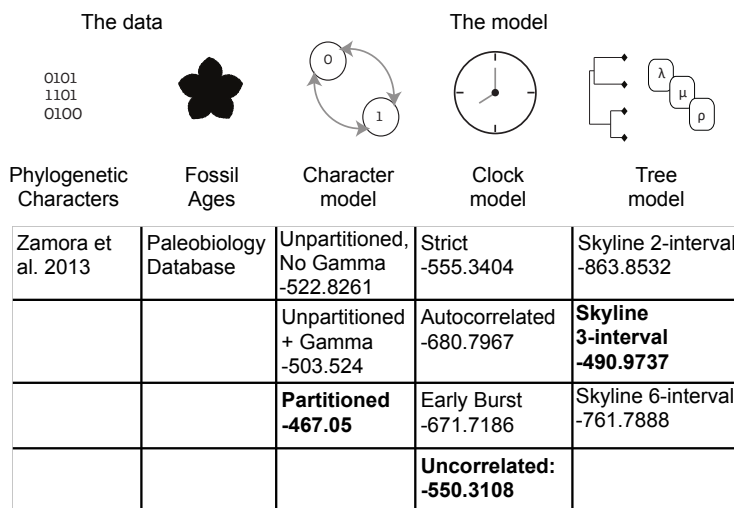


Figure 2: A table of the competed models for each component of the FBD process. Underneath each model component are the models competed for that component. The model indicated in bold text is the one that fit the data best, per Bayes Factor model selection.

parameter to the analyses, whereas relaxed clock models require many additional parameters.

- An uncorrelated lognormal clock: This clock treats each branch as an independent draw from a distribution (Drummond et al., 2006; Drummond and Rambaut, 2007). In this case, we used a lognormal distribution, which says most evolutionary rates are likely to be low, but with allowances for some branches to have very high rates. It should be noted that because each branch is a separate draw, the rate of an ancestral branch’s evolution may be very different than its descendants - either greater or lesser. In terms of macroevolutionary theory, an uncorrelated clock model is consistent with there being no shifts in intrinsic constraints on rates of change within a clade, but where there is considerable heterogeneity in the effects of ecology on rates of change among different lineages.
- Autocorrelated clock: These clocks assume that the rate of evolution on a descendent branch is drawn from a distribution centered on the rate of evolution of that branch’s ancestor (Aris-Brosou and Yang, 2002). Here, the rate is heritable, but constantly changing in a manner analogous to a morphometric character under continuous change. Theoretically, we might expect this if rates are affected by some other variable that is continuously changing (e.g., climate variables or biological variables such as metabolism or size, see further discussion in Bromham et al. (1996); Gaut et al. (1992); Thomas et al. (2006); Bromham et al. (2015)). Thus, this will favor smaller

rate shifts than those seen in an uncorrelated clock. We modeled the distributions per branch as lognormally-distributed.

- **Early Burst:** This clock models rates of character change as exponentially decaying over time. This assumes that rates of evolution are fastest near the base of the tree, and decline into the present. As illustrated above, disparity patterns within the clade also suggest this (Fig. 1). Prior work has sought the question of detecting radiation in a phylogenetic context (Liow et al., 2010). This model builds on this idea while explicitly including fossils (Quental and Marshall, 2009, 2010).

Each of these models has a different number of parameters and took a different amount of time to converge. Therefore, for each model, we first ran an exploratory MCMC to see how long convergence takes. Then we used the convergence value to choose the number of iterations per stepping stone. A table of compared models can be seen in Figure 2.

4.3 Tree Models

In all of our comparisons of tree models, we used variants of the fossilized birth-death model. We compared several models, reflecting different scenarios of diversification and sampling in the group. The simplest model treats these rates as constant over time. Of course, innumerable paleobiological studies indicate that origination, extinction and sampling all vary over time within clades. Shifts in these rates means that the prior probability of a branch spanning 1 million years is not constant throughout clade history (Wagner, 2019). Skyline models (e.g., Stadler et al., 2013) treat this as a possibility within FBD analyses by allowing all three rates to vary in different time-intervals. We contrasted several skyline models. These models assume that the parameters of the FBD analysis can vary between discrete time bins. The cinctan fossil record spans three geological stages of the middle Cambrian: the Wulian, Drumian, and Guzhangian. Most of the known species appear in the late Wulian and Drumian, with a marked decrease in the Guzhangian (Table 1). This suggests temporal variation in origination and/or extinction rates. Therefore, we allowed all analytical parameters to vary between all three geological stages. It should be noted that for all skyline models, there is an additional interval of time from the origin to the first interval with its own possible origination, extinction and sampling rates.

- **Time-homogeneous:** The first FBD model is a time-homogeneous model in which it is assumed that one rate of speciation, extinction, fossil sampling and sampling at the last occurrence time apply to the whole tree.
- **Two intervals:** We tested a model in which the Drumian stage is split into two stages, for a total of two skyline categories (Wuliuan & Drumian 1, Drumian 2 & Guzhangian).

- Three intervals: We tested a model in which each stage is given its own set of FBD parameters, for a total of three skyline categories.
- Six intervals: In this model, we allowed each stage-slice to have its own rates.

Although most prior FBD analyses treat origination and extinction as independent variables, paleobiological studies show that the two are closely correlated (e.g., Marshall, 2017). Over long periods of time, the relationship is nearly linear within large clades (e.g., Figure 3A, illustrating prominent Cambrian to Silurian clades). There is more variation with clades over short periods of time such as the stage-slices that we use for these analyses (Figures 3B-C). However, there is still a distinct lognormal relationship between origination and extinction. Moreover, the lognormal relationship for only Cambrian stage-slices or for echinoderms by stage-slice fits the same overall lognormal relationship well. Thus, our MCMC searches vary origination as an independent variable, and vary turnover (extinction/origination) as a variable dependent on origination. For the time-homogeneous model, this reflects the linear relationship shown in Figure 3A (where turnover varies from 0.90 to 1.05 times origination). For skyline models, turnover follows the lognormal relationships illustrated in Figures 3B-C.

Finally, for all competed models, the best-fit character change model and clock model were used as the other model subcomponents.

5 Results

5.1 Model fitting

Model selection supported the choice of substitution model with feeding and non-feeding characters (posterior probability: -467.053) modeled separately (log Bayes Factor: -1.078; substantial evidence). An uncorrelated lognormal clock (posterior probability: -550.311) was favored over an autocorrelated clock (posterior probability: -680.797) or a strict clock (posterior probability: -555.3404) with a log Bayes Factor of 1.562 (substantial evidence). Finally, the three-time interval model (posterior probability: -490.9737) was supported over the two-interval (-863.8532) and six-interval (-761.7888) models. A schematic of the model competed with the best-fit models highlighted can be seen on Fig. 2. Parameters of the best-fit FBD model can be seen on Table 2.

5.2 Cinctan phylogeny

The phylogeny estimated can be seen in Fig. 2. As expected, *Ctenocystis* is sister to the Cinctans. *Protocinctus*, which has been recovered in some recent studies as nested deep within the cinctans (Smith and Zamora, 2009), is recovered here as sister to the rest of the clade. The genus *Gyrocyrtis* is monophyletic, with several species placed as sampled ancestors within the clade. However,

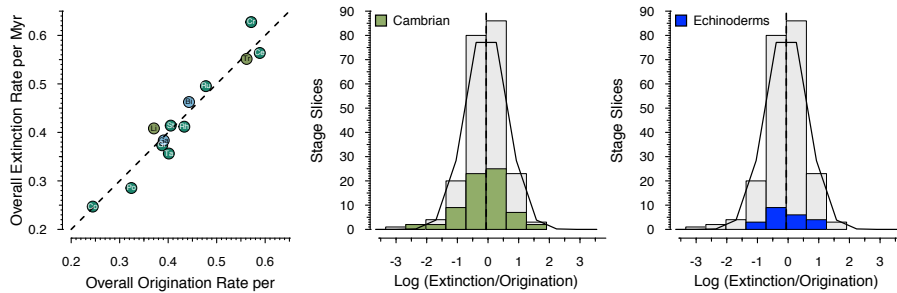


Figure 3: Relationships between origination and extinction rates given birth-death-sampling analyses for 12 clades prominent from the Cambrian through the Silurian. A. Most-likely origination and extinction rates for species over entire histories: trilobites (Tr), linguliform brachiopods (Li), conodonts (Co), poriferans (Po), tabulate corals (Ta), rugosan corals (Ru), rhynchonellate brachiopods (Rh), strophomenate brachiopods (St), cephalopods (Ce), crinoids (Cr), gastropods (Ga) and bivalves (Bi). Olive green denotes members of Sepkoski’s Cambrian fauna (Sepkoski, 1981) dark green denotes members of Sepkoski’s Paleozoic fauna, and light blue denotes members of Sepkoski’s Modern fauna. B & C. Distribution of logged turnover rates for individual clades and individual stage-slices (see Bergström et al. (2009), Cramer et al. (2011) and Rasmussen et al. (2019) for stage-slice definitions.) The best-fit lognormal distribution also is illustrated. B) Turnover rates for Cambrian time-slices separated. C) Turnover rates for crinoids separated. We use 3A to inform the MCMC analyses of range of turnover to consider for time-homogeneous models; we use 3B and 3C to inform analyses of the range of interval-specific turnovers to consider.

Table 2: Diversification parameters of the FBD model. Rates presented as the median of the 95% HPD of the Bayesian posterior sample for the best-fit model, the model in which each geological stage has its own speciation (λ), extinction (μ) and fossil sampling intensity (ψ) in parameters. Turnover = μ/λ ; Diversification= $\lambda-\mu$

Stage	ψ	Diversification	λ	μ	Turnover
Guzhangian	0.188	-0.193	0.493	0.687	2.148
Drumian	0.260	0.317	1.28	0.964	0.714
Wuliuan	0.224	-2.657	0.916	3.574	4.936

Progyrocystis appears in a clade with *Asturicystis* and *Graciacystis*. This clade is sister to the *Gyrocystis*, albeit with low posterior support. *Trochocystoides* and *Trochocystites* are nested deeper in the tree than in prior analyses, and are more closely related to species within the Sucocystidae. Similar to prior studies (Smith and Zamora, 2009; Zamora et al., 2013b), we also recover a clade comprising *Sucocystis*, *Lignanicystis* and *Ellipticintus*, though in this analysis *Sucocystis acrofora* groups with the *Undatacinctus-Ludwigicinctus* clade. The HPD on the age of the origin of this clade is between 505.365 and 507.724 Ma.

6 Discussion

6.1 Model-fitting for complex hierarchical models

When the fossilized birth-death model was first implemented for divergence time estimation, one of the noted benefits was avoiding incoherent fossil calibration points on nodes (Heath et al., 2014). “Incoherent” in here has multiple meanings: first, that fossils are not data under node calibration methods. In a node calibration framework, fossils constrain the possible ages a split can have. The fossil ages ranges are not data under this framework (Gavryushkina et al., 2017). The researcher parameterizes what they believe to be the waiting time between the divergence and this fossil subtending it. This waiting time is capturing two different quantities - the uncertainty around the age of the fossil and how long since the divergence the fossil took to arise. In practice, choice of prior is often subjective, and not based on any one criterion or method, though methods for doing this do exist (Nowak et al., 2013).

The second way in which this practice can result in incoherence is through the collision of priors on different nodes. Depending on the shape of the prior chosen, the upper age bound of an ancestor split may conflict with the lower age bound of its descendant splits. For example, if a researcher has little intuition for when a fossil arose in relation to the split that it subtends, they may place a uniform distribution specifying the longest and shortest the distance between the split and the fossil subtending it may be. Imagine this split and fossil are the descendants of an earlier node, which also has a fossil associated with it. Perhaps the researcher has an intuition that this older fossil is likely close to

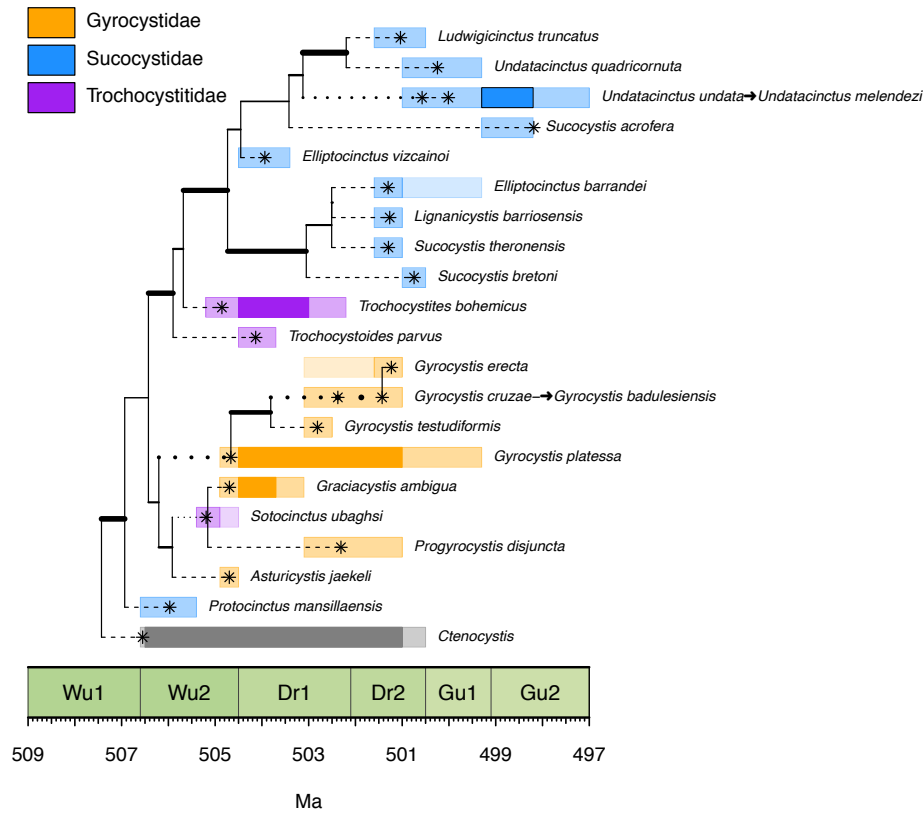


Figure 4: A dated phylogeny of the Cinctans. Branch widths of nodes are proportional to the posterior probability of the branch, with wider branches reflecting higher posterior probability. Branch durations preceding sampled species are dashed lines; dashed nodes denote cases where we reconstruct a sampled species as ancestral to the others. We reconstruct instances where the implied ancestor was still extant when the daughter lineage appears as evidence for budding cladogenesis (see, e.g., Eldredge, 1971). Asterisks denote most probable ages of first appearances. For taxon ranges, solid colors reflect ages for which species have definitely older and definitely younger finds; transparent bars represent range of possible first and last appearances. Extra pale bars represent cases where some of the candidates for oldest/youngest occurrence might be that age. For example, some occurrences of *Gyrocystis erecta* might be 503.1-501.0 Ma whereas others might be 501.6-501.0 Ma.

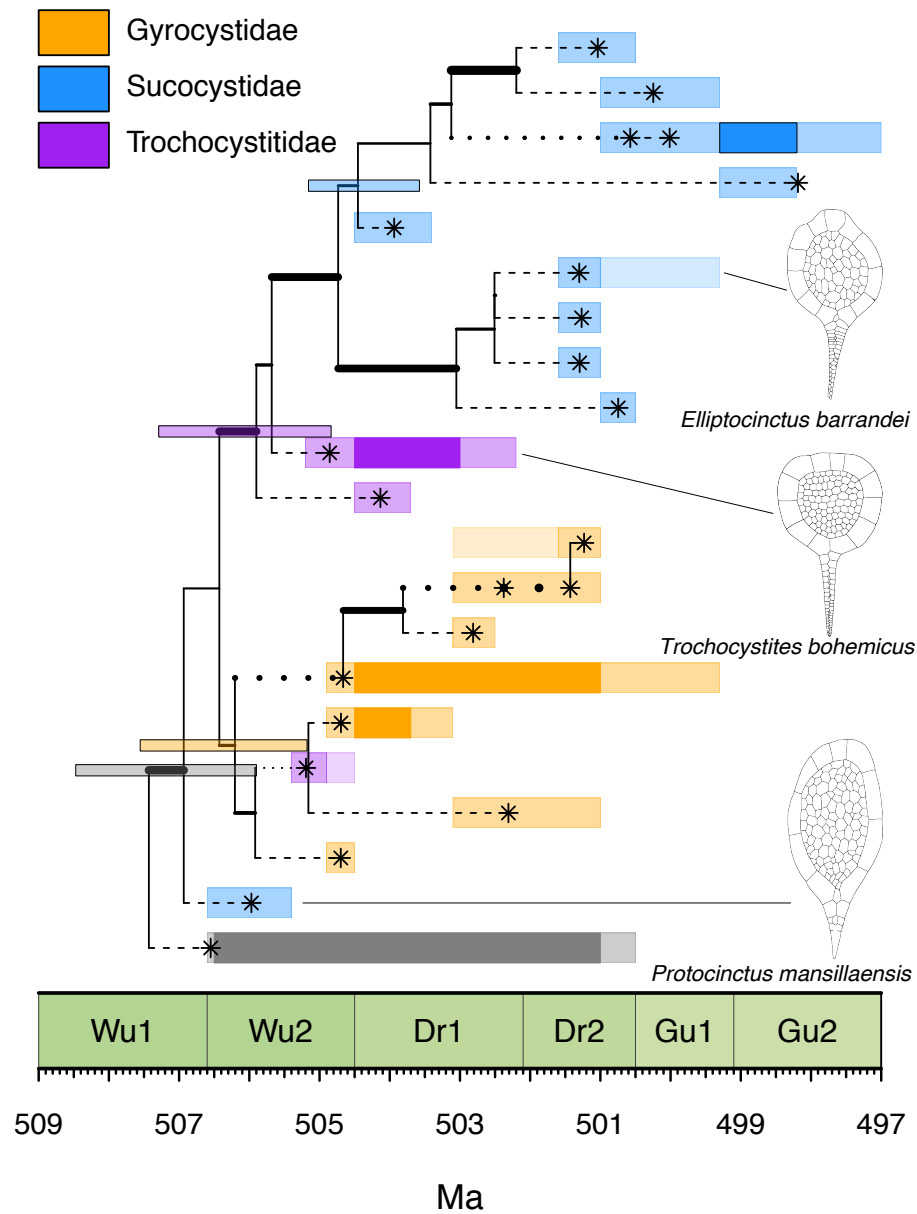


Figure 5: The same phylogeny but with credible intervals given for a few major divergences, include: 1) cinctans from ctenocystoids (gray); 2) gyrocytids from other cinctans (orange); 2) trochocystitids and sucocystids from other cinctans (purple); 4) sucocystids from trochocystitids. (Because this tree reconstructs all prior familial definitions as polyphyletic, the clades corresponding to these divergences represent plausible groupings for future revisions to cinctan higher taxa.)

its ancestor node. And so the researcher places a lognormal prior on the fossil waiting time, saying the fossil is likely close to the node, but allowing for it to possibly be much younger. If incorrectly parameterized, the upper bound of the lognormal could overlap the lower bound of the uniform, implying in those cases that the descendant split could occur before the ancestor split.

This is obviously undesirable. The fossilized birth-death model does not use node calibrations, instead parameterizing the uncertainty of the age associated with each tip. This is done by placing a uniform prior on each fossil tip that begins with the first occurrence of the fossil and ends with the last occurrence. For some taxa, this will mean a fairly wide uncertainty per tip. For example, in the terrestrial realm, fossil insect occurrences are often dated based on the type of amber they were found in (LaPolla et al., 2013). Some ambers can be precisely dated, as the trees that generate the amber has a narrow range. For others, the range of dates might be quite broad as the amber type could be made from multiple trees, or in tree types with long geological persistences (Poinar and Mastalerz, 2000). In fossils that have been individually dated, this uncertainty may correspond to the uncertainty on the radiometric dating. Similarly, fossil age uncertainty is ubiquitous in the marine fossil record, even for well-sampled fossil taxa. For example, few marine fossils are sandwiched between rock units available for fine-scale radiometric dating. Instead, these layers must be correlated to other units using chronostratigraphic methods, which always involve an envelope of uncertainty. Sometimes, the oldest fossil belong to a particular species may occur just above an unconformity (e.g., a sequence boundary), or stratigraphic correlations of fossil-bearing formations may otherwise be highly uncertain and contentious. Moreover, some fossils, particularly those from historical collections, may have been collected from a locality with low-precision stratigraphic data (i.e., “Silurian”), with no further information available to narrow its stratigraphic or temporal precision. Regardless of the manner in which the uncertainty is derived, the meaning is clear and singular: the uncertainty on a fossil tip represents the minimum and maximum plausible age of the fossil. This is a far clearer quantity to describe than uncertainty on a fossil, plus the waiting time between the fossil and the speciation that generated it. Critically, it is important to account for fossil age uncertainty in FBD studies, as not doing so can lead to inaccurate inference of tree topologies, divergences times, or both (Barido-Sottani et al., 2019, 2020b)

However, the fossilized birth-death model still contains parameters for which it may be difficult to choose parameters. It is generally known that a small proportion of life that has ever existed has fossilized. But what should the fossilization rate in any particular clade be? Should it change over time? Model selection has long been considered an important part of phylogenetic inference (Zwickl and Holder, 2004; Allman and Rhodes, 2008; Baele and Lemey, 2013). But in the absence of easy to use selection software (such as the seminal software for molecular model testing, modelTest; Posada and Crandall (1998)), has not been as widely used in other areas systematic research, particularly for divergence time estimation (Duchêne et al., 2015). Here, we have used heirarchical model selection to fit a model for each of the FBD’s component submodels. For

each subcomponent, we competed plausible models. The winning models were combined into a final analysis. Using stepping-stone model estimation, we were able to calculate precise model likelihoods and use them to compare models using the log Bayes Factor.

While this methodology is computationally intensive, it was also tractable. Because no time tree can be inferred without the means to infer a tree first, we first chose our model of morphological evolution. This is also the least computationally-intense part of the estimation, and can be completed in a few hours. Using this model, we then chose a clock model, testing four different models (see the next section for a discussion of these models). Finally, using our morphological evolution and clock models, we competed several versions of the FBD model, including three skyline models. Scoring a precise marginal likelihood for the total tripartite model is the most computationally intensive part of the work. By saving this for last, and first fitting the less parameter-rich morphological evolution and clock models, this estimation is made far more tractable for an average researcher to conduct on a laptop or desktop computer.

6.2 What does the chosen set of models tell us?

Being able to fit a model doesn't mean that fitting that model tells us anything about biology. Ideally, we will use our knowledge of the system to turn that model fit into knowledge. As shown on Fig. 2, we competed several different models of morphological evolution, clock rate distribution across the tree, and the tree model. Each of these models and parameters has meaning in terms of evolution. As the biological or geological interpretation of these phylogenetic models may not be intuitive to geologist readers, we will now examine what we have learned about evolution from this exercise.

The model of morphological evolution is intended to capture how the phylogenetic characters have evolved over time. It is the chief source of information about the topology. The models of morphological evolution we used were all based on the Mk model (Lewis, 2001). In this model, it is assumed that characters can be in any one of k known character states, that each character can change instantly along a branch, and that probability of change between any two states is equiprobable. As shown on Fig. 2, among-character rate variation fits the dataset better than a single rate of evolution across a dataset. This is somewhat unsurprising, as most work in this group has been conducted under parsimony, a model which assumes each character has its own rate of evolution. We also investigated partitioning in this dataset. Prior work has been equivocal about whether "ecological" traits such as feeding structures evolve at higher rates than do other characters, with nearly equal numbers of studies contradicting the notion (Foote, 1994; Sánchez-Villagra and Williams, 1998; Ciampaglio, 2002) and supporting it (Wagner, 1995; Blomberg et al., 2003; Hopkins and Smith, 2015). Here, we find strong support for this hypothesis.

We examined four clock models. The first was a strict clock. These types of clocks rarely are supported in molecular systematics. Molecular evolution rates are impacted by generation times, metabolic rate, and mutation rate. For

a more in-depth review of this concept, see Wright and Warnock (2020) in this issue. How this translates to rates of morphological evolution over time is not as well-studied, but all the above factors are also likely impact the evolution of anatomical form. Therefore, the lack of support for a strict clock in our data (Fig. 2) is unsurprising.

The remaining three clock models describe more biologically interesting scenarios. An autocorrelated clock model implies that a descendant branch will have a rate of evolution that is related to the rate of evolution of its ancestor. This is an appealing model, as we would expect that life history traits that effect possible rates of change may accumulate variation slowly, and be similar to their ancestors. We also examined an early burst model, in which the rate of evolution slows over time. This, too, is an interesting biological hypothesis that is testable given our data. However, both models were less well-supported than the uncorrelated lognormal clock, a model in which large changes in rates of evolution can be seen among ancestor-descendent pairs. It should be noted that while more flexible in terms of the rate variation allowed between ancestors and descendants, the uncorrelated lognormal is not necessarily the most complex model parametrically. The strength of support for the most flexible model suggests that perhaps there is a substantial amount of variation that is not being currently captured by our current generations of clock models. There may be a universe of models awaiting description that could be tapped into to fill this need.

The final model subcomponent is the tree model. We were able to easily reject a time-homogeneous FBD model in which one rate of speciation, extinction, and fossil sampling applies across the whole tree. The entirety of the tree is only a 12 million year span of evolutionary history. Being able to reject one model over a relatively small amount of time implies that variable-rate models might be appropriate for a great many systems. The cinctans appeared in a three-stage slice of the Miaolingian Epoch. One competed model looked at having the Wuliuan, Drumian, and Guzhangian stages having different sets of FBD parameters. Another was a two-stage model in which time was split down the middle in the Drumian. And a third, most complex model in which each stage was split into two intervals was also examined. It is worth noting that discriminate power between these models was fairly good, and that the most complex model was not simply chosen. The three-stage model fit best, followed by the strict clock, seven-stage model, and finally the two-stage model. This is somewhat comforting: if the most complex model had been chosen for each component model, one would wonder if we were not simply choosing from a candidate set of under parameterized models. The Bayes Factor is a reasonably conservative test, and did reject more parameter-rich models in favor of simpler ones.

Together, these models paint a picture of evolution in which trophically-important characters evolve according to different mechanisms than non-trophic characters. We find evidence for a world in which at times of notable transitions of the Earth (geological stages), we see change in the fundamental processes of diversification and sampling. And we come to understand that from ancestor

to descendant, different life history pressures lead to changes in the rate at which evolutionary change accumulates. These first forays into hierarchical model fitting call attention to notable pieces, such as the clock model, where we may be able to examine sources of heterogeneity and improve our models even further.

6.3 Cinctan phylogeny: implications for systematics and macroevolution

The origin time of the cinctan-*Ctenocystis* group was 507.52 mya (HPD 505.808 - 508.11 mya), with the ingroup originating at 505.747 mya (HPD 507.27 - 504.699 mya). As we note in our discussion of the chronostratigraphic data that we use, each tip (i.e., species) has uncertainty associated with its first appearance: we might know that the first occurrence (or possible first occurrences) are in a particular trilobite zone, but that typically restricts the age to a 1-3 million year window. This might sound trivial if we are thinking about divergences for modern taxa, but here it represents a significant proportion of expected species lifetimes, and thus a potentially large amount of time to accumulate (or not accumulate) character change. In RevBayes' implementation of the FBD model, tip uncertainty is typically treated as a uniform prior between the first and last appearance on the tip taxon (Barido-Sottani et al., 2020a). The uniform prior says that no age within the bounds is *a priori* more likely than any other. Nonetheless, we do see significant structure in the distributions for each tip (Fig. S1). Some tips, such as *Ctenocystis* and *Elliptocinctus vizcainoi* show strong skew towards the older or younger ages within their uniform tip range. Others, such as *Asturicystis jaekeli* show less signal, retrieving more-or-less the input uniform prior. This suggests that FBD analyses may be useful in the future for helping to narrow tip age ranges. In clades where tip uncertainty tends to be quite high, this could be an analytical path to higher precision on tip ages.

The topology of the tree is fully-resolved but poorly-supported on many nodes (Fig. 4). The tree in Fig. 4 displays the width of the branch scaled by the posterior support. As can be seen on that figure, the posterior probability of many nodes is quite low. This is unsurprising, as bootstrap support values in prior work have also been low (Smith and Zamora, 2009; Zamora et al., 2013a). The placement of *Protocinctus* is interesting on this phylogeny. Although it is the oldest cinctan, however, *Protocinctus* also possesses some derived character states if we assume that *Ctenocystis* is the appropriate outgroup (Rahman and Zamora, 2009). Accordingly, although prior phylogenetic studies place it as a basal member of the Sucocystidae, but evolving after the Sucocystidae diverged from both the Gyrocystidae and Trochocystitidae (Smith and Zamora, 2009; Zamora et al., 2013b). The placement of *Protocinctus* as sister to the rest of the cinctans is likely not solely due to the age of the fossil. This fossil is younger than its parent's next several ancestor nodes, meaning it could have been plausibly placed in a more derived position nested within the cinctan clade, but was not. The split between *Protocinctus* and the rest of the cinctans is also one of the most well-supported nodes on this tree.

In the in-group topology, *Asturicystis*, *Progyrocystis*, and *Graciacystis* form a weakly-supported clade that is sister to the rest of the *Gyrocystis*. *Trochocystoides* and *Trochocystites* do not form a monophyletic grouping. Our phylogeny also reflects a closer relationship between *Undatacinctus* and *Ludwigicinctus*. Neither *Elliptocinctus* nor *Sucocystis* are monophyletic in this analysis. Some of these differences may represent differences between the model applied here and in previous work. We used the Mk model (Lewis, 2001), which is more robust to superimposed or homoplasious changes than parsimony (Felsenstein, 1978; Wright and Hillis, 2014).

But differences may also reflect the inclusion of age information. For example, *Elliptocinctus* is a genus with two species on this tree. Prior analyses have reflected these as monophyletic. We did not recover these as monophyletic, with *Elliptocinctus barrandei* descending from a node that is 501.865 million years old (age HPD 500.326 - 503.626 Ma). This node is younger than the earliest appearance of *Elliptocinctus vizcainoi*. In order for these two taxa to be monophyletic, the strength of evidence in the character data would have to be strong enough to either move *Elliptocinctus vizcainoi* into that grouping (which is a canonical position for *Elliptocinctus*), thereby moving the age of the whole group back several million years, or would have to move *Elliptocinctus barrandei* out of it. Cinctans have notable homoplasy in the group and a small amount of characters. Fossil age information are treated as data under the FBD model, which has historically not been true of calibration methods, in which fossil age information was used to set constraints on clade ages. This means that the age information does not directly constrain the topology, but does still affect the distribution of topologies that are plausible given those ages. It will be worth further exploration to find out when we expect age information to exert a stronger influence than character information in determining a dated tree.

Interestingly, *Gyrocystis* has a number of sampled ancestors in the genus. In this genus, there are a total of three sampled ancestors, one pair of which (*G. erecta* and *G. badulesiensis*) likely represent budding cladogenesis, i.e., a case of speciation where the ancestral species persists. Its sister group also has one, which may also represent evidence for budding speciation. We emphasize these data show evidence for budding speciation despite the fact that we did not explicitly model stasis nor distinguish between “punctuational” changes associated with speciation vs. continuous change within lineages (e.g., Eldredge and Gould, 1972; Wagner and Marcot, 2010). However, the differences between character changes associated with speciation vs. continuous background change for these data might be small enough to be subsumed by the uncorrelated relaxed clock model we employed, which models rate shifts as occurring between branches but constant across a given branch’s duration. Nevertheless, future analyses employing more complex approaches to modeling speciation dynamics in fossil cinctans may outperform the models considered herein.

Evidence for sampled ancestors in the cinctan fossil record might seem surprising given that the echinoderm record is less complete than many other marine invertebrates (Foote and Sepkoski, 1999), although recent FBD studies find strong evidence for their occurrence in the particularly well-sampled record of

Paleozoic crinoid echinoderms (Wright, 2017b; Wright and Toom, 2017). However, four of the five cases for cinctans are from the Drumian, for which skyline models imply the highest sampling rate for cinctans. The probability of sampling ancestor-descendant pairs reflect the probability of sampling two species (i.e., [completeness]², see Foote, 1996c) is:

$$Pr[\text{sampling two species}] = \left(\frac{\psi}{\psi + \mu}\right)^2$$

where ψ is the fossilization rate and μ is the extinction rate (see Foote, 1997, equation 1b).

Still, even at the peak during the Drumian, we expect completeness of 0.21. This in turn predicts that we should find direct ancestor-descendant pairs only 4% of the time. However, sampling of ancestors and descendants often should be more probable than global sampling rates imply because sampling rates vary geographically as well as temporally. Because ancestors and descendants usually occur in the same geographic regions and environments, and because ancestors and descendants must at least abut temporally, factors favoring the sampling of any one species often favor the sampling of close relatives, including ancestor(s) (Wagner and Erwin, 1995).

Cinctans have other paleobiological characteristics that make discovery of sampled ancestors relatively probable: (1) the group occurs over a small window of time, allowing for the assessment of taxonomic completeness, (2) they are marine taxa, allowing for better fossilization potential than many groups, such as terrestrial vertebrates, (3) they have mineralized skeletons, and are preserved well enough (often as either molds or recrystallized calcite) to code morphological features, (4) they are numerically abundant fossils, particularly in rocks from the Iberian Chains of Spain and the Montagne Noire of southern France, which enables the collection of multiple specimens and assessment of more complete material, and (5) they are small, making it more tractable to score characters from relatively complete specimens.

One final reason why we might not be surprised to find as many cinctan sampled ancestors as we do stems from the fact that reconstructed ancestors co-occur with reconstructed descendants in some cases (Figs. 4-5). Budding cladogenesis is consistent with a variety of allopatric speciation models in which biological traits enhancing preservation probabilities (e.g., broad geographic ranges and long durations) also enhance the probability of leaving daughter taxa (Wagner and Erwin, 1995), all else being equal. This becomes particularly relevant because both kinds of occupancy patterns and sampling among contemporaneous species are typically exponential (Liow, 2013; Wagner and Marcot, 2013; Foote, 2016), with common species having individual sampling probabilities much greater than average. Therefore, if high occupancy is linked with the propensity for generating more daughter species, then we expect a fossil record biased in favor of species that had daughter species. This in turn means that we expect ancestor-descendant pairs to be more common than completeness metrics would predict. Our results suggest cinctans conform to this general model,

and other taxa with similar preservation rates and sampling intensity may also contain sampled ancestors.

We hope our case study of cinctan echinoderms illustrates how the methodological approaches to phylogenetic paleobiology discussed herein provide useful tools for a diverse range of paleontological interests and pursuits. Not only does our phylogenetic analysis of the Cincta provide insight into patterns of their evolutionary relationships and highlight potential taxonomic issues, but it also provide a wealth of data for addressing a wide variety of far-reaching broader issues in macroevolutionary theory, including fundamental questions regarding the tempo and mode of evolutionary change and the origin of species.

7 Conclusion

In this contribution, we have laid forth a framework for fitting complex, hierarchical phylogenetic models. We also draw attention to the relationship between macroevolutionary models on which many paleobiological studies focus and their corresponding phylogenetic models. The fossilized birth-death represents a significant leap forward in terms of the integration of fossils in Bayesian phylogenetic analyses. Under this model, fossils are data, not mere clade constraints. However, to leverage this framework involves fitting multiple submodels to a particular dataset. In doing so, we also inferred a new dated phylogeny for cinctan echinoderms, and provide some insight as to how and why this phylogeny differs from prior work, and highlighted its systematic and macroevolutionary implications for cinctan paleobiology. We have highlighted several theoretical and empirical concerns, such as how age information impacts topology and how common sampled ancestors are in empirical datasets, which have major implications for discerning models of speciation in the fossil record. It is our hope that in describing how complex model fitting can be carried out in a tractable way, we will empower more taxonomic empiricists to use the FBD approach we utilize herein with their data. We believe the interplay between theoretical phylogenetics and deep taxonomic knowledge of empirical paleontologists is critically important for generating models that not only help us better understand the peculiarities of our favorite taxonomic groups, but also help unravel generalities in the history of life.

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