Supplement to "Rate heterogeneity, time scale separation and non-equilibrium evolution explain fat-tailed fluctuations in marine biodiversity of the Phanerozoic"

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Limit distribution of a time-averaged homogeneous S1origination-extinction process

- Fossil taxa gain and lose taxa according to an origination-extinction process. We assume that
- most fossil occurrences of a taxon come from the period of its history when it is dominant
- and in steady state. In a time slice of duration τ during such a period of steady state
- the latent per capita rates of origination and extinction would be equal (i.e. $\lambda = \mu \equiv \rho$)
- and the number of origination or extinctions events (call such events Y) each follow an
- inhomogeneous Poisson process with rate ρN_t where N_t is the number of species or general
- in the taxon of interest at time t. Allowing N_t to vary smoothly with time, and invoking the
- communicative property of the Poisson distribution, we arrive at the number Y of extinction
- or origination events in τ being distributed

$$Y \sim \text{Pois}(\rho \int_{t=0}^{\tau} N(t)dt).$$
 (1)

Under the steady state assumption we can approximate N(t) by \bar{N} , the steady state diversity, leading to

$$Y \sim \text{Pois}(\rho \bar{N}\tau).$$
 (2)

Assuming the τ of each time period in the Paleobiology Database or Sepkoski's com-14 pendium to be approximately equal (i.e. equal durations of major stratographic units) then the distribution of fluctuations within taxa will be asymptotically Gaussian.

The Gaussian assymptotics of time-avereaged birth-death processes have been proven and explored elswhere as well (s1, s2).

S2Additional super-statistical analyses

To evaluate the sensitivity of our super-statistical analysis on the particular data used and we tested our predictions on different data sets (see below). The fact that it works in all 21 different applications indicates that it is robust to vagaries of different recording strategies and bias corrections in paleobiology. This could mean that much of the raw signal in massive 23 fossil datasets, at least signals regarding fluctuations, are not artifacts of sampling, as has been proposed before (s3). 25

S2.1Raw PBDB data

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We calculated the super-statistical prediction at the order level from raw genus diversity 27 recorded in the PBDB without correcting for taphonomic or sampling bias (Fig. S2). The 28 super-statistical calculation also closely fits the raw data as in the case of sampling and publication bias-corrected data.

S2.2Different taxonomic ranks in PBDB data

As noted in the main text, the super-statistical prediction predictably breaks down at higher taxonomic scales. In Figure S3 we present this worsening fit graphically using class level data with three-timer and publication corrected PBDB data

35 S2.3 Sepkoski's compendium

- Sepkoski's compendium (s4) provided the first hypothesis of Phanerozoic diversification. As
- such, it has served as a benchmark for further investigation into large-scale paleobiological
- patterns (s5). We conducted the same super-statistical analysis as in the main text and find
- 39 comparable results. Specifically, the super-statistical prediction far out preforms the null
- 40 Gaussian model (Fig. S4) and worsens with increasing taxonomic scale (Fig. S4), again
- implying the uniqueness of orders.

⁴² References

- s1. J. Keilson, S. S. Rao, Journal of Applied Probability 7, 699 (1970).
- s2. W. K. Grassmann, Annals of Operations Research 8, 165 (1987).
- 45 s3. B. Hannisdal, S. E. Peters, Science **334**, 1121 (2011).
- s4. J. J. Sepkoski, *A compendium of fossil marine animal families* (Milwaukee Public Museum, Milwaukee, WI, 1992).
- 48 s5. J. Alroy, et al., Science **321**, 97 (2008).
- ⁴⁹ s6. J. Alroy, *Science* **329**, 1191 (2010).

50 Supplemental Figures

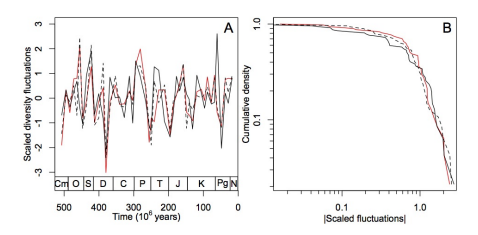


Figure S1: Comparison of SQS method (s6) (solid black line) with the raw data (dashed black) and our three-timer and publication bias correction method (red). The time-series of all marine invertebrate genera shows general agreement with the only major deviations toward the modern (A). Despite these differences the distribution of fluctuations in genus diversity across all marine invertebrates show good agreement (B).

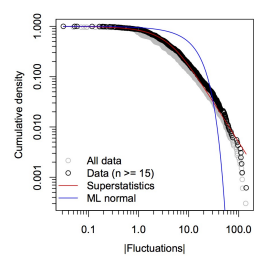


Figure S2: Super-statistical prediction of raw (i.e. not bias corrected) order-level fluctuations in genus diversity recorded in the PBDB. Grey dots are the full data of orders, while black ones are orders with more than 15 points. The red line is our theoretical prediction and the blue line the best Gaussian fit to the data.

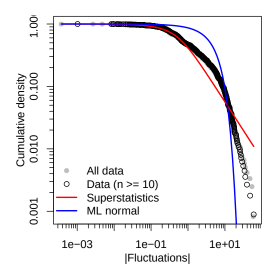


Figure S3: Super-statistical prediction of bias corrected class-level fluctuations in genus diversity recorded in the PBDB. Grey dots are the full data of orders, while black ones are orders with more than 15 points. The red line is our theoretical prediction and the blue line the best Gaussian fit to the data. Note at the class level the fit is predictably worse, see main text for discussion.

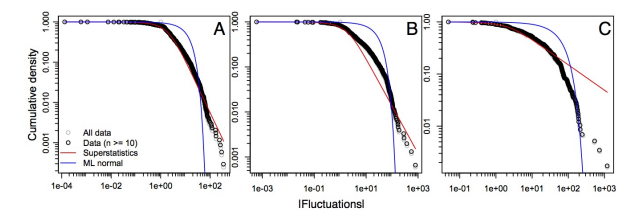


Figure S4: Super-statistical prediction (red line) of fluctuations in genus diversity recorded in Sepkoski's compendium of marine invertebrates compared to maximum likelihood normal distribution (blue line). Super-statistical theory explains order level fluctuations well (A) with increasingly poorer fits at the class (B) and phylum (C) levels.