



Dataset of antiarch placoderms (the most basal jawed vertebrates) throughout Middle Paleozoic



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Abstract. Antiarch placoderms, the most basal jawed vertebrates, have the potential to enlighten the origin of the last common ancestor of jawed vertebrates. Quantitative study based on credible data is more convincing than qualitative study. To reveal the antiarch distribution in space and time, we created a comprehensive structured dataset of antiarchs comprising 64 genera and 6025 records. This dataset, which includes associated chronological and geographic information, has been digitalized from academic publications manually into the DeepBone database as a dataset. We implemented the paleogeographic map marker to visualize the biogeography of antiarchs. The comprehensive data of Antiarcha allow us to generate its biodiversity and variation rate changes throughout its duration. Structured data of antiarchs has tremendous research potential, including testing hypotheses in the fields of the biodiversity changes, distribution, differentiation, population and community composition. Also, it will be easily accessible by the other tools to generate new understanding on the evolution of early vertebrates. The data file described in this paper is available on <https://doi.org/10.5281/zenodo.5639529> (Pan and Zhu, 2021).



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25 1 Introduction

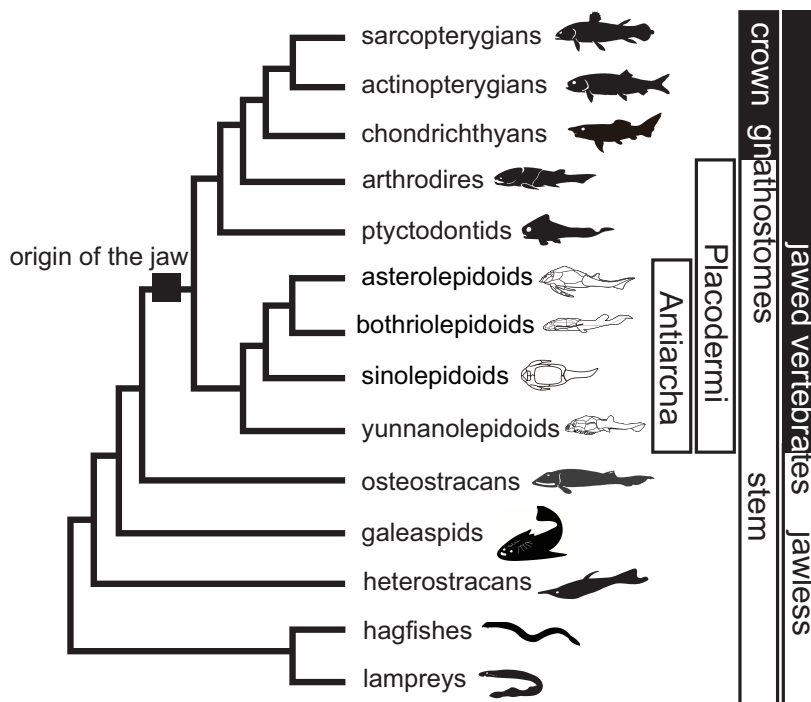
Placodermi is an extinct grade of jawed vertebrates that first occurred in the Silurian and then dominated the Devonian (Carr, 1995; Young, 2010). Recent prevailing phylogenetic hypotheses placed Placodermi as jawed stem-Gnathostomata that is sister to crown-Gnathostomata or modern jawed vertebrates (Brazeau, 2009; Davis et al., 2012; Dupret et al., 2014; Giles et al., 2015; King, 2021; Long et al., 2015;





30 Qiao et al., 2016; Trinajstić et al., 2015; Zhu et al., 2013, 2016). In this scenario, Antiarcha has usually been placed at the most basal position in the Placodermi (Fig. 1), representing the most basal jawed vertebrates. The geospatial-temporal distribution of Antiarcha will thus help us understand the origin and early evolution of jawed vertebrates. Moreover, Antiarcha was a widely distributed, diverse, and successful group within Placodermi from the late Silurian to the end of Devonian (Denison, 1978; Janvier, 35 1996; Zhao et al., 2016; Zhu, 1996) to allow the inference of biogeographic assumptions (Ritchie et al., 1992; Young, 1984b). As a successful vertebrate group during the Devonian, Antiarcha has much contributed to the Devonian stratigraphic correlation. For instance, the biozonation of East Baltic and southern East Antarctica Devonian succession is partly based upon the antiarchs *Bothriolepis*, *Asterolepis*, and *Pambulaspis* (Young, 1974, 1988). Lukševičs (1996) identified 14 bothriolepid species 40 (12 *Bothriolepis* and 2 *Grossilepis*) in the Frasnian-Famennian formations of the East European Platform, proposed nine antiarch assemblages, and set up the most detailed zonation of the Main Devonian Field. However, Blicek et al. (2000) reminded that *Asterolepis*, *Bothriolepis*, and *Remigolepis*, which have different stratigraphic distributions on different continents, must be used with caution for long-distance correlation.





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Figure 1 Phylogenetic relationships of major early vertebrate groups from Qiao et al. (2016) and Pan et al. (2018). Line drawings indicate groups of the Antiarcha. Silhouettes indicate the other vertebrate groups. The black cube on the cladogram points to the origin of the jaw.

Explaining the spatial and temporal distribution of early vertebrates is the prerequisite to understand their biogeographic exchange. Although Zigaite and Blicek (2013) advocated a quantitative analysis to define biogeographic patterns of early vertebrates, there is still lacking efficient quantitative analysis to understand the dispersal of early vertebrates. This occurs mostly because no comprehensive data collection of early vertebrates was accomplished.

The main objective of this study is to present an unprecedented structured dataset of Antiarcha that is potentially facilitate the research of the geospatial distribution patterns for antiarchs and the quantitative study on early vertebrates. This dataset is the first step to accomplish the global coverage of the vertebrate fossil dataset to analyze the Middle Paleozoic biogeography and paleogeography. Revealing the distribution of antiarchs in the paleomap background with reference to the global paleogeographic reconstructions of Scotese (2002), our preliminary results can be used to test the hypothesis of paleogeographic reconstructions.



2. Method

2.1 Overview

To facilitate the quantitative study and simulation analysis on early vertebrates, we are looking forward to a comprehensive database. Sallan et al. (2018) pointed out that a lack of early vertebrate fossil data has limited quantitative approaches, and hindered the resolution of issues in vertebrate evolution. With the implementation of a project entitled “Big Earth Data Science Engineering (CASEarth)” in the Strategic Priority Research Program since 2018 (Guo, 2017), we build the DeepBone database. Via three years of efforts conscientiously, the data of DeepBone database already has reached a certain scale. With continuously refining data, the Antiarcha dataset in the DeepBone database is the most comprehensive dataset endorsed by Chinese researchers in the Institute of Vertebrate Paleontology and Paleoanthropology, Chinese Academy of Sciences. The Antiarcha dataset differs from the other datasets in its basic unit, which is the specimen ID coupled with the occurrence and other detailed data. All the specimens are referenced to taxa and literature. Because the Antiarcha dataset was designed as a vertebrate paleontological dataset and its input format was designed as specimen-based, data entry assistants input the metadata according to the published specimen or virtual specimen.

Since there is no satisfactory approach that could automatically extract the paleontological data from literature, we have recruited several data entry assistants including relevant researchers, master’s and PhD students to collect and curate the data. In order to guarantee the quality of the data, we have designed a four-step data processing procedure (Fig. 2):

1. Data source by an experienced expert who got his PhD degree on early vertebrates.
2. The data entry assistants read the related references, extract the antiarch placoderm data and fill them into the record file under the supervision of experts.
3. Experts review and clean the data according to the references to implement quality control.
4. Further data examinations by senior experts to guarantee the data quality.
5. Data visualization and open access. This step is to provide better user interface to help disseminate the dataset.

Next, we provide more details of the data processing and visualization.



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Figure 2 Workflow of the data processing. Even the web data entry system could be accessed by any registered users, we have recruited several data entry assistants to enrich the data following the standardized workflow.

1, collecting and sifting literature by experienced experts. 2, data entry assistants digitalize paleontological descriptions from the page into the DeepBone database. 3, experts who are accountable for data review and clean the data according to the references to implement quality control. 4, associated senior researchers review the data. 5, data managers release the data to the public

2.2 Data Source

We invite researchers who have a good reputation in the research scope to sift published literature from the bibliography of the paleontology journals and books. The main journals includes *Alcheringa*, *Acta Geologica Polonica*, *Bulletin of the Geological Society of China*, *Estonian Journal of Earth Sciences*, *Journal of Vertebrate Paleontology*, *Journal of Paleontology*, *Palaeontologia Electronica*, *Palaeontology*, *Palaeoworld*, and *Vertebrata Palasiatica*. Totally, we have collected 126 publications, which span from the year 1939 to the year 2021. The satisfactory literature should include an accurate description or revision on specimen and taxon. We accepted the latest peer-reviewed literature to deal with the inconsistent descriptions.



2.3 Data Processing and Quality Control

We have made a tailored web page that provides a better user interface for them to fill in the rows of paleontological data. After that, the data would be reviewed by the other related experts so that a researcher could quickly access them to perform quantitative analysis reliably (Fig. 2). This workflow was learned from the Geobiodiversity Database (GBDB) (Xu et al., 2021). Almost all antiarch literature was published in English, Russian, French, German, and Chinese. Data entry assistance structure could handle the literature in Chinese and English well. Literature in French, Russian, and German was dealt with by paleontology postgraduates who know well these languages.

2.4 Data Visualization

The analysis is conducted on the raster tile form paleomaps (Scotese, 1998, 2016). We first convert the excavation locations from current GPS to paleo-GPS using TrackPoint software (Ke et al., 2016; Scotese, 2002). However, the construction of paleontological maps and networks for visual analysis is based on paleomaps. Thus, we convert these latitude and longitude coordinates into pixel coordinates of raster tile maps using the Web Mercator algorithm. The Web Mercator algorithm is a variant of the Mercator projection and the de facto standard for Web mapping applications (Battersby et al., 2014). Specifically, the projection can be modelled as:

$$\begin{cases} x = a * \theta \\ y = a * \ln \tan\left(\frac{\pi}{4} + \frac{\varphi}{2}\right) \end{cases}$$

Where a is the long axis of the earth; θ is the longitude in radians, the value range is $[-\pi, \pi]$, the east longitude is positive and the west longitude is negative; φ is the latitude, the value range is $[-\pi/2, \pi/2]$, north latitude is positive and south latitude is negative.

As for the paleomap, we also need to consider the map scaling and canvas size, so the final formula used is as follows:

$$\begin{cases} x_i = l * \theta * 2^{n-1} \\ y_i = l * \ln \tan\left(\frac{\pi}{4} + \frac{\varphi}{2}\right) * 2^{n-1} + m * (i - 1) \end{cases}$$

Where l is the length of the plane map; n is the size of the zoom scale, $n \in \mathbb{N}^+$; in the analysis of geological periods, i is taken as 1; in the analysis of cross-geological periods, i represents the i -th geological period, (x_i, y_i) are the pixel coordinates of the node in the i -th geological period, and m is the fixed pixel distance between the map centers of adjacent geological periods.



2.5 Biodiversity Visualization

135 We calculate the genus and species biodiversity and plot it as a **bar map**. To better view the results, we utilize the kernel density estimation (KDE) algorithm to smooth the curve and estimate the biodiversity. In statistics, KDE is a fundamental data smoothing algorithm that can help inferences about the population based on finite data samples.

140 The counts of genus or species of **various time slots** (visualized as the bins in the charts) are defined as (x_1, x_2, \dots, x_n) . In statistics, they can be defined as samples drawn from a univariate distribution with an unknown density f . To estimate the shape of this distribution f , its kernel density estimator is defined as

$$\hat{f}_h(x) = \frac{1}{n} \sum_{i=1}^n K_h(x - x_i) = \frac{1}{nh} \sum_{i=1}^n K\left(\frac{x - x_i}{h}\right)$$

145 where K is the kernel function for estimation, we use the Epanechnikov kernel for better smoothing; $h > 0$ is a smoothing parameter named as bandwidth. For better assessment of the biodiversity changes, we adopt the first derivative of the Continuous Probability Distributions function to estimate the rate of variation.

3. Results

3.1 Data Overview

150 This dataset, which was extracted from 126 published papers or books manually, consists of 64 genera and 6025 records, covering all antiarch lineages. The 6025 records include 5867 fossil specimens that had been systematically described and documented, and 158 virtual specimens, which were introduced to describe the taxon information when no specimen was assigned for the referred records. The quantities distribution of specimens are given in the supplement. Each record has at least one reference within our dataset, and the specimens lacking precise age are excluded. We transferred the unstructured data from literature to structured data for further research as detailed as possible. Table 1 shows the framework of our dataset. Among all the referred specimens, 6.51% belong to Yunnanolepidoidei, 2.86% belong to Sinolepidoidei, 78.92% belong to 'Bothriolepidoidei', and 11.71% belong to Asterolepidoidei. All the fossil sites of the constituent groups are plotted in Figure 3 and the phylogeny of Antiarcha follows that in Pan et al. (2018).



Specimen record		Reference
Formation	Specimen ID	Literature type
Member	Genus	Title
Fossil Locality	Species	Keywords
Discovery country	Custodian Institute	Volume
Latitude	System / Period	Issue
Longitude	Series / Epoch	Pages
Paleo-latitude	Stage / Age	Authors
Paleo-longitude	Group	Year
		Journal
		Doi

Table 1 The structure of Antiarcha dataset.

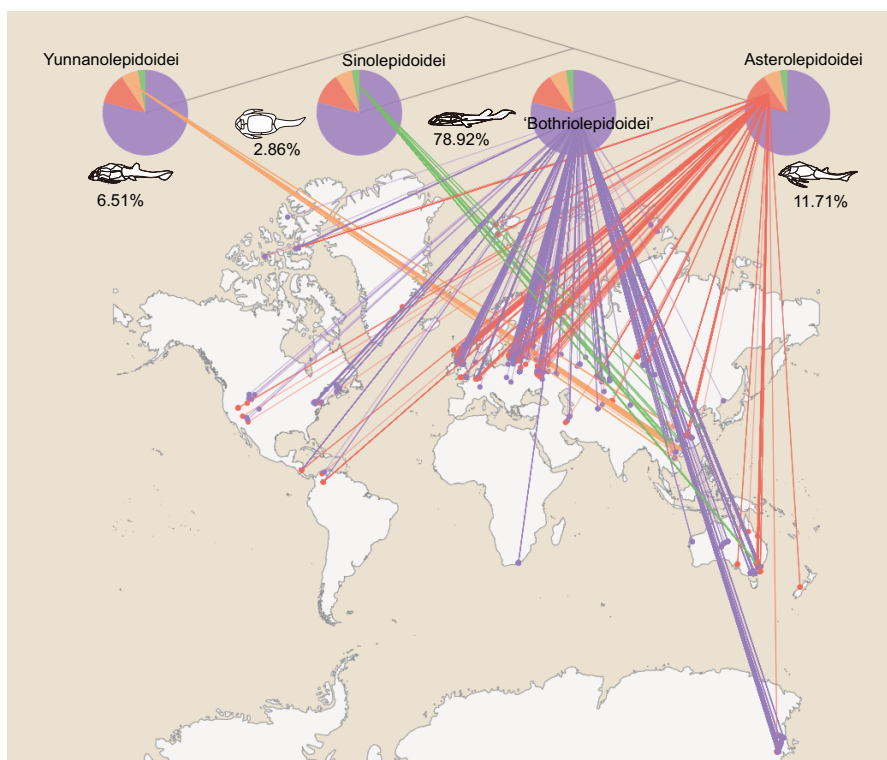


Figure 3 Graphic abstract of Antiarcha data. The tree topology is simplified from the phylogenetic result in Pan et al. (2018). Numbers under the pie chart represent the relative amounts of each group. Terminal groups are linked with their geographic distributions.

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No antiarch record was in GBDB. Compared to the 138 records of Antiarcha in the Paleobiology Database (PBDB, 2021-08-12), this is the most comprehensive dataset of Antiarcha up to now (Table 2). Moreover, only taxon rank, reference, and occurrence location are available in PBDB. More structured information of Antiarcha is available in DeepBone. Every record of DeepBone was assigned to an age with the latest reference. It is open accessible through the website of the DeepBone database or <https://doi.org/10.5281/zenodo.5639529>.



	DeepBone Database	Paleobiology Database(PBDB)
Type	specimen-based	fossil-occurrence-based
No. of references	355	19
No. of genera	64	26
No. of species	187	98
No. of Specimens/ occurrences	6025	138
Found in	2018	1998
Websit (last access: 22 October 2021)	www.deepbone.org	https://paleobiodb.org/#/

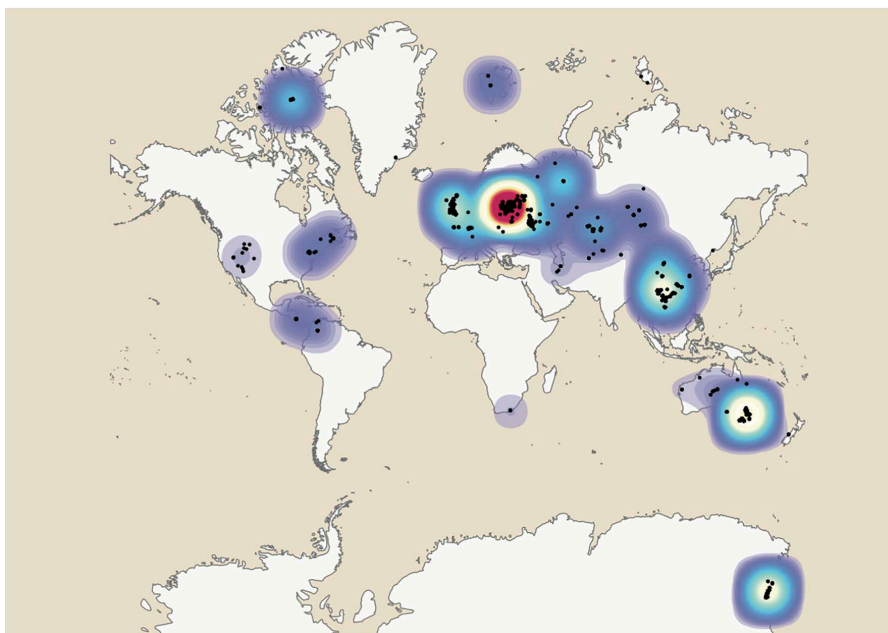
Table 2 The comparison of data of Antiarcha in two paleontological databases.



175 3.2 The Geospatial Distribution of the Antiarcha Dataset

As for the fossil site distribution, Yunnanolepidoidei is endemic in the South China block (comprising southern China and northern Vietnam). Sinolepidoidei is limited in South China and Australia (East Gondwana). In contrast, 'Bothriolepidoidei' and Asterolepidoidei are cosmopolitan, especially *Bothriolepis*. The faunistic elements in the communities are used herein at the genus level for their distributions because many *Bothriolepis* species were described based on isolated plates lacking diagnostic characters (Blicek and Janvier, 1993; Downs, 2011). The heat map of fossil records (Fig. 4) shows that Europe, Australia, and China account for the most records in the world, partly due to their long research history.





185 **Figure 4 Heat map of Antiarcha records.**



As Young (1990a) mentioned that biogeographic data must be interpreted in the context of paleogeographic hypotheses, we plot our data on paleomap (Fig. 5) to generate an outline of their past.

The analysis of biogeographic trends in Paleozoic vertebrates is highly dependent on fossil data and paleocontinent reconstruction. Due to the easy access of the paleo-geographic coordinates calculator



190 (PointTrack version 7.0) (Ke et al., 2016; Scotese, 2002), we decided to use Scotese's paleocontinent reconstruction to perform the plot map, although many paleogeographic reconstructions were proposed



(Heckel and Witzke, 1979; Li and Powell, 2001; Scotese et al., 1985). The continental reconstructions of Scotese place Baltica, China and Australia in the tropic and subtropic near the equator from Llandovery to Famennian. The timescale follows the International Commission on Stratigraphy



195 International Chronostratigraphic Chart version 2021/07.

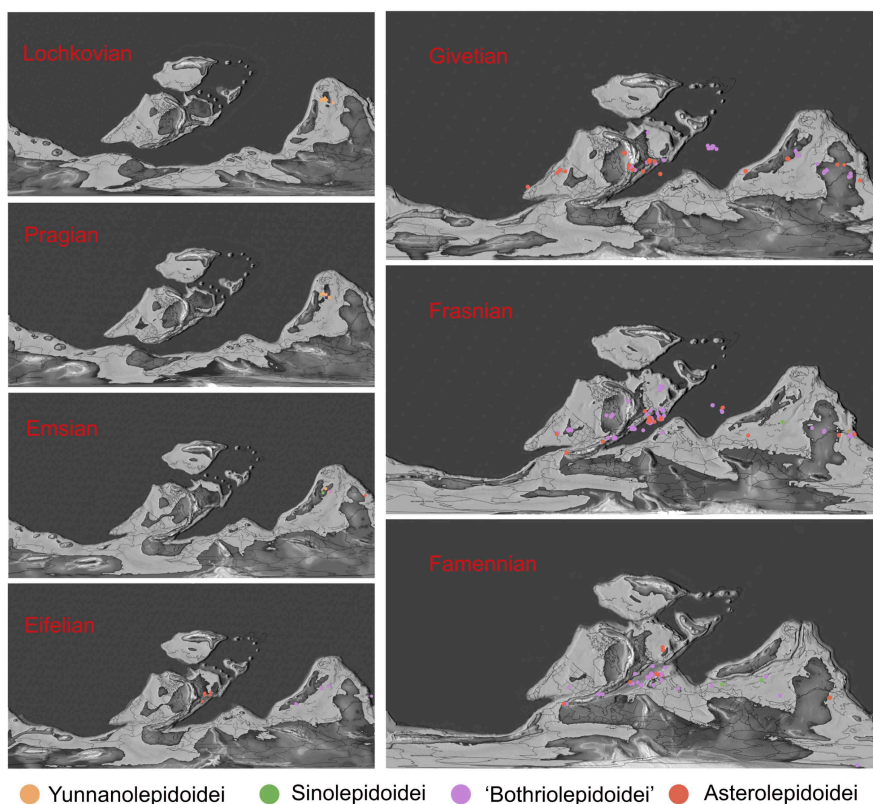
3.3 The Paleogeographic Distribution of the Antiarcha Dataset

We plotted these fossil sites on the paleogeographic map (Fig. 5) except the Silurian *Shimenolepis*, which is the earliest record of Yunnanolepidoidei and the only documented antiarch specimen before the Devonian (Wang, 1991; Zhao et al., 2016). Most of the fossil sites were positioned around the equator.

200 In the present scenario, the suborder Yunnanolepidoidei apparently originated as early as Silurian in the



205 South China block, forming a highly endemic fauna. All fossil sites of Yunnanolepidoidei lied in southern
China and northern Vietnam (Wang et al., 2010). From Ludlow (Silurian) to Early Devonian,
Yunnanolepidoidei formed dominant antiarchs. Sinolepidoidei and ‘Bothriolepidoidei’ first appeared in
Pragian in South China, and Asterolepidoidei first evolved in Emsian in Australia or East Gondwana.
210 During Middle Devonian, along with lessened isolation of South China, Yunnanolepidoidei became
extinct. Euantiarcha (‘Bothriolepidoidei’ + Asterolepidoidei) dominated the antiarchs in Middle and Late
Devonian, and only a few members of Sinolepidoidei coexisted with them in China and Australia. In
Eifelian, Asterolepidoidei suddenly bloomed in Baltica without any clue from the older horizons. The
distribution and diversity of Antiarcha reached a peak in Givetian. ‘Bothriolepidoidei’ and
Asterolepidoidei represent the main groups of Antiarcha in Givetian, comprising 5 bothriolepidoid
genera with 42 fossil locations and 9 asterolepidoid genera with 49 fossil locations. The records of
antiarchs decreased in the Pan-Cathaysian and East Gondwana in Late Devonian, contrary to those
increased in Baltica. Antiarchs in Baltica went through an independent evolution in Late Devonian.





215 **Figure 5** The distributions of *Antiarcha* during Devonian. The paleo-coordinates are calculated by TrackPoint.
Colors denoting respective groups follow Figure 3.



4 Discussion

4.1 First Appearance Record

220 First appearance record of a taxon or a lineage is important in paleontology and evolutionary biology as
it renders a hard minimum constrain on molecular clock calibration for a taxon (Benton and Donoghue,
2007; Benton et al., 2009; Donoghue and Benton, 2007). Based on our dataset, the oldest record of
yunnanolepidoids or antiarchs is *Shimenoalepis graniferus* from the Xiaoxi Formation at Shanmen
Reservoir, Lixian County, Hunan, China. *Shimenoalepis* was first described as the oldest known
placoderm, dated as Telychian of Llandovery (Janvier, 1996; Wang, 1991). However, after a detailed
225 stratigraphic work, Zhao et al. (2016) suggested that the age of *Shimenoalepis* is late Ludlow rather than
late Llandovery. Janvier and Tông-Dzuy (1998) also documented an indeterminate yunnanolepidoid
(*Antiarcha* gen. sp. indet.) from the Do Son Formation of northern Vietnam, which could be another
earliest antiarch potentially.

230 The oldest sinolepid is *Liujiangolepis suni*, from the Nakaoling Formation (Pragian), Xiangzhou,
Guangxi, China (Wang, 1987). The oldest bothriolepidid is *Houershanaspis zhangi*, documented from
the Danlin Formation (Pragian) in Mt. Houershan, Guizhou, southwestern China, on the basis of a
bothriolepid-like anterior median dorsal plate (Lu et al., 2017). The earliest asterolepidoid records are
represented by *Wurungulepis* and some disarticulated specimens, which had been documented from the
Broken River Formation, Broken River, Australia. The age of the Broken River Formation was first
235 referred to Eifelian, then reassigned to Emsian (*serotinus* Zone) (Burrow, 1996; De Pomeroy, 1996;
Young, 1984a, 1990b).

4.2 The Origin and Differentiation of Euanteriarcha

240 *Houershanaspis*, *Wudinolepis*, and *Wufengshania* from Pragian and Emsian represent the earliest
members of ‘Bothriolepidoidei’ (Chang, 1965; Lu et al., 2017; Pan et al., 2018), highlighting the origin
and early diversification of this group in South China. In Eifelian, the distribution range of
‘Bothriolepidoidei’ extended from South China to Iran that was positioned at the northern margin of East
Gondwana with the discovery of *Hyrceanaspis* (Janvier and Pan, 1982). ‘Bothriolepidoidei’ became
cosmopolitan during Givetian and Late Devonian. The earliest asterolepidoid records are from Australia

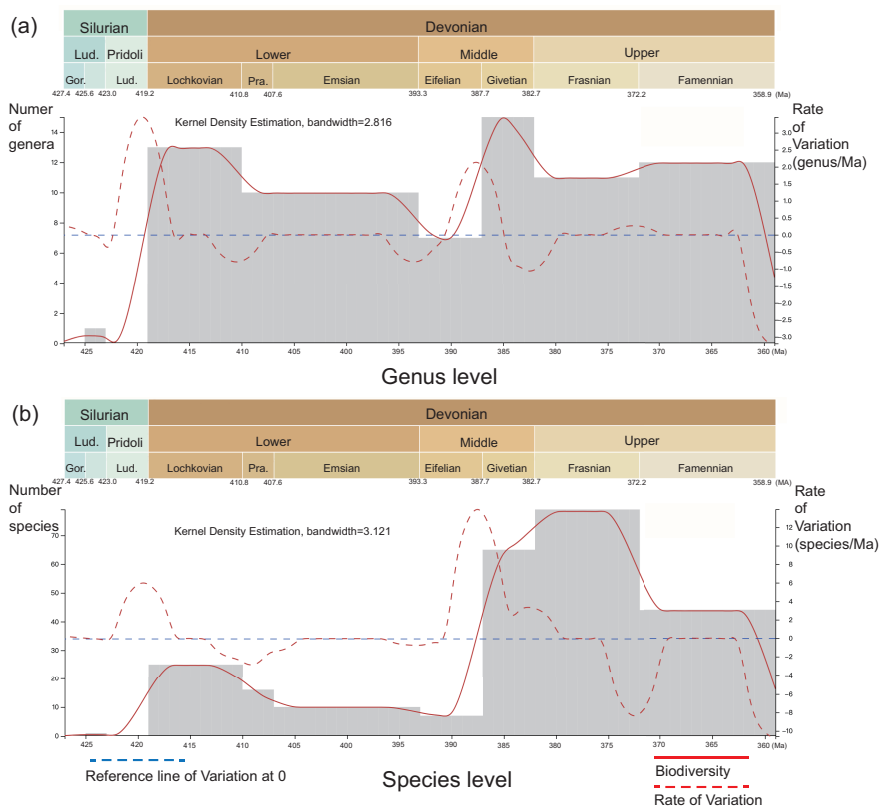


or East Gondwana in Emsian (Burrow, 1996; Young, 1984a, 1990b). In Eifelian, asterolepidoids
245 occurred in Baltica, in addition to East Gondwana. Compared to the abundant Early and Middle Devonian
fish faunas of in South China, asterolepidoids were late arrivals to South China (Young, 1984a) since no
asterolepidoid was known from this province before Givetian. It could be inferred that Asterolepidoidei
originated and differentiated in East Gondwana.

4.3 The Diversity Changes of Antiarcha

250 The comprehensive data of Antiarcha allow us to generate its diversity changes through its life span. We
used Kernel Density Estimation to calculate the curves of its biodiversity and variation rate (Fig. 6). The
curves of biodiversity at genus and species levels were similar except for their peaks. The curve of genus-
level reached its peak at Givetian, but the curve of species-level reached its peak at Frasnian. Both curves
could identify two sharp risings in diversity in Pridoli and Eifelian. The first rise in Pridoli coincided
255 with the rising of the jawed vertebrate and resulted in Yunnanolepidoidei reaching its diversity peak in
Lochkovian. The velocity rate of the genus level reached its peak at the end of Pridoli. After the E'Em
Event, the second rising was recognized by a faunal turnover from endemics to cosmopolitans (Zhu,
2000). Before this event, Petalichtyida and Antiarcha were endemic to China (Pan et al., 2015).
Wurungulepis was the only documented antiarch outside China in Early Devonian (Young, 1990b).
260 Yunnanolepidoidei was extinct after the E'Em Event, coinciding with the differentiation of euantriarchs.
The variation rate of species-level reached its peak at the end of Eifelian. Zhu (2000) has proposed the
Late Eifelian Event (Walliser, 1996) for the Chinese placoderm extinction. Although decreasing in Pan-
Cathaysian, the genus and species diversity of Antiarcha in other parts of the world kept rising after the
Late Eifelian Event. Walliser (1995) proposed the Mid-Famennian Event is characterized by the increase
265 of the species diversity of Antiarcha. On the contrary, our data shows that the species diversity of
Antiarcha underwent the sharpest decrease from Frasnian to Famennian except for the end of the
Devonian. Only the genus diversity of Antiarcha increased slightly at the beginning of Famennian.
Another difference between the two biodiversity curves is that the decreased genus diversity was in
marked contrast to the increased species diversity from Givetian to Frasnian.





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Figure 6 Biodiversity curves of Antiarcha. A, biodiversity curve at genus level. B, biodiversity curve at species level. Histograms indicate the diversity of Antiarcha in each time interval. The continuous red line indicates the biodiversity. The dash red line indicates the rate of variation. The dash blue line is the reference line of Variation at 0. Both red lines are calculated by Kernel Density Estimation.



275 5 Data Availability

The current dataset achieved via Zenodo represents a static version of the dataset in 3 November 2021: <https://doi.org/10.5281/zenodo.5639529> (Pan and Zhu, 2021). The latest version of the dataset is always freely available via <https://deepbone.org/> (last access: 3 November 2021).

6 Conclusions

280 Data are significant for quantitative analysis and contribute to data-driven scientific research. Previous works of early vertebrates usually focused on anatomy and phylogeny because of their significant impact on the origin of key characters. Nowadays, new collaborations have focused on biostratigraphy and biogeography, which throw new lights on the study of early vertebrates. Even if the fossil record is



incomplete, it is well demonstrated that the biogeographic evolution pattern of antiarchs can be
285 summarized from the compiled data and new analysis methods.

Author contributions. MZ supervised the project. ZHP and ZBN developed the model and performed
the simulations. ZHP prepared and revised the manuscript with contributions from MZ and ZBN. ZHP
and ZMX prepared the data.

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Competing interests. The authors declare that they have no conflict of interest.

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Reference



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Battersby, S. E., Finn, M. P., Usery, E. L., and Yamamoto, K. H.: Implications of web Mercator and its
use in online mapping, *Cartographica: J. Geogr. Inf. Geovis.*, 49(2), 85-101, doi:
10.3138/carto.49.2.2313, 2014.

310 Beals, E. W.: Bray-Curtis ordination: an effective strategy for analysis of multivariate ecological data,
Adv. Ecol. Res., 14, 1-55, doi: 10.1016/S0065-2504(08)60168-3, 1984.

Benton, M. J. and Donoghue, P. C. J.: Paleontological evidence to date the tree of life, *Mol. Biol. Evol.*,
24(1), 26-53, doi:10.1093/molbev/msl150, 2007.



- Benton, M. J., Donoghue, P. C. J., and Asher, R. J.: Calibrating and constraining molecular clocks, in: The Timetree of Life , edited by: Hedges, S. B. and Kumar, S., pp.35-86, Oxford Univ. Press, Oxford, 315 2009.
- Blicek, A., and Janvier, P.: Silurian-Devonian vertebrate biostratigraphy of Siberia and neighbouring areas, in: Palaeozoic Vertebrate Biostratigraphy and Biogeography, edited by: Long J. A., pp. 87-103, Bellhaven Press, London, 1993.
- Blicek, A., Turner, S., and Young, G. C.: Devonian vertebrate biochronology and global marine/non-marine correlation, Cour. Forschungsinst. Senckenberg, 220, 161-193, 2000.
- Brazeau, M. D.: The braincase and jaws of a Devonian ‘acanthodian’ and modern gnathostome origins. Nature, 457, 305-308, doi:10.1038/nature07436, 2009.
- Burrow, C. J.: Placoderm scales from the Lower Devonian of New South Wales, Australia, Mod. Geol., 20, 351-369, 1996.
- 325 Carr, R. K.: Placoderm diversity and evolution. Bull. Mus. Natl. Hist. Nat., Sect. C, 17(1-4), 85-125, 1995.
- Chang, K. J.: New antiarchs from the Middle Devonian of Yunnan, Vertebr. Palasiat., 9(1), 1-9, 1965.
- Davis, S. P., Finarelli, J. A., and Coates, M. I.: *Acanthodes* and shark-like conditions in the last common ancestor of modern gnathostomes, Nature, 486(7402), 247-250, doi:10.1038/nature11080, 2012.
- 330 De Pomeroy, A. M.: Biostratigraphy of Devonian microvertebrates from the Broken River region, north Queensland, West. Aust. Mus. Rec., 17, 417-437, 1996.
- Denison, R.: Placodermi. Handbook of Paleoiichthyology, vol. 2, 128, 1978.
- Donoghue, P. C. J. and Benton, M.: Rocks and clocks: calibrating the tree of life using fossils and molecules, Trends Ecol. Evol., 22(8), 424-431, doi:10.1016/j.tree.2007.05.005, 2007.
- 335 Downs, J. P.: Mass mortality of juvenile antiarchs (*Bothriolepis* sp.) from the Catskill Formation (Upper Devonian, Famennian Stage), Tioga County, Pennsylvania, Proc. Acad. Nat. Sci. Phila., 161, 191-203, 2011.
- Dupret, V., Sanchez, S., Goujet, D., Tafforeau, P., and Ahlberg, P. E.: A primitive placoderm sheds light on the origin of the jawed vertebrate face, Nature, 507, 500-503, doi:10.1038/nature12980, 2014.
- 340 Giles, S., Friedman, M., and Brazeau, M. D.: Osteichthyan-like cranial conditions in an Early Devonian stem gnathostome., Nature, 520, 82-85, doi:10.1038/nature14065, 2015.



- Guo, H. D. : Big Earth data: A new frontier in Earth and information sciences, *Big Earth Data*, 1(1-2), 4-20, doi: 10.1080/20964471.2017.1403062, 2017.
- Heckel, P. H., and Witzke, B. J.: Devonian world palaeogeography determined from distribution of carbonates and related lithic palaeoclimatic indicators, in: *The Devonian System*, edited by: House M. R., pp. 99-123, Vol. Spec. Pap. Paleontol. 23, 1979.
- Janvier, P. (Ed.): *Early Vertebrates*, Clarendon Press, Oxford, United States, 1996.
- Janvier, P. and Pan, J.: *Hyrceanaspis bliecki* n.g. n.sp., a new primitive euanthrop (Antiarcha, Placodermi) from the Middle Devonian of northeastern Iran, with a discussion on antiarch phylogeny, *Neues. Jahrb. Geol. Palaontol. Abh.*, 164(3), 364-392, 1982.
- Janvier, P. and Tông-Dzuy, T.: The Silurian and Devonian vertebrates of Viêt Nam: a review, *J. Geol., Series B*, 1998(11-12), 18-28, 1998.
- Ke, Y., Shen, S. Z., Shi, G. R., Fan, J. X., Zhang, H., Qiao, L., and Zeng, Y.: Global brachiopod palaeobiogeographical evolution from Changhsingian (Late Permian) to Rhaetian (Late Triassic), *Paleogeogr. Paleoclimatol. Paleoecol.*, 448, 4-25, doi:10.1016/j.palaeo.2015.09.049, 2016.
- King, B.: Bayesian tip-dated phylogenetics in paleontology: topological effects and stratigraphic fit, *Syst. Biol.*, 70(2), 283-294, doi: 10.1093/sysbio/syaa057, 2021.
- Kohonen, T.: The self-organizing map, *P. IEEE*, 78(9), 1464-1480, doi: 10.1109/5.58325, 1990.
- Kohonen, T.: The self-organizing map, *Neurocomputing*, 21(1-3), 1-6, doi: 10.1016/S0925-2312(98)00030-7, 1998.
- Li, Z. X. and Powell, C. M.: An outline of the palaeogeographic evolution of the Australasian region since the beginning of the Neoproterozoic, *Earth-Sci. Rev.*, 53(3-4), 237-277, doi: 10.1016/S0012-8252(00)00021-0, 2001.
- Long, J. A., Mark-Kurik, E., Johanson, Z., Lee, M. S., Young, G. C., Zhu, M., Ahlberg, P. E., Newman, M., Jones, R., Blauwen, J., Choo, B., and Trinajstić, K.: Copulation in antiarch placoderms and the origin of gnathostome internal fertilization, *Nature*, 517, 196-199, doi:10.1038/nature13825, 2015.
- Lu, L., Tan, K., and Wang, X.: A New Antiarchi (Placoderm Fishes) from Devonian Strata of Dushan, Guizhou Province, *Acta Geosci. Sin.*, 38(2), 144-148, 2017.
- Lukševičs, E.: Bothriolepid antiarchs (Placodermi, Bothriolepididae) from the north-western part of East European platform, Ph.D. thesis, Ģeogrāfijas un Zemes zinātņu fakultāte, University of Latvia, Latvia, 196 pp., 1996.



- Pan, J., Lu, L., and Ji, S. A.: A brief review of studies on palaeogeography of Middle Paleozoic vertebrates in China, *J. Southeast Asian Earth Sci.*, 13(3-5), 185-190, 1996.
- Pan, Z. H., Zhu, M., Zhu, Y. A., and Jia, L. T., A new petalichthyid placoderm from the Early Devonian of Yunnan, China, *C. R. Palevol*, 14(2), 125-137. doi:10.1016/j.crpv.2014.10.006, 2015.
- Pan, Z. H., and Zhu, M. VPPDB: Hosting the data for vertebrate paleoanthropology, *Acta Geol. Sin.-Engl.*, 93(supp.1), 61-63, doi: 10.1111/1755-6724.14246, 2019.
- Pan, Z. H., Zhu, M., Zhu, Y. A., and Jia, L. T.: A new antiarch placoderm from the Emsian (Early Devonian) of Wuding, Yunnan, China, *Alcheringa*, 42, 10-21, doi:10.1080/03115518.2017.1338357, 2018.
- Pan, Z. H. and Zhu, M, Dataset of antiarch placoderms (the most basal jawed vertebrates) throughout Middle Paleozoic (Version 31-Aug-2021), Zenodo, 10.5281/zenodo.5347006, 2021.
- Qiao, T., King, B., Long, J. A., Ahlberg, P. E., and Zhu, M.: Early gnathostome phylogeny revisited: multiple method consensus, *PLoS One*, 11(9), e0163157, doi:10.1371/journal.pone.0163157, 2016.
- Ricotta, C. and Podani, J.: On some properties of the Bray-Curtis dissimilarity and their ecological meaning, *Ecol. Complex.*, 31, 201-205, doi: 10.1016/j.ecocom.2017.07.003, 2017.
- Ritchie, A., Wang, S. T., Young, G. C., and Zhang, G. R.: The Sinolepidae, a family of antiarchs (placoderm fishes) from the Devonian of South China and eastern Australia, *Rec. Aust. Mus.*, 44(3), 319-370, doi:10.3853/j.0067-1975.44.1992.38, 1992.
- Sallan, L., Friedman, M., Sansom, S. R., Bird, M. C., and Sansom, J. I.: The nearshore cradle of early vertebrate diversification, *Science*, 362, 460-464, doi: 10.1126/science.aar3689, 2018.
- Scotese, C. R.: The PALEOMAP Project: paleogeographic atlas and plate tectonic software, *Oceanographic Literature Review*, 3(45), 606-607, 1998.
- Scotese, C. R.: Palaeo Map Project, <http://www.scotese.com>, last access: 31 August 2021.
- Scotese, C. R.: Tutorial: PALEOMAP paleoAtlas for GPLates and the paleoData plotter program, Technical Report, 56, <https://www.earthbyte.org/paleomap>, last access: 31 August 2021.
- Scotese, C. R., Voo, R. V., and Barrett, S. F.: Silurian and Devonian base maps [and Discussion], *Philos. Trans. R. Soc. Lond. Ser. B. Sci.*, 309(1138), 57-77, 1985.
- Trinajstić, K., Boisvert, C., Long, J., Maksimenko, A., and Johanson, Z.: Pelvic and reproductive structures in placoderms (stem gnathostomes), *Biol. Rev.*, 90(2), 467-501, doi:10.1111/brv.12118, 2015.



- Vesanto, J., and Alhoniemi, E.: Clustering of the self-organizing map, *IEEE Trans. Neural Netw. Learn. Syst.*, 11(3), 586-600, doi: 10.1109/72.846731, 2000.
- Walliser, O.H.: Global events in the Devonian and Carboniferous, in: *Global events and event stratigraphy in the Phanerozoic*, edited by: Walliser O.H., pp. 225-250, Springer, Berlin, Heidelberg, 405 New York, 1996.
- Wang, J. Q.: The Antiarchi from Early Silurian of Hunan, *Vertebr. Palasiat.*, 29(3), 240-244, 1991.
- Wang, S. T.: A new antiarch from the Early Devonian of Guangxi, *Vertebr. Palasiat.*, 25, 81-90, 1987.
- Wang, W., Qu, Q. M., and Zhu, M.: A brief review of the Middle Palaeozoic vertebrates from Southeast Asia, *Palaeoworld*, 19(1-2), 27-36, doi:10.1016/j.palwor.2009.12.005, 2010.
- 410 Xu, H. H., Niu, Z. B., and Chen, Y. S.: A status report on a section-based stratigraphic and palaeontological database the Geobiodiversity Database, *Earth Syst. Sci. Data*, 12, 3443–3452, doi: 10.5194/essd-12-3443-2020, 2020.
- Young, G. C.: Stratigraphic occurrence of some placoderm fishes in the Middle and Late Devonian, *Newsl. Stratigr.*, 243-261, 1974.
- 415 Young, G. C.: An asterolepidoid antiarch (placoderm fish) from the Early Devonian of the Georgina Basin, central Australia, *Alcheringa*, 8, 65-80, 1984a.
- Young, G. C.: Comments on the phylogeny and biogeography of antiarchs (Devonian placoderm fishes), and the use of fossils in biogeography, *Proc. Linn. Soc. N. S. W.*, 107(3), 443-473, 1984b.
- Young, G. C.: Antiarchs (placoderm fishes) from the Devonian Aztec siltstone, southern Victoria Land, 420 Antarctica, *Palaeontogr. Abt. A*, 202, 1-125, 1988.
- Young, G. C.: Devonian vertebrate distribution patterns and cladistic analysis of palaeogeographic hypotheses, *Geol. Soc. Lond., Mem.*, 12(1), 243-255, doi:10.1144/gsl.mem.1990.012.01.23, 1990a.
- Young, G. C.: New antiarchs (Devonian placoderm fishes) from Queensland, with comments on placoderm phylogeny and biogeography, *Mem. Queensland Mus.*, 28(1), 35-50, 1990b.
- 425 Young, G. C.: Placoderms (armored fish): dominant vertebrates of the Devonian period, *Annu. Rev. Earth Pl. Sc.*, 38(1), 523-550, doi:10.1146/annurev-earth-040809-152507, 2010.
- Young, G. C., Burrow, C. J., Long, J. A., Turner, S., and Choo, B.: Devonian macrovertebrate assemblages and biogeography of East Gondwana (Australasia, Antarctica), *Palaeoworld*, 19, 55-74, doi: 10.1016/j.palwor.2009.11.005, 2010.



- 430 Young, G. C. and Lu, J.: Asia-Gondwana connections indicated by Devonian fishes from Australia: palaeogeographic considerations, *J. Palaeogeogr.*, 9(8), 1-22, doi:10.1186/s42501-020-00057-x, 2020.
- Zhao, W. J., Zhu, M., Liu, S., Pan, Z. H., and Jia, L. T.: A new look at the Silurian fish-bearing strata around the Shanmen Reservoir in Lixian, Hunan province, *J. Stratigr.*, 40(4), 341-350, 2016.
- Zhu, M.: The phylogeny of the Antiarcha (Placodermi, Pisces), with the description of Early Devonian antiarchs from Qujing, Yunnan, China, *Bull. Mus. Natl. Hist. Nat., Sect. C*, 18, 233-347, 1996.
- 435 Zhu, M.: Catalogue of Devonian vertebrates in China, with notes on bio-events, CFS, Cour. Forschungsinst. Senckenberg, 223, 373-390, 2000.
- Zhu, M., Ahlberg, P. E., Pan, Z. H., Zhu, Y. A., Qiao, T., Zhao, W. J., Jia, L. T., and Lu, J.: A Silurian maxillate placoderm illuminates jaw evolution, *Science*, 354(6310), 334-336,
- 440 doi:10.1126/science.aah3764, 2016.
- Zhu, M., Yu, X. B., Ahlberg, P. E., Choo, B., Lu, J., Qiao, T., Qu, Q. M., Zhao, W. J., Jia, L. T., Blom, H., and Zhu, Y. A.: A Silurian placoderm with osteichthyan-like marginal jaw bones, *Nature*, 502(7470), 188-193, doi:10.1038/nature12617, 2013.
- Zigaite, Z. and Bliciek, A.: Chapter 28 Palaeobiogeography of Early Palaeozoic vertebrates, *Geol. Soc. Lond., Mem.*, 38(1), 449-460, doi:10.1144/m38.28, 2013.
- 445