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Main Manuscript for

Fossilized cell structures identify an ancient origin for the teleost whole-genome duplication.

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Author Contributions: D.D., M.F. P.E.A., S.S. and R.B.J.B. conceived and designed the project. D.D., M.F., G.C. and R.B.J.B. collected and prepared the material. D.D., A.D.S, V.F., S.S. and R.B.J.B. performed the synchrotron experiments. V.F. and S.S. prepared the PPC-SRµCT scan data and provided training for their reconstruction. A.D.S. segmented the scan data. D.D., A.D.S., S.S. and R.B.J.B. analysed the data. All authors discussed the interpretations. D.D., M.F. and R.B.J.B. developed the main text and made the figures. D.D., V.F. and G.C. wrote the SI Appendix. All authors provided a critical review of the manuscript and approved the final draft.

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Abstract

Teleost fishes comprise half of all vertebrate species, and possess a duplicated genome. This whole-genome duplication (WGD) occurred on the teleost stem-lineage, in an ancient common ancestor of all living teleosts, and is hypothesized as a trigger of their exceptional evolutionary radiation. Genomic and phylogenetic data indicate that WGD occurred in the Mesozoic, after the divergence of teleosts from their closest living relatives, but before the origin of the extant teleost groups. However, these approaches cannot pinpoint WGD among the many extinct groups that populate this 50- to 100-million-year lineage, preventing tests of the evolutionary effects of WGD. We infer patterns of genome size evolution in fossil stem-group teleosts using high-resolution synchrotron X-ray tomography to measure the bone cell volumes, which correlate with genome size in living species. Our findings indicate that WGD occurred very early on the teleost stemlineage, and that all extinct stem-group teleosts known so far possessed duplicated genomes. WGD therefore substantially pre-dates both the origin of proposed key innovations of the teleost skeleton and the onset of substantial morphological diversification in the clade. Moreover, the early occurrence of WGD allowed considerable time for post-duplication reorganization prior to the origin of the teleost crown-group. This suggests at most an indirect link between WGD and evolutionary success, with broad implications for the relationship between genomic architecture and large-scale evolutionary patterns in the vertebrate Tree of Life.

Significance Statement

Some lineages of organisms have undergone major evolutionary radiations, while others have not. Establishing why is a central goal of evolutionary research. Whole-genome duplication (WGD) is often proposed as having caused the spectacular evolutionary radiation of teleost fishes. However, due to the absence of genetic data for fossil species, it has been impossible to pinpoint precisely when WGD occurred during teleost history. We use 3D measurements of fossilized bone cell spaces to estimate genome sizes in extinct species, observing a near-doubling of size during earliest teleost ancestry. This suggests that WGD occurred very early, substantially pre-dating the dramatic radiation of teleosts. These findings suggest at most and indirect link between WGD and teleost diversification.

Main Text

Introduction

Whole-genome duplication (WGD) has occurred independently in multiple lineages of plants, fungi, and animals (1–3). This represents a major change to genomic architecture, with hypothesized impacts on evolutionary diversification (4, 5), caused by the origin of new gene functions from duplicate copies, expanding the genetic toolbox available for evolutionary 'tinkering' (6). However, in spite of its mechanistic plausibility, this hypothesis is so far supported by only limited and contradictory empirical evidence (7–10). Teleost fishes – comprising more than half of modern vertebrates – are a key example, with their spectacular variety of form and kind (ranging from eels to seahorses) often viewed as *prima facie* evidence for the role of WGD in triggering evolutionary diversification (6, 11). Teleost also show an incredible diversity of genome biology, demonstrating particularly high rates of evolution of protein-coding genes (12) and non-

coding elements (13), a broad range of genome sizes including the smallest known in vertebrates (14), and multiple polyploid lineages (15).

The genome of all living teleosts derives from an ancient WGD event that occurred before the last common ancestor of modern species (16). Additional duplication events occurred more recently in several teleost subgroups (9, 17), but are not generally proposed as drivers of diversification (9). Studies of the role of WGD in contributing to teleost diversity so far have analyzed the distribution of species richness among extant lineages and morphometric data for fossil phenotypes, with potentially conflicting results: extant teleosts have high rates of lineage diversification compared to other ray-finned fishes (7), but early fossil members of the teleost crown-group do not show increased rates of morphological evolution (18).

Molecular phylogenetic studies indicate that WGD occurred on the teleost stem-lineage: after the divergence of teleosts from their extant sister taxon (Holostei), but before the most recent common ancestor of all living teleosts (19, 20). However, these bounds encompass a large phylogenetic diversity of extinct groups that diverged during an interval of 50–100 million years, from the initial divergence of the teleost total-group from the holostean total-by the earliest Triassic (21), up to the first appearance of crown-group teleosts in the Late Jurassic (18, 22). Molecular-clock estimates provide only broad constraints on the precise timing of duplication (316-226 Ma (23); ~310 Ma (24)), and offer no information on its phylogenetic position on the teleost stem lineage. The imprecision of these estimates and the sometimes considerable incongruence of molecular clocks with the teleost fossil record questions the reliability of these inferences in the absence of further evidence.

Patterns of genome size evolution on the teleost stem lineage could provide alternative and independent evidence on the timing and phylogenetic position of the teleost WGD. However, stem lineages by definition comprise entirely extinct species that are known only from fossils, for which genomic data are absent. Nevertheless, some information about vertebrate genome size is preserved within fossil bone (25–27). Living organisms show a positive correlation between cell size and genome size (28–30), such that the volumes of bone cell spaces (osteocyte lacunae) allow estimates of genome size. This relationship has been demonstrated in ray-finned fishes, including teleosts, and is predictive for large-scale variation in genome size (31). The precision of this approach is sufficient for inferring the large change (presumably, doubling) of genome size involved in WGD (31). Here, we use this relationship to trace the evolution of genome size in extinct ray-finned fishes, using osteocyte lacuna volumes as a proxy for genome size. Our sample includes a broad range of stem- and crown-group teleosts, providing information on patterns of teleost genome-size evolution during the deep evolutionary history of the teleost total-group.

Three-dimensional measurement of fossil bone cell spaces with µm-scale diameters presents considerable technical challenges. We used propagation phase contrast synchrotron radiation X-ray micro-computed tomography (PPC-SRµCT) to address this, collecting standardized measurements of osteocyte lacuna volumes for 61 fossil ray-finned fish species ranging from 2.5 to 252 million years in age (*SI Appendix, Part I*). This fossil evidence is complemented by data from a previous study including 34 modern ray-finned fish species with known genome sizes (31). Our fossil sample includes all major groups of stem-group teleosts, members of both living and extinct lineages within the teleost crown group, and several non-teleost ray-finned fishes. This sample allows us to estimate relative genome sizes in extinct groups, providing information on the absolute timing and specific phylogenetic position of the teleost WGD as well as the timescale of post-duplication reductions in genome size (24). Both statistical analysis and qualitative observations demonstrate the effectiveness of lacuna size for inferring large evolutionary increases in genome size: known polyploid lineages such as catostomids and salmonids, which

underwent additional rounds of WGD, both show large osteocyte lacuna volumes compared to their close relatives (31).

Results

Our results suggest that WGD occurred early on the teleost stem lineage. Osteocyte lacuna volumes of early to mid-Mesozoic (Early Jurassic-Early Cretaceous) teleosts overlap with those of extant polyploid taxa and exceed values for non-polyploid taxa (Figs. 1-2, S1-3). Phylogenetic ancestral trait estimates attribute this to a substantial increase in lacuna volume (from 148 to 240 µm³) immediately following the split of teleosts from holosteans, and therefore before the deepest-known divergences of the teleost total group (Fig. 1). The oldest stem-group teleosts in our sample with measurable lacuna volumes hail from the Early Jurassic (~195 Ma) and have large lacuna volumes similar to those of recent polyploids such as salmonids and catostomids (Fig. 2). Late Jurassic and Early Cretaceous members of the teleost crown-group generally also have large lacunae compared to extant teleosts. In contrast, late Mesozoic and Cenozoic teleosts mostly have lacuna volumes comparable to those of their closest modern relatives (Figs. 1-2, S1). Osteocyte lacuna volumes therefore decrease through time along the teleost stem-lineage, with further reductions within major lineages of crown teleosts (such as elopomorphs and clupeocephalans) or within some members of the stem group (such as ichthyodectiforms). Aspects of these patterns of lacuna size evolution are consistent with those inferred from molecular data. Nevertheless, our study provides valuable additional evidence, placing important constraints on the timing and phylogenetic placement of WGD on the extinct teleost stem lineage as a test of its relationship to phenotypic diversification.

Discussion

A large, almost two-fold increase in osteocyte sizes at the base of the teleost stem lineage suggests a phylogenetically-early occurrence of WGD. Large genome sizes can also occur via other genomic mechanisms such as the accumulation of transposable elements, introns, or tandem repeats (32–34) (e.g. within vertebrates, lungfishes and salamanders). Nevertheless, the timing, phylogenetic distribution (spanning early members of the teleost stem-lineage and their crown group), and observation of subsequent decreases in cell size all suggest that the patterns of genome size variation inferred here result from WGD and not from other forms of evolutionary genome size expansion. Furthermore, our previous work has demonstrated that polyploidy provides a strong statistical explanation for large osteocyte size in actinopterygians, on par with genome size (31).

The oldest stem-group teleosts in our sample (*Pholidophoroides* and *Pholidophoropsis*) are of Early Jurassic age. These taxa represent lineages that diverged much earlier from the teleost stem-group, and therefore constrain the timing of the teleost WGD to no later than the Late Triassic (~235 Ma) using phylogenetic ancestral state estimation (Fig. 1). Direct sampling of Triassic stem teleosts will provide a further test of this hypothesis (35), but this has not so far been possible in spite of multiple attempts, due to poor preservation of bone microstructure (*SI Appendix, Part II*). Our phylogenetically-inferred age for the teleost WGD falls within the timeframe estimated by some molecular studies (316-226 Ma (23); ~310 Ma (24)). The subsequent decrease in lacuna volumes toward the crown and within individual crown-group lineages indicates post-duplication genome-size reduction, spanning tens of millions of years. This confirms previous inferences of post-duplication genome size reduction based on clock analysis of paralogous genes,mathematical modelling, and the phylogenetic distribution of genome size in extant teleosts (24). Ichthyodectiforms, a long-lived clade of Mesozoic stemteleosts, show a similar pattern of smaller volumes in more recent species compared to earlier

ones, providing evidence of parallel reduction in an extinct lineage closely related to the crown (Fig. 1).

Our findings are inconsistent with hypotheses that propose that spectacular evolutionary diversification of teleosts is an immediate consequence of the WGD. Because of the phylogenetically-early occurrence of WGD, all currently recognized members of the teleost stemgroup have inferred duplicate genomes. Therefore, WGD did not coincide with the origin of celebrated teleost functional innovations like the mobile premaxilla, symmetrical caudal-fin and aspects of vertebral geometry implicated in their later evolutionary success (36, 37) (Fig. 1). These traits arose later — conceivably many millions of years after the inferred origin of WGD (35). WGD occurred during or before the Triassic, and therefore also predates more general phenotypic diversification, which occurred from the Late Jurassic onwards (38). The earliest known (i.e., Triassic and Early Jurassic) stem teleosts conform to a conservative bodyplan, being small, fusiform fishes with ganoid scales. Quantitative analyses show consistently low levels of morphological diversity for at least ~50 Myr after the origin of the teleost total-group (38). The conservative early history of teleosts cannot be attributed to inadequacies of the Triassic record, which yields an abundance of fish fossils (39) including anatomically diverse holosteans (38). The morphological diversity of teleosts increased only gradually in the later stages of the Mesozoic, especially in the Cretaceous with the origin of fundamentally new bodyplans (e.g., eel-like, deepbodied (40, 41)) and amplified by prolific morphological diversification in the early Paleogene (42, 43). Triassic-Early Cretaceous teleosts, including early crown-group taxa, show comparable or lower rates of body-shape evolution than contemporary holosteans (18), which are known to lack duplicated genomes (20). Early teleosts also show delayed patterns of taxonomic and ecological diversification compared to the timing of WGD inferred here: teleosts comprise only a minority of actinopterygian genera in marine and freshwater settings throughout the Triassic (39) and much of the Jurassic (44), even after the origin of their crown-group. Only in the Late Jurassic — at least 80 Myr after the WGD — does the taxonomic diversity of teleosts approach or exceed that of other actinopterygians, as evidenced by the marine fossil assemblages of famous Lagerstätten like Cerin and Solnhofen (43).

This temporal pattern suggests an alternative hypothesis: that reorganization and integration of genomic architecture after the WGD, and not WGD itself, drove the phenotypic diversification and origin of key morphological traits during later stages of teleost evolution. Consistent with this hypothesis, the 50 Myr-interval between the origin of the teleost total-group and their observed increase in morphological diversity (38) is similar to the estimated duration of post-WGD genome size reductions to modern teleost-like levels (20)(Fig. 2). Decrease in genome size is associated with profound changes in genome organization, through gene loss, neo-functionalization, and changes in expression.

Because of recent focus on genomic hypotheses, relatively little attention has been given to alternative explanations for the phenomenal diversity of extant teleosts. However, the temporal incongruence between a Triassic WGD and the late Mesozoic and early Cenozoic events of teleost diversification (18, 38, 43) also demands broader consideration of non-genomic hypotheses. One possibility is that teleosts were the beneficiaries of biotic turnover during major environmental changes of the late Mesozoic. For example, the mid-Cretaceous interval saw substantial environmentally-driven alterations to marine ecosystems induced by large igneous province volcanism (45), regional ocean anoxic events (46), and other climatic variations. Among these, the Turonian thermal maximum c. 92 million years ago (47) was a major climate perturbation associated with biotic turnover at all levels of the marine trophic chain (46, 48). The early Late Cretaceous coincided with a major burst of cladogenesis in marine fishes, the diversity of which is strongly and positively correlated with sea surface temperature during the Jurassic and Cretaceous (44, 49). Although teleosts dominate this mid-Cretaceous radiation, other ray-finned fish groups show similar patterns (e.g., pycnodonts; (50)), arguing for a general—rather than clade-specific—mechanism. The Cretaceous concluded 66 million years ago with a major

mass extinction event. This induced a major reorganization of aquatic ecosystems, particularly in higher trophic levels of marine environments (51, 52). Paleontological and molecular data indicate major morphological diversification among marine teleosts in the early Paleogene (42, 53). This was associated with the colonization of different environments and the refilling of vacated functional roles (54–57), and included short-lived "evolutionary experiments" found alongside familiar living lineages (58, 59). As with the mid-Cretaceous, the early Paleocene represented a "hothouse" period in Earth's climate history, with evidence that higher temperatures might have led to increased fish production in some marine settings (60). Collectively, these observations suggest a role for environmental change and ecosystem reorganization in facilitating the diversification of teleosts, either in addition to or instead of the effects from WGD.

Our results provide direct fossil evidence of phylogenetic placement of the teleost WGD, and a more robust window for the timing and consequences of teleost genome duplication. WGD occurred early on the teleost stem-lineage, with considerable time for post-duplication reorganizations of genomic architecture before the morphological and taxonomic radiation of teleosts. This suggests an indirect and temporally-offset link between WGD and evolutionary diversification, if any, and is consistent with genomic studies of recent polyploids among fishes (e.g. salmonids (9)) and vascular plants (10). More broadly, our data demonstrate the potential of the fossil record of cell structures (26) as an independent source of evidence on genomic events that may have underpinned major diversification events, not just in fishes and other vertebrates, but also in plants and across the Tree of Life (2, 3).

Materials and Methods

Specimen collection

Our taxon sample builds on a previous study (31) that collected osteocyte lacuna volumes for 34 species of extant actinopterygians (including 28 teleost species). The present study adds 61 fossil species displaying osteocytic bone (61) to this sample: 8 non-teleost actinopterygians, 26 stemgroup teleosts, and 27 crown teleosts. They range from 2.5 (Pliocene) to 252 (Early Triassic) million years in age (*SI Appendix, Part I*, Dataset S1). However, no representative of the teleost lineage is older than 200 million years (Early Jurassic; *SI Appendix, Part II*). As fossil specimens have a higher density than extant ones, capturing a decent signal of transmitted X-ray beam could be challenging should the specimens be too large. Thus, we took small (~ 1-5 mm) bone samples from the fossils, targeting areas that reduced the loss of anatomical information from the specimens (e.g., areas affected by pre-existing cracks or broken bone). Since lacunae vary in volume from one bone to another (30, 31), the same skeletal element has to be used consistently across the sample. Therefore, all samples were taken from the dentary bone (lower jaw) as it is found in all target taxa and is often preserved in the fossil record. Moreover, the already-existing dataset of extant actinopterygians (31), used dentary bones for each included species.

Data acquisition

The data presented in this study were acquired using propagation phase contrast synchrotron radiation X-ray micro-computed tomography (PPC-SRµCT). Acquisition was performed over the course of three sessions in two synchrotrons: two sessions were done at the ID19 beamline of the European Synchrotron Radiation Facility (ESRF, Grenoble, France) and the remaining session was done at the I13-2 Diamond Manchester Imaging beamline of the Diamond Light Source (DLS, Didcot, UK). The setups on the two beamlines were generally similar, the main difference being that much higher energies were available at the ID19 beamline (up to 112 keV in this study) compared to I13-2 beamline (here, 21.39 keV). Both setups produced high-resolution data, with recorded voxel sizes of 0.357 µm at the I13-2 beamline and near 0.7 µm at the ID19 beamline. A voxel size of 0.7 µm or smaller is empirically adequate for imaging osteocyte lacunae (31, 62–64). Detailed information on the synchrotron experimental setups is found in the *SI Appendix, Part III* and Dataset S2. The tomograms (digital 'slices') obtained from our specimens

were processed with the software VGSTUDIO MAX v. 3.0 and 3.1 (Volume Graphics, Heidelberg, Germany), to segment osteocyte lacunae from the fossil bone matrix. We used specimen-specific grey-value thresholds to segment osteocyte lacunae from a selected region of interest, focusing on well-defined lacunae from primary bone, excluding objects such as specimen edges, cracks and abiotic inclusions manually. Grey-value thresholds were selected to ensure that the boundaries of segmented osteocytes coincided with the boundaries of osteocytes in the image volume, attempting to maximize fit of the segmented object to the underlying data. We did not segment the canaliculi that accommodate cytoplasmic projections of the osteocytes, as they were not visible in every tomogram and thus could bias our estimates of the osteocyte lacuna volumes. We used the 'Porosity/Inclusion' module of VGSTUDIO MAX, to measure the individual lacuna volumes. This module also provides visualization, coloring lacunae according to their volume with a consistent color range across our sample (Figs. 1-2, S1). We analyzed median osteocyte volumes computed from the entire population of segmented objects, excluding objects smaller than 25 µm³, which did not generally represent osteocyte lacunae. The whole protocol followed the one we applied previously for extant actinopterygians (31), allowing us to incorporate extant and fossil samples together in downstream analyses. The complete measurement and positional data for populations of osteocyte lacunae in all specimens analyzed are available in Dryad (https://doi.org/10.5061/dryad.bcc2fqzcc), and our scan image volumes are available on MorphoSource (https://www.morphosource.org/projects/0000C1125). We used objective quality control criteria to exclude some specimens from the analysis after

We used objective quality control criteria to exclude some specimens from the analysis after obtaining PPC-SRμCT tomograms (*SI Appendix, Part I.3*). Datasets were excluded based on the following criteria: [1] When the data clearly showed the absence of osteocyte lacunae from the sample, either because the taxon has anosteocytic bone (61, 65), or because osteocytes were not fossilized. [2] When osteocyte lacunae were present but not sufficiently well-resolved to allow an accurate segmenting of the lacunae or measurement of their volume. [3] After segmenting the osteocyte lacunae, we excluded more specimens in which the number of osteocyte lacunae we were able to measure was too small to warrant confident estimates of average size (n<~50). Prior to the segmentation and measurement, our sample of tomograms included 77 fossil specimens in total representing 71 species, and notably included five stem-group teleosts from the Triassic. Of these, 16 specimens were excluded based on quality control criteria (see the *SI Appendix, Part I.1 and I.2* for the complete list of specimens).

• Reference tree

We used a composite phylogeny based on a consensus of published hypotheses on the interrelationships of fossil actinopterygians, and divergence ages inferred from earliest known occurrences in the fossil record (*SI Appendix, Part I.2*, Dataset S1). The framework for extant taxa is a recently published molecular timetree (66) that has the best species sample and taxonomic coverage of any actinopterygian phylogeny so far published. Fossil taxa were stitched to the resulting tree using a custom script that makes extensive use of the R package ape version 5.0 (67) and is available in Dryad (https://doi.org/10.5061/dryad.bcc2fqzcc). Fossil age ranges, specifier taxa for their sister clades, and minimum divergence times from those sister clades based on fossil occurrences are available as Dataset S1. Phylogenetic position and age data for the fossil taxa and their divergence times from other species were taken from the most recent available paleontological and stratigraphic information (see the *SI Appendix, Part I.2* for the justification of phylogenetic and stratigraphic placement for each species in the sample). Extant terminals were pruned, retaining the 34 extant species for which we measured osteocyte volumes, prior to analysis. The resulting tree is available as Fig. S1 and Dataset S3.

Data analysis

To visualize patterns of genome size evolution, we mapped osteocyte lacuna volume (Dataset S4) to our composite phylogeny with branch lengths of units in time, using ancestral character state estimation (68) via the ace function of ape version 5.0 (67). Osteocyte lacuna volume was log₁₀-transformed prior to analysis and is used as a proxy for genome size (31). WGD predicts an approximate doubling of osteocyte size somewhere on the teleost stem-lineage.

Data availability

The SRµCT data (tomograms and 3D reconstructions) generated during this study have been deposited online in MorphoSource: www.morphosource.org/projects/0000C1125 (fossil specimens) and www.morphosource.org/projects/00000C959 (extant specimens). The median osteocyte lacuna volumes for each specimen are available in Dataset S4. The complete volume measurement data for populations of osteocyte lacunae in all specimen analysed, as well as the custom R script used to build the reference tree and reconstruct osteocyte lacuna volume evolution are available in Dryad (https://doi.org/10.5061/dryad.bcc2fqzcc). All other data files are included in the SI Appendix and Datasets S1-S4.

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References

- 1. Y. Van De Peer, S. Maere, A. Meyer, The evolutionary significance of ancient genome duplications. *Nat. Rev. Genet.* **10**, 725–732 (2009).
- 2. J. W. Clark, P. C. J. Donoghue, Whole-genome duplication and plant macroevolution. *Trends Plant Sci.* **23**, 933–945 (2018).
- 3. O. Simakov, *et al.*, Deeply conserved synteny resolves early events in vertebrate evolution. *Nat. Ecol. Evol.* (2020) https://doi.org/10.1038/s41559-020-1156-z.
- 4. S. Ohno, Evolution by Gene Duplication (Springer, 1970).
- K. D. Crow, G. P. Wagner, What is the role of genome duplication in the evolution of complexity and diversity? *Mol. Biol. Evol.* 23, 887–892 (2006).
- 6. S. M. K. Glasauer, S. C. F. Neuhauss, Whole-genome duplication in teleost fishes and its evolutionary consequences. *Mol. Genet. Genomics* **289**, 1045–1060 (2014).
- 7. F. Santini, L. J. Harmon, G. Carnevale, M. E. Alfaro, Did genome duplication drive the origin of teleosts? A comparative study of diversification in ray-finned fishes. *BMC Evol. Biol.* **9**, 194 (2009).
- 8. R. Ren, *et al.*, Widespread whole genome duplications contribute to genome complexity and species diversity in angiosperms. *Mol. Plant* **11**, 414–428 (2018).
- 9. D. J. Macqueen, I. A. Johnston, A well-constrained estimate for the timing of the salmonid whole genome duplication reveals major decoupling from species diversification. *Proc. R. Soc. B Biol. Sci.* **281**, 20132881 (2014).

- 10. I. Mayrose, *et al.*, Recently formed polyploid plants diversify at lower rates. *Science (80-.).* **333**, 1257–1257 (2011).
- 11. V. Ravi, B. Venkatesh, Rapidly evolving fish genomes and teleost diversity. *Curr. Opin. Genet. Dev.* **18**, 544–550 (2008).
- 12. V. Ravi, B. Venkatesh, The divergent genomes of teleosts. *Annu. Rev. Anim. Biosci.* **6**, 47–68 (2018).
- 13. T. Desvignes, J. Sydes, J. Montfort, J. Bobe, J. H. Postlethwait, Evolution after whole genome duplication: teleost microRNAs. *Mol. Biol. Evol.*, 0–2 (2021).
- 14. E. M. Smith, T. R. Gregory, Patterns of genome size diversity in the ray-finned fishes. *Hydrobiologia* **625**, 1–25 (2009).
- 15. R. A. Leggatt, G. K. Iwama, Occurrence of polyploidy in the fishes. *Rev. Fish Biol. Fish.* **13**, 237–246 (2003).
- 16. A. Amores, *et al.*, Zebrafish *hox* clusters and vertebrate genome evolution. *Science (80-.).* **282**, 1711–1714 (1998).
- 17. C. Berthelot, *et al.*, The rainbow trout genome provides novel insights into evolution after whole-genome duplication in vertebrates. *Nat. Commun.* **5**, 3657 (2014).
- J. T. Clarke, G. T. Lloyd, M. Friedman, Little evidence for enhanced phenotypic evolution in early teleosts relative to their living fossil sister group. *Proc. Natl. Acad. Sci.* 113, 11531–11536 (2016).
- 19. J. S. Taylor, I. Braasch, T. Frickey, A. Meyer, Y. Van de Peer, Genome duplication, a trait shared by 22,000 species of ray-finned fish. *Genome Res.* **13**, 382–390 (2003).
- 20. I. Braasch, *et al.*, The spotted gar genome illuminates vertebrate evolution and facilitates human-teleost comparisons. *Nat. Genet.* **48**, 427–437 (2016).
- 21. M. Friedman, The early evolution of ray-finned fishes. *Palaeontology* **58**, 213–228 (2015).
- 22. G. Arratia, "The Jurassic and the early history of teleosts" in *Mesozoic Fishes Systematics and Paleoecology*, G. Arratia, G. Viohl, Eds. (Verlag Dr. Friedriech Pfeil, 1996), pp. 243–259.
- 23. I. A. Hurley, *et al.*, A new time-scale for ray-finned fish evolution. *Proc. R. Soc. B Biol. Sci.* **274**, 489–498 (2007).
- 24. J. Inoue, Y. Sato, R. Sinclair, K. Tsukamoto, M. Nishida, Rapid genome reshaping by multiple-gene loss after whole-genome duplication in teleost fish suggested by mathematical modeling. *Proc. Natl. Acad. Sci.* **112**, 14918–14923 (2015).
- C. L. Organ, A. M. Shedlock, A. Meade, M. Pagel, S. V. Edwards, Origin of avian genome size and structure in non-avian dinosaurs. *Nature* 446, 180–184 (2007).
- 26. P. C. J. Donoghue, Fossil cells. *Curr. Biol.* **30**, R485–R490 (2020).
- 27. C. L. Organ, A. Canoville, R. R. Reisz, M. Laurin, Paleogenomic data suggest mammal-like genome size in the ancestral amniote and derived large genome size in amphibians. *J. Evol. Biol.* **24**, 372–380 (2011).

- 28. T. Cavalier-Smith, Skeletal DNA and the evolution of genome size. *Annu. Rev. Biophys. Bioeng.* **11**, 273–302 (1982).
- 29. E. Olmo, Nucleotype and cell size in vertebrates: a review. *Basic Appl. Histochem.* **27**, 227–256 (1983).
- 30. M. D. D'Emic, R. B. J. Benson, Measurement, variation, and scaling of osteocyte lacunae: a case study in birds. *Bone* **57**, 300–310 (2013).
- 31. D. Davesne, A. D. Schmitt, V. Fernandez, R. B. J. Benson, S. Sanchez, Three-dimensional characterisation of osteocyte volumes at multiple scales, and its relationship with bone biology and genome evolution in ray-finned fishes. *J. Evol. Biol.* **33**, 808–830 (2020).
- 32. T. Lefébure, *et al.*, Less effective selection leads to larger genomes. *Genome Res.* **27**, 1016–1028 (2017).
- H. C. Liedtke, D. J. Gower, M. Wilkinson, I. Gomez-Mestre, Macroevolutionary shift in the size of amphibian genomes and the role of life history and climate. *Nat. Ecol. Evol.* 2, 1792–1799 (2018).
- 34. A. Kapusta, A. Suh, C. Feschotte, Dynamics of genome size evolution in birds and mammals. *Proc. Natl. Acad. Sci.*, 201616702 (2017).
- 35. G. Arratia, Morphology, taxonomy, and phylogeny of Triassic pholidophorid fishes (Actinopterygii, Teleostei). *J. Vertebr. Paleontol.* **33**, 1–138 (2013).
- 36. B. Schaeffer, D. E. Rosen, Major adaptive levels in the evolution of the actinopterygian feeding mechanism. *Am. Zool.* **1**, 187–204 (1961).
- 37. L. Sallan, Evolution: Spinal innovation enabled by genome duplication. *Curr. Biol.* **30**, R1006–R1008 (2020).
- 38. J. T. Clarke, M. Friedman, Body-shape diversity in Triassic–Early Cretaceous neopterygian fishes: sustained holostean disparity and predominantly gradual increases in teleost phenotypic variety. *Paleobiology* **44**, 402–433 (2018).
- 39. C. Romano, *et al.*, Permian-Triassic Osteichthyes (bony fishes): diversity dynamics and body size evolution. *Biol. Rev.* **91**, 106–147 (2016).
- 40. J. G. Maisey, Santana Fossils: an Illustrated Atlas, J. G. Maisey, Ed. (T.F.H., 1991).
- 41. P. L. Forey, L. Yi, C. Patterson, C. E. Davies, Fossil fishes from the Cenomanian (Upper Cretaceous) of Namoura, Lebanon. *J. Syst. Palaeontol.* **1**, 227–330 (2003).
- 42. M. Friedman, Explosive morphological diversification of spiny-finned teleost fishes in the aftermath of the end-Cretaceous extinction. *Proc. R. Soc. B Biol. Sci.* **277**, 1675–1683 (2010).
- 43. G. Guinot, L. Cavin, 'Fish' (Actinopterygii and Elasmobranchii) diversification patterns through deep time. *Biol. Rev.* **91**, 950–981 (2016).
- 44. L. Cavin, P. L. Forey, C. Lécuyer, Correlation between environment and Late Mesozoic ray-finned fish evolution. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* **245**, 353–367 (2007).
- 45. S. E. Bryan, et al., The largest volcanic eruptions on Earth. Earth-Science Rev. 102, 207-

- 229 (2010).
- 46. R. M. Leckie, T. J. Bralower, R. Cashman, Oceanic anoxic events and plankton evolution: biotic response to tectonic forcing during the mid-Cretaceous. *Paleoceanography* **17**, 1–29 (2002).
- 47. C. Linnert, *et al.*, Evidence for global cooling in the Late Cretaceous. *Nat. Commun.* **5**, 4194 (2014).
- V. Fischer, N. Bardet, R. B. J. Benson, M. S. Arkhangelsky, M. Friedman, Extinction of fish-shaped marine reptiles associated with reduced evolutionary rates and global environmental volatility. *Nat. Commun.* 7, 10825 (2016).
- 49. G. Guinot, L. Cavin, Distinct responses of elasmobranchs and ray-finned fishes to long-term global change. *Front. Ecol. Evol.* **7**, 513 (2020).
- 50. J. J. Cawley, *et al.*, Rise and fall of †Pycnodontiformes: Diversity, competition and extinction of a successful fish clade. *Ecol. Evol.* (2021) https://doi.org/10.1002/ece3.7168.
- 51. N. Bardet, Extinction events among Mesozoic marine reptiles. *Hist. Biol.* **7**, 313–324 (1994).
- M. Friedman, Ecomorphological selectivity among marine teleost fishes during the end-Cretaceous extinction. *Proc. Natl. Acad. Sci.* 106, 5218–5223 (2009).
- 53. E. C. Sibert, R. D. Norris, New Age of Fishes initiated by the Cretaceous-Paleogene mass extinction. *Proc. Natl. Acad. Sci.* **112**, 8537–8542 (2015).
- 54. E. Ribeiro, A. M. Davis, R. A. Rivero-Vega, G. Ortí, R. Betancur, Post-Cretaceous bursts of evolution along the benthic-pelagic axis in marine fishes. *Proc. R. Soc. B Biol. Sci.* **285**, 20182010 (2018).
- 55. M. Friedman, *et al.*, A phylogenomic framework for pelagiarian fishes (Acanthomorpha: Percomorpha) highlights mosaic radiation in the open ocean. *Proc. R. Soc. B Biol. Sci.* **286**, 20191502 (2019).
- 56. D. R. Bellwood, C. H. R. Goatley, O. Bellwood, The evolution of fishes and corals on reefs: form, function and interdependence. *Biol. Rev.* **92**, 878–901 (2017).
- 57. S. A. Price, *et al.*, Two waves of colonization straddling the K-Pg boundary formed the modern reef fish fauna. *Proc. R. Soc. B Biol. Sci.* **281**, 20140321 (2014).
- 58. D. Davesne, G. Carnevale, M. Friedman, *Bajaichthys elegans* from the Eocene of Bolca (Italy) and the overlooked morphological diversity of Zeiformes (Teleostei, Acanthomorpha). *Palaeontology* **60**, 255–268 (2017).
- 59. A. Capobianco, *et al.*, Large-bodied sabre-toothed anchovies reveal unanticipated ecological diversity in early Palaeogene teleosts. *R. Soc. Open Sci.* **7**, 192260 (2020).
- 60. G. L. Britten, E. C. Sibert, Enhanced fish production during a period of extreme global warmth. *Nat. Commun.* **11**, 1–6 (2020).
- 61. D. Davesne, *et al.*, The phylogenetic origin and evolution of acellular bone in teleost fishes: insights into osteocyte function in bone metabolism. *Biol. Rev.* **94**, 1338–1363 (2019).

- 62. S. Sanchez, *et al.*, 3D microstructural architecture of muscle attachments in extant and fossil vertebrates revealed by synchrotron microtomography. *PLoS One* **8**, e56992 (2013).
- 63. S. Sanchez, P. Tafforeau, P. E. Ahlberg, The humerus of *Eusthenopteron*: a puzzling organization presaging the establishment of tetrapod limb bone marrow. *Proc. R. Soc. B Biol. Sci.* **281**, 20140299 (2014).
- 64. S. Sanchez, P. E. Ahlberg, K. M. Trinajstic, A. Mirone, P. Tafforeau, Three-dimensional synchrotron virtual paleohistology: a new insight into the world of fossil bone microstructures. *Microsc. Microanal.* **18**, 1095–1105 (2012).
- 65. D. Davesne, F. J. Meunier, M. Friedman, R. B. J. Benson, O. Otero, Histology of the endothermic opah (*Lampris* sp.) suggests a new structure–function relationship in teleost fish bone. *Biol. Lett.* **14**, 20180270 (2018).
- 66. D. L. Rabosky, *et al.*, An inverse latitudinal gradient in speciation rate for marine fishes. *Nature* **559**, 392–395 (2018).
- 67. E. Paradis, K. Schliep, ape 5.0: an environment for modern phylogenetics and evolutionary analyses in R. *Bioinformatics* **35**, 526–528 (2019).
- 68. D. Schluter, T. Price, A. Ø. Mooers, D. Ludwig, Likelihood of ancestor states in adaptive radiation. *Evolution (N. Y).* **51**, 1699–1711 (1997).

Figures

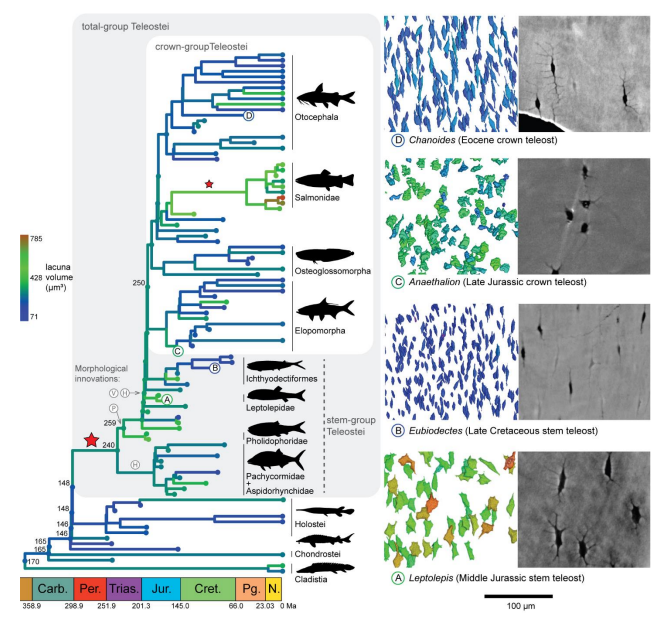


Figure 1. Evolution of osteocyte lacuna volume in fossil and modern teleosts revealed by synchrotron microtomography. Timescaled composite phylogeny of actinopterygians with mapped cell-size volumes for fossil and modern lineages. Red stars visualise the inferred occurrences of whole-genome duplications: the teleost WGD (large star), and the salmonid-specific WGD (small star). Renderings of osteocyte lacunae shown for highlighted fossil taxa branching close in time to the inferred teleost-specific WGD (A, C) and those substantially postdating it (B, D). Letters correspond to key teleost synapomorphies, mapped to the corresponding node based on morphological phylogeny (35): P, mobile premaxilla; H, homocercal caudal fin; A, vertebral autocentra.

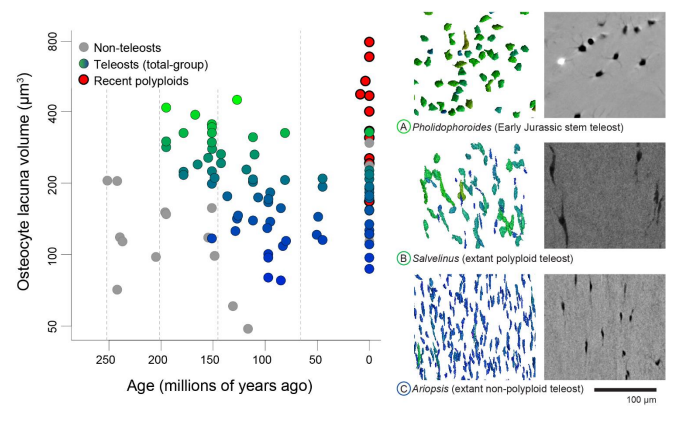


Figure 2. Evolution of osteocyte lacuna volume in fossil and modern teleosts revealed by synchrotron microtomography, plotted against time. We show cell size in an Early Jurassic stem-group teleost (A), within the range of modern polyploid species (exemplified by the salmonid *Salvelinus*; B), and indicative of a WGD. A steady decrease in cell size through time is evident in the fossil record, and is reflected by the relatively small cells of many extant teleosts, which lack polyploidy events subsequent to the ancestral teleost WGD (e.g. the catfish *Ariopsis*; C). Note semi-log axes (osteocyte lacuna volume values are log-transformed). Color scale follows Fig. 1.

Supplementary Information for

Fossilized cell structures identify an ancient origin for the teleost whole-genome duplication.

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This file includes:

Supplementary text
Figures S1 to S3
Legends for Datasets S1 to S4
SI References

Other supplementary materials for this manuscript include the following:

Datasets S1 to S4

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Supplementary Information Text

I. Complete list of specimens used in the study

This list includes all fossil specimens that we SRµCT-scanned for the present study. Some of the specimens were not included in the downstream analysis because they did not yield exploitable data in the form of osteocyte lacuna volumes.

I.1. Institutional abbreviations for fossil specimens

IRSNB – Institut Royal des Sciences Naturelles de Belgique, Brussels, Belgium; MB.f – Museum für Naturkunde, Berlin, Germany; MCSNB – Museo Civico di Scienze Naturali, Bergamo, Italy; MNHN – Muséum National d'Histoire Naturelle, Paris, France; NHMUK – Natural History Museum, London, UK; OUMNH – Oxford University Museum of Natural History, Oxford, UK; PIMUZ – Paläontologisches Institut und Museum, Zurich, Switzerland; UMMP – University of Michigan Museum of Paleontology, Ann Arbor, USA.

I.2. Justification of the phylogenetic and stratigraphic placement of the fossil species used in the analyses

We used the Rabosky *et al.* (1) phylogeny as our reference tree for extant taxa, since it was the most complete molecular tree of actinopterygians available at the date of our study. Moreover, it provides estimates of divergence times that are amongst the closest to what is suggested by the fossil record (i.e. 192 Ma for the origin of crown-group teleosts).

- †*Paleolox larsoni* Kimmel, 1975

Specimen number: UMMP V74157

Position: stem-group *Salvelinus* according to (2, 3)

Position in the reference tree: sister to the clade (crown *Salvelinus*) comprising *Salvelinus leucomaenis leucomaenis*, *Salvelinus confluentus*, their most recent common ancestor and all its descendants

Age / Locality: Miocene / Chalk Hills Formation, USA – 9.8-7.4 Ma (2, 4)

Minimum divergence time: † Paleolox larsoni itself is the oldest representative of the total-group

Salvelinus.

- †Prosopium prolixus Smith, 1975

Specimen number: UMMP 21728

Position: in total-group *Prosopium* according to (2)

Position in the reference tree: sister to the clade (crown *Prosopium*) comprising *Prosopium coulterii*, *Prosopium spilonotus*, their most recent common ancestor and all its descendants

Age / Locality: Pliocene / Glenns Ferry Formation, USA – 5.0-2.5 Ma (2, 4)

Minimum divergence time: †*P. prolixus* itself is the oldest representative of the total-group

Prosopium.

- †Spaniodon elongatus Pictet, 1850

Specimen number: NHMUK PV OR 44831

Position: crown Euteleostei, sister to "Osmeroidei" (i.e. Osmeriformes in the reference

phylogeny) according to (5)

Position in the reference tree: sister to the clade (crown Osmeriformes) comprising *Retropinna retropinna*, *Osmerus eperlanus*, their most recent common ancestor and all its descendants

Age / Locality: Late Santonian / Sahel Alma, Lebanon – 86.3-83.6 Ma (6, 7)

Minimum divergence time: † Spaniodon itself is the oldest representative of the total-group

Osmeriformes.

- †Pattersonella formosa (Traquair, 1908)

Specimen number: IRSNB Vert-01680-00240 ce

Position: crown Euteleostei, is an "argentinoid" (i.e. Argentiniformes in the reference phylogeny)

according to (8)

Position in the reference tree: sister to the clade (crown Argentiniformes) comprising *Argentina silus. Bathylagus antarcticus*, their most recent common ancestor and all its descendants

Age / Locality: Barremian-Aptian / Bernissart, Belgium – 128.0-126.0 Ma (9, 10)

Minimum divergence time: † *Pattersonella* itself is the oldest representative of the total-group Argentiniformes.

- †Leptolepides sprattiformis (Blainville, 1818)

Specimen number: MB.f.18203

Position: total-group Euteleostei according to (11–14)

Position in the reference tree: sister to the clade (crown Euteleostei) comprising Lepidogalaxias

salamandroides, Oncorhynchus mykiss, their most recent common ancestor and all its

descendants

Age / Locality: Kimmeridgian-Tithonian / Solnhofen Limestone, Germany - 150.94 Ma (15) **Minimum divergence time**: †*Leptolepides* itself is the oldest representative of the total-group

Euteleostei.

- †Chanoides macropoma Agassiz, 1843

Specimen number: NHMUK PV P 37226 **Position:** stem Otophysi according to (16, 17)

Position in the reference tree: sister to the clade (crown Otophysi) comprising Gymnotus

carapo. Danio rerio, their most recent common ancestor and all its descendants

Age / Locality: late Ypresian / Bolca, Italy – 49.11 Ma (15)

Minimum divergence time: †Santanichthys diasi, oldest representative of the total-group Otophysi, probably within the clade formed by †Chanoides and crown Otophysi – 113.4-110 Ma (18).

- †Tharrias araripis Jordan & Branner, 1908

Specimen number: NHMUK PV P 54675b

Position: stem Chanidae (crown Gonorhynchiformes), sister to Chanos chanos according to (10)

Position in the reference tree: sister to Chanos chanos

Age / Locality: early Albian / Romualdo Member of the Santana Formation, Brazil - 113.4-110

Ma (10)

Minimum divergence time: † Tharrias itself is the oldest representative of the clade it forms with Chanos.

- † Dastilbe elongatus Silva-Santos, 1947

Specimen number: NHMUK PV P 63381

Position: stem Chanidae (crown Gonorhynchiformes), sister to [† *Tharrias* + *Chanos*] according to (10)

Position in the reference tree: sister to the clade comprising †*Tharrias araripes, Chanos chanos*, their most recent common ancestor and all its descendants

Age / Locality: late Aptian / Nova Olinda Member of the Crato Formation, Brazil – 123.0-113.0 Ma (10)

Minimum divergence time: † Dastilbe itself is the oldest representative of the clade it forms with † Tharrias and Chanos.

- † Aethalionopsis robustus (Traquair, 1911)

Specimen number: IRSNB Vert-01680-00556

Position: stem Chanidae (crown Gonorhynchiformes), sister to [†Dastilbe + †Tharrias + Chanos] according to (10, 19)

Position in the reference tree: sister to the clade comprising † *Dastilbe elongatus, Chanos chanos,* their most recent common ancestor and all its descendants

Age / Locality: Barremian-Aptian / Bernissart, Belgium – 128.0-126.0 Ma (9, 10) Minimum divergence time: †Aethalionopsis itself is the oldest representative of the clade it forms with †Tharrias, †Dastilbe and Chanos.

- †Knightia sp.

Specimen number: UMMP Tmp-1008

Position: crown Clupeidae (crown Clupeiformes), Pellonulinae according to (20)

Position in the reference tree: sister to the clade (crown Pellonulinae) comprising Sierrathrissa

leonensis, Pellonula vorax, their most recent common ancestor and all its descendants

Age / Locality: Eocene / Green River Formation, USA - 55.8-33.9 Ma (20)

Minimum divergence time: †Knightia itself is the oldest representative of the total-group

Pellonulidae.

- † Armigatus namourensis Forey et al., 2003

Specimen number: NHMUK PV P 63151a **Position:** stem Clupeiformes according to (21)

Position in the reference tree: sister to the clade (crown Clupeiformes) comprising *Denticeps*

clupeoides, Pellonula vorax, their most recent common ancestor and all its descendants Age / Locality: early to middle Cenomanian / Namoura, Lebanon – 99.7-94.4 Ma (7) Minimum divergence time: †Ezkutuberezi carmeni from the Valanginian, oldest †Ellimichthyiformes (within the clade that †Armigatus forms with crown Clupeiformes) – 132.9-

139.8 Ma (21)

- †Ornategulum sardinioides (Pictet, 1850)

Specimen number: NHMUK PV P 52508

Position: stem Clupeiformes according to (20, 22), possibly as sister to [† Armigatus + crown

Clupeiformes1

Position in the reference tree: sister to the clade comprising † *Armigatus namourensis*. Denticeps clupeoides, their most recent common ancestor and all its descendants Age / Locality: Cenomanian / Hakel and Namoura, Lebanon (7) - 99.7-94.4 Ma Minimum divergence time: †Ezkutuberezi carmeni from the Valanginian, oldest †Ellimichthyiformes (within the clade that † Ornategulum forms with † Armigatus and crown Clupeiformes) - 132.9-139.8 Ma (21)

- †Plethodus expansus (Dixon, 1850)

Specimen number: NHMUK PV P 49895

Position: member of †Tselfatiiformes, which are crown Teleostei, sister to crown Clupeocephala

according to (23)

Position in the reference tree: sister to the clade (crown Clupeocephala) comprising *Denticeps* clupeoides, Oncorhynchus mykiss, their most recent common ancestor and all its descendants

Age / Locality: Cenomanian-Turonian / English Chalk, UK - 100.5-89.8 Ma (24)

Minimum divergence time: †Leptolepides from the Kimmeridgian-Tithonian Solnhofen Limestone is the oldest representative of crown Clupeocephala - 150.94 Ma (15)

- †Rhacolepis buccalis Agassiz, 1841

Specimen number: UMMP 101952

Position: within the †Crossognathiformes clade of (25), sister to crown Clupeocephala (that includes Clupeomorpha, Ostariophysi and Euteleostei). According to (23), †Crossognathiformes are sister to [†Tselfatiiformes + crown Clupeocephala]. Sister to Euteleostei (represented by Salmo salar) according to (26)

Position in the reference tree: We adopt the phylogenetic position proposed by (25), as it is a recent phylogeny that incorporates many †crossognathiform and crown teleost taxa: sister to the clade (crown Clupeocephala + †Tselfatiiformes) comprising †Plethodus expansus, Oncorhynchus mykiss, their most recent common ancestor and all its descendants

Age / Locality: Albian / Santana Formation, Brazil – 113.4-110 Ma (10, 27)

Minimum divergence time: †*Leptolepides sprattiformis* from the Kimmeridgian-Tithonian Solnhofen Limestone is the oldest representative of crown Clupeocephala - 150.94 Ma (15)

- †Apsopelix anglicus (Dixon, 1850)

Specimen number: NHMUK PV P 4246

Position: within the †Crossognathiformes clade of (25) **Position in the reference tree:** sister to †*Rhacolepis buccalis*

Age / Locality: Cenomanian-Turonian / English Chalk, UK – 100.5-89.8 Ma (24)

Minimum divergence time: †Bavarichthys incognitus (13) from the Kimmeridgian-Tithonian Solnhofen Limestone is the oldest representative of the †Crossognathiformes clade of (25) -

150.94 Ma (15)

- †Phareodus encaustus Cope, 1871

Specimen number: NHMUK PV P 64636I

Position: stem Osteoglossidae according to (28)

Position in the reference tree: sister to the clade (crown Osteoglossidae) comprising Osteoglossum bicirrhosum, Arapaima gigas, their most recent common ancestor and all its

descendants

Age / Locality: Eocene / Laney Member, Green River Formation, USA - 55.8-33.9 Ma **Minimum divergence time:** †*Cretophareodus alberticus* from the Campanian is the oldest representative of the clade formed by †*Phareodus* and crown Osteoglossidae – 83.6-72.1 Ma

- † Brychaetus muelleri Woodward, 1901

Specimen number: NHMUK PV OR 28424 **Position:** sister to †*Phareodus* according to (29)

Position in the reference tree: sister to †*Phareodus encaustus* **Age / Locality:** Ypresian / Sheppey, London Clay, UK – 50.5 Ma (24)

Minimum divergence time: †Brychaetus itself is the oldest representative of the clade it forms

with †Phareodus.

- † Eohiodon falcatus Grande, 1979

Specimen number: NHMUK PV P 61245

Position: stem Hiodontidae according to (28, 30, 31)

Position in the reference tree: sister to the clade (crown Hiodontidae) comprising *Hiodon alosoides, Hiodon tergisus*, their most recent common ancestor and all its descendants

Age / Locality: Eocene / Green River Formation, USA - 55.8-33.9 Ma

Minimum divergence time: †Eohiodon itself is the oldest representative of the clade it forms

with Hiodon.

Elopomorpha have two alternative topologies in the reference phylogenetic study of (6). Here we chose the hypothesis presented in the Fig. 3 of (6) as it is better resolved for the fossil taxa of our sample.

- † Hayenchelys germanus (Hay, 1903)

Specimen number: NHMUK PV P 62726

Position: stem Anguilliformes according to (32)

Position in the reference tree: sister to the clade (crown Anguilliformes) comprising *Diastobranchus capensis* + *Anguilla anguilla*, their most recent common ancestor and all its descendants

Age / Locality: under the †*Mantilliceras mantelli* Ammonite zone, Cenomanian / Hadjula, Lebanon (6) – 98.0 Ma

Minimum divergence time: †*Hayenchelys* itself is the oldest representative of the total-group

Anguilliformes.

- †Istieus grandis Agassiz, 1842

Specimen number: NHMUK PV P 3886

Position: stem Pterothrissidae according to (6)

Position in the reference tree: sister to *Pterothrissus gissu*

Age / Locality: younger than the †Scaphites hippocrepis III Zone but older than the †Baculites

mclearni Zone (6) / Coesfeld Member of Westphalia, Germany – 81.28-80.67 Ma **Minimum divergence time:** † *Istieus* is the oldest representative of the total-group

Pterothrissidae.

- †Lebonichthys gracilis (Davis, 1887)

Specimen number: NHMUK PV P 49076 **Position:** stem Albulidae according to (6)

Position in the reference tree: sister to the clade (crown Albulidae) comprising *Albula glossodonta, Albula vulpes*, their most recent common ancestor and all its descendants

Age / Locality: Late Santonian / Sahel Alma, Lebanon - 86.3-83.6 Ma (6, 7)

Minimum divergence time: †Lebonichthys itself is the oldest representative of the clade it forms

with crown Albulidae.

- †Brannerion vestitum Jordan, 1920

Specimen number: NHMUK PV P 60923

Position: sister to [Anguilliformes + Notacanthiformes] according to (6)

Position in the reference tree: † Brannerion is found as sister to [Anguilliformes + Notacanthiformes] in (6), but this clade does not exist in the reference phylogeny, unless Albuliformes are also included. We chose the conservative approach to position it as sister to [Anguilliformes + Notacanthiformes + Albuliformes]: the clade comprising Anguilla anguilla, Albula vulpes, their most recent common ancestor and all its descendants

Age / Locality: early Albian / Romualdo Member of the Santana Formation, Brazil – 113.4-110

Ma (6)

Minimum divergence time: †Baugeichthys caeruleus, earliest representative of the total-group Albuliformes – 134.5-129.4 Ma (1, 6)

- †Osmeroides sp. (presumably †Osmeroides lewesiensis)

Specimen number: OUMNH K.64151

Position: sister to [Albuliformes + Notacanthiformes + Anguilliformes] according to (6) **Position in the reference tree:** sister to the clade comprising †*Brannerion vestitum, Albula vulpes*, their most recent common ancestor and all its descendants [following the proposed position of †*Brannerion*]

Age / Locality: †*Hoplites dentatus* through †*Mortoniceras inflatus* zones of the Albian / Gault Clay Formation, UK – 110.22 Ma

Minimum divergence time: †Baugeichthys caeruleus, earliest representative of the total-group Albuliformes – 134.5-129.4 Ma (1, 6)

- †Flindersichthys denmaedi Longman, 1932

Specimen number: NHMUK PV P 3886

Position: stem Megalopidae (crown Elopiformes) according to (33)

Position in the reference tree: sister to the clade (crown Megalopidae) comprising *Megalops cyprinoides, Megalops atlanticus*, their most recent common ancestor and all its descendants **Age / Locality:** Albian / Stewart Creek, Rolling Downs Formation, Australia – 113.0-100.5 Ma **Minimum divergence time:** † *Flindersichthys* itself is the oldest representative of the clade it forms with *Megalops*.

- † Arratiaelops vectensis (Woodward, 1890)

Specimen number: NHMUK PV P 42013

Position: stem Megalopidae (crown Elopiformes) according to (33)

Position in the reference tree: sister to the clade comprising † Flindersichthys denmaedi.

Megalops cyprinoides, their most recent common ancestor and all its descendants

Age / Locality: Barremian-Aptian / Bernissart, Belgium - 126.0 Ma

Minimum divergence time: † *Arratiaelops* itself is the earliest representative of the total-group Megalopidae.

- † Ichthyemidion vidali Poyato-Ariza, 1995

Specimen number: MNHN.F.MSE356a

Position: stem Elopiformes according to (6)

Position in the reference tree: sister to the clade (crown Elopiformes) comprising Elops saurus,

Megalops cyprinoides, their most recent common ancestor and all its descendants

Age / Locality: Late Berriasian – Early Valanginian / El Montsec, Spain – 131.3-125.9 Ma (6) **Minimum divergence time:** † *Ichthyemidion* itself is the oldest representative of the clade it forms with crown Elopiformes.

- † Anaethalion angustus (Münster, 1839)

Specimen number: NHMUK PV OR 37926

Position: stem-Elopidae (crown Elopiformes) according to (34), but may be a stem Elopiformes

according to (12, 25)

Position in the reference tree: since the position of †*Anaethalion* varies from one phylogenetic analysis to the other, we chose the conservative approach to place it as a stem Elopiformes: sister to the clade comprising †*Ichthyemidion vidali*, *Megalops cyprinoides*, their most recent common ancestor and all its descendants

Age / Locality: †*Neochetoceras rebouletianum* Horizon within the †*Lithacoceras ulmense* Subzone of the †*Hybonoticeras beckeri* Ammonite Zone, Late Kimmeridgian / Eichstätt, Germany – 151.2 Ma (6)

Minimum divergence time: †*Anaethalion* itself is the oldest representative of the total-group Elopiformes, and of crown Elopomorpha.

†Ichthyodectiformes are a clade of stem Teleostei, which position varies from one study to another. Some studies find them as the sister to crown Teleostei (12, 25) while others find some stem Teleostei closer to the crown group than †Ichthyodectiformes (26, 35). We chose to follow (25) since it is a recent phylogenetic analysis including several †ichthyodectiform representatives alongside a comprehensive sample of stem- and crown Teleostei. Other recent large-scale studies either did not include any †ichthyodectiform representative (34, 36) or only one †ichthyodectiform (26). We thus interpret †Ichthyodectiformes as sister to crown Teleostei.

- †Xiphactinus audax Leidy, 1870

Specimen number: UMMP 11003

Position: in †Ichthyodectiformes, family †Ichthyodectidae (37)

Position in the reference tree: sister to the clade (crown Teleostei) comprising *Megalops cyprinoides*, *Oncorhynchus mykiss*, their most recent common ancestor and all its descendants

Age / Locality: Turonian-Maastrichtian / USA – 93.9-66 Ma (37)

Minimum divergence time: † Occithrissops wilsoni (38) is the oldest representative of †Ichthyodectiformes, and of the clade they form with crown Teleostei – Bathonian, 168.3-166.1 Ma

- †Ichthyodectes ctenodon Cope, 1870

Specimen number: UMMP V56318

Position: in †Ichthyodectiformes, family †Ichthyodectidae (37) **Position in the reference tree:** sister to †*Xiphactinus audax* **Age / Locality:** Turonian-Campanian / USA – 93.9-72.1 Ma (37)

Minimum divergence time: † Ichthyodectes and † Xiphactinus themselves are the oldest

representatives of the clade (†Ichthyodectidae) they form together.

- † Eubiodectes libanicus (Pictet & Humbert, 1866)

Specimen number: NHMUK PV P 62691a

Position: in †Ichthyodectiformes, family †Cladocyclidae (37)

Position in the reference tree: sister to the clade (†Ichthyodectidae) comprising †*Xiphactinus audax*, †*Ichthyodectes ctenodon*, their most recent common ancestor and all its descendants **Age / Locality:** Cenomanian / Hakel, Lebanon (7) – 99.7-94.4 Ma

Minimum divergence time: †*Chiromystus mawsoni* is the oldest representative of †Cladocyclidae (and of the clade †*Eubiodectes* forms with †Ichthyodectidae) – Hauterivian-Barremian, 132.9-125 Ma (37).

- †Thrissops formosus Agassiz, 1833

Specimen number: NHMUK PV OR 35013

Position: in †Ichthyodectiformes, outside of the clade that includes †Cladocyclidae and

†Ichthyodectidae (37)

Position in the reference tree: sister to the clade comprising †*Xiphactinus audax*, †*Eubiodectes*

libanicus, their most recent common ancestor and all its descendants **Age / Locality:** Kimmeridgian / Eichstätt, Germany – 151.2 Ma (6)

Minimum divergence time: †Thrissops itself is the oldest representative of the clade it forms

with †Ichthyodectidae and †Cladocyclidae.

- † Allothrissops regleyi (Thiollière, 1854)

Specimen number: NHMUK PV P 921

Position: in †Ichthyodectiformes, outside of the clade that includes †Thrissops, †Cladocyclidae

and †Ichthyodectidae (37)

Position in the reference tree: sister to the clade comprising †*Thrissops formosus*, †*Xiphactinus*

audax, their most recent common ancestor and all its descendants

Age / Locality: Kimmeridgian / Eichstätt, Germany – 151.2 Ma (6)

Minimum divergence time: † Occithrissops wilsoni (38) is the oldest representative of the clade

†Allothrissops forms with other †Ichthyodectiformes (37) – Bathonian, 168.3-166.1 Ma.

- †Pachythrissops laevis Woodward, 1919

Specimen number: NHMUK PV P 41859

Position: (37) considers the inclusion of †*Pachythrissops* inside †lchthyodectiformes to be dubious. However, several analyses, including the extensive one of (25) find it to be an †ichthyodectiform, sister to all other members of the clade.

Position in the reference tree: sister to the clade comprising †*Allothrissops regleyi*, †*Xiphactinus audax*, their most recent common ancestor and all its descendants

Age / Locality: Valanginian / Purbeck Group, UK – 139.8-132.9 Ma

Minimum divergence time: † *Occithrissops wilsoni* (38) is the oldest representative of the clade † *Pachythrissops* forms with other † Ichthyodectiformes – Bathonian, 168.3-166.1 Ma (37).

Stem-group teleosts (outside of †Ichthyodectiformes): Several recent phylogenetic studies have tried to resolve the relationships between stem teleost taxa (25, 26, 34, 36, 39). They all recover different topologies for our sampled taxa, but there are several areas of agreement, most notably:

- 1) a clade including †*Leptolepis coryphaenoides*, †*Tharsis*, †*Ascalabos*, †Ichthyodectiformes (when included) and crown Teleostei;
- 2) a clade including the above, plus † Dorsetichthys, † Siemensichthys (and † Ankylophorus when included), and † Eurycormus;
- 3) †Pholidophoridae sensu stricto, as defined by (34) as sister to this clade:
- 4) †Aspidorhynchidae and †Pachycormiformes outside of the clade formed by all of the above. †*Ichthyokentema* is a wildcard taxon and has a different position in every study where it is included.

Since the positions of the different taxa constitutive of these various clades vary from one study to another, we cannot choose which topology is the most likely to reflect reality. In that context, we have chosen to follow the study that has the most comprehensive taxon sample and the best resolution for each portion of the total-group teleost tree: (25) for the clade described in (1), and (34) for the rest of the tree.

- †Tharsis dubius (Blainville, 1818)

Specimen number: NHMUK PV OR 37852b

Position: sister to [†Ascalabos + †Varasichthyidae + †Ichthyodectiformes + crown Teleostei]

according to (25)

Position in the reference tree: sister to the clade comprising †*Xiphactinus audax*, *Megalops*

cyprinoides, their most recent common ancestor and all its descendants

Age / Locality: Kimmeridgian-Tithonian / Solnhofen, Germany -157.3-145 Ma

Minimum divergence time: † Occithrissops wilsoni (38) is the oldest representative of the clade

†Ascalabos forms with crown teleosts (25) - Bathonian, 168.3-166.1 Ma.

- †Leptolepis coryphaenoides (Bronn, 1830)

Specimen number: NHMUK PV P 947c

Position: sister to [all of the above, including crown Teleostei] according to (25, 34, 36, 39) **Position in the reference tree:** sister to the clade comprising † *Tharsis dubius, Megalops*

cyprinoides, their most recent common ancestor and all its descendants

Age / Locality: Toarcian / Curcy, France – 182.7-174.1 Ma (26)

Minimum divergence time: †Leptolepis coryphaenoides itself is the oldest representative of the

clade it forms with crown teleosts.

- †Leptolepis disjectus Woodward, 1895

Specimen number: NHMUK PV P 3793a

Position: †Leptolepis disjectus is not included in any phylogenetic analysis, therefore we

tentatively consider it to be closely related to the congeneric †L. coryphaenoides

Position in the reference tree: sister to †*Leptolepis coryphaenoides* **Age / Locality:** Bathonian / Thaynton Limestone, UK – 168.3-166.1 Ma

Minimum divergence time: †Leptolepis coryphaenoides is the oldest representative of the clade

†Leptolepis forms with crown teleosts – 182.7-174.1 Ma.

- †Ichthyokentema purbeckensis (Davies, 1887)

Specimen number: NHMUK PV P 7807

Position: sister to [†Leptolepis coryphaenoides + all of the above, including crown Teleostei],

according to (34)

Position in the reference tree: sister to the clade comprising †Leptolepis coryphaenoides,

Megalops cyprinoides, their most recent common ancestor and all its descendants

Age / Locality: Berriasian / Purbeck Group, UK - 145-139.8 Ma

Minimum divergence time: †Leptolepis coryphaenoides is the oldest representative of the clade

†Ichthyokentema forms with crown teleosts – 182.7-174.1 Ma.

- †Dorsetichthys bechei (Agassiz, 1837)

Specimen number: OUMNH J.3369

Position: sister to [†Ichthyokentema + all of the above, including crown Teleostei] according to

(34)

Position in the reference tree: sister to the clade comprising † Ichthyokentema purbeckensis,

Megalops cyprinoides, their most recent common ancestor and all its descendants

Age / Locality: Sinemurian / Lyme Regis, UK – 199.3-190.8 Ma (26, 34)

Minimum divergence time: † Dorsetichthys itself is the oldest representative of the clade it forms

with crown teleosts.

- †Siemensichthys macrocephalus (Agassiz, 1834)

Specimen number: NHMUK PV P 9135

Position: sister to [†Dorsetichthys + all of the above, including crown Teleostei] according to (34)

Position in the reference tree: sister to the clade comprising † *Dorsetichthys bechei, Megalops*

cyprinoides, their most recent common ancestor and all its descendants

Age / Locality: Kimmeridgian-Tithonian / Solnhofen, Germany -157.3-145 Ma

Minimum divergence time: †Dorsetichthys bechei is the oldest representative of the clade

†Siemensichthys forms with crown teleosts - 199.3-190.8 Ma.

- †Ankylophorus similis (Woodward, 1895)

Specimen number: NHMUK PV P 1083

Position: sister to † Siemensichthys according to (34, 36)

Position in the reference tree: sister to †Siemensichthys macrocephalus **Age / Locality:** Kimmeridgian-Tithonian / Solnhofen, Germany –157.3-145 Ma

Minimum divergence time: †Ankylophorus and †Siemensichthys themselves are the oldest

representatives of the clade they form together (†Ankylophoridae).

- † Eurycormus speciosus Wagner, 1863

Specimen number: MB.f.7016

Position: sister to [†Ankylophoridae + all of the above, including crown Teleostei] according to

(34)

Position in the reference tree: sister to the clade comprising † Siemensichthys macrocephalus,

Megalops cyprinoides, their most recent common ancestor and all its descendants **Age / Locality:** Kimmeridgian-Tithonian / Solnhofen, Germany –157.3-145 Ma

Minimum divergence time: †Dorsetichthys bechei is the oldest representative of the clade

†Eurycormus forms with crown teleosts – 199.3-190.8 Ma.

- †Pholidophoropsis caudalis (Woodward, 1895)

Specimen number: OUMNH J.3363

Position: according to (34), part of †Pholidophoridae *sensu stricto*, a clade of Triassic and Early Jurassic stem Teleostei, represented in the phylogenetic analysis by Triassic genera and not by †*Pholidophoropsis*. The clade is sister to [†*Eurycormus* + all of the above, including crown Teleostei], according to (26, 34)

Position in the reference tree: sister to the clade comprising †*Eurycormus speciosus*, *Megalops cyprinoides*, their most recent common ancestor and all its descendants

Age / Locality: Sinemurian / Lyme Regis, UK - 199.3-190.8 Ma

Minimum divergence time: †*Pholidophoretes salvus* and †*Knerichthys bronni* are the oldest representatives of the †*Pholidophoridae sensu stricto*, and thus of the clade †*Pholidophoropsis* forms with crown teleosts – early Carnian, 237-227 Ma (34).

- †Pholidophoroides crenulata Egerton, 1843

Specimen number: NHMUK PV OR 36313

Position: according to (34), part of †Pholidophoridae *sensu stricto*, a clade of Triassic and Early Jurassic stem Teleostei, represented in the phylogenetic analysis by Triassic genera and not by †*Pholidophoroides*. The clade is sister to [†*Eurycormus* + all of the above, including crown Teleostei], according to (26, 34)

Position in the reference tree: sister to †Pholidophoropsis caudalis

Age / Locality: Sinemurian / Lyme Regis, UK - 199.3-190.8 Ma

Minimum divergence time: as †*Pholidophoroides* and †*Pholidophoropsis* were never included in a formal phylogenetic analysis, their position within †Pholidophoridae is unknown. Since they are both of the same age and locality, we chose the conservative approach to give their own age as the minimum divergence time for the clade they form together.

- †Protosphyraena sp.

Specimen number: NHMUK PV P 9644b

Position: in †Pachycormiformes (40, 41), the latter forming a clade with †Aspidorhynchiformes that is sister to all other total-group Teleostei, according to (26, 34, 36)

Position in the reference tree: sister to the clade comprising †*Prohalecites porroi, Megalops cyprinoides*, their most recent common ancestor and all its descendants

Age / Locality: Coniacian-Campanian / Smoky Hill Member, Niobrara Group, USA – 89.8-72.1 Ma

Minimum divergence time: †*Prohalecites porroi* is the oldest representative of the total-group Teleostei – Ladinian-Carnian, 237 Ma (34, 42).

- †Hypsocormus sp.

Specimen number: NHMUK PV P 46348b

Position: in †Pachycormiformes, sister to the 'macrocarnivore clade' comprising

†Protosphyraena (40, 41, 43)

Position in the reference tree: sister to †*Protosphyraena* sp. **Age / Locality:** Callovian / Oxford Clay, UK – 166.1-163.6 Ma

Minimum divergence time: †Hypsocormus itself is the oldest representative of the clade it forms

with †Protosphyraena (40, 43).

- †Rhinconichthys taylori Friedman et al., 2010

Specimen number: NHMUK PV OR 33219

Position: in †Pachycormiformes, sister to the clade comprising †Hypsocormus and

†Protosphyraena (40, 41, 43)

Position in the reference tree: sister to the clade comprising †*Protosphyraena* sp., †*Hypsocormus* sp., their most recent common ancestor and all its descendants **Age / Locality:** Lower Chalk, Cenomanian / Burnham, UK – 100.5-93.3 Ma (41) **Minimum divergence time:** †*Pachycormus*, †*Euthynotus* and other taxa are the oldest representatives of the clade †*Rhinconichthys* forms with †*Protosphyraena* and †*Hypsocormus* (40, 43) – late Toarcian, 182.7-174.1 Ma.

- †Pachycormus macropterus (Blainville, 1818)

Specimen number: MNHN.F.STC103 (as. †*P. curtus*)

Position: in †Pachycormiformes, sister to the 'filter-feeding clade' comprising †*Rhinconichthys* (40, 41, 43). †*Pachycormus curtus* is considered a junior synonym of †*P. macropterus* by (43)

Position in the reference tree: sister to †*Rhinconichthys taylori* **Age / Locality:** late Toarcian / Curcy, France – 182.7-174.1 Ma

Minimum divergence time: † *Pachycormus* itself is the oldest representative of the clade it forms

with †Rhinconichthys (40, 43).

- † Euthynotus incognitus (Blainville, 1818)

Specimen number: NHMUK PV P 2044

Position: in †Pachycormiformes, sister to all other members of the clade (40, 41, 43)

Position in the reference tree: sister to the clade comprising †*Protosphyraena* sp., †*Pachycormus macropterus*, their most recent common ancestor and all its descendants Age / Locality: Toarcian / *Posidonia* shale, Holzmaden, Germany – 182.7-174.1 Ma

Minimum divergence time: †*Euthynotus* itself is the oldest representative of †Pachycormiformes (40, 43).

- † Vinctifer comptoni (Agassiz, 1841)

Specimen number: UMMP 101950

Position: in †Aspidorhynchiformes, themselves sister to †Pachycormiformes according to (26, 34, 36)

Position in the reference tree: sister to the clade (†Pachycormiformes) comprising †*Euthynotus incognitus*, †*Pachycormus macropterus*, their most recent common ancestor and all its descendants

Age / Locality: early Albian / Romualdo Member of the Santana Formation, Brazil – 113.4-110 Ma (44)

Minimum divergence time: † *Euthynotus incognitus* is the oldest representative of the [†Pachycormiformes + †Aspidorhynchiformes] clade (26, 40) – Toarcian, 182.7-174.1 Ma.

- †Belonostomus sp.

Specimen number: NHMUK PV P 4029

Position: in †Aspidorhynchiformes, sister to †Vinctifer according to (44)

Position in the reference tree: sister to † Vinctifer comptoni

Age / Locality: early to middle Cenomanian / Namoura, Lebanon – 99.7-94.4 Ma (7, 44)

Minimum divergence time: †Belonostomus tenuirostris and †B. dorsetensis are the oldest representatives of †Belonostomus – Kimmeridgian, 157.3-152.1 Ma (44).

- †Belonostomus muensteri (Agassiz, 1834)

Specimen number: NHMUK PV OR 37796
Position: in †Belonostomus according to (44)

Position in the reference tree: sister to †*Belonostomus* sp.

Age / Locality: Tithonian / Solnhofen Limestone, Germany – 152.1-145 Ma (44)

Minimum divergence time: †Belonostomus tenuirostris and †B. dorsetensis are the oldest

representatives of †Belonostomus – Kimmeridgian, 157.3-152.1 Ma (44).

- † Aspidorhynchus fisheri Egerton, 1854

Specimen number: NHMUK PV P 17317

Position: in †Aspidorhynchiformes, sister to [†Belonostomus + †Vinctifer] according to (44)

Position in the reference tree: sister to the clade comprising † *Vinctifer comptoni*, † *Belonostomus* sp., their most recent common ancestor and all its descendants

Age / Locality: Berriasian / Purbeck Group, UK – 145-139.8 Ma

Minimum divergence time: †Aspidorhynchus crassus is the oldest representative of

†Aspidorhynchiformes – Bathonian, 168.3-166.1 Ma (44).

- † Aspidorhynchus acutirostris (Blainville, 1818)

Specimen number: NHMUK PV P 97

Position: in † *Aspidorhynchus* according to (44)

Position in the reference tree: sister to †*Aspidorhynchus fisheri* **Age / Locality:** Kimmeridgian / Kimmeridge Clay, UK– 157.3-152.1 Ma

Minimum divergence time: †Aspidorhynchus crassus is the oldest representative of

†Aspidorhynchiformes – Bathonian, 168.3-166.1 Ma (44).

- †Semionotus elegans Newberry, 1888

Specimen number: UMMP 13664

Position: stem Gynglimodi according to (26)

Position in the reference tree: sister to the clade (crown Gynglimodi) comprising *Atractosteus tropicus*, *Lepisosteus osseus*, their most recent common ancestor and all its descendants **Age / Locality:** Hettangian-Sinemurian / Newark Supergroup, USA – 201.3-190.8 Ma (26, 45) **Minimum divergence time:** †*Semionotus bergeri* is the oldest representative of the clade †*S. elegans* forms with crown Gynglimodi – Carnian, 237-227 Ma (26).

- †Heterolepidotus dorsalis Kner, 1866

Specimen number: NHMUK PV P 10290

Position: stem Halecomorphi, potentially close to † Ophiopsis according to (46) but never

included in a phylogenetic analysis

Position in the reference tree: sister to Amia calva (crown Halecomorphi)

Age / Locality: Rhaetian / Hallein, Germany - 208.5-201.3 Ma

Minimum divergence time: †Heterolepidotus itself is the oldest representative of the clade it

forms with crown Halecomorphi.

- †Eoeugnathus megalepis Brough, 1939

Specimen number: PIMUZ T344

Position: stem Halecomorphi, close to †*Panxianichthys* and thus outside of the clade formed by †*Ophiopsis* (and thus †*Heterolepidotus*) and *Amia* according to (26, 47)

Position in the reference tree: sister to the clade comprising †Heterolepidotus dorsalis, Amia

calva, their most recent common ancestor and all its descendants

Age / Locality: Ladinian / Monte San Giorgio, Switzerland – 242-237 Ma

Minimum divergence time: †*Panxianichthys imparilis* is the oldest representative of the clade †*Eoeugnathus* forms with crown Halecomorphi – Anisian, 247.2-242 Ma.

- †Dapedium cf. politum Leach, 1822

Specimen number: OUMNH J.3323

Position: in †Dapediiformes, stem Holostei according to (26, 48)

Position in the reference tree: sister to the clade (crown Holostei) comprising Lepisosteus

osseus, Amia calva, their most recent common ancestor and all its descendants

Age / Locality: Sinemurian / Lyme Regis, UK – 199.3-190.8 Ma (49)

Minimum divergence time: † Watsonulus eugnathoides is the oldest representative of crown Holostei, and of the clade † Dapedium forms with them – Induan-Olenekian, 251.9-247.2 Ma (26, 50).

- †Perleidus sp.

Specimen number: NHMUK PV P 26456

Position: stem Neopterygii, according to (51, 52)

Position in the reference tree: sister to the clade (crown Neopterygii) comprising Amia calva,

Megalops cyprinoides, their most recent common ancestor and all its descendants

Age / Locality: Proptychites beds, Induan / Stensiö Plateaus, Greenland – 251.9-251.2 Ma Minimum divergence time: †Perleidus itself is the oldest representative of the clade it forms

with crown Neopterygii (53).

- †Pholidopleurus typus Bronn, 1858

Specimen number: NHMUK PV P 19314

Position: in †Pholidopleuridae, stem Neopterygii, outside of [†Perleidus + crown Neopterygii]

according to (51, 52)

Position in the reference tree: sister to the clade comprising † Perleidus sp., Megalops

cyprinoides, their most recent common ancestor and all its descendants

Age / Locality: Anisian-Ladinian / Besano, Italy - 247.2-237 Ma

Minimum divergence time: † Perleidus is the oldest representative of the clade † Pholidopleurus

forms with crown Neopterygii – 251.9-251.2 Ma.

- †Proscinetes elegans (Agassiz, 1833)

Specimen number: NHMUK PV P 1626

Position: in †Pycnodontiformes, that are stem Neopterygii according to (48). †Pycnodontiformes, †*Pholidopleurus* and †*Perleidus* have never been included in the same phylogenetic analysis, so we choose the conservative approach and place †*Proscinetes* outside of the clade [†*Perleidus* + †*Pholidopleurus* + crown Neopterygii]

Position in the reference tree: sister to the clade comprising †*Pholidopleurus typus, Megalops cyprinoides*, their most recent common ancestor and all its descendants

Age / Locality: lower Tithonian / Solnhofen, Germany - 152.1-145 Ma (54)

Minimum divergence time: †*Perleidus* is the oldest representative of the clade †*Proscinetes* forms with crown Neopterygii – 251.9-251.2 Ma.

- †Birgeria stensioei Aldinger, 1931

Specimen number: PIMUZ T2188

Position: traditionally viewed as a stem Chondrostei (50), † *Birgeria* is a stem Actinopteri

according to (55)

Position in the reference tree: sister to the clade (crown Actinopteri) comprising *Acipenser transmontanus*, *Megalops cyprinoides*, their most recent common ancestor and all its descendants

Age / Locality: Anisian-Ladinian / Besano, Italy – 247.2-237 Ma (56)

Minimum divergence time: † Discoserra pectinodon is the oldest representative of the total-

group Neopterygii, and crown Actinopteri (50, 55) - Bashkirian, 323.2-315.2 Ma.

I.3. Quality control criteria and specimens excluded from the analysis

We performed an initial quality-control evaluation of our scan data based on a list of three objective criteria. Specimens that did not pass any one of these criteria were not included in our analysis of evolution of osteocyte volumes and inferred genome size through time:

- (1) It was possible to confidently identify osteocyte lacunae in the PPC-SRµCT tomograms. Specimens lacking osteocytes, or in which osteocytes could not be confidently identified were not included in the analysis:
- (2) The osteocyte lacunae were well-preserved or well-defined enough in the PPC-SRµCT tomograms to guarantee that their volumes would be accurately measured. Poor definition or preservation could result from taphonomic factors (e.g. infilling of the lacunae by mineral-rich fluids) or issues with the scans themselves causing low contrast between fossil bone and its lacunae;
- (3) Sufficiently large numbers of osteocyte lacunae were present in the scan (count > ~50) to provide quantitative confidence in estimates of their average volumes. Specimens that had fewer osteocyte lacunae, even if they were well-defined, were excluded from the analysis.

- †Oncorhynchus lacustris (Cope, 1870)

Specimen numbers: UMMP 47839, UMMP 52619, UMMP 21756.

Position: in Salmonidae, close to *Oncorhynchus myki*ss according to (2, 3) **Age / Locality:** Miocene / Chalk Hills Formation, USA – 9.8-4.8 Ma (2, 4)

Failed criterion 2.

- †*Paleolox larsoni* Kimmel, 1975 Specimen number: UMMP 50352.

Failed criterion 2.

- †Lycoptera middendorffi Müller, 1848 Specimen number: NHMUK PV P 6728.

Position: stem Osteoglossomorpha according to (28)

Age / Locality: Early Cretaceous / Mt Uksuk, Mongolia - 136.4-125.5 Ma

Failed criterion 2.

- †*Anaethalion angustus* (Münster, 1839) Specimen number: NHMUK PV P 3575.

Specimen number. Namor PV

Failed criteria 2 and 3.

- †Cladocyclus gardneri Agassiz, 1841 Specimen number: UMMP 117484.

Specimen number. Owner 117404.

Position: in †Ichthyodectiformes, family †Cladocyclidae (37)

Age / Locality: Aptian-Albian / Crato and Santana Formations, Brazil - 123-110 Ma (37)

Failed criteria 2 and 3.

- † Ascalabos voithii Münster, 1839

Specimen number: NHMUK PV P 3673a.

Position: sister to [†Ichthyodectiformes + crown Teleostei] according to (25)

Age / Locality: Kimmeridgian / Kelheim, Germany - 157.3-152.1 Ma

Failed criteria 2 and 3.

- †Lombardichthys gervasuttii (Zambelli, 1980)

Specimen number: MCSNB 6662.

Position: part of †Pholidophoridae sensu stricto (36)

Age / Locality: Norian / Ponte Giurino, Italy – 210 Ma (34, 36)

Failed criterion 1.

- †Pholidorhynchodon malzannii Zambelli, 1980

Specimen number: MCSNB 11015.

Position: part of †Pholidophoridae *sensu stricto* (34, 36) **Age / Locality:** Norian / Cene, Italy – 210 Ma (34, 36)

Failed criterion 1.

- †Parapholidophorus nybelini Zambelli, 1975

Specimen number: MCSNB 6530c.

Position: part of †Pholidophoridae *sensu stricto* (34, 36) **Age / Locality:** Norian / Cene, Italy – 210 Ma (34, 36)

Failed criterion 1.

- †Pholidoctenus serianus Zambelli, 1977

Specimen number: MCSNB 4827.

Position: part of †Pholidophoridae *sensu stricto* (34, 36) **Age / Locality:** Norian / Cene, Italy – 210 Ma (34, 36)

Failed criterion 1.

- †Prohalecites porroi (Bellotti, 1857)

Specimen number: NHMUK PV P 19471.

Position: sister to [†Pholidophoridae sensu stricto + all other total-group Teleostei] according to

(34, 36)

Age / Locality: Ladinian-Carnian boundary / Perledo, Italy – 237 Ma (42)

Failed criteria 2 and 3.

- †Pachycormus macropterus (Blainville, 1818)

Specimen number: NHMUK PV P 32433.

Failed criterion 2.

- † Aspidorhynchus acutirostris (Blainville, 1818)

Specimen number: NHMUK PV P 56466.

Failed criterion 2.

- †Cheirolepis canadensis Whiteaves, 1881

Specimen number: UMMP 3453.

Position: stem Actinopterygii according to (48, 55, 57, 58)

Age / Locality: Escuminac Formation, Frasnian / Miguasha, Canada - 382.7-372.2 Ma (59)

Failed criterion 2.

II. Comments on Triassic stem-group teleosts

Stem-group teleosts are known since the Middle Triassic, the oldest ones being the pholidophorids † Malingichthys nimaiguensis and † M. wanfenglinensis from the late Ladinian (239-238 Ma) of southern China (60). These species are slightly older than † Prohalecites porroi from the latest Ladinian (237 Ma) of the Perledo-Varenna Formation (Grigna Mountains, Italy) and Kalkschieferzone (Meride Limestone) of Monte San Giorgio (42). In order to reflect as accurately as possible the temporal distribution of total-group teleosts, we included five Triassic stem-group teleosts in our sample: † Prohalecites porroi (NHMUK PV P 19471), as well as the Norian †Lombardichthys gervasuttii (MCSNB 6662), †Pholidorhynchodon malzannii (MCSNB 11015), †Parapholidophorus nybelini (MCSNB 6530c) and †Pholidoctenus serianus (MCSNB 4827) from Northern Italy (34, 36). All of these five Triassic taxa were represented by one sample from the dentary (lower jaw), and SRµCT-scanned at the Diamond Light Source I13-2 beamline, U.K. They were no issues with the SRuCT data acquisition itself, however, none of the four Norian specimens apparently preserved osteocyte lacunae. The apparent absence of preserved osteocyte lacunae may be related to the presence of abundant synsedimentary pyrite in fossils from these localities (61), which may have affected the skeletal tissue obliterating any evidence of cellular structure, but additional data on the taphonomy of the localities would be needed to substantiate this hypothesis. There is no reason to believe that osteocytes were absent in these taxa (i.e. anosteocytic bone (62, 63)), because other histological structures (e.g., growth lines, lines of reversal) were not visible either. Conversely, osteocyte lacunae have been detected and segmented in †Prohalecites porroi. However, only very few of them were preserved in the examined sample for this species and their preservation was poor, limiting their statistical power and potentially introducing a bias. For this reason, and because of the strong influence of this taxon on the results, we chose not to include it in the analyses. Due to the impossibility to obtain information from Triassic stem-group teleosts, the oldest total-group teleosts present in our sample are the Early Jurassic (Sinemurian) † Pholidophoroides crenulata and † Pholidophoropsis caudalis from Lyme Regis, U.K. Future investigations should aim to include Triassic stem-teleosts in order to more precisely define the age of the teleost WGD. Additional taxa not sampled in this study which could be targeted are †Annaichthys pontegiurinensis, †Eopholidophorus forojuliensis, †Jiangilichthys sp., †Knerichthys bronni, †Parapholidophorus caffi, †Pholidophoretes salvus, †Pholidophorus gervasuttii, †Pholidophorus latiusculus and †Zambellichthys bergamensis (34, 64). In addition, a few enigmatic Middle Triassic neopterygians may represent the earliest stem-group teleosts, including the middle Anisian (about 242 Ma) †Marcopoloichthys ani from the Upper Member of the Guangling Formation, Yunnan, South China (65). This taxon is characterized by a unique combination of features, including an apparently mobile premaxilla, a dorso-ventrally symmetrical caudal fin with a reduced number of hypurals, and an extreme reduction of the cover of ganoid scales (65, 66). A reevaluation of the phylogenetic position of this taxon would be desirable to address key questions about the earliest stages of teleost evolution, the origin of teleost morphological innovations and of their WGD.

III. Synchrotron beamline setups

III.1. DLS I13-2 Diamond-Manchester Imaging beamline setup

The X-ray beam was generated by a 2 m long in vacuum undulator (gap 5 mm), filtered with pyrolytic graphite (1340 μ m), aluminium (2100 μ m) and molybdenum (84 μ m). Additionally, a rhodium mirror was used to cut high energies. It resulted in a weighted mean detected energy of 21.39 keV. Radiographs were recorded using an indirect detector comprising a 34 μ m GGG scintillator, a 10x microscope lens, a 2x eyepiece and a PCO.edge 5.5 sCMOS camera (PCO AG, Kelheim, Germany). The effective recorded pixel size for this system was 0.357 μ m. Based on the pixel size and the detected energy, the sample-detector propagation distance was set to 17 mm. The acquisition consisted of 2500 projections recorded over a 180° rotation of the sample, with an exposure time of 0.5 s per projection. Additionally, 20 flatfield images (sample out

of the beam) were recorded before and after the scan as well as 20 dark images before the beginning of the scan (X-ray beam off to record the noise of the camera). Tomographic reconstruction was performed using the Savu processing pipeline (67) and the I13 analysis Python package. The workflow consisted of a dezinger correction (i.e. removing the scattering spikes), an optical distortion correction, ring removal on sinogram, single distance phase retrieval using the Paganin filter (68), tomographic reconstruction using TomoPy (69), and a conversion of the 32-bits HDF5 volume as a 16-bits tiff stack.

III.2. ESRF ID19 beamline setup

The ID19 beamline could be tuned for a much broader range of energies and the setup was regularly changed to adjust to the sample (see Supplementary Data). The X-ray beam was generated either by a wiggler or a single harmonic undulator, depending on the required energy. Filters were also adjusted depending on the energy. Beryllium X-ray lenses were used to focus the beam and increase flux on the sample (number of lenses adjusted with energy). Finally, we used a 1 mm graphite rotating disk as a decoherer and reduce potential ring artefact (70). Radiographs were recorded using an indirect detector comprising a scintillator (10 µm GGG or 25 µm LuAG), a 10x microscope lens and a PCO.edge camera (5.5 or 4.2 gold). The effective pixel size for this system was ca. 0.7 µm (as small variations were induced by changing the propagation distance). Based on the pixel size and the detected energy, the sampledetector propagation distance was adjusted, ranging from 20 to 220 mm. The acquisition consisted of 2999 projections recorded over a 360° rotation of the sample. Exposure time was ranging from 0.03 to 0.2 seconds, as transmitted signal was getting lower as the energy increased. Additionally, 21 flatfield images (sampled out of the beam) were recorded before and after the scan as well as 20 dark images before the beginning of the scan (X-ray beam off to record the noise of the camera). Tomographic reconstruction was performed using PyHST2 (71). The workflow consisted of a tomographic reconstruction using the single distance phase retrieval approach of PvHST2, followed by a conversion of the 32-bits EDF stack as a 16-bits tiff stack. Additional processing included ring correction (72).

III.3. Note on the resolution

Outside of the higher energies available at the ESRF ID19 beamline, the second main difference between both beamlines was the magnification setups, generating data with a recorded voxel size of $0.357~\mu m$ at the I13-2 beamline and near $0.7~\mu m$ at the ID19 beamline. It is important to note that these values indicate the magnification only and not the resolution, i.e. the smallest feature observable in the resulting tomographic data. Several parameters will influence the resolution, including notably the brilliance and spatial coherence of the incoming X-ray beam, the thickness and composition of the scintillator, the optical lenses and the digital camera. As the aim of this project is to measure osteocyte volumes, the variation in magnification and possibly in resolution could affect: 1) the smallest osteocyte detected; 2) the volume of osteocytes by adding or removing voxels at the periphery of the volume. The smallest osteocytes measured have a volume of ca. $70~\mu m$ 3; such a volume would be defined by ca. 1500~voxels and 195~voxels for voxel sizes of $0.357~\mu m$ and $0.7~\mu m$ respectively. The variation in voxel size hence could not influence the detection of the smallest osteocytes in our dataset.

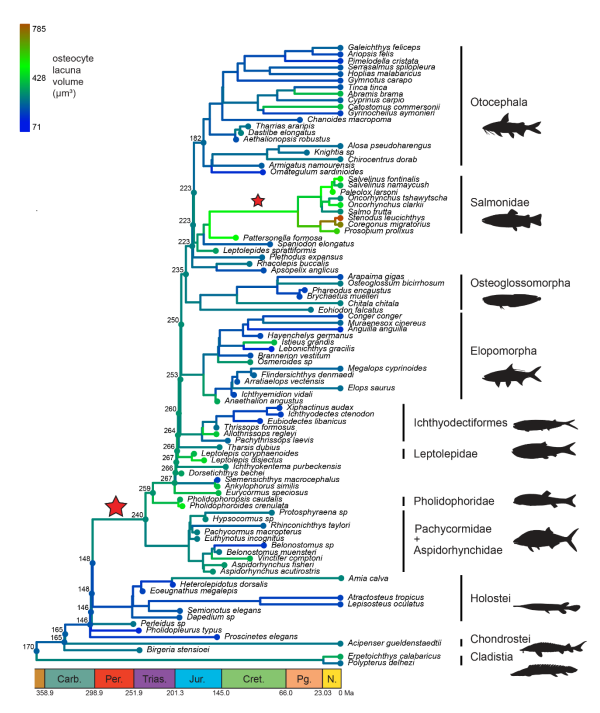


Figure S1. Evolution of osteocyte lacuna volume in fossil and modern teleosts revealed by synchrotron microtomography. Timescaled composite phylogeny of actinopterygians with mapped cell-size volumes for fossil and modern lineages. This version includes terminal taxon names.

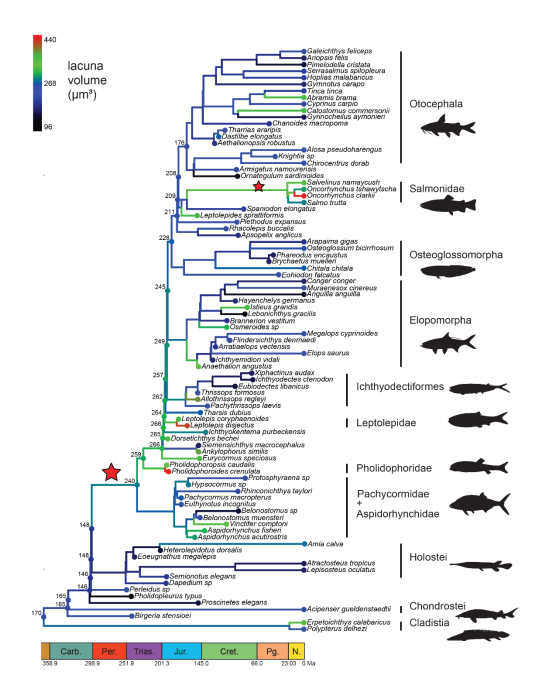


Figure S2. Evolution of osteocyte lacuna volume in fossil and modern teleosts revealed by synchrotron microtomography. Alternative version of Fig. 1 where *Salvelinus fontinalis*, *Coregonus migratorius*, *Stenodus leucichthys*, *Prosopium prolixus*, *Paleolox larsoni*, and *Pattersonella formosa* were omitted due to their very large osteocyte lacuna volumes. Therefore, the color scale distinguishes better the variation in volume of most of the taxon sample. This scale is different to that of Figs. 1-2.

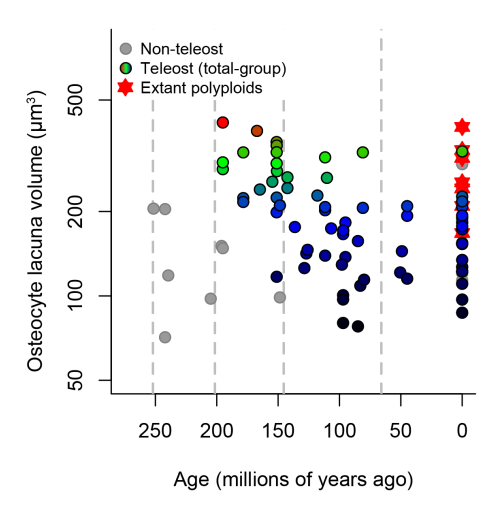


Figure S3. Evolution of osteocyte lacuna volume in fossil and modern teleosts revealed by synchrotron microtomography, plotted against time. Alternative version of Fig. 2, omitting the same taxa and using the same color scale as Fig. S2.

Dataset S1 (separate file). Full list of specimens used in the study, with their age ranges, the specifier taxa for their sister clades, and minimum divergence times from those sister clades based on fossil occurrences.

Dataset S2 (separate file). Full list of specimens for which PPC-SRµCT data were obtained, with the corresponding detailed experimental setups.

Dataset S3 (separate file). Tree file used in our analysis, obtained from stitching fossil taxa to the reference tree (1).

Dataset S4 (separate file). Full list of specimens, with the corresponding median osteocyte volumes using the minimum and maximum realistic grey value thresholds. We only used the "max" values for the analyses.

SI References

- 1. D. L. Rabosky, *et al.*, An inverse latitudinal gradient in speciation rate for marine fishes. *Nature* **559**, 392–395 (2018).
- 2. R. F. Stearley, G. R. Smith, Fishes of the Mio-Pliocene Western Snake River plain and vicinity I. Salmonid fishes from Mio-Pliocene lake sediments in the Western Snake River plain and the Great Basin. *Misc. Publ. Museum Zool. Univ. Michigan* **204**, 1–43 (2016).
- 3. R. F. Stearley, G. R. Smith, Phylogeny of the Pacific trouts and salmons (*Oncorhynchus*) and genera of the family Salmonidae. *Trans. Am. Fish. Soc.* **122**, 1–33 (1993).
- 4. M. A. Alexandrou, B. A. Swartz, N. J. Matzke, T. H. Oakley, Genome duplication and multiple evolutionary origins of complex migratory behavior in Salmonidae. *Mol. Phylogenet. Evol.* **69**, 514–523 (2013).
- 5. L. Taverne, A. Filleul, Osteology and relationships of the genus *Spaniodon* (Teleostei, Salmoniformes) from the Santonian (Upper Cretaceous) of Lebanon. *Palaeontology* **46**, 927–944 (2003).
- 6. A. Dornburg, M. Friedman, T. J. Near, Phylogenetic analysis of molecular and morphological data highlights uncertainty in the relationships of fossil and living species of Elopomorpha (Actinopterygii: Teleostei). *Mol. Phylogenet. Evol.* **89**, 205–218 (2015).
- 7. P. L. Forey, L. Yi, C. Patterson, C. E. Davies, Fossil fishes from the Cenomanian (Upper Cretaceous) of Namoura, Lebanon. *J. Syst. Palaeontol.* **1**, 227–330 (2003).
- 8. L. Taverne, Sur *Pattersonella formosa* (Traquair, 1911) et *Nybelinoides brevis* (Traquair, 1911), téléostéens salmoniformes argentinoides du Wealdien inférieur de Bernissart, Belgique, précédemment attribués au genre *Leptolepis* Agassiz, 1832. *Bull. l'Institut R. Sci. Nat. Belgique, Sci. la Terre* **54**, 1–27 (1982).
- 9. S. Olive, L. Taverne, L. Cavin, U. Deesri, Systematic revision of the Cretaceous actinopterygian fauna from Bernissart, Belgium. *Res. Knowl.* **3**, 43–46 (2017).
- T. J. Near, A. Dornburg, M. Friedman, Phylogenetic relationships and timing of diversification in gonorynchiform fishes inferred using nuclear gene DNA sequences (Teleostei: Ostariophysi). *Mol. Phylogenet. Evol.* 80, 297–307 (2014).
- 11. G. Arratia, Basal teleosts and teleostean phylogeny. *Palaeo Ichthyol.* **7**, 5–168 (1997).
- 12. G. Arratia, Remarkable teleostean fishes from the Late Jurassic of southern Germany and their phylogenetic relationships. *Mitteilungen aus dem Museum für Naturkd. zu Berlin, Geowissenschaftliche R.* **3**, 137–179 (2000).
- 13. G. Arratia, H. Tischlinger, The first record of Late Jurassic crossognathiform fishes from Europe and their phylogenetic importance for teleostean phylogeny. *Foss. Rec.* **13**, 317–341 (2010).
- 14. L. B. Bean, G. Arratia, Anatomical revision of the Australian teleosts *Cavenderichthys talbragarensis* and *Waldmanichthys koonwarri* impacting on previous phylogenetic interpretations of teleostean relationships. *Alcheringa An Australas. J. Palaeontol.* (2019) https://doi.org/10.1080/03115518.2019.1666921.
- 15. M. J. Benton, *et al.*, Constraints on the timescale of animal evolutionary history. *Palaeontol. Electron.* **18.1.1FC**, 1–106 (2015).
- 16. C. Patterson, *Chanoides*, a marine Eocene otophysan fish (Teleostei: Ostariophysi). *J. Vertebr. Paleontol.* **4**, 430–456 (1984).
- 17. D. Mayrinck, P. M. Brito, O. Otero, Review of the osteology of the fossil fish formerly attributed to the genus †*Chanoides* and implications for the definition of otophysan bony characters. *J. Syst. Palaeontol.* **13**, 397–420 (2015).
- 18. D. Mayrinck, "Relações Filogenéticas dos Otophysi (Actinopterygii, Teleostei), com ênfase em Characiformes, incluindo as formas fósseis," Université de Poitiers Universidade do Estado do Rio de Janeiro. (2010).
- L. Taverne, Ostéologie et position systématique d'Aethalionopsis robustus (Pisces, Teleostei) du Crétacé inférieur de Bernissart (Belgique) et considérations sur les affinités des Gonorhynchiformes. Bull. la Cl. des Sci. Académie R. Belgique 5, 958–982 (1981).
- 20. L. Grande, Recent and fossil clupeomorph fishes with materials for revision of the subgroups of clupeoids. *Bull. Am. Museum Nat. Hist.* **181**, 231–372 (1985).
- 21. J. Alvarado-Ortega, E. Ovalles-Damián, G. Arratia, "A review of the interrelationships of

- the order Ellimmichthyiformes (Teleostei: Clupeomorpha)" in *Mesozoic Fishes 4 Homology and Phylogeny*, G. Arratia, H.-P. Schultze, M. V. H. Wilson, Eds. (Verlag Dr. Friedriech Pfeil, 2008), pp. 257–278.
- 22. P. L. Forey, A primitive clupeomorph fish from the middle Cenomanian of Hakel, Lebanon. *Can. J. Earth Sci.* **10**, 1302–1318 (1973).
- L. Taverne, Tselfatia formosa, téléostéen marin du Crétacé (Pisces, Actinopterygii), et la position systématique des Tselfatiiformes ou Bananogmiiformes. Geodiversitas 22, 5–22 (2000).
- 24. M. Friedman, H. T. Beckett, R. A. Close, Z. Johanson, The English Chalk and London Clay: two remarkable British bony fish *Lagerstätten. Geol. Soc. London, Spec. Publ.* **430**, 165–200 (2016).
- 25. E. Sferco, A. López-Arbarello, A. M. Báez, Phylogenetic relationships of †*Luisiella feruglioi* (Bordas) and the recognition of a new clade of freshwater teleosts from the Jurassic of Gondwana. *BMC Evol. Biol.* **15**, 268 (2015).
- 26. A. López-Arbarello, E. Sferco, Neopterygian phylogeny: the merger assay. *R. Soc. Open Sci.* **5**, 172337 (2018).
- 27. P. L. Forey, The osteology of *Notelops* Woodward, *Rhacolepis* Agassiz and *Pachyrhizodus* Dixon (Pisces: Teleostei). *Bull. Br. Museum (Natural Hist. Geol.* **28**, 123–204 (1977).
- 28. A. M. Murray, D. K. Zelenitsky, D. B. Brinkman, A. G. Neuman, Two new Palaeocene osteoglossomorphs from Canada, with a reassessment of the relationships of the genus † *Joffrichthys*, and analysis of diversity from articulated versus microfossil material. *Zool. J. Linn. Soc.* **183**, 907–944 (2018).
- 29. J. Zhang, Phylogeny of Osteoglossomorpha. Vertebr. Palasiat. 44, 43–59 (2006).
- 30. L. Grande, *Eohiodon falcatus*, a new species of hiodontid (Pisces) from the late Early Eocene Green River Formation of Wyoming. *J. Paleontol.* **53**, 103–111 (1979).
- 31. E. J. Hilton, Comparative osteology and phylogenetic systematics of fossil and living bonytongue fishes (Actinopterygii, Teleostei, Osteoglossomorpha). *Zool. J. Linn. Soc.* **137**, 1–100 (2003).
- 32. A. Belouze, M. Gayet, C. Atallah, Les premiers Anguilliformes : II. Paraphylie du genre *Urenchelys* Woodward, 1900 et relations phylogénétiques. *Geobios* **36**, 351–378 (2003).
- 33. L. Taverne, Ostéologie et position systématique d'*Arratiaelops vectensis* gen. nov., téléostéen élopiforme du Wealdien (Crétacé inférieur) d'Angleterre et de Belgique. *Bull. l'Institut R. Sci. Nat. Belgique, Sci. la Terre* **39**, 77–96 (1999).
- 34. G. Arratia, Morphology, taxonomy, and phylogeny of Triassic pholidophorid fishes (Actinoptervaii, Teleostei). *J. Vertebr. Paleontol.* **33**, 1–138 (2013).
- 35. C. Patterson, D. E. Rosen, Review of ichthyodectiform and other Mesozoic teleost fishes and the theory and practice of classifying fossils. *Bull. Am. Museum Nat. Hist.* **158**, 81–172 (1977).
- G. Arratia, New Triassic teleosts (Actinopterygii, Teleosteomorpha) from northern Italy and their phylogenetic relationships among the most basal teleosts. *J. Vertebr. Paleontol.* 37, e1312690 (2017).
- 37. L. Cavin, P. L. Forey, S. Giersch, Osteology of *Eubiodectes libanicus* (Pictet & Humbert, 1866) and some other ichthyodectiformes (Teleostei): phylogenetic implications. *J. Syst. Palaeontol.* **11**, 115–177 (2013).
- 38. B. Schaeffer, C. Patterson, Jurassic fishes from the western United States, with comments on Jurassic fish distribution. *Am. Museum Novit.* **2796**, 1–86 (1984).
- 39. G. Arratia, New remarkable Late Jurassic teleosts from southern Germany: Ascaloboidae n. fam., its content, morphology, and phylogenetic relationships. *Foss. Rec.* **19**, 31–59 (2016).
- 40. M. Friedman, Parallel evolutionary trajectories underlie the origin of giant suspension-feeding whales and bony fishes. *Proc. R. Soc. B Biol. Sci.* **279**, 944–951 (2012).
- 41. M. Friedman, *et al.*, 100-million-year dynasty of giant planktivorous bony fishes in the Mesozoic seas. *Science* (80-.). **327**, 990–993 (2010).
- 42. A. Tintori, The actinopterygian fish *Prohalecites* from the Triassic of Northern Italy. *Palaeontology* **33**, 155–174 (1990).

- 43. L. Wretman, H. Blom, B. P. Kear, Resolution of the Early Jurassic actinopterygian fish *Pachycormus* and a dispersal hypothesis for Pachycormiformes. *J. Vertebr. Paleontol.*, e1206022 (2016).
- 44. P. M. Brito, Révision des Aspidorhynchidae (Pisces, Actinopterygii) du Mésozoïque : ostéologie, relations phylogénétiques, données environnementales et biogéographiques. *Geodiversitas* **19**, 681–772 (1997).
- 45. P. E. Olsen, A. R. McCune, Morphology of the *Semionotus elegans* species group from the Early Jurassic part of the Newark Supergroup of Eastern North America with comments on the family Semionotidae (Neopterygii). *J. Vertebr. Paleontol.* **11**, 269–292 (1991).
- 46. L. Grande, W. E. Bemis, A comprehensive phylogenetic study of amiid fishes (Amiidae) based on comparative skeletal anatomy. An empirical search for interconnected patterns of natural history. *J. Vertebr. Paleontol.* **18**, 1–696 (1998).
- 47. Z. Sun, *et al.*, A new non-parasemionotiform order of the Halecomorphi (Neopterygii, Actinopterygii) from the Middle Triassic of Tethys. *J. Syst. Palaeontol.* (2016) https://doi.org/10.1080/14772019.2016.1181679.
- 48. A. E. Latimer, S. Giles, A giant dapediid from the Late Triassic of Switzerland and insights into neopterygian phylogeny. *R. Soc. Open Sci.* **5**, 180497 (2018).
- 49. D. Thies, J. Waschkewitz, Redescription of *Dapedium pholidotum* (Agassiz, 1832) (Actinopterygii, Neopterygii) from the Lower Jurassic Posidonia Shale, with comments on the phylogenetic position of *Dapedium* Leach, 1822. *J. Syst. Palaeontol.* **4**, 339–364 (2016).
- 50. M. Friedman, The early evolution of ray-finned fishes. *Palaeontology* **58**, 213–228 (2015).
- 51. G.-H. Xu, K.-Q. Gao, M. I. Coates, Taxonomic revision of *Plesiofuro mingshuica* from the Lower Triassic of northern Gansu, China, and the relationships of early neopterygian clades. *J. Vertebr. Paleontol.*, e1001515 (2015).
- 52. G.-H. Xu, X.-Y. Ma, A Middle Triassic stem-neopterygian fish from China sheds new light on the peltopleuriform phylogeny and internal fertilization. *Sci. Bull.* **61**, 1766–1774 (2016).
- 53. G.-H. Xu, L.-J. Zhao, K.-Q. Gao, F.-X. Wu, A new stem-neopterygian fish from the Middle Triassic of China shows the earliest over-water gliding strategy of the vertebrates. *Proc. R. Soc. B Biol. Sci.* (2012) https://doi.org/10.1098/rspb.2012.2261.
- 54. F. J. Poyato-Ariza, S. Wenz, A new insight into pycnodontiform fishes. *Geodiversitas* **24**, 139–248 (2002).
- 55. S. Giles, G. H. Xu, T. J. Near, M. Friedman, Early members of "living fossil" lineage imply later origin of modern ray-finned fishes. *Nature* (2017) https://doi.org/10.1038/nature23654.
- 56. C. Romano, W. Brinkmann, Reappraisal of the lower actinopterygian *Birgeria stensioei* Aldinger, 1931 (Osteichthyes; Birgeriidae) from the Middle Triassic of Monte San Giorgio (Switzerland) and Besano (Italy). *Neues Jahrb. für Geol. und Paläontologie, Abhandlungen* **252**, 17–31 (2009).
- 57. T. Argyriou, *et al.*, Internal cranial anatomy of Early Triassic species of † *Saurichthys* (Actinopterygii: †Saurichthyiformes): implications for the phylogenetic placement of †saurichthyiforms. *BMC Evol. Biol.* **18**, 161 (2018).
- 58. J. Lu, S. Giles, M. Friedman, J. L. den Blaauwen, M. Zhu, The oldest actinopterygian highlights the cryptic early history of the hyperdiverse ray-finned fishes. *Curr. Biol.* **26**, 1602–1608 (2016).
- 59. S. Giles, *et al.*, Endoskeletal structure in *Cheirolepis* (Osteichthyes, Actinopterygii), an early ray-finned fish. *Palaeontology* (2015) https://doi.org/10.1111/pala.12182.
- 60. A. Tintori, *et al.*, Oldest stem Teleostei from the Late Ladinian (Middle Triassic) of Southern China. *Riv. Ital. Di Paleontol. E Stratigr.* **121**, 285–296 (2015).
- 61. A. Tintori, Fish taphonomy and Triassic anoxic basins from the Alps: a case history. *Riv. Ital. di Paleontol. e Stratigr.* **97**, 393–407 (1992).
- D. Davesne, *et al.*, The phylogenetic origin and evolution of acellular bone in teleost fishes: insights into osteocyte function in bone metabolism. *Biol. Rev.* **94**, 1338–1363 (2019).
- 63. D. Davesne, F. J. Meunier, M. Friedman, R. B. J. Benson, O. Otero, Histology of the endothermic opah (*Lampris* sp.) suggests a new structure–function relationship in teleost

- fish bone. Biol. Lett. 14, 20180270 (2018).
- 64. R. Zambelli, Note sui Pholidophoriformes: VII contributo *Eopholidophorus forojuliensis* n.g., n.sp. *Gortania* **11**, 63–76 (1990).
- 65. A. Tintori, *et al.*, New specialized basal neopterygians (Actinopterygii) from Triassic of the Tethys Realm. *Geol. Insubrica* **10**, 13–20 (2008).
- 66. A. Tintori, C. Lombardo, D.-Y. Jiang, Z.-Y. Sun, "*Pholidophorus*" faccii Gortani 1907: nuovi dati tassonomici. *Gortania* **32**, 45–52 (2010).
- 67. N. Wadeson, M. Basham, Savu: a Python-based, MPI framework for simultaneous processing of multiple, N-dimensional, large tomography datasets. *arXiv*, arXiv:1610.08015 (2016).
- 68. D. Paganin, S. C. Mayo, T. E. Gureyev, P. R. Miller, S. W. Wilkins, Simultaneous phase and amplitude extraction from a single defocused image of a homogeneous object. *J. Microsc.* **206**, 33–40 (2002).
- 69. D. Gürsoy, F. De Carlo, X. Xiao, C. Jacobsen, TomoPy: a framework for the analysis of synchrotron tomographic data. *J. Synchrotron Radiat.* **21**, 1188–1193 (2014).
- S. Sanchez, P. E. Ahlberg, K. M. Trinajstic, A. Mirone, P. Tafforeau, Three-dimensional synchrotron virtual paleohistology: a new insight into the world of fossil bone microstructures. *Microsc. Microanal.* 18, 1095–1105 (2012).
- 71. A. Mirone, E. Brun, E. Gouillart, P. Tafforeau, J. Kieffer, The PyHST2 hybrid distributed code for high speed tomographic reconstruction with iterative reconstruction and a priori knowledge capabilities. *Nucl. Instruments Methods Phys. Res. Sect. B* **324**, 41–48 (2014).
- 72. A. Lyckegaard, G. Johnson, P. Tafforeau, Correction of ring artifacts in X-ray tomographic images. *Int. J. Tomogr. Stat.* **18**, 1–9 (2011).