Marine Early Triassic Actinopterygii from Elko County (Nevada, USA): implications for the Smithian equatorial vertebrate eclipse

Carlo Romano,1 James F. Jenks,2 Romain Jattiot,1,3 Torsten M. Scheyer,1 Kevin G. Bylund,4 and Hugo Bucher1

1Paleontological Institute and Museum, University of Zurich, Karl Schmid-Strasse 4, 8006 Zurich, Switzerland (carlo.romano@pim.uzh.ch); (romain.jattiot@pim.uzh.ch); (tscheyer@pim.uzh.ch); (hugo.fr.bucher@pim.uzh.ch)
21134 Johnson Ridge Lane, West Jordan, Utah 84084, USA (jenksjimruby@gmail.com)
3UMR CNRS 6282 Biogéosciences, Université de Bourgogne, 6 Boulevard Gabriel, 21000, Dijon, France (romain.jattiot@pim.uzh.ch)
4140 South 700 East, Spanish Fork, Utah 84660, USA (kevin@ammonoid.com)

Abstract.—The Early Triassic vertebrate record from low paleolatitudes is spotty, which led to the notion of an ‘equatorial vertebrate eclipse’ during the Smithian. Here we present articulated ray-finned fishes (Actinopterygii), collected from the marine Lower Triassic Thaynes Group at three new localities in Elko County (Nevada, USA), which were deposited within the equatorial zone. From the Smithian of the Winecup Ranch, we describe two partial skulls of the predatory actinopterygian Birgeria (Birgeriidae), attributed to B. americana new species and Birgeria sp. Birgeria americana n. sp. is distinguished from other species by a less reduced operculogular series. With an estimated total length of 1.72–1.85 m, it is among the largest birgeriids. We confirm that Birgeria encompasses species with either two or three rows of teeth on the maxilla and dentary, and suggest that species with three well-developed rows are restricted to the Early Triassic. From the latest Smithian of Palomino Ridge, we present a three-dimensional, partial skull of the longirostrine predator Saurichthys (Saurichthyidae). This and other occurrences indicate that saurichthyids were common in the western USA basin. From the early late Spathian of Crittenden Springs, we describe a posterior body portion (Actinopterygii indet.). This find is important given the paucity of Spathian osteichthyan sites. We provide a summary of Early Triassic vertebrate occurrences in the United States, concluding that vertebrate fossils remain largely unstudied. The presence of predatory vertebrates in subequatorial latitudes during the Smithian confirms that Early Triassic trophic chains were not shortened and contradicts the ‘equatorial vertebrate eclipse’.

Introduction

The end-Permian mass extinction event (~251.9 Ma; Burgess et al., 2014) was the most severe biotic crisis of the Phanerozoic, marking a significant change within marine and continental biocenoses (e.g., Raup, 1979; Sepkoski, 1984). In the aftermath of this catastrophic event, the surviving and newly evolved clades went through a series of subsequent crises during the Early Triassic, which selectively affected their recovery (e.g., Galfetti et al., 2007b; Orchard, 2007; Brayard et al., 2009, 2011, 2017; Song et al., 2011; Hofmann et al., 2013a, b; Hochuli et al., 2016).

Bony fishes (Osteichthyes) displayed relatively lower diversity during the Paleozoic, but radiated extensively after the end-Permian mass extinction event (e.g., Tintori et al., 2014a; Friedman, 2015; Romano et al., 2016a). The Triassic fossil record bears testimony to the first diversification event of the Neopterygii, the group to which over half of all living vertebrate species belongs. Whereas most neopterygians were small during the Triassic, the ‘Palaeopterygii’ (non-neopterygian actinopterygians) were predominantly represented by large species, suggesting that this group was important at higher trophic levels (Romano et al., 2016a).

Despite recent advances, the detailed pattern and tempo of recovery of the various clades of fishes remain unsettled because Early Triassic taxa are not as well studied as Middle Triassic ones (Scheyer et al., 2014; Tintori et al., 2014a; Romano et al., 2016a, b). Although Early Triassic fish assemblages have been described from numerous localities around the world (Brinkmann et al., 2010), some large paleogeographical domains and time intervals still suffer from a scanty record (e.g., López-Arbarello, 2004; Romano et al., 2016a, b). For example, little is known about low-latitude Early Triassic vertebrate faunas (e.g., Scheyer et al., 2014; Romano et al., 2016a)—a circumstance that led Sun et al. (2012) to speculate about an ‘equatorial vertebrate eclipse’, which they linked to extreme temperatures during the Smithian. Furthermore, the fossil record of Osteichthyes is marked by an extended interval with only a few, scattered occurrences during the Spathian (late Early Triassic). This Spathian gap in the osteichthyan fossil record potentially overlaps with the onset of the first neopterygian radiation (Romano et al., 2016a).

Here we describe for the first time articulated fish remains from the Early Triassic of Nevada (USA)—a paleoecographic domain situated near the paleoequator. The presented material
notably improves the Early Triassic record of fishes from the United States and provides new information on low-latitude vertebrate faunas during the essentially warm Early Triassic (Gouedemand et al., 2013; Romano et al., 2013). One of the presented fossils is derived from strata of Spathian age, thus adding a new occurrence to this interval.

**Localities, ages, and depositional settings**

The fossils described below were collected from three distinct localities in Elko County, northeastern Nevada, USA: (1) Winecup Ranch, (2) Palomino Ridge, and (3) Crittenden Springs.

**Winecup Ranch.**—Two specimens (NMMNH P-66225, NMMNH P-77117) were found near the Winecup Ranch (Elko County, Nevada, USA; Fig. 1). The Winecup Ranch (e.g., Oversby, 1972), or Wilkins Ranch of the older literature (e.g., Clark, 1957), is located ~43 km north of Wells, and ~7.2 km southeast of Wilkins Junction, on the south side of Thousand Springs Valley (Fig. 1.2). Lower Triassic exposures, designated as ‘Wilkins Ranch (V)’ in Clark (1957), crop out in two separate, but nearly contiguous, large areas encompassing ~7.7 km² each, and beginning ~1.6 km and ~3.2 km south of the ranch, respectively. These exposures occur in T40N, R64E and 65E and extend southward for ~5.5 km, reaching into the northern end of the Windermere Hills (Clark, 1957; Oversby, 1972; Coats, 1987).

Compared to most other Lower Triassic exposures in northeastern Nevada, relatively little is known regarding this particular section, especially the portion that represents strata of Smithian age. According to Clark (1957), the section, consisting...
of vertical and overturned beds in isolated outcrops, is highly faulted, with outcrops abutting against other outcrops with differing attitudes, making detailed logging impossible. Contact with Permian rocks is not visible and alluvial material covers much of the upper beds (Clark, 1957). Approximately 460 m of probable Spathian sediments, consisting of massive gray limestone and gray shaly limestone yielding bivalves, brachiopods, and ammonoids crop out in the higher hills ~6 km south of the ranch (Clark, 1957).

A succession of Smithian, dark-brown limestone overlain by 60–90 m of black shaly limestone is repeated several times on the east side of an unimproved road ~3 km south of the ranch (Clark, 1957). These rocks are relatively incompetent and, consequently, the topography in this particular area consists of sagebrush flats and a few, very low rolling hills. Outcrops are few—weathered shale covers most of the area and small pieces of dark-brown limestone occasionally found lying on the surface usually indicate the position of subsurface beds. A few well-preserved middle Smithian ammonoids (e.g., Inoities oweni) belonging to the ‘Meekoceras zone’ of Clark (1957) have been found by JJ and HB in these limestone beds of the Thaynes Group at two sites located 340 m apart. Oversby (1972) mentioned the occurrence of bones (large vertebral and fish remains) in the study area, but neither described nor depicted them.

The discovery sites of NMMNH P-66225 (N41°22′58.2″, W114°40′11.9″) and NMMNH P-77117 (N41°22′49.5″, W114°40′24.8″) are located ~2.75 km south-southeast of the Winecup Ranch. P-77117 was found at the most productive locality together with Pseudaspidites, Procurtcoreoceras, propycithid indet., and prionitid indet., which, combined with Inoities oweni, indicate a late early Smithian to early late Smithian age (Brayard and Bucher 2008; Brühwiler et al., 2010; Brayard et al., 2013). P-66225 was found more or less on strike ~300 m northeast of the P-77117 locality. Conodonts or ammonoids have not been detected in the matrix surrounding P-66225, but a few exceptionally well-preserved Smithian ammonoids, many of which retain relict color bands, and other fossils have been collected from the site since it was discovered in the early 1950s (Mullen, 1985; Jenks et al., 2010). Disarticulated vertebrate bones are occasionally found in association with the ammonoids of the ‘Meekoceras beds’ (personal observation, JJ, KGB, TMS, HB, 2010–2016).

Lower Triassic marine sediments, consisting of the Dinwoody Formation and units belonging to the overlying Thaynes Group, crop out in the hills immediately north of the Long Canyon road and extend to the northeast for ~8 km, covering an area of ~33 km² (Clark, 1957; Mullen, 1985). Good exposures of Spathian shales are limited to a few seasonal storm drainage channels. One such channel cuts numerous, gently northward dipping layers with early diagenetic limestone nodules, most of which are barren, but a few yield ammonoids (e.g., Stacheites) typical of the early late Spathian Fengshanshi/ Prohungarites fauna (Bucher, 1989; Guex et al., 2010). A float concretion found in the same channel contains a partial skeleton of an actinopterygian (NMMNH P-77357), which is described herein. Additional, articulated cranial material (PIMUZ A/I 4641; Birgeria?) has recently been recovered from the Spathian of Crittenden Springs and is currently undergoing preparation for eventual study.

**Materials and methods**

The described material is curated in part by the NMMNH, and in part by the PIMUZ. PIMUZ A/I 4397 from Palomino Ridge and one specimen from the Winecup Ranch (NMMNH P-66225)
were prepared using air pens under a binocular microscope. The fragile bones were consolidated with liquid glue (low-viscosity cyanoacrylate) during preparation.

For consistency, the anatomical terminology used herein follows that of previous publications (e.g., Stensiö, 1921, 1925, 1932; Nielsen, 1949; Lehman, 1952; Ørvig, 1978; Mutter et al., 2008; Romano and Brinkmann, 2009). However, no homology with similarly named bones of other vertebrates is necessarily implied (cf. Schultze, 2008). When describing the scales, we refer to length as their longitudinal (anteroposterior) extent and depth as their dorsoventral dimension. Open nomenclature is used in accordance with the recommendations of Bengtson (1988).

The following comparative material was consulted: ZMUC VP 3176 (Birgeria groenlandica, holotype; Stensiö, 1932); MNHN.F MAE 605, MAE 606 (B. nielseni; Beltan, 1980); PMU P 1421 (B. aldingeri, holotype; Stensiö, 1932; Schwarz, 1970); PMU P 349, P 1422 (B. cf. aldingeri; Stensiö, 1932); PIMUZ A/I 4301 (Birgeria sp.; Scheyer et al., 2014); PIMUZ material of B. stensioei (as listed in Romano and Brinkmann, 2009); PIMUZ A/I 3900 (Saurichthys cf. elongatus; Romano et al., 2012); and PIMUZ A/I 4135, A/I 4144 (S. madagascariensis; Kogan and Romano, 2016a). Additionally, a photograph of MPUM 9334 (Birgeria sp.; Lombardo and Tintori, 2005, fig. 1b) was obtained for this study.

We employ Tozer’s (1965) stage subdivisions of the Early Triassic: Griesbachian (early Induan), Dienerian (late Induan), Smithian (early Olenekian), and Spathian (late Olenekian). These stages are well defined by global biotic events (e.g., Galfetti et al., 2007b; Hochuli et al., 2016). Recent biochronostratigraphic datings indicate a duration of maximally $1.4 \pm 0.4 \text{ myr}$ (million years) for the Griesbachian–Dienerian interval, $~0.7 \pm 0.6 \text{ myr}$ for the Smithian, and $~3 \text{ myr}$ for the Spathian (Ovtcharova et al., 2006; Galfetti et al., 2007a).

Repositories and institutional abbreviations.—Types, figured, and other specimens examined in this study are stored in the following institutions: New Mexico Museum of Natural History and Science (NMMNH), Albuquerque, New Mexico, USA; Paleontological Institute and Museum, University of Zurich (PIMUZ), Zurich, Switzerland; Museum National d’Histoire Naturelle (MNHN.F), Paris, France; Museo del Dipartimento di Scienze della Terra ‘A. Desio’ dell’Università degli Studi di Milano (MPUM), Milan, Italy; Museum of Evolution (PMU, formerly Paleontological Museum), Uppsala University, Uppsala, Sweden; Zoological Museum, Natural History Museum of Denmark (ZMUC), Copenhagen, Denmark.

Systematic paleontology

Anatomical abbreviations.—Af, anal fin (lepidotrichia); An, angular (external plate); Aop, antoperculum; Ar, Meckel’s cartilage (ossified as the articular); Br, branchiostegal ray;
Remarks.—The family includes only Birgeria Stensiö, 1919. Two other genera formerly referred to Birgeriidae, the Early Jurassic Ohmdenia Hauff, 1953 and the Early Cretaceous Psilichthys Waldman, 1971; Friedman, 2012, have been shown to be unrelated (Waldman, 1971; Friedman, 2012).

Genus Birgeria Stensiö, 1919, emend. Romano and Brinkmann, 2009

Type species.—Birgeria mougeoti (Agassiz, 1843) from the Middle Triassic of Bayreuth, Germany.

Remarks.—Stensiö (1919) erected the genus based on a maxilla from the Middle Triassic of Germany that contains teeth resembling those of the type material of 'Saurichthys' mougeoti Agassiz, 1843, which henceforth became the type species of Birgeria. The type material from the Muschelkalk of France and Germany is composed of several isolated teeth and dentigerous maxilla fragments, which clearly cannot be attributed to Saurichthys Agassiz, 1834, because in this taxon the maxilla lacks such large teeth (e.g., Stensiö, 1925; Rieppel, 1985; Mutter et al., 2008; Maxwell et al., 2015; Kogan and Romano, 2016a). The material of Agassiz (1833–1843) and Stensiö (1919) is too fragmentary for identification at the species level, but bears close resemblance to material of other, much better known species referred to Birgeria. Additional, more complete material from the Early Triassic of Spitsbergen (Svalbard, Arctic Norway), supposedly belonging to B. mougeoti (Agassiz, 1843), was figured and described by Stensiö (1919, 1921, 1932). One specimen of Stensiö’s (1932) material, however, was later considered as a separate species, B. aldingeri Schwarz, 1970. Although Schwarz (1970) did not explicitly comment on the taxonomic status of other material from Spitsbergen (Stensiö, 1919, 1921, 1932), we agree that it is likely not conspecific with that from the Middle Triassic Germanic Basin, due to notable differences in the angle of the anterior margin of the postorbital maxillary blade (see Discussion). Pending a proper revision of the type species and the species from Spitsbergen, we follow previous authors and treat B. mougeoti from the Muschelkalk Sea as a valid species, whereas Stensiö’s (1919, 1921, 1932) material from Spitsbergen is herein provisionally referred to as B. cf. aldingeri (except for the holotype of B. aldingeri).

Occurrence.—Marine Triassic, global: from the Griesbachian (earliest Triassic) of Greenland to the Rhaetian (latest Triassic) of Europe.

Birgeria americana new species

Figures 3, 4

Holotype.—NMMNH P-66225 (Figs. 3, 4), from upper lower Smithian to lower upper Smithian beds (Thaynes Group), ~2.75 km south-southeast of the Winecup Ranch, east-central Elko County, Nevada, USA (Fig. 1). P-66225 is a partial skull preserved within a large limestone nodule, with its right side exposed. A digital 3D surface scan of P-66225 is available at MorphoMuseuM (Romano et al., 2017).

Differential diagnosis.—Very large (>150 cm) birgeriid (larger than B. groenlandica Stensiö, 1932, B. nielseni Lehman, 1948, B. liui Jin, 2001, B. guizhouensis Liu, Yin, and Luo in Liu et al., 2006, and B. acuminata [Agassiz, 1843]); postorbital blade of maxilla elongate, anteriorly low, posteriorly high, with inclined anterior border (relatively shorter, anteriorly about as high as posteriorly, with steep anterior margin in B. mougeoti [Agassiz, 1843], B. stensioei Aldinger, 1931, and B. acuminata); antoperculum present; suboperculum dorsoventrally elongate (absent or weakly ossified in B. stensioei); dorsoventrally shorter in B. groenlandica and B. nielseni); five to six postmandibular branchiostegal rays (four to five in B. nielseni, three to five in B. groenlandica, maximum one in B. stensioei); teeth of the intermediate row on the maxilla and dentary mostly widely spaced, varying in size, but mostly high (small, equal-sized, close-set intermediate teeth in B. stensioei and B. acuminata); labial teeth distinct (smaller in B. stensioei, absent in B. acuminata).

Description.—NMMNH P-66225 (Figs. 3, 4) shows the portion between the cleithrum posteriorly, and the level of the hind margin of the orbital opening anteriorly. The preserved part has a length of 26 cm. Most bones are still in situ. Parts of the maxilla, the preoperculum, the suborbitals, and the lower jaw were damaged due to weathering.

Upper jaw.—The maxilla is the dominant bone of the upper jaw (Fig. 3). It consists of a cleaver-shaped external plate (composed of a low suborbital portion and a high, elongate postorbital blade) and an internal lamina. The postorbital blade of the maxillary bone is confined by a slanted, concave anterior margin, a nearly straight dorsal border, a posterior margin that is straight in its upper segment and distinctly sigmoid in its lower part, and a dentigerous ventral margin, which is largely straight except for its posteriormost portion, which is concave. The postorbital plate is mostly flat except for the posteroventral part, which is laterally convex. The ornamentation of the maxilla is only preserved in proximity to the tooth-bearing margin, where
Figure 3. *Birgeria americana* n. sp. (NMMNH P-66225, holotype) from the Smithian of the Winecup Ranch, Elko County, Nevada, USA. (1) Fossil in dorsal view (above) and in right aspect (below); (2) schematic drawing of 3.1 with interpretations of skeletal features; anterior is right. Scale bar = 50 mm (total).
it consists of minute teeth (odontodes; Ørvig, 1978). The internal lamina extends from the rostral end of the maxilla until the level of the posterior end of the dentigerous margin of the dentary (Fig. 3).

Lower jaw.—The mandible is largely complete, with only the anteriormost part missing (Fig. 3). Most of the lateral surface is weathered, meaning that the margins between the dentary, angular, and surangular represent the medial ones (the dentary usually covers a large area of the angular laterally; e.g., Romano and Brinkmann, 2009).

The dentary is composed of an external plate as well as a dorsal and a ventral internal lamina. The external plate is bounded by a gently convex ventral margin and a nearly straight, tooth-bearing dorsal border. The dorsal internal lamina runs close to the upper confinement of the external plate, whereas the ventral internal lamina parallels the lower margin of the bone. The angular forms the posterior and posteroventral borders of the lower jaw. This bone, too, is composed of an external plate and a large internal lamina. The plate-like part of the angular is confined by a long, convex border ventrally, and by an S-shaped margin posteriorly, which is dorsally concave and ventrally convex. The posterior and ventral margins run suborthogonally and together form a rounded posteroventral corner on the mandible. The suture between the external plate of the dentary and the surangular runs obliquely from anterodorsal to posteroventral, except in the most dorsal segment; there, the posterior margin of the dentary forms a recess, which houses the pointed anterior part of the surangular (Fig. 3). The upper portion of the boundary between the external plates of the angular and dentary is vertical, whereas the lower portion is distinctly inclined, running from posterodorsal to anteroventral.

The wedge-shaped surangular exhibits a coronoid process (Fig. 3), without contribution of the dentary. The mandibular sensory canal traverses the external plate of the angular near its caudal and ventral borders and continues through the external plate of the dentary near its ventral margin, probably along the base of the ventral internal lamina.

The conspicuous internal lamina of the angular projects from the medial side of the plate-like part (Fig. 3). The base of this internal lamina follows the ventral and posterior borders of the external plate, but is offset with regard to these margins. In the posterior region of the lower jaw, the internal lamina is oriented mediocaudally, forming an obtuse angle with the external plate. In contrast, the external plate and internal lamina run suborthogonally within the ventral part of the angular. The ventral component of the medial lamina is lateromedially less broad than the posterodorsal portion of the lamina. The posterior part of the lamina becomes successively broader from the posteroventral angle of the angular to the level of the jaw joint, forming a sizeable postero medial projection at the caudal end of the mandible (Fig. 3). The internal lamina is also curved medially.

Dentition.—Macroscopic teeth are developed on the dentary and the maxilla, whereas the surangular lacks such teeth. The dentition of the maxilla and dentary consists of conical teeth that are arranged in three longitudinal rows: a lingual, an intermediate, and a labial one. The lingual row consists of fairly large, stout teeth growing on the internal lamina of the maxilla, and the dorsal internal lamina of the dentary, respectively. In the

---

**Figure 4.** *Birgeria americana* n. sp. (NMMNH P-66225, holotype) from the Smithian of the Winecup Ranch, Elko County, Nevada, USA. (1) Tentative restoration of the skull of P-66225; (2) close-up view (position indicated in 4.1) of the dentition (above), and drawing thereof (below); the large lingual teeth growing on the internal lamina of the maxilla and the dorsal internal lamina of the dentary, respectively, are highlighted in dark gray; teeth of the labial row (white arrows) and intermediate row are colored white; (3) close-up view of a lingual tooth of the maxilla (position shown in 4.1) illustrating the surface ornamentation of the base and acrodin cap (photographed using a Leica MZ16F camera mounted on a stereomicroscope, contrast enhanced with ammonium chloride); the white arrow in 4.3 points to the demarcation between the tooth base and the acrodin cap; anterior is right (1–3). Scale bars = 10 mm (total) (2), or 5 mm (total) (3).
rostral part of these bones, the large lingual teeth are fully exposed (but partly weathered), whereas farther posteriorly only the apical parts are visible (Figs. 3, 4). The lingual teeth are aligned almost perfectly, forming a palisade (although gaps may occur where teeth fell out in vivo). The largest lingual teeth of the maxilla are located near the rostral end of the bone. From there, they slowly decrease in size in caudal direction. Conversely, the lingual teeth of the dentary slightly increase in height in posterior direction. The posteriormost lingual teeth of the dentary could not be reached with the air pen.

The intermediate series of teeth line the ventral margin of the external plate of the maxilla, and the dorsal margin of the external plate of the dentary, respectively (they are lost in the anterior parts of both of these bones due to weathering). The teeth of the intermediate row are always smaller than the neighboring teeth of the lingual row. They are irregularly distributed, but consecutive teeth are mostly separated by wide interspaces. The intermediate teeth of the maxilla decrease in size posteriorly, becoming minute at the level of the caudal end of the dentigerous margin of the dentary. Farther posteriorly, within the convex part of the maxilla that laterally covers the lower jaw, the intermediate teeth of the maxilla are larger again (Fig. 3). The intermediate teeth in the anterior segment of the dentary are about the same size as their antagonists on the maxilla, whereas those in the posterior segment of this bone are distinctly larger (Fig. 4). Close to the caudal end of the dentigerous margin of the dentary, the intermediate teeth rapidly decrease in size.

The teeth of the labial row are smaller than those of the intermediate row (Fig. 4). Like the intermediate teeth, the labial teeth are irregularly distributed along the jaw margins. They are quite frequent in the posterior part of the dentary, but a few labial teeth are also intermittently developed on the maxilla, especially in the anterior segment of this bone. The labial teeth are bordered by minute teeth laterally (odontodes; Ørvig, 1978), forming the ornamentation of the maxilla and the dentary.

Whereas the apicobasal axes of the lingual teeth are inclined medially (and gently caudally), those of the intermediate teeth are more or less oriented dorsoventrally (except in the caudal part of the maxilla, where they are medially tending). The apicobasal axes of the labial teeth are inclined laterally, whereas the lateral leaning varies between individual labial teeth. The apicobasal axis of some labial teeth is nearly perpendicular to that of the lingual teeth (Fig. 4.2). The teeth of the three longitudinal rows are all linguolabially compressed and curved medially.

All teeth comprise an acrodin cap (Ørvig, 1978) and a high base (Fig. 4.2, 4.3). The acrodin cap usually makes up only a small part of the total tooth height in lingual teeth (probably about one-fifth to one-sixth), whereas in teeth of the intermediate and labial rows the acrodin cap measures at least one-third of the total height. The surface of the acrodin cap is mostly smooth, except for its basal segment, which shows a few widely spaced, meandering, apicobasally oriented ridges (Fig. 4.3). Anterior and posterior cutting edges are present on teeth of all three rows. The surface of the tooth base, where preserved, is ornamented with very fine, subvertical striae. The striae are close-set and frequently anastomose (Fig. 4.3).

Preoperculum and operculogular series.—The incompletely preserved preoperculum (Fig. 3) adjoins the postorbital plate of the maxilla dorsally and posteriorly. The boomerang-shaped preoperculum is composed of a long anterior shank and a short posterovertral branch, which meet in an obtuse angle. The rostral margin of the anterior shank is deeply concave and the anterodorsal part of the bone protrudes rostrally. The posterovertral branch of the preoperculum adjoins the straight upper part of the posterior margin of the maxilla, but does not reach farther ventrally, thus exposing the quadratum laterally. The caudal margin of the preoperculum is convex. A section of the preopercular sensory canal is visible in the lower shank. The canal traverses the bone centrally (Fig. 3).

The bones of the operculogular series (Fig. 3) are arranged in the characteristic manner for Birgeria. The operculum is situated dorsally to the anterior shank of the preoperculum. It is tilted in a way that its external side now faces dorsally. The operculum is incompletely preserved, but had an oblong, ovoid outline with acute anterior and posterior ends. Rostral to the operculum is another small, plate-like bone, interpreted as the antoperculum. Caudally, the operculum reaches to the knee of the preoperculum, where it is still articulated with the suboperculum (sensu Romano and Brinkmann, 2009). The vertically arranged suboperculum is elongate and slender, being morphologically indistinguishable from the posteriorly adjoining branchiostegal rays. The suboperculum has a triangular outline and is confined by very long anterior and posterior borders and a short dorsal margin contacting the operculum. The lateral side of the suboperculum is convex.

The branchiostegal series is divided into a postmandibular and a submandibular series, which are separated by a gap (Fig. 3). Five postmandibular radii branchiostegii are preserved ventrocaudal to the suboperculum. The postmandibular branchiostegals are coalesced neither with each other nor with the suboperculum. The postmandibular rays are slender, elongate elements. The first postmandibular branchiostegal ray is dorsoventrally almost as long as the suboperculum. The caudally following postmandibular rays are much shorter and their length consecutively decreases towards the last of these bones. Moreover, whereas the first and second rays are dorsoventrally oriented, like the suboperculum, the subsequent ones are more and more caudally inclined, with the fourth and fifth ray being somewhat anteroposteriorly arranged. Each ray has an acute ventral/anterior end and a rounded dorsal/posterior termination (except for the posteriormost of these elements, which has a pointed caudal end). Their lateral surface is convex. In addition to the five postmandibular rays situated behind the suboperculum, a small, thin, sixth postmandibular radium branchiostegium is seen isolated about halfway between the postmandibular and the submandibular series (Fig. 3). Mediolaterally, this ray is situated at the same level as the other branchiostegals.

Nine slender submandibular branchiostegals are preserved medioventral to the lower jaw (Fig. 3), though their original number was higher. Each of these elements is plate-like and has a quadrangular outline, whereas their anterolateral corner protrudes rostrally. The anteroposterior width of these bones subsequently decreases from the first (anterior) to the last one. Furthermore, while the anterior submandibular branchiostegals are only
moderately posteriorly inclined, the caudally following ones are more distinctly so.

Suborbital and infraorbital series.—The bones of the suborbital series are preserved in situ, albeit mostly in a poor state of preservation. Their morphology and arrangement agree with those of other species of Birgeria, meaning that they are elongate, obliquely and serially arranged bones with a broad posteroventral part and a slender anterodorsal portion. About three suborbitals are visible (Fig. 3), the posterodorsal parts of which are still intact and nestled below the anterior process of the preoperculum. The anteroventral parts of the two, posteriormost suborbitals of P-66225 are fragmentarily preserved, but it is evident that they extended rostrad until the level of the third lingual tooth of the maxilla (which would mark the level of the posterior border of the orbital opening; Fig. 4.1).

A subtriangular bone fragment located anterodorsal to the suborbitals (Fig. 3) may belong to the dermosphenotic (this bone is usually subdivided into two or more elements in Birgeria, see Stensiö, 1932; Romano and Brinkmann, 2009). Other elements of the circumorbital series and the sclerotic ring are perhaps visible at the rostral end of the fossil, but they are too poorly preserved for accurate description.

Splanchnocranium.—The quadratum and articular are preserved in situ (Fig. 3). The quadratum is partly covered by the maxilla laterally, but the articulation condyle is exposed. The articular is mostly hidden by the angular and only its dorsalmost portion can be seen. The hyomandibula is only partly visible, being exposed where the preoperculum is damaged (Fig. 3). Some additional bones situated just medial to the postmandibular branchiostegal rays possibly belong to the hyomandibula or the symplectic, but they are not well exposed. The dermohyal is preserved in situ between the preoperculum, the operculum, and the antoperculum (Fig. 3). This bone is elongate and wedge-like, with a pointed posterior end, and a rounded, club-shaped anterior termination. The dermohyal is marked by a longitudinal crest running along its dorsal side. A few elements of the branchial skeleton are partly exposed (Fig. 3). For instance, a portion of an epibranchial is probably visible medially to the operculum. A segment of a possible ceratobranchial is seen at the posterior end of the concretion. Several rod-like bones preserved anterodorsal to the lower jaw likely belong to the hyoid arch or the branchial arches.

Pectoral girdle.—The right cleithrum is the only bone of the shoulder girdle that is visible (Fig. 3). Only the large lower branch of the cleithrum is preserved, the anterodorsal portion of which could not be prepared. The cleithrum fragment has a roughly triangular outline, being confined by a long, gently convex ventral border, and a long dorsal margin, being deeply concave in its caudalmost segment. The posterodorsal corner of the cleithrum fragment tapers, marking the transition to the missing upper branch of this bone. The external surface of the cleithrum is divided into a large lower portion, facing laterally, and a subhorizontally oriented upper portion.

Etymology.—The species name refers to its provenance.

Remarks.—The morphology of NMMNH P-66225 agrees well with that of the Triassic genus Birgeria Stensiö, 1919. Species of Birgeria are essentially differentiated by cranial features, but also by a few postcranial traits, such as the arrangement pattern of the dorsal fin pterygophores (Romano and Brinkmann, 2009). Diagnostic characters justifying the erection of a new species for the Nevada material are the less reduced dermal gill cover (i.e., presence of an antoperculum), very elongate suboperculum, and relatively numerous postmandibular branchiostegal rays, including a rudimentary ray situated within the gap of the branchiostegal series (see Discussion).

Based on the length of the preserved part (26 cm) of P-66225, corresponding to ~70–75% of the snout to shoulder girdle length in birgeriids with elongate postorbital skull portions (e.g., Nielsen, 1949), a snout to shoulder girdle length of ~34.5–37 cm for P-66225 is estimated. In large species of Birgeria, the skull plus shoulder girdle length usually corresponds to ~20% of the total length (see Liu et al., 2006; Romano and Brinkmann, 2009), suggesting a total length of ~172–185 cm for P-66225.

Birgeria sp.

Figure 5

Occurrence.—From upper lower Smithian to lower upper Smithian strata (Thaynes Group), ~2.75 km south-southeast of the Winecup Ranch, east-central Elko County, Nevada, USA (Fig. 1).

Description.—NMMNH P-77117 (Fig. 5) is a cranial fragment with a length of 17 cm. The fossil is seen in left aspect and preserved in a limestone nodule as part (P-77117 a) and counterpart (P-77117 b), with the fracture surface going through the bones of the left cheek. The specimen is associated with bivalves, one of which resembles Crittendenia (personal communication to CR, M. Hautmann, 2016).

Palatoquadrum and its dermal bones.—Several upper jaw bones are exposed: the maxilla, dermopalatine, entopterygoid, ectopterygoid, and ‘dermometapterygoid’ (Fig. 5). The quadrum is probably present, though not very well visible. The maxilla is only fragmentarily preserved, with portions of the postorbital blade and the internal lamina discernible. The dermopalatine is an elongate, low element situated in the anterodorsal region of the upper jaw. This bone exhibits its maximum depth rostrally. The height of this element tapers caudally. Only one dermopalatine is visible. Dorsal to the dermopalatine is the entopterygoid, of which only the rostral portion is preserved. The ectopterygoid lies posterior to the dermopalatine and posterodorsal to the entopterygoid. The ectopterygoid is a large, rostrocaudally elongate element that extends posteriorly until the jaw joint area. The boundary between the ectopterygoid and the entopterygoid runs obliquely from anterodorsal to posterodorsal. Another, albeit incompletely preserved, element situated posterior to the entopterygoid and posteroventral to the ectopterygoid, represents the ‘dermometapterygoid’ (terminology of Nielsen, 1949; but see Arratia and Schultz, 1991).

Lower jaw.—The mandible is incompletely preserved. Its anterior tip is missing and the lateral surface is broken off (Fig. 5). The exposed margins between the dentary, angular, and surangular represent the medial ones, and the morphologies of these bones are similar to those of NMMNH P-66225 (holotype of B. americana n. sp.; Fig. 3). The angular of P-77117 (Fig. 5) is equipped with a well-developed internal lamina. The dorsal and
ventral internal laminae of the dentary are well visible, delimiting the space occupied by Meckel’s cartilage. A fragment of the dentigerous upper margin of the prearticular is preserved.

**Dentition.**—Teeth are preserved on several jaw bones (Fig. 5). The basal portions of close-set lingual teeth are preserved in the posterior segment of the dentary. One tooth of the maxilla and one of the dentary is each visible in transverse section on part b of the fossil. In addition, longitudinal rows of close-set, macroscopic teeth are present on the dermopalatine and the ectopterygoid in the upper jaw, and on the prearticular in the lower jaw. At least one row of large teeth is observed on both the dermopalatine and the ectopterygoid. These teeth are smaller than the lingual teeth of the maxilla. The medial surface of the dermopalatine and ectopterygoid is covered with minute teeth, cross sections of which are seen along the oral margins of these bones. At minimum one row of teeth lines the dorsal margin of the prearticular. These teeth are smaller than the larger teeth of the dermopalatine and the ectopterygoid. Macroscopic teeth are not developed on the surangular.

**Preoperculum, operculogular series, and suborbitals.**—The fragmentarily preserved preoperculum is composed of two shanks—an anterior and a posteroroventral one (Fig. 5). Mainly the dorsal and posterior bone margins are visible. The dorsal border is straight and subvertical, whereas its caudal confinement is convex. At the level of the knee of the preoperculum, a weak notch is discernible. A section of the preopercular sensory canal traverses the posteroroventral shank centrally in anterodorsal direction, then continues rostrad near the straight dorsal margin of the bone. The most anterior part of the canal is not preserved. The operculum is situated dorsal to the anterior shank and has an elongate, ovoid shape. The rostral end of this bone is pointed. Nine submandibular branchiostegal rays are preserved ventral to the dentary. The anterior rays are rostrocaudally broader than the posterior ones. The anterolateral corner of each ray protrudes rostrally. One or two suborbitals are possibly seen near the anterior end of the fossil.

**Remarks.**—The overall anatomy of NMMNH P-66225 supports its referral to *Birgeria* Stensiö, 1919. However, attribution at the species level is not possible due to the absence of diagnostic characters. Based on comparison with *B. groenlandica* Stensiö, 1932 (cf. Nielsen, 1949), a total length of ~145–165 cm is estimated for P-77117.

**Family Saurichthyidae Owen, 1860, emend. Stensiö, 1925**

**Genus Saurichthys Agassiz, 1834**

**Type species.**—*Saurichthys apicalis* Agassiz, 1834 from the Ladinian (Middle Triassic) of Bayreuth, Germany, by original designation.

**Occurrence.**—From the latest Permian of China to the Late Triassic of Europe and possibly China.

*Saurichthys* sp. Figure 6

**Occurrence.**—Bed PLR 35 (Jattiot et al., in press), latest Smithian *Pseudosageceras augustum* Zone, Thaynes Formation (Fig. 2), Palomino Ridge, Section 2 (Jattiot et al., in press), southern Elko County, Nevada, USA (Fig. 1).

**Description.**—PIMUZ A/I 4397 (Fig. 6) is a three-dimensional skull fragment preserved in a limestone slab. The posterior part of the specimen, as well as the maxilla, preoperculum, lower jaw, and operculogular series are missing. The superficial layer of the
dermal bones, carrying the ornamentation, is mostly weathered and the suture lines are visible.

Dermal skull roof.—The dermal cranial roof is mainly formed by the large, paired frontals, and the large, paired dermopterotics. The medial boundaries between these bones are intermittently visible from near the caudal end of the skull fragment up to a short distance posterior to the tip of the rostrum. The frontals are oriented mostly horizontally, except for the part anterodorsal to the orbital opening, where the bone exhibits a small, laterally facing flange. The ossification centers of both frontals are well visible and located at about the level of the hind margin of the orbit. The supraorbital sensory canal runs from these ossification centers up until the level of the external narial openings, where the canal turns laterally.

The ossification center of the dermopterotic is situated close to the posterolateral margin of the dermal skull roof (Fig. 6). The boundary between the frontal and the dermopterotic is scarcely visible, but judging from the circumferences of the growth lines radiating from the ossification centers of both of these bones, their mutual boundary runs sinuously from anterolateral to posteromedial. Anteriorly, the dermopterotic forms the posterior part of the dorsal margin of the orbit. The dermopterotic forms an extended lateral flange posterodorsal to the orbital opening. The remains of a poorly preserved dermosphenotic may be visible along the anterior margin of this lateral flange. A sensory canal is traceable along the lateral border of the postorbital portion of the dermal skull roof, running through the dermopterotic in the rostrocaudal direction. Posterodorsal to the orbital opening, the canal turns ventrolaterally. Some tubercles are visible along the sensory canal in the posterolateral part of the left dermopterotic. A pair of small parietals is developed medially to the dermopterotics and...
posteriorly to the frontals, but the boundaries between these elements are mostly unclear.

Rostrum.—The margins between the bones composing the rostrum are only partly visible. The tip of the rostrum is formed by the large, unpaired rostropremaxilla (Fig. 6). Posterolaterally, the rostropremaxillary bone tapers on both sides. Caudal to the rostropremaxilla lies the small lachrymal, which is possibly preserved on the left side. The nasaloantorbital forms the anterior border of the orbit. This bone is wedged between the lachrymal and the postero lateral part of the rostropremaxilla ventrally, and the frontal dorsally. Its anterior end is elongate and probably acute. The nasaloantorbital contains both external narial openings, which are well visible on the right side of the skull (Fig. 6). The anterior naris is deeper than the posterior one. The nasaloantorbital also includes the junction of three sensory canals, which is situated ventral to the nares. The supraorbital canal enters the nasaloantorbital dorsally and runs between the narial openings. The ethmoidal canal passes laterally through the rostropremaxilla and pierces the nasaloantorbital through its anterovelventral margin. The infraorbital canal enters the nasaloantorbital coming from the lachrymal.

Remarks.—PIMUZ A/I 4397 is a saurichthyid but, identification to the species level is complicated due to its incomplete preservation. The majority of diagnostic traits differentiating species of Saurichthys pertain to the postcranium, which is not preserved in the specimen from Palomino Ridge (see Discussion).

The skull has a preserved length of ~132 mm, suggesting that it belonged to a medium-sized saurichthyid (sensu Tintori, 2013). The rostrum is subcomplete and long (~100 mm from the tip of the snout until the anterolateral margin of the orbit).

Actinopterygii indet.

Figure 7

Occurrence.—Early late Spatian Fengshanites/Prohungarites beds (Thaynes Group), Crittenden Springs, northeast Elko County, Nevada, USA (Fig. 1).

Description.—NMMNH P-77357 (Fig. 7) is a posterior body portion of a medium-sized actinopterygian (estimated total length: ~25–30 cm). Preserved are the scales, most of the caudal fin, and the basal portions of some lepidotrichia of the dorsal and anal fins. The skeletal elements are largely in situ and mostly preserved as imprints, which have been affected weakly by surface erosion.

Squamation.—The squamation consists of rhombic scales arranged in oblique vertical rows (Fig. 7). Scales are also developed on the proximal part of the dorsal caudal lobe. About 31 scale rows are counted along the lateral midline of the trunk, from the anterior end of the fossil to the caudal inversion. Dorsally, some of the vertical scale rows seem to split into two rows. The scales along the flank are about as long as they are deep, whereas those in the dorsal and ventral areas are less deep. The scale ornamentation and the lateral line sensory canal are not preserved, and a peg-and-socket articulation is not visible.

Fins.—The proximal parts of at least six lepidotrichia of the dorsal fin are preserved in situ near the rostral end of P-77357 (Fig. 7). The fin rays are subdivided into short segments. The anal fin, although incomplete as well, seemingly inserts a few scale rows posterior to the dorsal fin. At least seven anal fin rays are discernible. Each fin ray is segmented into several short, close-set units, and some of the lepidotrichia bifurcate at least once.

The caudal fin is abbreviated-heterocercal, exhibiting a reduced, scaled body lobe (Fig. 7). The distal ends of the dorsal and ventral caudal lobes are not preserved. Most of the caudal fin web is not well visible, but the lepidotrichia are clearly segmented into short elements and at least the central fin rays are distally branched. The leading margin of the dorsal caudal fin ramus is preceded by one lanceolate scute, followed posteriorly by three unpaired basal fulcra and ~12 paired basal fulcra (Pattern II of Arratia, 2009). The first (anterior) unpaired basal fulcrum has a weakly concave anterior margin, whereas the two caudally following ones have distinctly concave anterior borders. Small fringing fulcra are observed along the leading margin of the ventral caudal lobe.

Remarks.—Preservation of NMMNH P-77357 is not sufficient for an attribution at low taxonomic rank, and it is thus left in open nomenclature.

Taxonomy and comparative anatomy

Birgeria Stensiö, 1919.—Birgeria is known from most marine Triassic fish localities, with up to 11 nominal species (Stensiö, 1919, 1921, 1932; Boni, 1937; Lehman, 1948, 1952; Nielsen, 1949; Savage and Large, 1966; Schwarz, 1970; Beltan, 1980; Bürgin and Furrer, 1992, 1993; Jin, 2001; Liu et al., 2006; Romano and Brinkmann, 2009; this study). Its occurrence in the Permian of Bolivia is questionable (Beltan et al., 1987; Cione et al., 2010). Contrary to Saurichthys (Romano et al., 2012), species richness of Birgeria remained relatively low and steady during the Triassic. Fossils of Birgeria are often (but not always) relatively rare within fish assemblages (e.g., Lombardo and Tintori, 2005; Scheyer et al., 2014), a fact that has been ascribed to the anatomically inferred offshore habitat of this taxon (e.g., Schwarz, 1970).

From the eastern Panthalassan rim, Birgeria has thus far been described from the Early Triassic of western Canada (Schaeffer and Mangus, 1976; Neuman, 2015), Greenland (Stensiö, 1932; Nielsen, 1949; Jessen, 1972; Ørvig, 1978; Bartsch, 1988), Spitsbergen (Stensiö, 1921, 1932; Scheyer et al., 2014), and Arctic Russia (Berg et al., 1964). The Canadian and Russian birgeriids are poorly known. Birgeria was also reported from Lower Triassic exposures near Bear Lake, Idaho, USA (Dunkle cited in Schaeffer and Mangus, 1976, p. 552), but this material has not been described in the literature and the repositories are unknown. Lastly, cranial remains from the Late Triassic of California described as Xenestes velox Jordan, 1907 have been reassigned to Birgeria by Stensiö (1932); however, this poorly known species needs revision. The specimens described herein represent the first fossil evidence for the occurrence of Birgeria in the Early Triassic of the western USA.

Birgeria encompasses species of large size that, together with some species of Saurichthys Agassiz, 1834, pertained to the actinopterygian apex predators of the Triassic (Lombardo and Tintori, 2005). One of the largest individuals of Early

Downloaded from https://www.cambridge.org/core. IP address: 54.70.40.11, on 31 May 2019 at 17:07:24, subject to the Cambridge Core terms of use, available at https://www.cambridge.org/core/terms. https://doi.org/10.1017/jpa.2017.36
Triassic age is PIMUZ A/I 4301 from Spitsbergen, a few fragments of which were illustrated by Scheyer et al. (2014). This specimen had an estimated total length of >2 m (skull without shoulder girdle length is ~35 cm). Specimens NMMNH P-66225 and NMMNH P-77117 from Nevada are both large individuals. With an estimated total length of 1.72–1.85 m, P-66225 had a size comparable to the birgeriids from the Smithian of Spitsbergen (Stensiö, 1921, 1932; Scheyer et al., 2014), the Middle Triassic of Monte San Giorgio and Besano (Swiss-Italian borderland; Schwarz 1970; Romano and Brinkmann, 2009), the Middle–Late Triassic of China (Jin, 2001; Liu et al., 2006; Sun et al., 2016), and the Late Triassic of Slovenia (Jurkovšek and Kolar-Jurkovšek, 1986), Italy (e.g., Boni, 1937), and Switzerland (Bürgin and Furrer, 1992). A clear evolutionary trend in body size is not apparent, except that very large birgeriids (>120 cm) are as yet unknown from strata older than Smithian.

As a member of the ‘Palaeopterygii,’ Birgeria mostly retained the plesiomorphic ‘paleoniscoid’ bauplan, being
characterized, for instance, by the possession of an extended, cleaver-shaped maxilla (reflecting the large gape size and the far forward location of the eyes), the arrangement of jaw teeth in parallel rows, the remote position of the dorsal fin, and the heterocercal caudal fin (e.g., Aldinger, 1931, 1937; Nielsen, 1949; Lehmán, 1952). Like many other early actinopterygians, Birgeria possesses a dermohyal (Romano and Brinkmann, 2009; this study), but the presence of this bone remained undetected in earlier descriptions. In B. americana n. sp., B. nielseni, B. aldingeri, and B. stensioei, the dorsal side of the dermohyal is marked by a longitudinal crest (Stensiö, 1932; Schwarz, 1970; Beltan, 1980; Romano and Brinkmann, 2009; this study).

Apart from the aforementioned plesiomorphic traits, Birgeria stands out by a set of derived features, such as the advanced reduction of the squamation, the presence of an unpaired rostrormaxilla, the posterior extension of the frontals medially separating the parietals, or the large internal lamina on the angular (e.g., Stensiö, 1921, 1932; Nielsen, 1949; Lehmán, 1952; Schwarz, 1970; Jin, 2001; Liu et al., 2006; Romano and Brinkmann, 2009). A further, autapomorphic condition is the arrangement of the elements of the opercular series. The gill cover of Birgeria consists of a low, anteroposteriorly elongated operculum, located dorsal to the maxillary blade, a narrow, vertically arranged suboperculum (sensu Romano and Brinkmann, 2009; i.e., first ray of the ‘suboperculum’ of Nielsen, 1949; Beltan, 1980), and a branchiostegal series that is divided into separate postmandibular and submandibular series (e.g., Nielsen, 1949; Lehmán, 1952; Beltan, 1980; Romano and Brinkmann, 2009). These highly specialized modifications are attributable to the extreme obliquity of the suspensorium (Aldinger, 1937; Schwarz, 1970), related with the tremendous enlargement of the gape size (Nielsen, 1949), pushing the jaw joint close to the pectoral girdle.

Interspecific variation within the opercular series concerns chiefly the number of postmandibular branchiostegal rays (sensu Romano and Brinkmann, 2009; i.e., second to last ray of the ‘suboperculum’ of Nielsen, 1949; Beltan, 1980). Whereas three to five postmandibular branchiostegals are developed in B. groenlandica (cf. Nielsen, 1949) and four to five in B. nielseni (cf. Beltan, 1980), none or at most one is usually preserved in B. stensioei (Romano and Brinkmann, 2009). In these species, all postmandibular branchiostegals are vertically arranged, like the suboperculum, whereas in B. americana n. sp. the posterior ones are caudally tending (Fig. 3). The opercular series of B. americana n. sp. also displays a rudimentary branchiostegal ray between the main postmandibular and the submandibular series, which has thus far not been observed in other species. In addition, an antoperculum is developed in the Nevada taxon, a bone that is present in several other early actinopterygians, but until now was unknown in Birgeria (e.g., Aldinger, 1937; Lehmán, 1952; Gardiner and Schaeffer, 1989). The gill cover of B. americana n. sp. is less reduced compared to other birgeriids.

Other interspecific differences within the skull concern, among others, the outlines of the parasphenoid and maxilla (Boni, 1937; Schwarz, 1970). Many Early Triassic birgeriids possess a maxilla with a relatively low, elongate postorbital blade (e.g., B. groenlandica, B. aldingeri, B. cf. aldingeri, and B. americana n. sp.). Furthermore, the anterior margin of the postorbital blade is much more oblique (e.g., B. groenlandica, B. aldingeri, B. cf. aldingeri, B. nielseni, and B. americana n. sp.) than, for instance, in the Middle Triassic B. mougoueti or B. stensioei (Agassiz, 1834–1843; Stensiö, 1919, 1932; Aldinger, 1931; Nielsen, 1949; Schwarz, 1970; Romano and Brinkmann, 2009; Fig. 3). Additionally, the dorsal and ventral margins of the postorbital blade are subparallel in Middle and Late Triassic species (B. mougoueti, B. stensioei, and B. acuminata) and some Early Triassic species (B. nielseni and Birgeria sp. from British Columbia; Lehmán, 1952; Schaeffer and Mangus, 1976), whereas in most Early Triassic forms they clearly converge rostrad (e.g., B. americana n. sp., B. groenlandica, B. aldingeri, B. cf. aldingeri, Birgeria sp. from Russia, and some specimens referred to B. nielseni; Stensiö, 1919, 1932; Boni, 1937; Nielsen, 1949; Lehmán, 1952; Berg et al., 1964; Schwarz, 1970; Beltan, 1980; Romano and Brinkmann, 2009; Fig. 3). In general, Griesbachian–Smithian birgeriids have low, elongate skulls, whereas Middle–Late Triassic species possess higher, shorter crania (see Schwarz, 1970). A reduction of the postorbital skull length has also been documented in Saurichthys during the Early–Middle Triassic transition (Mutter et al., 2008; Romano et al., 2012), being a possible case of parallelism between these two predatory actinopterygians.

Some comments are necessary concerning the number of tooth rows and the size distribution pattern of teeth on the maxilla and dentary. Stensiö (1921, 1932), Nielsen (1949), Lehmán (1952), Savage and Large (1966), and Bürgin and Furrer (1992) described two rows of teeth on the maxilla and/or dentary of B. groenlandica, B. nielseni, B. aldingeri, B. cf. aldingeri, and B. acuminata, whereas Schwarz (1970) observed three rows in B. stensioei. Three discrete rows of teeth are also developed in B. americana n. sp. (Fig. 4) and in the poorly known Middle–Late Triassic B.? costata (Münster, 1839), whose generic attribution has been questioned (Boni, 1937; Bürgin and Furrer, 1993). Based on unpublished computer tomography generated images of the holotype of B. groenlandica (ZMUC VP 3176; personal communication to CR, T. Argyriou, 2016), it is evident that most of the large lingual teeth are not exposed on the surface, but two rows of teeth are confirmed here. Regarding B. aldingeri, only the lateral imprints of the intermediate and labial teeth of the maxilla are visible, with the internal lamina and lingual teeth not preserved (personal observation, C. Romano, 2016; Fig. 8; Stensiö, 1932), thus three rather than two rows are developed in this species.

Notably, only some Early Triassic birgeriids (B. aldingeri and B. americana n. sp.) possess intermediate teeth that are variable in size but predominantly high, and chiefly widely spaced, whereas in the Middle Triassic B. stensioei and the Late Triassic B. acuminata and Birgeria sp. they are close-set and of relatively small, uniform size throughout the length of the jaw (Woodward, 1889; Boni, 1937; Savage and Large, 1966; Schwarz, 1970; Bürgin and Furrer, 1992; Lombardo and Tintori, 2005; Fig. 8). The lateral teeth are small but distinct in B. aldingeri and B. americana n. sp., whereas in stratigraphically younger taxa they are either very small or absent (Fig. 8), with the exception of the problematic B.? costata (Bürgin and Furrer, 1993). Ørvig (1978), confirming observations by previous authors (e.g., Stensiö, 1919), showed that the
odontodes on the external side of the jaw bones become larger and tooth-like in vicinity to the marginal teeth, exhibiting both an acrodin cap and a pulp cavity. It is conceivable that the labial teeth developed from such odontodes.

The present paper provides additional evidence that Birgeria encompasses both species with three rows of teeth and species with two rows of teeth along the oral margins of the maxilla and dentary. Based on the current state of knowledge, it appears that birgeriids with three well-developed tooth rows as well as widely spaced intermediate teeth with varying height are restricted to the Early Triassic. On the other hand, species with only two principal rows of teeth and small, close-set, equal-sized intermediate teeth occur throughout the Triassic. Whether the reduction of the intermediate and labial tooth rows on the maxilla and dentary represents an evolutionary trend in Birgeria requires further comparative studies.

Compared to the intermediate teeth, the size distribution pattern of the lingual teeth is seemingly more conservative between species. The same size distribution pattern is found in Birgeria americana n. sp., B. stensioei, and B. acuminata (Aldinger, 1931; Boni, 1937; Schwarz, 1970; Romano and Brinkmann, 2009; Figs. 3, 4). In both B. stensioei and B. acuminata, the lingual teeth in the caudal part of the dentary are curved towards the anterior (Boni, 1937; Bürgin and Furrer, 1992; Romano and...
although Nielsen (1949) noted that the lingual teeth of B. stensioei are more widely spaced than in B. groenlandica. Although this is true for some specimens of B. stensioei (e.g., the lectotype), the teeth are more densely distributed in other specimens (Aldinger, 1931; Schwarz, 1970; Fig. 8). Tooth loss cannot account for such regular distribution patterns, but more studies are necessary to better assess the taxonomic value of tooth spacing.

Previous workers (Stensiö, 1932; Lehman, 1952; Schwarz, 1970) suggested interspecific variation in the ornamentation of the acrodin cap, ranging from completely smooth, to only basally striated, to fully striated. Nevertheless, the taxonomic value of tooth ornamentation is doubtful, as this character also shows intraspecific variability. For instance, in B. stensioei the acrodin cap is fully striated in PIMUZ T 4780 (Schwarz, 1970), but only basally striated in PIMUZ T 1 (Fig. 8). Nielsen (1949) also mentions variability concerning this character in B. groenlandica. Crown ornamentation is also subjected to wear.

The dentitions of the rostrum of the maxilla, maxilla, and dentary are supplemented by at least one row of macroscopic teeth on the prearticular, coroid, ectopterygoid, and dermalpalatine, respectively, and myriads of minute teeth also cover the lingual surfaces of the prearticular, coroid, ectopterygoid, endopterygoid, parasphenoid, and the bones of the branchial arches (Stensiö, 1921; Nielsen, 1949; Lehman, 1952; Bürgin and Furrer, 1992; Romano and Brinkmann, 2009; this study). All species are characterized by a strong dentition, which together with the weakly developed bones of the operculogular series, suggest that Birgeria was a ram feeder (contra Lombardo and Tintori, 2013), meaning that prey was chased and bitten rather than engulfed through current action (Schaeffer and Rosen, 1961). The coroid process on the mandible—developed in convergence to holosteans—reduced torque on the jaw joint (Schaeffer and Rosen, 1961). Birgeria is often allied with saurichthyids and the Acipenseriformes (sturgeons and paddlefish), even though they share only a few characters (e.g., reduced squamation, posterior elongation of the parasphenoid; Bemis et al., 1997), some of which are also present in other actinopterygians. The suggested close affiliation goes mainly back to the ‘Stockholm school’ (Schultze, 2009), whose influential works repeatedly highlighted similarities between Birgeria and Acipenseriformes, some of which were later called into question (e.g., nerve sac groups, Órvig, 1978). Jessen (1972) also doubted a close affinity due to differences in the pectoral girdle skeleton, and according to Coates’ (1999) cladistic analyses, Birgeria is resolved as closely related to Acipenser only if endocranial characters are omitted. A close relationship among Birgeria, Saurichthys, and Acipenseriformes was, nonetheless, recovered in the cladistic analysis of Gardiner et al. (2005).

Nielson (1949), among others, compared the peculiar ray-like elements posteroventral to the operculum of B. groenlandica with the lobate suboperculum of Polyodon. However, as pointed out by Romano and Brinkmann (2009), the anteriormost subopercular ray of Nielsen (1949) borders on the operculum, and the same condition is also seen in B. nielseni, B. stensioei, and B. americana n. sp. (Beltan, 1980; Romano and Brinkmann, 2009; Fig. 3). Although Nielsen (1949) stated that the rays are proximally fused in B. groenlandica, like in Polyodon (e.g., Bemis et al., 1997), fusion is not evident in our material. The present study supports the view that the ‘suboperculum’ of Nielsen (1949) is a composite element and that it is not homologous with the suboperculum of the American paddlefish. The homology of the slender suboperculum of Birgeria with the suboperculum of other actinopterygians requires further study; a homology with the ‘accessory operculum’ of early ray-fins (e.g., Cheirolepis) would also be possible.

Saurichthys Agassiz, 1834.—Saurichthys is known from Triassic sites around the world, both marine and freshwater, and with over forty named species (Kogan and Romano, 2016a and references therein), it is much more speciose than Birgeria. After its first appearance in the latest Permian, Saurichthys rapidly reached global distribution and high species richness during the Early–Middle Triassic, but later became less diverse and geographically more restricted (Mutter et al., 2008; Romano et al., 2012). Within the United States, Saurichthys has previously been described from the early late Smithian Anasibirites beds west of Georgetown, Bear Lake County, Idaho (Romano et al., 2012; a second skull, PIMUZ A/I 4621, was found in 2013, and a cranial fragment, NMMNH P-77359, in 2015 by JJ). Saurichthys was also described from the Middle Triassic of Pershing County, Nevada (Sander et al., 1994; Rieppel et al., 1996).

Three-dimensionally preserved skulls or skull fragments of Saurichthys (preserved as either body fossils or external molds) are frequently found in strata of Early Triassic age (Stensiö, 1925; Lehman, 1952; Schaeffer and Mangus, 1976; Beltan and Janvier, 1978; Minikh, 1981, 1982; Mutter et al., 2008; Romano et al., 2012; Kogan and Romano, 2016a; this study) or Middle Triassic, but later became less diverse and geographically more restricted (Mutter et al., 2008; Romano et al., 2012; Kogan and Romano, 2016a; this study) or Middle Triassic age (Frech, 1903–1908; Hennig, 1909; Beltan et al., 1979; Rieppel, 1985; Wu et al., 2015), but are seemingly rare in younger deposits. Although saurichthyid crania contain some diagnostic features (e.g., Romano et al., 2012; Werneburg et al., 2014), species are predominantly differentiated by postcranial characters (e.g., squamation or fin segmentation pattern, morphology of the axial skeleton; e.g., Rieppel, 1985; Mutter et al., 2008; Tintori, 2013; Tintori et al., 2014b; Maxwell et al., 2015; Kogan and Romano, 2016a, b). Moreover, comparisons of dermal skull bone patterns between species are complicated due to the fusion of bones during ontogeny. However, PIMUZ A/I 4397 from the latest Smithian of Palomino Ridge shows some features useful for comparison, such as the elongate rostrum, the anterior extension of the lachrymal, or the distinct lateral flange of the dermopterotic.

In addition to Saurichthys sp. from Palomino Ridge, a dorsoventrally and anteroposteriorly expanded lateral flange on the dermopterotic is found in S. toxolepis Mutter, Cartanyà, and Basaraba, 2008 from the Early Triassic of Canada, S. obruchevi Minikh, 1981 from the Early Triassic of Russia, and some specimens ascribed to S. wimani Woodward, 1912, S. ornatus Stensiö, 1925, and S. elongatus Stensiö, 1925 from the Smithian of Spitsbergen. The same condition is possibly also seen in S. cf. elongatus from the early late Smithian of Idaho (Romano et al., 2012). In contrast, the lateral flange of the dermopterotic is more reduced in other Early Triassic species, such as S. madagascariensis Piveteau, 1945, S. cf. ornatus from Greenland, and S. wimani (Stensiö, 1925;
In addition to PIMUZ A/I 4397, S. cf. elongatus, S. ornatus, and S. toxolepis the lachrymal reaches anteriorly beyond the level of the posterior external naris, whereas in S. cf. ornatus and S. madagascariensis the lachrymal extends less far rostrally (e.g., Stensiö, 1925; Mutter et al., 2008; Romano et al., 2012; Kogan and Romano, 2016a). The significance of this character requires further study.

The rostrum is often incompletely preserved in Saurichthys. In addition to PIMUZ A/I 4397, largely complete, elongate rostra are preserved, for instance, in specimens ascribed to S. toxolepis, S. elongatus, and S. curionii (Stensiö, 1925; Rieppel, 1985; Mutter et al., 2008), as well as in an undescribed specimen from Idaho (PIMUZ A/I 4621).

**Early Triassic vertebrate occurrences in the western USA**

Localities yielding Early Triassic fishes are known from Greenland, Canada, and the USA (Schaeffer and Mangus, 1976; Wilson and Bruner, 2004; Brinkmann et al., 2010). Early Triassic paleichthysological sites are absent in South America (López-Arbarello, 2004; Brinkmann et al., 2010) and, to our knowledge, there are no such localities in Central America and the Caribbean Islands. Consequently, our knowledge of Early Triassic fishes from the former western Pangean margin relies mainly on sites that were then located at mid or high latitudes in the northern hemisphere (i.e., localities in present-day northern North America and Arctic Eurasia).

Marine Early Triassic fishes from Greenland are derived from several horizons, all of earliest Triassic age (Griesbachian–earthy Dienerian; Perch-Nielsen et al., 1974). They have been extensively researched especially during the twentieth century (e.g., Stensiö, 1932; Nielsen, 1949; Mutter et al., 2008; Kogan, 2011). From Canada, Early Triassic fishes have been reported from sites in western Alberta (Neuman, 2015 and references therein), Ellesmere Island (Nunavut; Schaeffer and Mangus, 1976), and Wapiti Lake area, British Columbia. Fishes from the latter locality in particular have been the focus of several recent studies (e.g., Mutter and Neuman, 2006; Mutter et al., 2008; Wendruff and Wilson, 2012, 2013). The western Canadian fish fossils come from various layers, but age constraints are limited (e.g., Mutter and Neuman, 2006; Neuman, 2015). In addition to the aforementioned Canadian sites, Wignall and Newton (2003) reported the occurrence of well-preserved remains of cf. Bobasatrania in the upper Grayling Formation (Dienerian?) of Ursula Creek, northwest of Wapiti Lake.

In contrast to Canada and Greenland, only a few, short papers were dedicated to the fishes from the Early Triassic of the United States. Exposures of Lower Triassic, marine rocks, which are found in Alaska, Montana, Wyoming, Idaho, Utah, Nevada, and California, have yielded a plethora of marine invertebrate fossils, including the earliest metazoan reefs of the Mesozoic (e.g., Silberling and Tozer, 1968; Nichols and Silberling, 1979; Brayard et al., 2011, 2013, 2017; Hofmann et al., 2013a, b). Several formations produce Early Triassic vertebrate remains, but these occurrences have often only been casually mentioned (e.g., Oversby, 1972; Silberling, 1973; Schaeffer and Mangus, 1976; Poole and Wardlaw, 1978). On the other hand, the marine and freshwater fishes from the Middle Triassic of the USA have received slightly more research interest (e.g., Wemple, 1906; Wells, 1947; Schaeffer and Gregory, 1961; Sander et al., 1994; Rieppel et al., 1996; Cuny et al., 2001).

One of the longest known collecting areas for Early Triassic marine vertebrate fossils is southeast Idaho, yielding ichthyloliths (Evans, 1904; Goddard, 1907; Mutter and Rieber, 2005; Romano et al., 2012; Brayard et al., 2017), but also articulated fishes (Tanner, 1936; Romano et al., 2012), and some ichthyopterygian reptile remains (Massare and Callaway, 1994; Scheyer et al., 2014). In addition to these published specimens, new vertebrate material (isolated bones or articulated partial/complete skeletons) was recently discovered at several sites in Idaho, for instance in the middle Smithian ‘Meekoceras beds’ (personal observation, CR, JJ, KGB, HB, 2013–2016), the late Smithian Anasibirites beds (Saurichthys: PIMUZ A/I 4621, NMMNH P-77359; several unlabelled specimens of Actinopterygii indet.), and the early Spathian Tirolites beds (personal communication, A. Brayard, 2016; personal observation, JJ, KGB, HB, 2013–2016).

Very little is known about marine vertebrate occurrences in Lower Triassic formations in the other western states. Aside from the Smithian and Spathian fish material from Nevada described herein, some ichthyopterygian remains from Spathian layers were documented by Mazin and Bucher (1987) and Kelley et al. (2016). Isolated vertebrate remains have also been found in the Smithian ‘Meekoceras beds’ at Crittenden Springs as well as in Spathian strata at several sites in Nevada (personal observation, JJ, RJ, TMS, KGB, HB, 2010–2016). Articulated osteichthyan fossils of Dienerian age have recently been discovered in the Candelaria Hills in southwest Nevada (e.g., Brinkmann et al., 2010; Romano et al., in preparation). From Wyoming, Case (1936) described a sauroptrypgian reptile of Spathian age (Lovelace and Doebert, 2015; Kelley et al., 2016).

Most vertebrates from the Early Triassic of the Western United States are Smithian or Spathian in age, whereas Griesbachian and Dienerian fossils are less well known. The material from Nevada presented herein is derived from equatorial shelfal series deposited on the North American Craton. Material from sites located farther to the East (Utah, Idaho) was deposited in a low-latitude, epicontinental sea (western USA basin; Brayard et al., 2013). The depth and areal extent of this Early Triassic sea were controlled by both regional tectonics and eustatic sea level changes (e.g., Paull and Paull, 1993; Brayard et al., 2013).

In conclusion, several localities in the western USA yield Early Triassic vertebrate remains, but for the most part these fossils received little interest by researchers. The specimens described herein, in combination with other published and unpublished material from the Smithian of the western USA, contradict Sun et al.’s (2012) notion of a near absence of vertebrates within low latitudes due to ‘lethally hot’ temperatures during this interval. This alleged ‘equatorial vertebrate eclipse’ largely ignored the contradictory data available from China (e.g., Tong et al., 2006), and was also based on a wrong conodont age (Goudemand et al., 2013). Sun et al. (2012)
further inferred too high sea surface temperatures (~38°C, possibly over 40°C), because the presence of fishes in the USA and China suggests that a 36°C physiological limit was not crossed (Motani and Wainwright, 2015).

Spathian Osteichthyes

Of interest is the presence of vertebrate fossils in Spathian strata in the western USA. The Spathian had a duration of ~3 myr, therefore longer than the in the western USA. The Spathian had a duration of ~3 myr, the first three stages of the Early Triassic combined (~2 myr; Ovtcharova et al., 2006, 2015; Galfetti et al., 2007a; Fig. 8). However, only a few, well-dated Spathian sites yield articulated osteichthyan fossils (Romano et al., 2016a).

Apart from the new material from Crittenden Springs (NMMNH P-77357, PIMUZ A/I 4641), articulated actinopterygian remains (a perleidid and a saurichthyid) have been described from Spathian deposits of Anhui Province, China (Sun et al., 2013; Tintori et al., 2014b). In addition, largely complete marine osteichthyan fossils have also been documented from beds close to the Smithian–Spathian boundary at localities in Anhui, Hubei and Jiangsu provinces, China (e.g., Tong et al., 2006 and references therein). A saurichthyid (Saurichthys elongatus?) with a short postorbital skull portion was found slightly above the Smithian ‘Fischniveau’ of Spitsbergen, thus being of possible Spathian age (Stensiö, 1925; Kogan and Romanov, 2016b). Two saurichthyids and a colobo-dontid were found in the Gogolin beds of Upper Silesia, Poland, whose age is close to the Spathian-Anisian boundary (e.g., Eck, 1865; Frech, 1903–1908; Nawrocki and Szulc, 2000; Szulc and Becker, 2007). Aldinger (1931) mentioned undescribed actinistian material from Gogolin. Marine and freshwater osteichthyan remains of presumable Spathian age have been described from localities in Canada, Russia, Kazakhstan, northern China, and South Africa (e.g., Jubb and Gardiner, 1975; Minikh, 1981; Mutter and Neuman, 2006), although often with limited biocrustonotigraphic constraints. Besides, there are several Spathian occurrences of isolated fish remains (e.g., Stensiö, 1921; Mutter and Rieber, 2005; Kogan and Romanov, 2016b; Brayard et al., 2017), but such material often provides little taxonomic information.

Griesbachian–Smithian fishes differ notably from post-Spathian ones. Earliest Triassic sites are characterized by cosmopolitan taxa, whereas Middle Triassic assemblages comprise many endemic taxa, with a distinct neopterygian component (Tintori et al., 2014a; Romano et al., 2016a). This suggests an ichthyofaunal turnover either during the Spathian or in conjunction with the end-Smithian event. This extinction event decimated nektobenthic clades such as ammonoids and conodonts (Galfetti et al., 2007b; Orchard, 2007) and saw the replacement of treme-tosauroid ‘amphibians’ by reptiles among the marine tetrapod apex predators (Scheyer et al., 2014). However, the impacts of the end-Smithian event on fishes remain elusive, particularly with regard to the timing of the radiation of the ‘Triassic Middle Fish Fauna’ of Tintori et al. (2014a). The sparse Spathian actinopterygians known thus far show more affinities with Middle Triassic taxa than with Griesbachian–Smithian ones (Sun et al., 2013; Tintori et al., 2014b). There is a need for more research on Spathian Osteichthyes, emphasizing the importance of the discovery of articulated osteichthyan fossils in the Spathian of Nevada and Idaho (Brayard et al., 2017; this study).

Conclusions

The presented material from three new sites in Elko County (Nevada, USA) notably improves the fossil record of Early Triassic fishes from the western USA. Although Early Triassic fish remains occur in several places in the United States, most of them have not been studied. We introduce a new Spathian fish occurrence. Spathian Osteichthyes are preserved in several sites in the western USA and their study may provide vital clues concerning the post-Paleozoic evolution of fishes.

The Smithian material studied here includes two large-sized individuals of the predatory actinopterygian Birgeria, providing the first fossil evidence that birgeriids expanded their distribution into the western USA Sea, even during the Smithian thermal maximum (Goudemand et al., 2013; Romano et al., 2013). One specimen can be ascribed to a new species, Birgeria americana n. sp., which is characterized by three rows of teeth on the maxilla and dentary, and a less reduced opercular series compared to other taxa. We further describe material of the ambush predator Saurichthys that, together with other late Smithian occurrences in Idaho (PIMUZ A/I 3900, PIMUZ A/I 4621, NMMNH P-77359), suggests that this taxon was relatively common in the western USA basin. The presented fishes from low paleolatitudes, along with other Smithian vertebrate occurrences in the United States and South China (Tong et al., 2006), invalidate the alleged ‘equatorial vertebrate eclipse’ of Sun et al. (2012). These low-latitude occurrences imply that a 36°C temperature tolerance was not permanently exceeded during the Smithian thermal maximum (Motani and Wainwright, 2015). The presence of the actinopterygian top predators Birgeria and Saurichthys also contradicts anew the claim of missing apex predators and truncated food chains during the Early Triassic made by Chen and Benton (2012).

Acknowledgments

We appreciate information from and insightful discussions with T. Argyriou, M. Leu, D. Ware, T. Brühwiler, M. Brosse, C. Klug, M. Hautmann, W. Brinkmann (all PIMUZ), I. Kogan (Technische Universität Bergakademie Freiberg, Germany), P. Skrzyczyk and R. Skrzycza (Krakow, Poland), A. Brayard (Université de Bourgogne, Dijon, France), R. Hofmann (Museum für Naturkunde Berlin), and A. Tintori (MPUM). E.E. Maxwell (Staatliches Museum für Naturkunde Stuttgart, Germany) and the editor are thanked for comments that improved this paper. I. Kogan and A. Tintori are also thanked for photographs. M. Leu, C. Kolb, and F. Blattmann (all PIMUZ) are thanked for lab support. CR thanks M. Hebeisen, R. Roth, and T. Argyriou for advice during preparation of fossils, and M. Véran, G. Clément (both MNHN.F), and J.O. Ebbestad (Bureau of Land Management Nevada) for swift administrative processing. We deeply acknowledge past and present support by


Accepted 4 May 2017