Olenekian (Early Triassic) Fossil Assemblage from Eastern Julian Alps (Slovenia)

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Abstract: New palaeontological and sedimentological data from the Lower Triassic strata of the eastern Julian Alps in Slovenia are presented. They are unusual for the Early Triassic of the Alps in representing a relatively deeper, unrestricted marine (mid-ramp) setting. There are two basic microfacies types in the section investigated (types A and B), which are organized as couplets with coarse-grained tempestitic deposits (microfacies A), overlain by laminated or bioturbated lime mudstones and/or marls (microfacies B), frequently containing ammonoids. This pattern is interpreted as storm deposition with occasional winnowing of bottom sediments and the formation of coarse-grained skeletal deposits (lags), followed by the slow settling of suspended particles, when the storm waned, in addition to background deposition. Dominantly lime mud deposition and the presence of ammonoids indicate deposition on a more distal, deeper ramp with an unrestricted connection to the open sea. Intense reworking of bottom skeletal-rich sediment and accumulation of storm lags suggest deposition above the storm wave base, possibly in a wide low-energy mid-ramp environment. Faunas from such settings have been reported relatively rarely from the Early Triassic of the Alps. The macrofauna contains ammonoids, bivalves and gastropods, whereas the microfauna is represented by foraminifer tests and conodont elements; rare fish remains also occur. In the foraminifer assemblages, species of Ammodiscus, Hoyenella, Glomospirella dominated, corresponding to the widespread “Glomospira-Glomospirella” foraminifer community, with some miliolids and nodosariids. The conodont fauna is characterized by Triassosphathodus hungaricus (Kozur et Mostler), indicating an early Spathian (Olenekian) age. The fossil assemblage highlights the wide distribution of Early Triassic taxa in the Tethys and facilitates its worldwide correlation. Its relatively low diversity by comparison with shallow marine settings is interpreted as an evolutionary proximal-distal trend in the wake of the end-Permian mass extinction. Re-diversification first occurred in nearshore settings and expanded into deeper/distal marine environments through geological time.

Key words: Olenekian, Early Triassic, foraminifers, bivalves, gastropods, conodonts, sedimentology, Julian Alps, Slovenia.

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INTRODUCTION

The end-Permian mass extinction event was pivotal in the Phanerozoic history of marine life, terminating the incumbency of the “Palaeozoic Evolutionary Fauna” that had dominated life in the sea for nearly 250 million years (Sepkoski, 1984). In the wake of the extinction event, the ecological structure of the present-day biosphere evolved from the stock of the surviving taxa, shaping the “Modern Evolutionary Fauna” that characterizes marine ecosystems until today (Sepkoski, 1997). The recovery from the end-Permian mass extinction is therefore crucial for understanding
how the present-day biosphere evolved. An indispensable prerequisite in this context is a census of the faunas that lived and evolved during the Early Triassic, i.e., the epoch that followed the crises. Thus, collecting and interpreting new field data from the Early Triassic has been a major research agenda during the past decade (e.g., Brayard and Bucher, 2008; Posenato, 2008; Kumagae and Nakazawa, 2009; Kaim et al., 2010, 2013; Hautmann et al., 2011, 2013; Brühwiler et al., 2012; Wasmer et al., 2012; Hofmann et al., 2013a, b). However, Early Triassic benthic faunas have been described mostly from relatively shallow marine sections, particularly in Europe, where deeper shelf settings are rare. The section investigated is also known as the location, where an isolated temnospondyl bone (Lucas et al., 2008), was found and therefore represents an important site of Slovenian natural heritage. In this study, new sedimentological and palaeontological data are presented from the Spithian (late Early Triassic) section in the Julian Alps (Slovenia). They contribute to a better understanding of how diversity and faunal composition changed along depth gradients during re-diversification from the greatest crisis in the history of life.

**GEOLOGICAL SETTING**

The study area is in the Julian Alps, NW Slovenia (Fig. 1). Structurally they form the easternmost continuation of the Southern Alps, where the South Alpine and the Dinaric structures now overlap (Placer, 1999). The largest overthrust unit in the Slovenian part of the Julian Alps is represented by the Julian Alps Overthrust (Jurkovšek, 1987a, b) or the Julian Nappe, which is composed of successions, ranging from Early Triassic to Cretaceous strata, though a major part of it is made up of Late Triassic carbonates. Early Triassic strata are scattered in narrow disconnected belts or patches in the wider Julian Alps area. They usually occur within smaller tectonic slices (Figs 2, 3).

The area investigated is located in the uppermost tectonic slice of the Julian Alps Overthrust, SE of Triglav Mt. (2863 m), the highest peak in Slovenia. The lower part of these strata is in tectonic contact with the Norian–Rhaetian Dachstein Limestone, whereas upwards they pass continuously into Anisian limestones and dolomites (Jurkovšek, 1987a; Celarc and Kolar-Jurkovšek, 2008).

The Studorski preval section is strongly tectonized (coordinates: N46°21'14.46'', E13°52'55.80''; WGS 84). Thick lime mudstones and marls and thin beds enriched in coarse-grained bioclastic material, predominate. The fauna is characterized by Early Triassic mollusks, dominated by bivalves and gastropods. The microfauna is diversified, but consists mainly of foraminifer assemblages in association with rare conodont elements and fish remains.

**Lithology**

The Studorski preval section mainly consists of thick mudstones (Fig. 4A) and marls (Fig. 4B), and thin beds, enriched in coarse-grained bioclastic material. These lithologies form units, in which coarse-grained bioclastic material...
is concentrated at the base and gradually passes upward into mudstones/marl. Thus, the microfacies in the Studorski
preval section correspond to two microfacies types: micro-
facies type A, fossiliferous floatstone/packstones, and mi-
icrofacies type B, dense or laminated lime mudstones that
may pass gradually into marl.

Centimetre- to decimetre-thick fossiliferous floatstone/
packstone beds (A) have distinctive sharp lower erosional
boundaries. The floatstone/packstone interbeds gradatio-
nally change upward to much thicker mudstone/marl inter-
vals, which locally were intensively reworked by organisms
(Fig. 4C).

Floatstones/packstones of microfacies type A consist of
coarse (usually > 2 mm) well preserved skeletal detritus
(Fig. 4F). Among the skeletal fragments bivalves, gastro-
pod, and ammonoid detritus predominate. Echinoderm ossi-
cles and foraminifer tests are rare. In the floatstone/pack-
stone a bimodal sorted biofabric is found, which consists of
large bioclasts and micritic matrix. Very frequently, the ma-
trix is non-homogenous and contains a high proportion of
peloidal particles and some micrite (Fig. 4E). Whole valves
are dominantly oriented parallel to the bedding surface, very
often in a convex-up position. Completely preserved bi-
valve or gastropod shells are geopetally infilled with sparry
calcite. Some shelter pores, at the concave side of bivalve
shells, are also infilled by sparite.

Microfacies type B consists of dense, laminated or bio-
turbated mudstones and/or marls. Sparse, well preserved fos-
sils can be dispersed in the mudstones (Fig. 4D). The mud-
stones consist of homogenous micritic mud or micrite-rich
laminae. Disturbance of the lamination is due to reworking
by infaunal organisms. Lime mud is slightly recrystallized. In
both microfacies types euhedral pyrite crystals and large ce-
lestite crystals (determined by EDS analysis) are present.

MATERIAL AND METHODS

The present study is based on field work, carried out in
2011 at Studorski preval in the Julian Alps. Approximately
10 m of the Lower Triassic strata was sampled just below
the contact with the Anisian dolomite. Nine samples (SP
1–9) were collected for examination from the Lower Trias-
sic strata; sample SP 1 is the lowest and the sample SP 9 is
the highest. Rock samples with an average weight of 4 kg
were processed for conodont study, using standard labora-
tory techniques. Several thin sections were made for the
study of foraminifers as well as for petrographic purposes.
Laboratory preparation was carried out at the Geological
Survey of Slovenia (Geološki zavod Slovenije) where all of
the micropalaeontological material is stored and inventoried
under repository numbers 4429, 4548–4555, 4901–4904
and abbreviated GeoZS. The collection of macrofossil spec-
imens: Jurkovšek Paleontological Collection, Kamnica 27,
Dol pri Ljubljani, Slovenia, has been registered with the
Natural History Museum of Slovenia, Ljubljana and abbre-
viated as BJ. The determinations of conodont elements pre-
sented here and the SEM/EDS analyses were carried out,
using the JEOL JSM 6490LV Scanning Electron Micro-
scope at the Geological Survey of Slovenia.

Fig. 2. Generalized stratigraphic section of Triassic strata in
Julian Alps. Thick dark bar indicates section studied in tectonic
slice at Studorski preval

Fig. 3. Palaeogeographic map for Early Triassic with position
of the Julian Alps marked (star), modified from Scotese (2001)
Fig. 4. Microfacies types from Early Triassic of Studorski preval, Slovenia. Hammer length 26.5 cm (A) and 29 cm (B, C). A. Laminated, thin-bedded mudstones. B. Homogenous marl interval. C. Bioturbated lime mudstone. D. Rare skeletal fragments in dominantly muddy sediment – microfacies type B. E. Floatstone (microfacies type A) with coarse skeletal detritus and inhomogeneous matrix containing high proportion of pelloidal particles and some micrite. F. Floatstone of microfacies type A consists predominantly of bivalves and gastropod detritus. Note convex-up position of most valves.
PALAEOENVIRONMENT

The Studorski preval section can be compared to the Werfen Formation of the Southern Alps that represents deposition on a storm-influenced ramp (Brandner et al., 2012). Sedimentation of the dominantly fine-grained microfacies type B can be interpreted as slow deposition in a deeper marine environment below the fair-weather wave base, which corresponds to the distal part of a ramp. The presence of ammonoid shells indicates a good connection with the open unconfined sea and also implies deposition in a deeper-ramp environment. Nevertheless, the interlayering with limestone of microfacies type A suggests that slow deposition of fine-grained sediments (microfacies type B) was periodically interrupted by the deposition of coarse skeletal detritus during high-energy events (microfacies type A). The sharp and erosive lower boundaries of microfacies type A beds imply the abrupt commencement of sedimentation. The well-preserved skeletal detritus in the floatstones/packstones and the upward-increasing proportion of matrix indicate deposition during storms, in a zone above the storm-weather wave base. As the beds of microfacies type A are overlain by laminated or bioturbated calcareous mudstones and/or marls (microfacies type B), their deposition reflects short storm events in a deeper low-energy ramp environment, where the deposition of lime mud prevails. The intense bottom-shear conditions during the storm peak concentrated shells of living and dead organisms from the sea bottom, either by burying them under a sudden influx of storm-suspended particles or by exhuminning previously buried shells by rippling up the underlying weakly consolidated sediments and forming skeletal concentrations (lag deposits). Therefore, well-preserved fossils within a fine micritic/peloidal matrix of microfacies type A are interpreted as the result of storm events, which preserve whole fossils by burying and protecting them from normal destructive processes. The presence of foraminifera indicates detritus derived by storms from the shallow proximal ramp. Valves oriented parallel to the bedding plane in a convex-up position also indicate deposition under short-term high-energy conditions. Furthermore, winnowing and suspension of fine sediment resulted in the deposition of the upward-finishing units above coarse lag deposits. As the storm waned, more mud was deposited from suspended material, resulting in fining-upward grading. Intense colonization by organisms and bioturbation could reflect the cessation of the storm and a return to normal low-energy ramp sedimentation below the fair-weather wave base. The presence of pyrite also indicates low-energy, and possibly partly anaerobic conditions, whereas the genesis of celestite is considered as diagenetic in deep-sea sediments, as explained by Baker and Bloomer (1988) although it also has been described from shallow marine settings (Hautmann, 1997).

Storms are frequently recorded both in the inner and outer shelf/ramp in the Induan/Olenekian sediments of the wider Dinaric region (Aljinović, 1995; Aljinović et al., 2006, 2011), while this investigation points to deposition in wide mid-ramp zone (the zone between the fair- and storm-weather wave bases), as defined by Burchette and Wright (1992).

There are only a few outcrops of Lower Triassic strata in the Julian Alps and they occur in small tectonically confined areas (e.g., Mavric, Lipanca). There is some similarity in their development and macrofossil content (Natiria costata and badly-preserved ammonoids). Conodont and other micro-paleontologic analyses were negative and therefore a detailed geological comparison of these strata had not been possible.

PALAEOONTOLOGY

Microfauna

The microfossil material recovered from nine conodont samples (SP 1–9) was examined in the present study. The associations consisted of conodont elements, rare fish remains, as well as some recrystallized free specimens of foraminifera from two samples (SP 1 and SP 5). In addition, several thin sections were prepared from each sample for foraminifera study. The conodonts recovered are white with CAI = 1 (Epstein et al., 1977).

The list of microfossils determined is presented in Table 1.

Foraminifera

The foraminifer classifications of Loeblich and Tappan (1988) and Mikhailovich (2000) are combined and used here. The foraminifer descriptions are short, because these foraminifers are mainly well-known and the preservation of these forms is not particularly good.

Genus Ammodiscus Reuss, 1862

Type species: Ammodiscus infimus Bornemann, 1874

Ammodiscus? parapriscus Ho, 1959

Fig. 5A–D

*1959 Ammodiscus parapriscus n. sp. – Ho: p. 408, pl. 2, figs 3–6.

Table 1

Distribution of microfossils in samples from Studorski preval

<table>
<thead>
<tr>
<th>Taxa</th>
<th>Sample</th>
</tr>
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<tbody>
<tr>
<td>Ammodiscus* parapriscus Ho</td>
<td>SP1</td>
</tr>
<tr>
<td>Ammodiscus* parapriscus Ho</td>
<td>SP2</td>
</tr>
<tr>
<td>Ammodiscus* parapriscus Ho</td>
<td>SP3</td>
</tr>
<tr>
<td>Ammodiscus* parapriscus Ho</td>
<td>SP4</td>
</tr>
<tr>
<td>Ammodiscus* parapriscus Ho</td>
<td>SP5</td>
</tr>
<tr>
<td>Ammodiscus* parapriscus Ho</td>
<td>SP6</td>
</tr>
<tr>
<td>Ammodiscus* parapriscus Ho</td>
<td>SP7</td>
</tr>
<tr>
<td>Ammodiscus* parapriscus Ho</td>
<td>SP8</td>
</tr>
<tr>
<td>Ammodiscus* parapriscus Ho</td>
<td>SP9</td>
</tr>
<tr>
<td>Glomospirella facilis Ho</td>
<td></td>
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<tr>
<td>Glomospirella shengi Ho</td>
<td></td>
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<tr>
<td>Glomospirella sp.</td>
<td></td>
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<tr>
<td>Pilammina? cl. praecdensa</td>
<td></td>
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<tr>
<td>Urosevic</td>
<td></td>
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<tr>
<td>Arenovidalina? sp.</td>
<td></td>
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<tr>
<td>Meandroidira? sp.</td>
<td></td>
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<tr>
<td>Agathammina? sp.</td>
<td></td>
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<tr>
<td>Hayennella sinensis (Ho)</td>
<td></td>
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<tr>
<td>Dentalina splendida Schleifer</td>
<td></td>
</tr>
<tr>
<td>Nodosaria? spp.</td>
<td></td>
</tr>
<tr>
<td>Triassocathodus hungaricus (Kozur &amp; Mostler)</td>
<td></td>
</tr>
</tbody>
</table>

Vr – very rare (1 specimen), r – rare (2–4 specimens), a – abundant (5–10 specimens), va – very abundant (more than 11 specimens)
1983  *Ammoniscus parapriscus* Ho – Salaj et al., p. 61, pl. 1, ?non fig. 2, pl. 6, figs 3–5, ?non fig. 6, non 7–8.
1992  *Ammoniscus parapriscus* Ho – Trifonova, p. 9, pl. 1, figs 3–5, pl. 6, fig. 10.

**Material:** 75 sections in samples SP 1, 2, 3, 4, 5, 6, 7, 8, 9 and 61 free specimens from the sample SP 1.

**Description:** The test is circular, discoid; periphery rounded and concave on both sides. The spherical proloculus is followed by a closely planispiral enrolled undivided tubular second chamber, gradually increasing in diameter. The number of whorls is 3–5. The wall is agglutinated and compact. The aperture is at the end of the tube.

**Remarks:** This species was assigned to *Cornospira* with a question mark by Rettori (1994), but the material of the present study is not well-preserved. Therefore, it is not possible to support his conclusion, but the question mark is retained here for definition of the genus. The preservation of the wall of the specimens studied is not good, because they are recrystallized and in some cases pyritized.

**Stratigraphic and geographic occurrence:** The species *Ammoniscus parapriscus* was first found in the Lower and Middle Triassic of China (Ho, 1959). Later this species was reported from the Lower and Middle Triassic (Anisian) of the Alps, Dinarides, Albania, Hungary, Bulgaria, Caucasus, Iran (Zaninetti, 1976; Pantić-Prodanović and Radošević, 1977b; Pisa et al., 1979; Oravecz-Scheffer, 1987; Pirdeni, 1988; Efimova, 1991; Trifonova, 1992; Rettori, 1995) and in the Upper Triassic of the West Carpathians (Salaj et al., 1983).

**Glomospirella facilis** was first described from the Lower and Middle Triassic of China (Ho, 1959). Later this species was reported from the Lower and Middle Triassic (Anisian) of the Alps, Dinarides, Albania, Hungary, Bulgaria, Caucasus, Iran (Zaninetti, 1976; Pantić-Prodanović and Radošević, 1977b; Pisa et al., 1979; Oravecz-Scheffer, 1987; Pirdeni, 1988; Efimova, 1991; Trifonova, 1992; Rettori, 1995) and in the Upper Triassic of the West Carpathians (Salaj et al., 1983).

Genus *Glomospirella* Plummer, 1945

**Type species:** *Glomospira umbilicata* Cushman & Waters, 1927

**Glomospirella facilis** Ho, 1959

*Fig. 5E, F*

1959  *Glomospirella facilis* n. sp. – Ho: p. 413, pl. 5, figs 20–25.
1983  *Glomospirella facilis* Ho – Salaj et al., p. 65, pl. 3, figs 1–10, 13-16, non 11–12, pl. 5, figs 10–11, 15–16, ?non figs 12–14.
1992  *Glomospirella facilis* Ho – Trifonova, p. 18, pl. 2, fig. 4, pl. 6, figs 20, 27, ?non fig. 28.

**Material:** 17 sections in the samples SP 1, 3, 5, 6, and 30 free specimens from the samples SP 1 and 5.

**Description:** The test is small, equatorial section oval, axial section mainly plain or convex on the both sides. The spherical proloculus is followed by a streptospirally enrolled undivided tubular second chamber and the last 2–3 whorls become planispirally coiled and gradually increasing in diameter. The streptospiral stage of coiling is wider in the axial section than the last whorl. The wall is agglutinated, thin, and smooth. The aperture is at the end of the tube.

**Remarks:** The characteristic features of this species are the oval form in the equatorial section and the convex or plain form in the axial section, the streptospiral stage of coiling is wider in the axial section than the last whorl. The above mentioned features of *Glomospirella shengi* are different from those of *Glomospirella facilis*. The preservation of the wall of the specimens studied is not good, owing to recrystallization or pyritization.

**Stratigraphic and geographic occurrence:** This species was first found in the Lower and Middle Triassic of China (Ho 1959). Later the species was described from Lower and Middle Triassic (Anisian) of the Dinarides, Albania, Hungary, Bulgaria, Greece, Turkey, Iran, China (Zaninetti, 1976; Pantić-Prodanović and Radošević, 1977a; Dager, 1978; Oravecz-Scheffer, 1987; Pirdeni, 1988; Salaj et al., 1988; Trifonova, 1992; Rettori, 1995) and in the Anisian and Rhaetian of the West Carpathians (Salaj et al., 1983).

Genus *Pilammina* Pantić, 1965

**Type species:** *Pilammina densa* Pantić, 1965

**Pilammina?* cf. praedensa* Urošević, 1988

*Fig. 5I*

1988  *Pilammina praedensa* n. sp. – Urošević: p. 372, pl. 1, figs 1–6.
1995  *Pilammina praedensa* Urošević – Rettori, p. 56, pl. 3, figs 1–3, 5, ?non fig. 4.

**Material:** One section in the sample SP 6.

**Description:** The test is free. It is close-coiled, consisting of a round proloculus and another inseptate, tubular chamberlet. The initial winding covers two and a half to three close coils. The last whorl is much wider than first whorls.

**Remarks:** The section studied is similar to the holotype. The preservation is not good, because the test is not complete. Therefore there is no possibility to describe the inner structure in detail. Usu-
ally the preservation of the wall of this species is not good and there is no confidence that this species belongs to *Pilammina* and this is true for the studied specimen too.

**Stratigraphic and geographic occurrence:** Species *Pilammina praedensa* was first described from the Lower Triassic of the Dinarides (Urošević, 1988) and after it was found in the upper part of Olenekian of Greece and the Lower Triassic of China (Rettori, 1995).

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**Genus Hoyenella Rettori, 1994**

**Type species:** *Glomospira sinensis* Ho, 1959

*Hoyenella sinensis* (Ho, 1959)

Fig. 5M–O

*1959*  
*Glomospira sinensis* n. sp. – Ho, p. 410, pl. 3, figs 16–34.

*1959*  
*Glomospira sinensis* var. *rara*, n. sp. n. var. – Ho, p.
type is broken, without a proloculus. The preservation of the specimens is not good, because they are mainly recrystallized, but there is proloculus present in all specimens.

Stratigraphic and geographic occurrence: Species *Dentalina splendida* was first described from the lower part of the Olenekian of the North Siberia (Gerke, 1961). Later this species was found in the Olenekian of the North Caucasus and Precaucasus (Efimova, 1991), and in the upper part of the Lower Triassic and in the lower part of the Anisian of Bulgaria (Trifonova, 1994).

Remarks on foraminifer fauna. The foraminifer assemblage consists of *Ammodiscus*? parapriscus Ho, *Glomospirella facilis* Ho, *Gl. shengi* Ho, *Gl. sp.*, *Pilaminina*? cf. *P. praeedens* Urošević, *Arenivalina*? *sp.*, *Meandrosoria*? *sp.*, *Agathammina*? *sp.*, *Hoyenella sinensis* (Ho), *Dentalina splendida* Schleifer, *Nodosaria*? *sp.* (Fig. 5A–R). Species of the genera *Ammodiscus*, *Glomospirella* and *Hoyenella* are dominant. An overview on the foraminifer taxa determined is given in Table 1.

The main feature of the taxonomic composition of this foraminifer assemblage from the Studorski preval locality in the Julian Alps of western Slovenia is the dominance of primitive agglutinated foraminifers (*Ammodiscus* and *Glomospirella* and some miliolids and nodosariids. The foraminifers are very small and not well-preserved. The generic composition of this foraminifer assemblage is mainly similar to the composition of the Early Triassic and Anisian assemblages from different parts of the Tethys, from the Alps to China. Several authors have indicated that there are “*Glomospira-Glomospirella*” foraminifer communities at different levels of the Lower Triassic and the lower Anisian in the Thayian Realm (Zaninetti, 1976; Salaj et al., 1983; Trifonova, 1992; Vaks, 2007). The foraminifer assemblage from the Studorski previal locality has some species common with the foraminifer assemblages from the Lower Triassic of China (He, 1993), the Olenekian of the Caucasus area (Efimova, 1991), the Lower Triassic and Anisian of Bulgaria (Trifonova, 1992) and Hungary (Oračev-Scheffer, 1987), and it can be correlated with foraminifer assemblages from these areas. In the Alps and the Dinardies the “*Glomospira-Glomospirella*” foraminifer communities with some miliolids and nodosariids are known mainly from the Lower Triassic (Dimitrijević et al., 1968; Pantić; 1970; Pantić-Prodanović and Radošević, 1977b; Pisa et al., 1979; Resch, 1979; Broglio Loriga et al., 1990).

Earlier in western Slovenia, the “*Glomospira-Glomospirella*” foraminifer assemblages with some nodosariids were reported from the Lower Triassic and Anisian (Ramovš, 1972). The foraminifer assemblage from western Slovenia is mainly similar to the foraminifer assemblages mentioned above, because they have a common generic composition. In eastern Slovenia “*Glomospira-Glomospirella*” foraminifer assemblages are known from the Anisian (Anićić and Dzet, 2000).

In only one thin-section a *Meandrosoria*-like specimen has been detected, but the preservation of the internal structure is poor. In Triassic time, the representatives of *Meandrosoria* are typical for the Early Triassic and the Anisian. It is reasonable to conclude that the foraminifer assemblage from the Studorski preval locality of the Julian Alps in western Slovenia occurred in Early Triassic–Anisian time, or that it was probably restricted to the upper part of the Early Triassic.

Conodonts

The conodont taxonomy follows the classification of Orchard (2005).


Type species: *Spathognathodus homeri* Bender, 1970
**Triassospathodus hungaricus** (Kozur and Mostler, 1970)  
Fig. 6A–G

*1970 Spathognathodus hungaricus* n. sp. – Kozur and Mostler: p. 453, pl. 4, figs 8–11.

**Material:** Five P-elements and eleven ramiform elements (samples SP 1, 3, 5) of moderate preservation.

**Description:** The spathodid P-elements examined are similar in length and bear three denticles. The basal cavity is wide and open. However, part of the posterior margin in the illustrated specimen is broken in the aboral view (Fig. 6D). It is obvious that the elements have an expanded basal cavity.

The ramiform elements are mostly fragmented and they do not permit reconstruction. Some of them are determined as ?M, ?S1 and S3 elements.

**Remarks:** *Triassospathodus* can be easily distinguished from *Neospathodus* Mosher, 1968, which has a terminally upward-directed lower margin of the basal cavity (Mosher, 1968). On the other hand, there exists a morphologic similarity in the P elements of *Triassospathodus* Kozur, 1998 (in Kozur et al., 1998) and *Novispathodus* Orchard, 2005 (Orchard, 2005); the latter has a shorter and more widened basal cavity and it has also a different apparatus composition (Heinz Kozur, pers. comm., 2012).

From the Spatian strata of the Thaynes Group in Nevada, Lucas and Orchard (2007) reported elements that are morphologically close or identical with “*Neospathodus* cf. hungaricus.”

**Stratigraphic and geographic occurrence:** Species *T. hungaricus* was first described from the *Tirolites* beds of Felsőrs in Hungary (Kozur and Mostler, 1970). *Triassospathodus* is an important Lower Triassic genus that comprises the majority of Upper Olenekian (Spathian) index species (Kozur et al., 1998).

In an integrated ammonoid and conodont zonation of the Triassic, Kozur (2003) presented *T. hungaricus* as the marker taxon for the basal Spathian equivalent to the *Tirolites cassianus* ammonoid Zone. According to H. Kozur (pers. comm., 2012), in the shallow western Tethys the *T. hungaricus* fauna lies within the lower Spathian, where the fauna with *Icriospathodus collinsoni* is missing.

There are several records of Early Triassic conodonts from Slovenia, but most of them are older than Spathian (see Kolar-Jurkovšek et al., 2011). Recovery of the spathodid Spathian fauna has been documented only from the borderland of the Ljubljana depression in central Slovenia (Dozet and Kolar-Jurkovšek, 2007). In Slovenia, a conodont fauna, high in spathodid elements and with few denticles, is abundant in the Spathian of Slovenia (Idrija-Žiri and Krško areas; unpublished data). From the External Dinarides, the Muč section in Croatia was proposed as a standard section for the European Upper Scythian. This section is well-known for its rich macrofauna and also contains *Triassospathodus triangularis* (Herak et al., 1983).

**Macrofauna**

The macrofauna collected contains bivalves and gastropods, as well as a few badly preserved specimens of ammonoids of the genus *Tirolites* that indicate an Early Spathian age.

**Bivalves**

Genus *Bakevellia* King, 1848

*Type species:* *Avicula antiqua* Graf zu Münster in Goldfuss, 1836 [non *A. antiqua* Defrance] = *Avicula binneyi* Brown, 1841, by original designation

*Bakevellia* cf. *incurvata* (Lepsius, 1878)  
Fig. 7A–C

*1878 Gervillia incurvata* sp. n. – Lepsius: p. 353, pl. 1, figs 3a–b.

*1908 Gervillia incurvata* Lepsius – Wittenburg, p. 31, pl. 4, fig. 7.

*1923 Gervillia incurvata* Lepsius – Diener, p. 91 (cum syn.).

**Material:** One well-preserved specimen with conjoined valves
(BJ 2766), one left valve and one specimen with conjoined valves (BJ 2763), ca. 40 moderately preserved specimens mostly with conjoined valves (BJ 2764).

**Description:** Relatively slender, posteriorly elongated *Bakevellia*, with left and right valve more or less equally convex. Beaks pointed, projecting above hinge margin in both valves.

**Remarks:** This species is unusual for its genus in having more or less equally convex left and right valves. On this basis, the material is tentatively assigned to *Bakevellia incurvata*, although there are slight differences in the outline by comparison with the single right valve, figured by Lepsius (1878, pl 1, figs 3a, b). Lepsius (1878, p. 353) indicated that the beak of the right valve did not project above the hinge line, whereas in the material described, it does. Another difference to the description by Lepsius (1878) is...
the apparent lack of a shell sulcus at the ventral margin, but the intraspecific variability of this feature is unclear.

A peculiarity of the present material is an unusual orientation of specimens that have both valves conjoined. Frequently, these specimens have their valves opened up to 30° and rest on the ventral valve margins, i.e. the hinge margin lies in a dorsal position, more or less parallel to the substrate. It is unlikely that this was the life position of the animal, because constructional morphology (elongated shape, long hinge axis, similar convexity of valves) indicates life as an endobystate mud sticker with the hinge axis oriented more or less vertically to the sediment surface (Seilacher, 1984, text-fig. 7; Muster, 1995, p. 98). Because the fauna occurs at the top of a tempestitic bed, as indicated by hummocky cross-stratification, a possible explanation is that the animals were excavated and displaced by storm waves and then smothered by the resettling silt-clay suspension. The observed orientation in the Studorsi preval material might thus be an unnatural dying position of this species.

Genus *Eumorphotis* Bittner, 1901

**Type species**: *Pseudomonotis telleri* Bittner, 1898 by subsequent designation (Cossmann, 1902)

*Eumorphotis cf. gronensis* (Wittenburg, 1908)

Fig. 7E

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**Material**: One left valve with a partly preserved external shell layer (BJ 2766), questionable three abraded left valves (BJ 2701).

**Description**: Very large *Eumorphotis* with about 70 radial ribs near the shell margin. Ribs nodose, increasing in number by intercalation during ontogeny. Outline of shell typical of the genus, but posterior wing relatively elongated.

**Remarks**: The largest specimen is about 8 cm long and more than 8 cm high. This is to the knowledge of the authors the largest *Eumorphotis* specimen, reported so far from the Early Triassic.

Although the type material of *E. gronensis* (Wittenburg, 1908) is incomplete, the specimen in the present study is tentatively assigned to that species on the basis of the similar ornamentation. The shell exterior of the specimen is dorsally eroded, but an increase in the number of ribs by intercalation of additional ribs is observed on the posterior wing. Therefore, it is assumed that the higher number of ribs in this specimen (ca. 70) by comparison with the holotype of *E. gronensis* (28) could be related to its larger size.

Species *E. kittlii* Bittner, 1901 differs from *E. gronensis* chiefly in its stronger development of the ten to twelve most posterior radial ribs, which cover the posterior wing and the adjacent part of the disc. Although the present authors currently treat both species separately, they wish to note that the difference in ornamentation might turn out to be a preservation effect or as lying within the range of intraspecific variation. In this case, *E. kittlii* would have priority over *E. gronensis* and species assignment of the specimen would change accordingly (see also Broglio Loriga and Mirabella, 1986).

Genus *Aviclamys* Allasinaz, 1972

**Type species**: *Pecten csopakensis* Frech, 1905 by original designation

*Aviclamys*? sp.

Fig. 7D

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**Material**: One complete left valve and two incomplete valves (BJ 2765).

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Fig. 8. Neritimorph gastropod *Natifia costata* (Münster, 1841) from Early Triassic of Studorsi preval, Slovenia, BJ 2480. A. General view of rock fragment with numerous poorly preserved individuals. B. Close-up of the specimen arrowed on A, showing mode of preservation

**Description**: Disc nearly circular, covered with about ten strong first-order radial ribs plus the same number of intercalated weaker second-order ribs. The posterior auricle is without a sinus.

**Remarks**: The state of preservation does not allow specific determination of this taxon. However, the circular disk, truncated posterior auricle and style of ornament are typical of *Avichlamys. Pecten vajolettensis* Wittenburg, 1908, subsequently referred to *Avichlamys* by Allasinaz (1972, p. 225), has a similar ornament. However, the presence of an elongated anterior auricle, as figured by Wittenburg (1908, pl. 1, fig. 9), cannot be verified in the material described.

**Gastropods**

Genus *Natifia* de Koninck, 1881

**Type species**: *Natifia lyrata* Phillips, 1836 by monotypy (de Koninck, 1881, p. 5)

*Natifia costata* (Münster, 1841)

Fig. 8A, B

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Genus *Naticella* Münster, 1841

**Type species**: *Naticella costata* sp. n. – Münster: p. 101, pl. 10, fig. 14.


2005 *Natifia costata* (Münster) – Nützel, p. 440, fig. 6.

**Type locality**: Most likely Gronehshof, South Tirol, Italy; Werfen Formation, Olenekian, Early Triassic (compare Kittl, 1892: 67).

**Material**: Single rock piece, composed of gastropod coquina from Studorsi preval. The association is monospecific, the shells are recrystallized and mostly weathered out from the visible surfaces. Sample BJ 2480.

**Remarks**: In spite of the mass occurrences of this gastropod in the Werfen Formation, its taxonomy – both generic and specific – status remains unclear. The type species of *Natifia* was described from the Visean (Lower Carboniferous) of Yorkshire (Phillips, 1836), while the species *Natifia costata* was defined by Münster (1841) in a monograph on gastropods from the Carnian (Upper

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DISCUSSION AND CONCLUSIONS

This paper presents detailed documentation and interpretation of sedimentological data on a remarkable Early Triassic fossil site in the Julian Alps of Slovenia. The report on the palaeontological material, collected at Studorski preval, confirms the wide distribution of mollusk and foraminifer taxa in the Early Triassic Tethys. The recovery of the earliest Spathian conodont fauna with *Triassospathodus hungaricus* indicates an Olenekian age of the fauna and is an important correlative element for comparison with coreval faunas in the Dinarid area and worldwide.

The sedimentological data presented here indicate that the strata of Studorski preval in the Julian Alps of western Slovenia were deposited in a more distal, storm-influenced ramp. A ramp setting was also mentioned for the deposition of the Werfen Formation in the Southern Alps (Brandner et al., 2012). Two types of beds were recognized, designated here as microfacies types A and B. These alternate vertically and form a sedimentary architecture that consists of coarse-grained skeletal lag deposits of microfacies A, overlain by the laminated or bioturbated mudstones and/or marls of microfacies B. Units with such organization imply the occasional winnowing of bottom sediments during storm peaks and the deposition of coarse-grained skeletal material as the storms waned, followed by the slow settling out of fine-grained material during intervals without storms. Ammonoids encountered in microfacies B imply a barrier-free connection of the depositional environment with the open sea, i.e. deposition in a more distal, deeper-ramp setting.

Numerous trace fossils in the mudstones and marls of microfacies B type beds suggest deposition in a low-energy deeper part of the ramp. A low-energy, poorly aerated environment is also indicated by the presence of pyrite and celestite. Periodic reworking of the bottom skeletal-rich sediment and the sedimentation of storm beds suggest deposition in a zone above the storm wave base, possibly in a wide low-energy mid-ramp environment.


The mollusk fauna includes taxa comparable to those of the *Tirolites* beds of the (Upper) Werfen Fm. Gastropods form monospecific coquinas that are most likely of the shells of *Natria costata*, a typical faunistic element of the Tethyan Early Triassic. The bivalve fauna includes three genera that differ in stratigraphical range and geographical distribution. *Bakevellia* is a long-lived genus, known worldwide from the Permian up to the Cretaceous. *Eumorphophis* was cosmopolitan in the Early Triassic and reached its acme in the Olenekian, although the genus might have persisted locally into the Middle Triassic. In contrast, *Avichlamys* was restricted to the Olenekian and is currently known only from the western Tethys. The bivalve species include *Bakevellia* cf. *incurvata, Eumorphophis* cf. *gromensis* and *Aviclamsys* sp. One of the specimens of *Eumorphophis* displays unusually large dimensions, seemingly constituting a record for the largest size of this genus, so far reported from the Early Triassic. The unusual position of articulated *Bakevellia* specimens is in accordance with storm-event deposition, inferred from sedimentological observations. By comparison with other recently described bivalve faunas from the Spathian (Neri and Posenato, 1985; Wasmer et al., 2012; Hautmann et al., 2013) and even from the Griesbachian (Kumage and Nakazawa, 2009; Hautmann et al., 2011), the bivalve diversity is low in the Studorski preval section, and no infaunal taxa are present. However, the normal to large growth sizes of the specimens indicate that this low diversity is not related to environmental stress. Rather, it may be related to the relatively greater water depth, provided that the specimens have not been transported over very large distances, which is unlikely given the low level of fragmentation. Miller (1988) demonstrated an evolutionary onshore-offshore trend in the Palaeozoic history of bivalves, in which taxa mainly evolved in shallow marine settings and expanded later into deeper/distal settings. This onshore-offshore or proximal-distal trend can be easily applied to the weakly inclined ramp situation, as depicted for
the site investigated. The low bivalve diversity in the relatively deeper distal setting described here may thus reflect an evolutionary delay of re-diversification in distal settings, compared to the more diverse faunas in shallower settings of the same age (e.g., Neri and Posenato, 1985; Hauumont et al., 2013). The data indicate a general evolutionary trend that onshore-offshore (see also Jablonski et al., 1983), seldom considered in analyses of the timing of biotic recovery after mass extinctions.

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