New Melissiodontinae (Mammalia, Rodentia) from the Paleogene of south-east Serbia

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Abstract Isolated teeth of Melissiodontinae from two Eocene and four Oligocene localities in southeastern Serbia are described. One new genus and two new species are named. The study of the derived morphology of the cheek teeth and of the contrastingly primordial microstructure of the tooth enamel of this diverse material provides a glimpse into the early history of the subfamily. The supposedly Asian murid ancestor of the Melissiodontinae seems to have reached the Serbian-Macedonian land area during the early or middle Eocene, which is shortly after the split up of the Muridae and Dipodidae and before the ‘Grande Coupure’ of central and Western Europe. We interpret the rapid consequent specialisation of the morphology of the chewing apparatus of the Melissiodontinae as an adaptation to feeding on small invertebrates on the floor of the Eocene forest.

Keywords Paleogene · Rodentia · Melissiodontinae · New genus and species · South-east Serbia

Introduction

The subfamily Melissiodontinae is in Europe represented by only one genus: Melissiodon Schaub, 1925, known from fossil assemblages dated between the late early Oligocene (MP23) to early Miocene (MN4) and includes nine species (Hrubesch 1957). It is the only murid that is present during the ‘Cricetid Vacuum’ (MN3 in the early Miocene) in south western Europe. Outside Europe, Melissiodon sp. is known from the Anatolian assemblage of Kargı-1 dated latest Oligocene-earliest Miocene (de Bruijn et al. 2013). The other genus belonging in this subfamily, Edirnella Ünay-Bayraktar, 1989, is known from the late Eocene of Süngülü (Lesser Caucasus; de Bruijn et al. 2003) and from two localities in the Thrace basin (MP25, Ünay-Bayraktar 1989).

The common presence of the Melissiodontinae in almost all of our Serbian localities was unsuspected. Moreover, the morphological diversity observed indicate that the Melissiodontinae were already diverse and highly specialised in the late Eocene and early Oligocene in this area (Table 1).

The samples of isolated teeth of melissiodontines were collected in the Babušnica-Korintica and Pčinja basins from two Eocene localities, Zvonice and Buštranje, and from five early Oligocene sites of Strelac-1, -2 and -3 Valniš and Raljin. The geological settings of these sites, their age assignment, overall
fossil content and methods of sample treatment is described in de Bruijn et al. (in press).

Material and methods

The terminology of parts of the cheek teeth basically follows Freudenthal et al. (1994) and is illustrated in Fig. 1. The melissiodontine material used for comparison consists of casts of Edirnella sinani Ünay-Bayraktar, 1989 cheek teeth and a fragment of a lower incisor from Kocayarma (Thrace basin, Turkey). In addition, casts of the cheek teeth of Edirnella kempeni de Bruijn et al. 2003 from Süngülü (Lesser Caucasus, Turkey) and original specimens of Melissiodon bernlochensis Hrubesch, 1957 from Bernloch (Germany) are available.

Abbreviations and terminology used in the description of the microstructure of enamel are enamel dentine junction (EDJ), Hunter-Schreger band (HSB), angle between the HSB and the normal to the EDJ (inclination), portio interna (PI), portio externa (PE), outer enamel surface (OES), external enamel layer without prisms (PLEX), inter prismatic matrix (IPM), enamel with parallel prisms that are at right angles to the EDJ (radial enamel) and basal ring of lamellar enamel in the molars (BRLE). The measurements of the teeth have been taken with a Leitz Ortholux measuring microscope with mechanical stage and measuring clocks. The pictures were made using a table-top and a high-resolution SEM. All specimens are figured as left ones. If the original is from the right side, this is indicated by underlining its number on the figure. Lower case letters refer to the lower dentition, upper case letters refer to the upper dentition. Abbreviations for measurements and descriptions are number of specimens (N), range of measurements (R), length (L), width (W), sinistral (sin) and dextral (dex).

The abbreviations used for of the localities are Zvonce (ZV), Buštranje (BUS), Strelac-1 (STR-1), Strelac-2 (STR-2), Strelac-3 (STR-3), Valniš (VA) and Raljin (RA). The fossil assemblages from southeastern Serbia are housed in the Natural History Museum in Belgrade (Serbia). Belgrade Museum locality codes are ZV = 037, BUS = 031, STR-1 = 024, STR2 = 015, STR3 = 026, VA = 027, RA = 028. A representative set of casts of rodents is kept in the collection of the department of Earth Sciences of Utrecht University, the Netherlands.

Taxonomy

Muridae Illiger, 1811
Melissiodontinae Schaub, 1925
Genera included: Melissiodon Schaub, 1920; Edirnella Ünay-Bayraktar, 1989 and Mogilia nov. gen.
Introduction

Schaub (1920) defined the genus *Melissiodon* on the combination of characteristics of the mandible and teeth that differentiate this genus from all other murids. In the same publication, he remarked that the group merits family rank, but the formal definition of the Melissiodontidae was published 5 years later (Schaub 1925). It is clear from these publications, as well as from a number of later studies (i.e. Freudenthal et al. 1992), that specialists hesitated to include *Melissiodon* into the Muridae (the family name Muridae is used here as the equivalent of Muroidea). This is understandable because it shows a number of aberrant characteristics in the skull: the lower incisor ends below the m2, the diastema is long and tubular, the scar of the masseter on the mandible is weak, the infraorbital foramen is large and the shape of the jugal is not like in other murids (Kristkoiz 1992). At the same time, the morphology of the cheek teeth is very derived, while the microstructure of the lower incisors remained primitive (Kalthoff 2000). Allocation to the Muridae was therefore rather induced by the lack of a suitable alternative than on similarity to members of that family. All species of *Melissiodon* of which the characteristics of skull and mandible are known share the characteristics listed above as well as characteristics of the cheek teeth such as the high ridges with steep walls (Hrubesch 1957), laterally compressed cusps and a square occlusal surface of the M2. Consequently the contents of the (sub)family remained clear cut and restricted to *Melissiodon* until Ünay-Bayraktar (1989) allocated her genus *Edirnella*, which is based on a few isolated upper cheek teeth of *E. sinani* from Kocayarma (Thrace basin, Turkey; ~MP 25), to the Melissiodontidae.

The classification of the Oligocene Muridae as suggested by Ünay-Bayraktar (1989), in which the subfamilies Paracricetodontinae (with *Paracricetodon* and *Trakmys*) and Melissiodontinae (with *Melissiodon* and *Edirnella*) are united in the family Melissiodontidae, has been criticised by Freudenthal et al. (1992), Kristkoiz (1992) and Kalthoff (2006) on two issues: (1) is *Edirnella* a member of the Melissiodontinae? (2) Are the Paracricetodontinae and Melissiodontinae as closely related as suggested by Ünay-Bayraktar (1989)?

Freudenthal et al. (1992) transferred *Edirnella* to the Paracricetodontinae on the basis of the morphology of its cheek teeth, while Kalthoff (2006) did so on the basis of the schmelzmuster of the lower incisor. However, the incisor

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**Fig. 1** Terminology of elements of the molars of melissiodonts used in the descriptions

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studied by Kalthoff (2006) was erroneously identified as *Edirnella* from Kavakdere (Thrace basin, Turkey), a locality that did not yield this species. Since this mistake was made, as well as discovered, by one of us (HdB) we apologise for causing confusion. Our recent analyses of a lower incisor of *E. sinani* from its type locality Kocayarma showed that it has the primitive type 1 schmelzmuster (Fig. 2) characteristic for melissiodontines (Kalthoff 2000).

The recognition that the enamel of the lower incisor of *E. sinani* shows the type 1 schmelzmuster in combination with the morphology of the upper cheek teeth, justifies the classification of *Edirnella* and *Melissiodon* in the same subfamily. Considering the second question, Freudenthal et al. (1992), Kristkoiz (1992) and Kalthoff (2006) have shown that the Melissiodontinae and Paracricetodontinae are different clades that do not have more in common than that they are both included in the Muridae.

Early diversification of the Melissiodontinae was also suggested by the allocation of *Edirnella kempeni* de Bruijn et al. 2003, from the late Eocene of eastern Turkey, to the Melissiodontinae on the basis of similarity in shape and morphology of its second and third upper and lower cheek teeth with those of the uncontested melissiodontine *Melissiodon* bernlochensis Hrubesch, 1957.

The presence of several melissiodontine species differing in the size of the cheek teeth, the height of the ridges relative to the cusps and in minor details of the microstructure of the enamel of the lower incisor in some of our Serbian localities came as a complete surprise (Table 1). The smaller species will be included in a new genus while the larger will tentatively be allocated to *Edirnella*. The uncertainty about the latter allocation is due to the unfortunate situation that, apart from one worn m2 (plate 7, Fig. 5 in Ünay-Bayraktar 1989), the lower dentition of the type species of *Edirnella* and the M1 of the two Serbian species is not known. The m2, m3, M2, and M3 that will be allocated to cf. *Edirnella* are very similar to those of *Melissiodon*. However, the anterior part of the m1 (described and figured below) is very different.

**Mogilia** nov. gen.

**Type species**: *Mogilia miloshi* nov. sp.

**Type locality**: Zvonce.

**Age**: Eocene.

**Derivatio nominis**: After the old Slav word ‘Mogila’, meaning a site where something lies buried.

**Included species**: *Mogilia miloshi* nov. sp. and *Mogilia lautus* nov. sp.

**Diagnosis**: *Mogilia* species have low-crowned cheek teeth with slender, but relatively high, cusps and a moderately to very complex network of low irregular ridges. The anterocone of the M1 is broad and situated on the central longitudinal axis of the occlusal surface. The small retracted anteroconid of the m1 is lower than the protoconid and metaconid. The anterior outline of the m1 is rounded and the anterosinusid of the m2 and m3 is wide as in all Melissiodontinae. The m3 has approximately the same length as the m1. The molars lack a basal ring of lamellar enamel. The outer surface of the thin enamel of the lower incisor is either smooth or shows a set of indistinct tangential ridges. The PI consists of transverse HSB with prism parallel IPM that make an angle of ~ 30° with the normal on the EDJ. The PE consists of radial enamel.

**Differential diagnosis**: *Mogilia* differs from *Melissiodon* in having lower-crowned cheek teeth with lower, more irregular, ridges. The almost symmetrical anterocone complex consisting of three cusps is situated on the central longitudinal axis in *Mogilia*, but has a labial position in *Melissiodon*. The
lingual part of the anteroconid of the m1 is connected to the protoconid in *Mogilia* as well as in *Melissiodon*, but the anterior outline of the m1 of these genera is quite different because a second, labially situated, anteroconid cusp developed in *Melissiodon*.

*Mogilia* differs from *Edirnella* by its more complex dental pattern and the structure and the central position of the anterocone complex of the M1. This complex consists of three cusps: a parastyl, the true anterocone and a cusp situated close to the lingual border between the anterocone and the protocone (= the *Melissiodon* cusp of Ünay-Bayaraktar 1989). In *Edirnella*, the anterocone has a labial position; it is situated on a straight line with the metacone and paracone. The *Melissiodon* cusp is not really part of the anterocone complex and the parastyl is absent. In *Mogilia* the anteroconid of the m1 is a small cusp that is connected to the protoconid, while it is developed as a cingulum in *Edirnella kempeni*. The m1 of the type species of *Edirnella* (*E. sinani*) is unfortunately not known.

*Mogilia miloshi* nov. sp.  
(Figs. 3d–i, 4d–i, 5d–i, and 6d–i).

**Derivatio nominis:** This species is named after our friend and colleague Miloš Milivojević, one of the most successful fossil hunters of the eastern hemisphere.

**Type locality:** Zvonce (coordinates 42°55′22″–23°34′43″).  
**Age:** Eocene.  
**Holotype:** Fragment of a right maxilla with M1-M2 (ZV-505) Fig. 3 g.  
**Material and measurements:** Tables 2 and 3, Fig. 7.

Upper dentition from the type locality shown in Figs. 3g–i and 4g–i.  
Lower dentition from the type locality in Figs. 5g–i and 6g–i. Other localities with *Mogilia miloshi*: Buštranje (late Eocene).

**Diagnosis:** Small species of *Mogilia*. Labial cusps of the upper cheek rounded, not strongly laterally compressed and ridges subordinate to the cusps. The anteroconid of the m1 is very small and, in the majority of the specimens, connected to the protoconid by a short oblique anterolophid.  
**Differential diagnosis:** *Mogilia miloshi* differs from *M. lautus* by its smaller size and simpler dental pattern of the M1 and m1. Moreover, the labial cusps of the upper cheek teeth are not laterally compressed as in *M. lautus*.

Description of the type material

All upper cheek teeth have three roots. In the M1, the anterocone proper is situated on the central longitudinal axis of the occlusal surface. Its size and height are about the same as in the four main cusps. The anterocone complex consists of the anteroconid and a parastyl on the labial border and the *Melissiodon* cusp on the lingual border of the occlusal surface. A peculiar feature shared by all M1 is the small, but well delimited, cusp that sits on the longitudinal axis behind the anterocone. It is not clear whether this cusp is the homologue of the protoconule or a neoformation. Among the many low, irregular, ridges connecting the cusps, remnants of the, more or less transverse, protoloph, metalophule, mesoloph and posteroloph and of the longitudinal ridge can be detected.

The occlusal surface of the M2 is about square as in all melissiodontines. The labial arm of the anteroloph is long and reaches the antero-labial base of the paracone. The lingual arm of the anteroloph is even longer and continues as a cingulum that reaches the base of the hypocone. The protolophule 1 and the metalophule are transverse and connected to the anterior arms of the protocone and hypocone. A complete protolophule 2 is present in 5 out of 10 specimens; in the others, this ridge is incomplete and developed as a second mesoloph. The length of the mesoloph itself shows strong individual variation. In only 1 out of 10 specimens it reaches the labial border of the occlusal surface. The posteroloph descends sharply from the hypocone to the metacone.

The anterior part of the M3 is very similar to that in the M2. The posterior part of the M3 is reduced and shows the V-pattern that characterizes most melissiodontine M3. The lingual arm of the anteroloph continues as a cingulum around the protocone and reaches the reduced hypocone. The protolophule 1 is connected to the anterior arm of the protocone. The length of the mesoloph shows much variation, but this ridge never reaches the labial border of the occlusal surface.

The lower cheek teeth have two roots. The m1 has a tiny anteroconid; it is not a true cusp in the majority of the specimens but the end of the anterior arm of the protoconid. Antero-labially of this ‘anteroconid’ there is an anterosinusid in four out of five specimens. The metalophulid 1 is directed obliquely forward and connects to the anterior arm of the protoconid. The strong posterior arm of the protoconid almost reaches the lingual border of the occlusal surface. An irregular, sometimes incomplete mesolophid is present in three out of five specimens. The hypolophulid is transverse or directed slightly forward and connects to the anterior arm of the hypoconid. The long posterolophid is connected to the postero-lingual slope of the entoconid.
Edimella kempeni (Süngülü)

Mogilia miloshi (Bustranje)

Mogilia miloshi (Zvonce)

Mogilia lautus (Valniš)

Melissiodon bernlochensis (Bernloch)
Fig. 5 Edirnella kempeni from Sünngili (type locality). a m1, b m2, c m3. Mogilia miloshi nov. sp. from Buštranje (code 031). d m1 (BUS-556), e m2 (BUS-561), f m3 (BUS-581). Mogilia miloshi nov. sp. from Zvonce, (type locality, code 036). g m1 (ZV-546), h m2 (ZV-552), i m3 (ZV-561). Mogilia lautus nov. sp. from Valniš (code 027), j m1 (VA-9264), k m2 (VA-937), l (VA-956) Melissaodon bernlochensis from Bernloch (type locality). Collection Bayerische Staatsammlung, München). m1 (nr.1536), n m2 (nr.65), o m3 (nr.25).

The m2 has a long straight lingual arm of the anterolophid that is separated from the metaconid by a notch, while the shorter labial branch encloses the anterosinusid. The parallel, slightly forwards directed, metalophulid and hypolophulid connect to the anterior arms of the protoconid and hypoconid. The posterior arm of the protoconid is long. The mesolophid is long in two out of six, short in two out of six and absent in two out of six specimens. The long posterolophid is connected to the postero-lingual base of the entoconid.

The morphology of the single m3 from Zvonce available (Fig. 5) is very similar to that of the m2. The metalopulid and hypolophulid are transverse and connect to the anterior arms of the protoconid and hypoconid. The posterior arm of the protoconid is long, but the mesolophid is absent.

Enamel structure of the lower incisor and the cheek teeth

The outer enamel surface of the lower incisor is either smooth (Buštranje) or shows an indistinct set of tangential ridges (Zvonce). The PI of the thin enamel (~30 μm) consists of transverse HSB with prism parallel IPM that make an angle of about 30° with the normal on the EDJ (Fig. 9). The PI of the specimen from Zvonce is about four times the thickness of the radial enamel of the PE, while the PI of the specimen from Buštranje is only slightly thicker than the PE. The enamel of the cheek teeth consists of radial enamel all the way to the base of the crowns (P-type, von Koenigswald 2004).

Comparison of the Mogilia miloshi cheek teeth from Buštranje and Zvonce

The teeth from Buštranje assigned to M. miloshi are metrically as well as morphologically very similar to those from Zvonce. The ridges of the few M1 available from Buštranje are somewhat higher and these teeth have a ridge connecting the ‘protoconule’ to the parastyl, a cusp that is less developed than in the M1 from Zvonce. The ridges of the M2 from Buštranje seem also to be slightly higher than in the specimens from Zvonce and the paracone and metacone are more laterally compressed. The morphology of the M3 from both localities overlaps. The ridges of the lower cheek teeth from Buštranje are slightly higher than in the ones from Zvonce and the anteroconid of the m1 is connected to the metaconid by the metalophulid in three out of four specimens. The m2 from the two localities differ sharply in the configuration of the metalophulid. In the six m2 from Zvonce the metalophulid connects labially with the anterior arm of the protoconid, while in the 11 specimens from Buštranje, the metalophulid and the anterior arm of the protoconid are connected to the anterolophid separately. The latter configuration is seen in most Melissaodon species as well as in Edirnella kempeni. The differences between the two samples of M. miloshi teeth observed above are in our opinion insufficient to define separate species. This is especially so because not all tooth positions would then allow identification to the species level. However, the morphological differences strongly suggest that Buštranje specimens are more derived and thus may be the younger of the two localities. Whether or not this age difference is at the basis of the sharp difference in composition between the rodent associations from these localities too (de Bruijn et al. in press) remains to be demonstrated.

Mogilia lautus nov. sp.
(Figs. 3j–l, 4j–l, 5j–l and 6j–l).

Derivatio nominis: Lautus has two meanings in Latin. The first is ‘well washed’ and the second is ‘luxurious’, both qualifications apply to the Mogilia teeth that will be described below.

Holotype: M1 dext. from Valniš (VA-878), Fig. 3j (shown reversed).

Type locality: Valniš.

Age: Early Oligocene.

Other localities with: Mogilia lautus Strelac-1, Raljin (both not illustrated).

Age: Early Oligocene.

Material and measurements: Tables 4 and 5, Fig. 7.

Diagnosis: Mogilia lautus is of medium size with upper cheek teeth that show an intricate pattern of low ridges. Among these is an exceptionally well-developed entomesoloph. The broad antercone complex of the M1 is situated on the longitudinal axis of the occlusal surface. The unicusp anteroconid of the m1 has a retracted position and a low cingulum at its anterior slope. A prominent extra cusp is present directly postero-labially of the metaconid.

Differential diagnosis: Mogilia lautus is larger than M. miloshi and the dental patterns of the upper teeth and the lower m1 are much more intricate than in that species. The elaborate structure of the anteroconid complex of the m1 of M. lautus is different from the small, simple anteroconid in the m1 of M. miloshi.
which is situated directly postero-labially of the metaconid. As the end of the posterior arm of the protoconid has formed a cusp unusual because the anteroconid proper is not situated close to the paracone to the protocone and the well-developed mesoloph is present in 11 out of 13 specimens. The protolophule 1 connects to the base of the paracone. An entomesoloph is separated from that cusp by a notch. The labial branch of the anterolophid encloses an oval anterosinusid, a feature shared by most Melissiodontinae. The posterior arm of the protoconid, the mesolophid and the ectomesolophid are well developed in all 10 m2.

The anterior portion of the m3 is very similar to that of the m2. A long posterior arm of the protoconid is present in all 12 m3, but the mesoloph is absent in 6 out of 12 specimens. The hypolophid inserts on the anterior arm of the hypoconid.

Discussion

The m2 from Raljin is somewhat larger than the ones from the type locality of M. lautus, while it is morphologically intermediate between M. lautus and cf. Edirnella nov. sp. 2 from Strelac-1. In spite of an effort undertaken in 2015 to enlarge the collection from Raljin, we did not find any more complete melissiodontine teeth, so the assignation of this single m2 remains uncertain.

The teeth in the associations of Mogilia from Zvonce, Buštranje and Valniš show in this sequence a trend towards increase in size and complexity of the dental pattern. Since the samples from Buštranje and Zvonce are small, the difference in stage of evolution between these associations is for most tooth positions not clear. However, the metalophid in the m2 from Zvonce connects to the anterior arm of the protoconid (the original murid configuration), while it bends forward and connects with the anterolophid in the specimens from Buštranje (the derived configuration). Although we have no independent age control that the locality of Zvonce is older than Buštranje, we interpret the observed difference between the Mogilia associations that way. Our

### Table 2

| Zvonce | Length (mm) | Width (mm) |
|--------|-------------|------------|
|        | Range | Mean | N | Mean | Range | N |
| M1    | 2.40–2.50 | 2.47 | 6 | 1.85 | 1.70–2.00 | 6 |
| M2    | 1.42–1.74 | 1.57 | 10 | 1.59 | 1.46–1.74 | 9 |
| M3    | 1.18–1.27 | 1.22 | 4 | 1.39 | 1.31–1.43 | 4 |
| m1    | 1.65–1.95 | 1.85 | 6 | 1.36 | 1.28–1.45 | 6 |
| m2    | 1.57–1.88 | 1.73 | 5 | 1.40 | 1.33–1.48 | 6 |
| m3    | –    | 1.77 | 1 | 1.37 | – | 1 |

### Table 3

| Buštranje | Length (mm) | Width (mm) |
|-----------|-------------|------------|
|           | Range | Mean | N | Range | Mean | N |
| M1        | 2.56–2.63 | 2.60 | 2 | 1.77 | 2 |
| M2        | 1.45–1.65 | 1.52 | 7 | 1.51 | 1.43–1.60 | 8 |
| M3        | 1.07–1.36 | 1.24 | 19 | 1.36 | 1.24–1.50 | 21 |
| m1        | 1.81–1.83 | 1.82 | 3 | 1.37 | 1.34–1.41 | 3 |
| m2        | 1.40–1.78 | 1.68 | 11 | 1.41 | 1.32–1.47 | 10 |
| m3        | 1.57–1.90 | 1.74 | 8 | 1.26 | 1.10–1.35 | 8 |
working hypothesis is that *M. miloshi* and *M. lautus* have an ancestor-descendant relationship.

*Edirnella* Ünay-Bayraktar, 1989

**Included species:** *Edirnella kempeni*, *Edirnella sinani*, cf. *Edirnella* nov. sp. 1 from Buštranje, cf. *Edirnella* nov. sp. 2 from Strelac 1.
The few specimens and fragments included in cf. *Edirnella* differ in size, cf. *Edirnella* nov. sp. 1 being larger than cf. *Edirnella* nov. sp. 2. The cf. *Edirnella* are different from those of *Mogilia* but more like the m1 of *E. kempeni*; hence, our assignation to cf. *Edirnella*. Unfortunately the m1 of the type species *E. sinani* is not known, thus some uncertainty will remain in the allocation of species to the genus. cf. *Edirnella* nov. sp. 1. (Figs. 8d–g).

**Locality** Buštranje.

**Age:** Late Eocene.

**Dental characters:** The morphology of these few upper teeth is very similar to the type material of *E. sinani*, but they are much larger. The M2 has three roots.

**Remark:** The M2 and M3 of cf. *Edirnella* nov. sp. 1 resemble those of *Melissiodon* in the shape of the cusps and the height of the ridges, but the associated anterior part of an m1 shows more similarity with that tooth in *Mogilia* than in *Melissiodon* (see Figs. 5, 6 and 8).

**Material and measurements:** Only the M2, M3 and a damaged m1 are available.

Table 6 Measurements of cf. *Edirnella* nov. sp. 1 from Buštranje.

| Description |
|---|

The morphology of the three-rooted M2 of *Edirnella* nov. sp. 2 resembles that of the M2 of *Melissiodon bernlochensis* in detail (Figs. 3n, 4n and 8i). The anteroloph is low and has a double connection to the transverse protolophule 1 and is weaker than in *Edirnella* nov. sp. 1. Characteristic is the small pit that is enclosed by the entoloph, the posterior arm of the protocone and the mesoloph. This pit seems to be precluded by the configuration of low ridges in the central part of the M2 of *Mogilia laetus* from Valníš (Fig. 3k).
Fig. 8  *Edirnella sinani* from Kocayarma (type locality).  a M1.  b M2.  c M3.  cf. *Edirnella* nov. sp. 1 from Buštranje (code 031).  d M2 (BUS-591).  e M2 (BUS-593).  f M3 (BUS-596).  g Anterior part of m1 (BUS-599).  cf. *Edirnella* sp. from Zvonce (code 036).  h (ZV-587).  cf. *Edirnella* nov. sp. 2 from Strelac-1 (code 024).  i M2 (STR1–302).  j M3 (STR1–308).  k M3 (STR1–310).  l anterior part of m1 (STR1–311).  m m2 (STR1–315).  n m3 (STR1–318)
The five M3 from Strelac-1 show considerable individual variation in the differences in degree of reduction of the posterior part of the occlusal surface. A peculiar characteristic that all these teeth share is the double connection between the protoloph and the anteroloph. The taxonomic value of this character is uncertain; it occurs also in the M3 of cf. Edirnella sp. indet. From Zvonce (Fig. 8h) and in the M3 of M. bernlochensis (Figs. 3o and 4o), but lacks in the M3 of Edirnella nov. sp. 1 from Buštranje (Fig. 8f).

The single m1 available from Strelac-1 consists of an anterior part only (Fig. 8i). This anteroconid complex resembles the configuration seen in the m1 of Mogilia lautus (Figs. 5j and 6j) in having a retracted anteroconid lingually which is connected by a short anterolophulid to a smaller, more ridge-shaped and a labially situated anteroconid cusp. Another characteristic shared by these two species is the development of an extra cusp near, or at the end of the posterior arm of the protoconid. The morphology of the rolled specimen from Strelac-2 is as described above.

The m2 shows a protoconid and hypoconid that are V-shaped and thus modified as in Melissiodon. The anterosinusid is wide. The metalophulid and the anterior arm of the protoconid reach the anterolophid separately. The posterior arm of the protoconid continues as a long thin mesolophid all the way to the lingual border of the occlusal surface as in the m2 of Melissiodon bernlochensis (Figs. 5n and 6n). The hypolophulid connects to the anterior arm of the hypoconid.

The long m3 has a dental pattern, slightly reduced, that is almost identical to that of the m2 (Fig. 8m, n).

cf. Edirnella sp. indet.
(Fig. 8h).

**Locality:** Zvonce.

**Age:** Late? Eocene.

**Material and measurements** 1 M3 (length 1.75 mm, width 1.93 mm).

**Description**

The single M3 from Zvonce is much too large to allocate it to *Mogilia miloshi* from that locality. Its morphology, showing a double connection between the protoloph and the anteroloph, and size are similar to the M3 of *Edirnella nov. sp. 2* from the late Oligocene site Strelac-1. Since the M3 of Melissiodontinae are very much alike this specimen cannot be identified to the species level.

### Table 6 Measurements of cf. *Edirnella* n. sp. 1 from Buštranje

|        | Length (mm) | Width (mm) |
|--------|-------------|------------|
| Buštranje | 2.42–2.77 | 2.55 | 2.29 | 2.17–2.38 | 3 |
| M2     | –           | 1.92 | 2.13 | –         | 1 |
| m1     | –           | 2.37 | –    | –         | – |

### Table 7 Measurements of cf. *Edirnella* n. sp. 2 from Strelac-1 and -2

|        | Length (mm) | Width (mm) |
|--------|-------------|------------|
| Strelac-1 | 2.42–2.62 | 2.52 | 2.15 | 2.19–2.05 | 2 |
| M3     | 1.58–1.79 | 1.66 | 1.82 | 1.66–1.93 | 5 |
| m2     | 1.94–2     | 1.82 | 1.89 | 1.84–1.94 | 2 |
| m3     | –           | 2.66 | 1.86 | –         | 1 |
| Strelac-2 | –           | 2.66 | 1.86 | –         | 1 |

### Table 8 Measurements of cf. *Edirnella* sp. indet. from Strelac-2

|        | Length (mm) | Width (mm) |
|--------|-------------|------------|
| M3 sup | 1.63        | 1.75       |

### The evolutionary history of the Melissiodontinae

Until Ünay-Bayraktar (1989) assigned her much disputed genus *Edirnella* from the early Oligocene of Turkish Thrace to the Melissiodontinae the geographic range of this, then monogenic, subfamily remained restricted to Europe where it is considered to be an immigrant from the East. The first occurrence of *Melissiodon* is in Bernloch (MP23), a site in southern Germany East of the Rhine Graben, while the first appearance west of the Rhine Graben is from localities assigned to MP24 (Russel et al. 1982). The stratigraphic range of the genus *Melissiodon* in Europe is from late early Oligocene (MP23) to well into the early Miocene (MN4). It is the only murid which range straddles the Cricetid Vacuum in south western Europe. Nine species have been formally named (Hrubesch 1957), which seems, considering the stability of their highly derived dental morphology, too many. Other than *Edirnella sinani* from the Thrace basin (MP25, Ünay-Bayraktar 1989) there are only two records of Melissiodontinae from outside of Europe: *E. kempeni* from the late Eocene of Sünğülü (Lesser Caucasus; de Bruijn et al. 2003) and *Melissiodon* sp. from the Oligo/Miocene transitional interval of Kargi-2 in central Anatolia (de Bruijn et al. 2013).

The discovery of the diverse array of Melissiodontinae from the late Eocene and early Oligocene of southeastern Serbia described above thus increases our knowledge of the early history of the subfamily substantially. We interpret the rapid consequent specialisation of the morphology of the chewing apparatus of the Melissiodontinae as an adaptation to feeding on small invertebrates.

It shows that its, presumably Asian, ancestor colonised the Serbian-Macedonian land area during the early or middle Eocene. That is, during the dawn of the origin of the Muridae, thus after the split up of the Muridae and
Dipodidae, and before the ‘Grande Coupure’, so the Melissiodontinae seem to be a very early branch of the Muridae. This conclusion is supported by the occurrence of two species in the late Eocene locality Buštranje that not only differ in the size and morphology of their cheek teeth, but also in details of the microstructure of their incisor enamel (Fig. 9). This radiation into several species probably occurred on or near the Serbian-Macedonian land area.

Fig. 9  a Longitudinal section of the lower incisor enamel of *Mogilia miloshi* nov. sp. from Buštranje. b Longitudinal section of the lower incisor enamel of *Mogilia miloshi* nov. sp. from Zvonce. c Longitudinal section of the lower incisor enamel of *Edirnella* nov. sp. 1 from Buštranje. d-e Transverse section of the m2 of *Mogilia miloshi* nov. sp. from Buštranje. f-g Transverse section of the m2 of *Edirnella* nov. sp. 1 from Buštranje.
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Compliance with ethical standards

Conflict of interest The authors declare that they have no competing interests.

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