Prolagus Pomel, 1853 (Lagomorpha, Mammalia) in the framework of the Pliocene faunal rearrangements in central Europe

Stanislav ČERMÁK, Chiara ANGELONE & Blanca MONCUNILL-SOLÉ
**Prolagus** Pomel, 1853 (Lagomorpha, Mammalia) in the framework of the Pliocene faunal rearrangements in central Europe

Stanislav ČERMÁK  
Institute of Geology of the Czech Academy of Sciences,  
Rozvojová 269, 165 00 Prague 6 (Czech Republic)  
cermaks@gli.cas.cz (corresponding author)

Chiara ANGELONE  
Department of Sciences, University Roma Tre,  
Largo S. L. Murialdo 1, 00146 Roma (Italy)  
and Institut Català de Paleontologia Miquel Crusafont, Edifici Z ICTA–ICP,  
Carrer de les Columnes s/n, Campus de la Universitat Autònoma de Barcelona,  
08193 Cerdanyola del Vallès, Barcelona (Spain)

Blanca MONCUNILL-SOLÉ  
Centro de Investigaciónes Científicas Avanzadas (CICA), As Caballeiras s/n,  
Campus de Elviña, Universidade da Coruña, 15071 A Coruña (Spain)  
and Department of Sciences, University Roma Tre, Largo S. L. Murialdo 1, 00146 Roma (Italy)

Submitted on 21 February 2020 | Accepted on 25 May 2020 | Published on 28 June 2021

**ABSTRACT**

Pliocene occurrences of *Prolagus* Pomel, 1853 in central Europe represent anomalies out of the peri-Mediterranean area, at that time the core distribution of the genus. Though known for several decades, those materials never underwent a general revision. The detailed analysis and comparison of all available materials performed here revealed two phenotypic entities: 1) *Prolagus bilobus* Heller, 1936 (Gundersheim localities, Raciszyn 1), for which were defined additional diagnostic characters and ontogenetic patterns of variation (d3/p3 and mandible); and 2) *Prolagus* sp. (Beremend 26/39), probably a new species. All the available occurrences are dated to MN15b. The morphological trends towards the reduction of p3 entoconid and of enamel folding evidenced in Pliocene *Prolagus* of western Europe cannot be recognized in coeval central European forms. Evidently, *P. bilobus* and *Prolagus* sp. do not pertain to the western European clade, whose separation is known since the early late Miocene. We hypothesize that at least *P. bilobus* originated from a dispersal of *Prolagus* from south-eastern regions of Europe rather than from an autochthonous speciation of isolated populations left as a relict after the southward displacement of *Prolagus* distribution area. The dispersal is likely to be related to the Pliocene global environmental changes during which extensive faunal rearrangements took place in Europe, in particular to those near the early/late Pliocene boundary.

**KEY WORDS**
Prolagidae, *Prolagus bilobus*, Gundersheim, Raciszyn 1, Beremend, palaeobiogeography.

urn:lsid:zoobank.org:pub:4FD1B8AF-7957-469E-A627-8126691743C3

Čermák S., Angelone C. & Moncunill-Solé B. 2021. — *Prolagus* Pomel, 1853 (Lagomorpha, Mammalia) in the framework of the Pliocene faunal rearrangements in central Europe. Comptes Rendus Palevol 20 (28): 597–617. https://doi.org/10.5852/cr-palevol2021v20a28
INTRODUCTION

Prolagus Pomel, 1853 represents a successful, long-lasting, species-rich lagomorph genus that during most of the Miocene was widely distributed throughout Europe and formed a significant part of the small mammal assemblages (López Martínez 1974, 1989, 2001). Probably, since the latest Miocene, its geographical range started to dwindle. The Pliocene is a critical moment in the evolution of Prolagus. The changes in its palaeobiogeographical distribution from pan-European to patchy and mainly peri-Mediterranean distribution become evident in this period. The Pliocene record of Prolagus in the peri-Mediterranean area is quite common and well documented, in contrast to central Europe, where the genus is very rare and occurs sporadically in a few localities.

Isolated Pliocene populations of Prolagus in central Europe have been known since the early 20th century (Heller 1936). The proven occurrences of the genus have been limited until now to Prolagus bilobus Heller, 1936 documented from Gundersheim sites (MN15b) in Germany and Raciszyn 1 (MN15b) in Poland (Heller 1936; Fejfar & Storch 1990; Fejfar et al. 2006; Fostowicz-Frelik 2010; Čermák & Angelone 2013). Apart from punctual taxonomic disputes (cfr. Fostowicz-Frelik 2010 vs Čermák & Angelone 2013), these remains have never been the subject of a detailed, dedicated study.

Some studies exist about the taxonomy and phylogeny of Pliocene Prolagus from western Europe (López Martínez & Thaler 1975; López Martínez 1989; Angelone 2008a), and as well, some enigmatic isolated remains have been reported from eastern Europe (Agadjanian & Erbajeva 1983; Erbajeva & Shushpanov 1988; Averianov & Tesakov 1998; Tesakov & Averianov 2002).

We decided to focus our attention to Pliocene central European Prolagus, in order to summarize the available materials, to clarify their taxonomic status, and to try to unravel the reasons that led to the palaeobiogeographical distribution observed after the Miocene.

THE LOCALITIES: HISTORICAL BACKGROUND AND GEOLOGICAL SETTING

The Pliocene record of Prolagus in Central Europe is limited to the following localities; their geographical locations are shown in Figure 1.

GUNDERSHEIM

Most of the fossil faunas at Gundersheim (Alzey-Worms district, Rheinland-Pfalz, Germany) come from three quarries, located SW from the homonymous village. A study of the fossil record from Gundersheim was first performed by Heller (1936). However, although the material originated from several fissures of quarries, F. Heller published the different faunas considering them as one assemblage, with only sporadic notes concerning their original provenance. Thus, the exact location, age, and name of each site/assemblage have since then remained unclear. Until the 1960s, all the samples were generally considered as pertaining to one faunal unit, and referred to simply as “Gundersheim”. Eventually, Kretzoi (1962) distinguished two faunal units in Heller’s original material: an older
Prolagus in the framework of the Pliocene faunal rearrangements in central Europe

one, correlated with the Csarnotanian (*sensu* Kretzoi 1962; c. MN15b), called Gundersheim 1 by Tobien (1980), and a younger one, correlated with the late Villányian (*sensu* Kretzoi 1962; c. MN17), called Gundersheim 2 by Tobien (1980).

Tobien (1980) also numbered particular Gundersheim fissures. Since the 1950s, he discovered and studied 22 fissures of which five yielded identifiable faunal remains. The most interesting was fissure no. 4, in which were recovered *Trilophomys schaubi*, *Bjoernkurtenia canterranensis*, *Baranomys longidens*, *Germanomys weileri*, and *Mimomys gracilis*, which indicate a late Ruscinian age (MN15b). This site, published in detail by Fejfar *et al.* (2006), is known as Gundersheim-fissure 4 (Gundersheim 3 *sensu* Koenigswald & Tobien 1990).

Tobien (1980) assumed that the most of Heller’s older faunal unit (Gundersheim 1) most likely originated from this fissure. The last fauna at Gundersheim was discovered in 1976 in an isolated block of calcified red loam found in the open field near the Rosengartenbruch quarry. The locality, known as Gundersheim-Findling (Gundersheim 4 *sensu* Koenigswald & Tobien 1990), yielded a faunal assemblage analogous to the one found in Gundersheim-fissure 4 described by Fejfar *et al.* (2006), and indicating a late Ruscinian (MN15b) age as well. Rodents from this locality were described in detail by Fejfar & Storch (1996), and prolacid lagomorphs partly by Čermák & Angelone (2013).

Apart from *Prolagus*, herein studied, Gundersheim localities yielded other lagomorph taxa: Gundersheim-Findling – *Pliopentalagus dietrichi* (Fejfar, 1961) published in detail by Čermák & Wagner (2013) and *Hypolagus petenyii* Čermák & Fladerer in Čermák, 2009 (p3 dex [morphotype I/a/A], 2 lower molariforms); Gundersheim-fissure 4 – Leporidae gen. et sp. indet. (fragmentary lower molariform); and “Hasenfundtelle I” (after label by Heller) – *Hypolagus* published by Heller (1936: 137-139; figs 1-2, pl. XI: 13-14) as *H. brachygynathus* and reassigned here as *H. cf. brachygynathus* (Kormos, 1930) (2 mandibles – sin with p3-p4 [morphotype II/c/D] and dex with p3 [II/II/c/C]).

**RACISZYN 1**
The locality Raciszyn 1 corresponds to a small abandoned quarry located c. 500 m W from the village of Raciszyn (district Gmina Działoszyn, province Łódź, central Poland). The fossil material comes from the terra rossa filling of a collapsed cave formed in Oxfordian limestones. Preliminary information about the locality and its fauna was provided by Kowalski (1990), Nadachowski (1990), Sulkowski & Szynkiewicz (1994), Szynkiewicz & Jagiełło (2015), and Nadachowski *et al.* (2015). The locality yielded a fossil assemblage consisting of about 40 mammalian taxa (Sulkowski in Szynkiewicz & Jagiełło 2015). The assemblage includes biostratigraphically valuable taxa such as a small *Mimomys of the gracilis-stehlini lineage* (with prevalence of primitive forms), *Mimomys bassiacus*, *Baranomys* or *Trilophomys* (Nadachowski 1990; Sulkowski & Jagiełło 2015; Nadachowski *et al.* 2015), indicating a late Ruscinian (MN15b) age (Kowalski 1990; Nadachowski *et al.* 2015). The remains of *Prolagus* addressed in this paper were first studied by Fotowicz-Frelik (2010) and described as a new species, *Prolagus*.
osmolskiae Fostowicz-Frelík, 2010. Eventually, Čermák & Angelone (2013) revised and synonymized the species with *Prolagus bilobus*. The only other lagomorph species found in Raciszyn 1 is *Hypolagus petenyii* (Nadachowski et al. 2015).

**BEREMEND**

The limestone quarry at Beremend is situated in the westernmost part of the Villány Hills (Baranya region, Hungary), an area palaeontologically investigated since the half of the 19th century. The first fossiliferous fissures in the quarry were discovered by S. J. Petényi in 1847 (Petényi 1864; Kordos 1991). During the 20th century, the quarry yielded several dozens of vertebrate-bearing sites, whose age span from the early Pliocene to the early Pleistocene (see Kretzoi 1956, 1959, 1962; Jánossy 1986; Kordos 1991; Pongrácz 1999; Császár & Kordos 2004 for details). The material, comprising remains of the genus *Prolagus* studied herein, was collected by L. Pongrácz from fissure fillings Beremend 26 and 39. The site no. 26 yielded a very extensive and diverse fossil assemblage of mammals including the biostatigraphical markers *Mimomys gracilis*, *Deromys nebrinius*, and *Propliomys hungaricus* (cf. Császár & Kordos 2004; Čermák 2007; Čermák unpubl. data). Such taxa indicate a late Ruscian (MN15b) age for Beremend 26, i.e., slightly younger than Csarnóta 2 (L. Kordos pers. comm.). Apart from *Prolagus*, the locality yielded also other lagomorphs, namely *Ochotonoma carnotana* Kretzoi, 1959 (partly studied by SC; see Čermák 2007) and *Hypolagus petenyii*.

In comparison with Beremend 26, the fossil record from Beremend 39 is much scantier and less diverse, consisting of “only” about 15 mammalian taxa (Pongrácz in litt.; Čermák & Wagner 2013). Nevertheless, the presence of *Beremendia fissidens*, *Elomys intermedius*, *M. cf. gracilis*, *Propliomys hungaricus*, *H. petenyii*, and *Trischizolagus dumitrescuae*, together with the evolutionary degree of the arvicOLIDs, indicate also a late Ruscian (MN15b) age, analogous to Beremend 26 (Čermák & Wagner 2013).

**OTHER LOCALITIES**

In addition to the above mentioned localities, there are other two relevant sites from which the presence of *Prolagus* was reported. Unfortunately, however, we were not successful to track down the lagomorph material from those localities. The taxonomic position of those samples remained thus unclear.

— **WÖLFERSHEIM** (Wetterau district, Germany; MN15b) — the occurrence of *P. bilobus* in this locality was reported in a faunal list by Tobien (1977). Dahlmann (2001) studied in detail small mammals from this locality, but not lagomorphs, which were reported only in a faunal list (*ibid.*: 95) with reference to Tobien’s list (1977). The collection of mammals from Wölfersheim is deposited in the SMF, but lagomorphs are missing.

— **KISLÁNG** (Polgárdi district, Hungary; MN15 or MN17 according to Mayhew 2012) — Kretzoi (1954) reports from this locality a sole left p3 of *Prolagus*. Based on the presence of the crochet (“Der Sporn am Mittelgraben der Kaufläche”), Kretzoi (1954: 247) hypothesized a taxonomic proximity of the Kisláng specimen to *Prolagus* from Gundersheim, and referred to it as *P. cf. bilobus*. The material of Kisláng collected and described by Kretzoi (1954) is currently curated in the collection of the MFGI, but the last surveys (SC in 2005) were not successful to track down the specimen. At any rate, the Kisláng mammalian assemblage is a mixture of material of different geological ages with a minimum estimated range between c. 1.7-3.5 Ma (Mayhew 2012). Thus, the age of the p3 reported by Kretzoi (1954) is unclear. Nevertheless, considering the presence of *Mimomys cf. bassiacus* in the locality (Mayhew 2012), the late Ruscian (MN15b) age cannot be excluded (see for details Čermák & Angelone 2013).

**MATERIAL AND METHODS**

All the material from the Gundersheim localities is stored in the collections of the SMF and the material from Raciszyn 1 is curated in the collections of the ZPAL. The material from Beremend 26 and 39 comes from the private collection of L. Pongrácz (Győr, Hungary) and is temporarily housed in the collection of the GLI. Following Čermák & Angelone (2013), we used Gundersheim 1 and Gundersheim 2 for Heller’s original localities/assemblages (sensu Tobien 1980), and Gundersheim-fissure 4 and Gundersheim-Findling for localities published by Fejfar et al. (2006) and Fejfar & Storch (1990).

For teeth nomenclature and metrics we follow López Martínez (1989) and Angelone & Sesé (2009), for jaws we follow Wible (2007) and Čermák (2009). Drawings and measures were taken with the aid of a Dino-Lite digital microscope and of a binocular microscope with ocular micrometer. Dental measurements were taken as maximum antero-posterior or bucco-lingual dimensions of the respective two-dimensional dental structures orthogonal to prismatic shaft. For consistency, only adult specimens (recognized by their prismatic tooth shape) were used for interspecific size comparisons. All measured data are given in millimeters [mm].

The biostratigraphic terminology used in this paper follows Fejfar & Heinrich (1983) and Fejfar et al. (1998). The term “Central Europe” is understood here as a geographic region in the center of Europe including Germany, Switzerland, Liechtenstein, Austria, Poland, the Czech Republic, Slovakia, and Hungary. Country abbreviations follow ISO 3166-1 alpha-2 codes. Following the recommendations of the International Commission of Stratigraphy, the first letter of informal subepochs is not capitalized.

Interspecific comparisons were made with the early Miocene to Pleistocene European species of *Prolagus* using original or additional materials (indicated by “*”), or on a bibliographical basis (unless otherwise stated, data were taken from the original descriptions of species): *P. oenigenisi* (König, 1825); *P. saurus* (Wagner, 1829); *P. vasconiensis* Viret in Roman & Viret, 1930; *P. savagei* Berzi, 1967; *P. crusafontii* López Martínez in López Martínez & Thaler, 1975; *P. depereti* López Martínez in López Martínez & Thaler, 1975; *P. michauxi* López Martínez in López Martínez & Thaler, 1975; *P. tobiens* López in López Martínez et al., 1977; *P. sorbinii* Masini, 1989; *P. forsi* López...
Martínez & Sesé in Álvarez-Sierra et al., 1990; P. caucasicus Averianov & Tesakov, 1998; *P. italicus* Angelone, 2008a; *P. latiuncinatus* Angelone & Čermák, 2015; and *P. pannonicus* Angelone & Čermák, 2015. Other metric data were taken from Álvarez-Sierra et al. (1990), Tesakov & Averianov (2002), and Angelone & Veitschegger (2015).

**ABBREVIATIONS**

**Institutions**
- GLI: Institute of Geology of the Czech Academy of Sciences, Prague;
- MFGI: Magyar Földtani és Geofizikai Intézet [Hungarian Institute of Geology and Geophysics], Budapest;
- SMF: Senckenberg Museum, Frankfurt;
- ZPAL: Institute of Palaeobiology, Polish Academy of Sciences, Warszawa.

**Teeth**
- D/d: Upper/lower deciduous teeth;
- I/i: Upper and lower incisors;
- M/m: Upper/lower molar;
- P/p: Upper/lower premolar.

**Measurements**
- AA: Partial width;
- L: Length;
- Ltrig: Trigonid length;
- PH: Hypofoxxius width;
- TH: Distal hyperloph length;
- W: Width;
- Wtl: Third lobe width of m2;
- Wtal: Talonid width;
- Wtrig: Trigonid width.

**Other abbreviations**
- Dp-d: (L of d3)-(L of p3) [mm];
- Kd/p: ((L of d3)/(L of p3))*100 [%];
- Md/l: ((L of alveolar row)/(L of diastema))*100 [%];
- M̄H: (((H of mandible at p3 + H of mandible at m2)/2)/L of alveolar row)*100 [%];
- N: Number of specimens;
- OR: Observed range;
- ̄X: Arithmetic mean.

**SYSTEMATIC PALAEONTOLOGY**

Class MAMMALIA Linnaeus, 1758
Order LAGOMORPHA Brandt, 1855
Family PROLAGIDAE Gureev, 1960
Genus Prolagus Pomel, 1853

_Prolagus bilobus_ Heller, 1936 (Figs 2-4)

**NAME-BEARING TYPE AND TYPE LOCALITY.** — Lectotype – left p3 (SMF 1996/136), illustrated in Heller (1936: fig. 3), designed by López Martínez & Thaler (1975), revised by Čermák & Angelone (2013). Gundersheim (probably Gundersheim 1 sensu Tobien 1980; age uncertain, most likely MN15b), Rheinland-Pfalz, Germany (see Čermák & Angelone 2013 for details).

**EMENDED DIAGNOSIS.** — See Čermák & Angelone (2013: 49).

**MATERIAL.** — Gundersheim (probably Gundersheim 1, original material of F. Heller, partly published in Heller (1936): 1 p3 sin, SMF 1996/136; 1 fragment of mandibular body dex with p4/m1, SMF 1994/895; 1 talonid of p4 dex, SMF 1994/896; 1 talonid of p4 or m1 dex, SMF 1994/897; 1 trigonid of p4 or m1 dex, SMF 1996/137; 1 mandibular body dex with p4-m2, SMF 1996/156; 1 mandibular ramus dex with articular head, SMF 1996/157; 1 mandibular ramus dex with p4-m1, SMF 1996/158. Gundersheim-fissure 4 (material published by Fejfar et al. 2006): 2 P2 sin, SMF 2005/1, 2; 1 P3 sin, SMF 2005/3; 1 d3 dex, SMF 2005/4; 1 p3 sin, SMF 2005/5; 1 fragment of mandibular body dex, SMF 2005/6. Gundersheim-Findling: 11 I1 sin, SMF 1985/171–181; 6 I1 dex, SMF 1985/182–187; 2 P2 sin, SMF 1985/70, 71; 5 P2 dex, SMF 1985/72–76; 5 P3 sin, SMF 1985/77, 80, 82, 84, 85; 7 P3 dex, SMF 1985/78, 79, 81, 83, 86–88; 8 P4 sin, SMF 1985/89–96; 12 P4 dex, SMF 1985/97–108; 1 fragment of P4 sin, SMF 1985/242; 4 fragments of P4 dex, SMF 1985/240, 241, 243, 244; 9 M1 sin, SMF 1985/109–112, 115, 124, 126, 247; 17 M1 dex, SMF 1985/113, 114: 116–123, 125, 141, 142, 245, 246, 249, 255; 4 M2 sin, SMF 1985/129, 133, 135, 140; 12 M2 dex, SMF 1985/127, 128, 130–132, 134, 136–139, 178, 179; 3 enamel fragments of upper molariforms, SMF 1985/252–254; 6 fragments of mandibular bodies dex with various teeth, SMF 1985/165–170; 1 i1 sin, SMF 1985/188; 6 p3 sin, SMF 1985/144–146, 160, 161, 163; 11 p3 dex, SMF 1985/143, 147–153, 159, 162, 164; 5 fragments of p3, SMF 1985/154–158; 9 p4/m1 sin, SMF 1985/189–196, 234; 6 p4/m1 dex, SMF 1985/215–220; 6 trigonids of p4/m1 sin, SMF 1985/204–206, 209, 223, 235; 4 trigonids of p4/m1 dex, SMF 1985/221, 225, 236, 237; 3 talonids of p4/m1 sin, SMF 1985/207, 208, 238; 2 talonids of p4/m1 dex, SMF 1985/222, 224; 1 m2 sin, SMF 1985/210; 1 m2 dex, SMF 1985/226; 1 trigonid of m2 sin, SMF 1985/211; 1 trigonid of m2 dex, SMF 1985/228; 3 talonids of m2 sin, SMF 1985/212–214; 4 talonids of m2 dex, SMF 1985/227, 229, 230, 239; 2 m2 sin, SMF 1985/232, 233; 2 D3/4 sin, SMF 1985/268, 269; 2 d3 sin, SMF 1985/256, 257; 5 d3 dex, SMF 1985/258–262; 4 d4 sin, SMF 1985/263–266; 1 d4 dex, SMF 1985/267. Raciszyn 1: 1 mandibular body dex with complete dentition, ZPAL M.10.

**MEASUREMENTS.** — See Tables 1 and 2, Fejfar et al. (2006), and Čermák & Angelone (2013).

**STRATIGRAPHIC AND GEOGRAPHIC DISTRIBUTION.** — Early Pliocene (late Ruscinian, MN15b) of central Europe: localities Gundersheim (probably Gundersheim 1 sensu Tobien 1980; original material of F. Heller), Gundersheim-fissure 4, Gundersheim-Findling, possibly Wölfersheim (all in Germany) and Raciszyn 1 (Poland).

**DESCRIPTION**

*d3* (Fig. 2A, B)

Three-lobe tooth; the anterocorid is always isolated (N = 7), shaped as an elongated triangle; it may bear an anterior and/or posterior undulation in the lingual side; in one case there is a connection of the anterocorid and the accessory cusp, marked by a deep anteroflexid. A small, roundish cusp is present in the antero-lingual side, between anterocorid and protocorid, in 86% of cases (N = 7). The trigonid is indented by an average-sized to deep centroflexid; the protoconulid is quite reduced, as well as the metaconid. A centroisthmus between trigonid and talonid is present in 40% of cases (N = 5). In
the talonid, the entoconid is much smaller than the globous hypoconid, which appears to be the largest cusp, sometimes even larger than the anteroconid.

d4 (Fig. 2C, D)
Trigonid and talonid are always well separated. The trigonid is of equal size or slightly smaller than the talonid; it bears a quite developed, flattened anterior lobe, separated from the main body of the trigonid by a simple, straight paraflexid (no centroflexid). The triangular talonid lacks or has a very reduced anterior isthmus.

p3 (Fig. 2E [in part], F-N)
The occlusal features of adult teeth (identified mainly by their prismatic shaft) are the following: large size; the anteroconid is as large as metaconid (92%, N = 12) – in one case, it is slightly smaller; the anteroconid is roughly shaped as a right-angled triangle and appears “tilted” (i.e., with the 90° angle pointing posteriorly); the main body of the anteroconid is always displaced towards the lingual side (N = 13), its posterior and labial sides with undulation or notch in 69% (N = 13); the metaconid is quadrangular, with enamel undulated and/or crenulated all around; the crenulation may interest also the anterior side of the entoconid, which is always quite thick (length > 1/5 of total L; N = 11) and without enamel hiatus; the mesoflexid is very variable, from relatively short and U-shaped to quite long and J-shaped (N = 12) – in one case, the mesoflexid connects to the centroflexid, isolating the metaconid, and in 2 cases it hosts an additional enamel islet in its lingual side (such islet in one case connects to the entoconid with wear, in the other case it remains isolated); the crochet is present in all adult specimens, predominantly with a very large size (in 50%; large in 36%, small in 14%; N = 14); the centroflexid has undulated sides (71%, N = 14), parallel in ¾ of its length, then widens in correspondence of the crochet – apart for the sporadical above mentioned connection with the mesoflexid, the centroflexid may be anteriorly “closed” by a connection of proto- and metaconid; the metaisthmus varies from relatively wide (2 cases) to very thin (N = 14), when it is thin, it may have a bottlenecked appearance; the protoisthmus is always thin to very thin (N = 13) and its connection with the protoconid has a “bottlenecked” appearance only in 23% (N = 13); the hypoconid is triangular in shape and predominantly of average size (only in two cases it is quite large); the hypoflexid is very deep, almost reaching the posterior side of the tooth; the anterior part of the protoconid is quite protruding lingually, and the passage to the thin posterior part is marked by an abrupt narrowing; the protoconid is connected to the protoconulid by a wide to very wide loph, at the junction protoconid/protoconulid there may be a spur in about half of the cases; the protoconulid is quite short, often very wide with a bulky appearance.

The occlusal features of juvenile teeth (identified mainly by their conical shaft; N = 6) are the following: the anteroconid, always displaced towards the lingual side, is medium-sized (smaller than metaconid in 67%), rather flattened oval in shape, distally smooth (in one case lingually with fold); the metaconid is mainly triangular, in one case is isolated; the crochet may be absent (1 case), very reduced or of average size, straight or inclined with respect to the antero-posterior axis of the tooth, and its position also varies from central to displaced (there is no univocal correspondence between its size, shape and position); the hypoconid is large compared to the size of the tooth, triangular or with rounded lingual edge; the hypoflexid is deep, as in adult individuals; the mesoflexid is predominantly V-shaped and short; the centroflexid shape is similar to that of adults, with undulated sides in 33%; the protoisthmus is thin to very thin, often bottlenecked (50%); the metaisthmus varies from thin to average wide; the protoconid has the same shape as in adults (developed anteriorly, reduced posteriorly), however the anterior part may be more or less developed towards the lingual side; the proportions protoconid-protoconulid are variable, but in general the protoconulid is usually thin, though not very long.

p4-m2 (Fig. 2E [in part], O)
The p4-m1 consist of two separate lobes of approximately similar width fused together by cement; posteriorly, both

---

### Lower and upper teeth

| Tooth | N | L | R | OR | CV |
|-------|---|---|---|----|----|
| p3    | 10| 2.47| 2.20-2.77| 7.96|
| p4/m1 | 13| 1.95| 1.69-2.13| 7.11|
|       | 22| 1.12| 0.90-1.32| 9.41|
|       | 23| 2.13| 1.88-2.47| 8.64|
| m2    | 3 | 2.65| 2.45-2.81| –   |
|       | 6 | 2.02| 1.71-2.19| 9.09|
|       | 7 | 1.99| 1.82-2.25| 7.36|
|       | 19| 2.09| 1.85-2.39| 7.73|
| m3    | 5 | 1.12| 1.00-1.21| 7.94|
|       | 11| 1.16| 1.05-1.25| 6.88|
| P2    | 5 | 1.69| 1.60-1.82| 5.52|
|       | 6 | 2.64| 2.50-2.87| 6.97|
| P3    | 10| 2.04| 1.71-2.21| 7.08|
|       | 9 | 3.45| 2.85-3.84| 8.92|
|       | 12| 2.36| 1.70-2.76| 12.77|
| P4    | 16| 1.65| 1.36-1.85| 8.73|
|       | 17| 0.98| 0.80-1.17| 10.79|
|       | 11| 3.23| 2.79-3.80| 9.98|
|       | 11| 2.87| 2.42-3.31| 11.01|
| M1    | 12| 1.45| 1.19-1.69| 11.61|
|       | 19| 1.66| 1.12-1.95| 13.64|
|       | 18| 3.07| 2.30-3.51| 12.39|
|       | 19| 2.70| 1.94-3.04| 11.76|
|       | 19| 2.42| 1.70-2.90| 12.70|
| M2    | 13| 1.51| 1.25-1.80| 11.03|
|       | 12| 2.84| 2.29-3.54| 13.32|
|       | 13| 2.44| 1.94-2.99| 13.33|
|       | 13| 2.21| 1.80-2.69| 12.99|
| D3/4  | 2 | –   | 1.21,1.29| –   |
|       | 2 | –   | 1.75,1.96| –   |

---

**Table 1.** Dental measurements of Prolagus bilobus Heller, 1936 from the early Pliocene (MN15b) locality Gundersheim-Findling. See Material and methods for abbreviations.
lobes have thickened enamel. The trigonids are of rhomboid shape, longer than talonids, with an accentuated anterior step; the talonid shows an antero-labial flexid, and a shallow notch in the labial edge; the m2 are three-lobed, the first two similar to p4-m1, whereas the last lobe is simple, triangular-shaped.

Lower jaw (Fig. 4)
Eleven fragments of mandibular bodies in different ontogenetic stages are available. In adult individuals, the mandible is rather robust. The diastema is long, but slightly shorter than the p3-m2 alveolar row (Table 2). The posterior mental foramen is large, oval, located close to the ventral edge of the mandibular body below m1 (in most cases, below the trigonid/talonid connection; c. 67%, N = 6), and remarkably maintains the same position both in juvenile and adult specimens. The area below p3-p4 is richly fenestrated. The lower incisor extends below the p4 (in most cases below the trigonid/talonid connection; c. 50%, N = 6). The ventral margin of the mandible is convex, and becomes more prominent and angular with increasing ontogenetic age. The mylohyoid line is moderately developed, and in ventral view it appears located near the buccal edge of the mandibular body. The coronoid process is weakly developed. The articular head is strongly convex in medio-lateral direction.

D3-4 (Fig. 3X, Y)
Three roots. The mesial hyperloph becomes longer with wear; in less worn specimens it covers the lagiloph, and then “grows” to cover the entire postcone. The mesial hyperloph has a notch on the posterior edge. The mesial hypercone is much less developed than the distal one, both in size and width. Consequently, the hypoflexus appears as a very open “V”. Both para- and mesoflexus are very deep, and their relative depth can vary; the paraflexus can be deeper than the mesoflexus or viceversa. The lagicone is small and thin; its connection with the very short lagiloph is bottlenecked. The postcone is rounded, with inclined axis and with a developed postlobule.
**P2 (Fig. 3A-E)**
Quite flattened in shape. The mesial hyperloph is long, covering 67-78% (N = 6) of the tooth width, often robust, and always without enamel hiatus. The lagicone is straight or very slightly inclined lingually, variable in thickness, and slightly longer than postcone in 83% (N = 6), in one case with an undulation in its postero-lingual side. Para- and mesoflexus are deep and curved, with the latter longer than the former.

**P3 (Fig. 3F-J)**
The mesial hyperloph is of average thickness, slightly widened at the distal end, always without enamel hiatus (N = 10). The mesial hypercone is lingually less protruding than the distal one. The hypoflexus is V-shaped, from very shallow to shallow. The para- and mesoflexus are deep, curved, of equal depth or sometimes with the paraflexus as the deepest of the two. The lagicone-lagiloph complex is very thin in most cases, the lagicone-lagiloph connection is marked by a sharp change of the curvature. The lagiloph covers half or the entire postcone (in any case, never reaching the labial side of the tooth). The postcone is rounded or quadrangular, connected posterolabially to a postlobule of very variable size.

**P4 (Fig. 3K-O)**
The mesial hyperloph is more protruding lingually than the distal one, but it is much shorter (TH, i.e., its average length, is 40% of tooth L). The hypoflexus reaches about 45% of tooth W (50% of AA) and never merged with parafossette (N = 21), it is V-shaped in its lingual part, then becomes narrow U-shaped in 62% (N = 21); the U-shaped part also has a different inclination with respect to the other part, i.e., pointing anteriorly. The parafossette is long to extremely long in its anterior part, slightly inclined towards the anterior side of the tooth in most cases, and its width remains constant all along, except for a few cases, in which the anterior end is slightly widened; sometimes the anterior end of the parafossettes covers and surpasses the mesofossette (18%; N = 22). The mesofossette is variable in size from large to very large, and in shape from V-shaped (predominant), to U-shaped and C-shaped.

**M1 (Fig. 3P-T)**
The mesial hypercone is more protruding than the distal one. The hypoflexus is very long (c. 79% of W; c. 90% of AA), curved posteriorly or straight. The fossettes are predominantly present; in 92% of cases (N = 26) there is one fossette, in 4%...
of cases two fossettes, and in the remaining cases (4%) fossettes are absent. The fossettes are positioned just under the distal end of the hypoflexus; they vary in shape from round to slightly oval (42%; N = 24) to elongated (58%).

**M2 (Fig. 3U-W)**

The precone is well-developed, the postlobule reduced. The mesial hypercone is slightly more protruding than the distal one. As with M1, the hypoflexus is very long (c. 78% of W; c. 91% of AA) and curved posteriorly or straight. The fossettes are predominantly absent (93%; N = 15); when present, they are round and rather small.

**Taxonomic comparison and remarks**

*Prolagus bilobus* has recently been partly revised by Čermák & Angelone (2013). The species is clearly differentiated by its very large p3 (Fig. 5) with lingually shifted, large, triangular, posteriorly, and buccally indented anteroconid; undulated sides of centroflexid; long crochet; metaconid of similar size as anteroconid; thick entoconid lacking enamel hiatus. The p3 of *P. bilobus* has a very homogeneous and stable morphology, as far as the anteroconid, crochet and entoconid are considered: all the adult individuals have a lingually displaced anteroconid with a unique “tilted” appearance, a voluminous crochet, and a thick, crenulated entoconid. These features seem unaffected by the intraspecific variability range. The p3 differs from that in: 1) *P. pannonicus*, *P. sorbinii*, *P. latiuncinatus*, *P. caucasicus*, *P. michauxi*, *P. ibericus*, and *P. savagei* in having a significantly larger size (Fig. 5); 2) *P. pannonicus* and *P. caucasicus* in having a larger and wider anteroconid; 3) *P. sorbinii*, *P. ibericus*, *P. depereti*, *P. savagei*, and *P. italicus* in having a larger crochet; 4) *P. caucasicus* in having a quadrangular metaconid; and 5) *P. michauxi* and *P. sorbinii* in having a triangular indented anteroconid, vertically aligned with the entoconid.

The availability of additional materials allowed us to highlight some peculiarities of the species also in other tooth positions:

- d3 with its isolated labial cusp and predominance of separated talonid also in advanced stages of wear. This feature, which recalls *P. crusafonti* (see Angelone 2007: table 2, p. 413), can be considered as quite primitive. In fact, latest Miocene and Pliocene species (as far as we know, taking into account the scarce availability of fossil lagomorphs deciduous teeth and of detailed studies on the subject) are characterized by a connected talonid.

- d4 with a very deep paraflexid but not a centroflexid, as in late Miocene congeneric populations of western Europe.

- P2 with generally flattened appearance and very long mesial hyperloph. A flattened P2 is a common and stable feature of *P. bilobus*, but it is instead quite variable in other Pliocene species. The mesial hyperloph is remarkably longer compared to the Pliocene species *P. pannonicus*, *P. latiuncinatus*, and *P. ibericus*.

- P3 with centrocone not reaching the labial edge and P4 with long, upturned parafossette. Both characters are in common with the sorbinii group, providing a hint about the possible phylogenetic affinity of *P. bilobus*.

- P4 with mesial hyperloph remarkably thin. The very thin dentine bridge between parafossette and hypoflexus in P4 of *P. bilobus* can be well observed in *P. pannonicus* and *P. latiuncinatus*; in *Prolagus* sp. from Beremend 39 these structures may even be confluent.

Tesakov & Averianov (2002) attributed to *P. bilobus* scantly materials of isolated p3s from Tanatary (MD, MN15; 1 p3, formerly attributed to *P. cf. oeningensis* by Erbajeva &
Čermák S. et al.

Shushpanov (1988) and Erbajeva (1988)), Tatareshty (MD, MN15; 1 p3), Kotlovina (UA, probably MN15 or MN16 according to Nesin & Nadachowski (2001); 1 p3, formerly attributed to *P. cf. oenensis* by Agadjanian & Erbajeva (1983)), and Kamenskoe (UA, age unclear, probably MN15; 1 p3, formerly attributed to *Prolagus* sp. by Topachevsky (1962)).

These p3 are characterized by: 1) a small, not tilted anteroconid shifted towards the lingual side of the tooth; 2) a reduced, diamond-shaped metaconid with unfolded enamel, and 3) a reduced entoconid. Thus, they are quite different from *P. bilobus*. In the specimens from Tatareshty and Kotlovina, the crochet is absent or quite reduced, contrarily to adult specimens of *P. bilobus*. The measurements, available only for Tatareshty (L × W = 1.95 × 1.93; Tesakov & Averianov 2002) and Tanatary (L × W = 1.80 × 1.70; Erbajeva & Shushpanov 1988), indicate that the Moldavian specimens are sensibly smaller in size than *P. bilobus*. The overall morphology of the p3 of Ukrainian and Moldavian materials described above reminds the appearance of some populations of *P. sorbinii* (see Angelone 2007), a species distributed in Greece and Italy in the late Miocene and earliest Pliocene (Angelone & Rook 2012). Thus, those remains are tentatively attributed here to *P. aff. sorbinii*.

For the sake of completeness, it is worth to mention that Tesakov & Averianov (2002) tentatively referred to *P. bilobus* the population from Polgárdi (HU, MN13), a hypothesis refuted by Angelone & Čermák (2015) who assigned the sample to the new species *P. latiuncinatus*.

To conclude, at present we include only the localities of Gundersheim and Raciszyn 1 in the proven *P. bilobus* distribution, as the remains from Wölfersheim and Kisláng are at present not available (see above for details).

### Table 2

Mandibular measurements of *Prolagus bilobus* Heller, 1936: 1, Gundersheim-Findling, SMF 1985/165; 2, Gundersheim (probably Gundersheim 1 sensu Tobien 1980), SMF 1996/158, unpublished material of F. Heller; 3, Raciszyn 1, ZPAL M.10.

| Measurements                                      | 1    | 2    | 3    |
|--------------------------------------------------|------|------|------|
| Lingual height of mandible at p3                 | 4.89 | 7.13 | 9.09 |
| Lingual height of mandible at p4                 | 6.50 | 9.01 | 12.98|
| Lingual height of mandible at m2                 | 6.23 | 8.66 | 12.25|
| Length of diastema                               | 5.75 | –    | 11.14|
| Alveolar length of p3-m3                          | 7.95 | 9.39 | 12.04|
| Mandible width at p4                              | 3.31 | 4.48 | 5.82 |
| Mandible width at m2                              | 2.53 | 3.56 | 4.29 |

**For the sake of completeness, it is worth to mention that**

Tesarov & Averianov (2002) tentatively referred to *P. bilobus* the population from Polgárdi (HU, MN13), a hypothesis refuted by Angelone & Čermák (2015) who assigned the sample to the new species *P. latiuncinatus*.

**To conclude, at present we include only the localities of**

Gundersheim and Raciszyn 1 in the proven *P. bilobus* distribution, as the remains from Wölfersheim and Kisláng are at present not available (see above for details).

### Prolagus bilobus

- Gundersheim (probably 1) - DE (?MN15b), Lt
- Gundersheim-Findling - DE (MN15b)
- Raciszyn 1 - PL (MN15b)

### Prolagus sp.

- *Prolagus italicus* (Montagnola Senese - IT, MN17)
- *Prolagus savagei* (Arondelli - IT, MN16a)
- *Prolagus depereti* (Perpignan - FR, MN15)
- *Prolagus ibericus* (Zeta - ES, MN15)
- *Prolagus michauxi* (Sète - FR, MN15a)
- *Prolagus caucasicus* (Kosyakino - RU, MN14)
- *Prolagus sorbinii* (Monte Castellaro - IT, MN13)
- *Prolagus latiuncinatus* (Polgárdi - HU, MN13)
- *Prolagus pannonicus* (Sümeg - HU, MN10/11)

**For the sake of completeness, it is worth to mention that**

Tesarov & Averianov (2002) tentatively referred to *P. bilobus* the population from Polgárdi (HU, MN13), a hypothesis refuted by Angelone & Čermák (2015) who assigned the sample to the new species *P. latiuncinatus*.

**To conclude, at present we include only the localities of**

Gundersheim and Raciszyn 1 in the proven *P. bilobus* distribution, as the remains from Wölfersheim and Kisláng are at present not available (see above for details).
is very narrow, bottleneck-shaped. The crochet is extremely feeble, in central position, symmetrical in shape. The centroflexid is straight, parallel to the antero-posterior axis of the tooth, with undulated sides, and enlarged at its distal end, in correspondence of the crochet. The hypoconid is small, triangular in shape, with flattened labial end. The protoconid is wide in its anterior part, narrowing towards the posterior part of the tooth in its distal part. The connection protoconid/protoconulid is marked by a spur. The protoconulid is relatively small, of average width.

p4-m2 (Fig. 6E [in part])
In p4-m1 trigonid and talonid have similar width and both show a thickening of the enamel on the posterior side. The trigonids are of rhomboid shape, longer than the talonids. In p4 the trigonid has an anterior step. The talonid shows an antero-labial flexid, more marked in p4, less marked in m1 and even less in m2. The three-lobed m2 have trigonid and talonid similar to other molariform teeth, plus a triangular-shaped posterior lobe.

Lower jaw (Fig. 6A-D)
The outline is slender in both lateral and ventral views; the diastema is long, and its length is almost equal to the alveolar one. The posterior mental foramen is large, oval, located close the ventral edge of mandibular body in the area below m1/ m2. Several accessory foramina are present below p3-p4. The lower incisor extends below the trigonid of p4. The ventral margin of the mandible is convex. The mylohyoid line is long, well-developed, and in ventral view appears located near the lingual edge of the mandibular body. The coronoid process is moderately developed. The articular head is strongly convex in medio-lateral direction.

P3 (Fig. 7E [in part])
The mesial hyperloph is thin, of constant width, without enamel hiatus, and covers part of the postcone. The mesial hypercone is less protruding than the distal one. The distal hyperloph, however, is very reduced in length. The V-shaped hypoflexus is short. Para- and mesoflexus have the same depth, and reach posteriorly almost the edge of the tooth. The lagicone is of average size and rounded, the lagiloph has no indentations or steps, and the centrocone does not reach the labial end.

P4 (Fig. 7D [in part], E [in part])
The mesial hyperloph is slightly longer than the distal one; the length of mesial- and distal hyperloph is similar. The hypoflexus is V-shaped in its lingual part, then takes a very narrow U-shape; in one individual the hypoflexus is connected to the parafossette, in the other, only a very thin enamel bridge separates the hypoflexus and the parafossette. The parafossette is very long, thin, and its anterior part may be upturned anteriorly; the mesofossette is C- or V-shaped, with the anterior tip slightly longer than the posterior one.
The sole specimen from Beremend 26, represented by taxOnOmic cOmParisOn and remarks

direction. The infraorbital canal is short; in relation to the
bital foramen is oval, prolonged only slightly in dorsoventral
masseteric spine is short and triangular in shape. The infraor-
posteriorly accompanied by several smaller foramina. The
the maxillopalatine suture, there is a large palatine foramen,
foramen is large, oval, positioned medially to P4. Next to
P . caucasicus in having a larger size; (Fig. 5); 2) P . depereti, P . savagei, and
P . bilobus in having a smaller size (Fig. 5); 3) P . pannonicus and
P . caucasicus in having a larger and wider p3 anteroconid; and
4) P . pannonicus, P . latiuncinatus, and P . bilobus in having a
significantly less developed crochet.

Surprisingly, Prolagus sp. from Beremend 26 is morphologi-
cally and dimensionally incompatible with the coeval P . bilobus.
The p3 of Prolagus bilobus has an extremely stable morphol-
yogy (see section “Taxonomic comparison and remarks” about
of P . bilobus). This characteristic makes impossible to “fit in” in
P . bilobus the lower jaw from Beremend 26, which p3 has a
central, not tilted anteroconid, an extremely small crochet,
and a medium-sized entoconid. Moreover, the lower jaw of
Prolagus sp. from Beremend 26 has a different outline with
respect to P . bilobus, especially in the ventral part (which is
regularly convex and not angular), and in the size and posi-
tion of the posterior foramen (smaller and positioned further
posteriorly). Also the measurements of the p3 and of the lower jaw of Prolagus sp. from Beremend 26 are incompatible with
an attribution to P . bilobus. In fact, though pertaining to an
old individual, the measurements of p3 and lower jaw of Prola-
gus sp. from Beremend 26 fall below the distribution of adult
P . bilobus (Fig. 5; Table 3). However, the upper teeth pertaining
to a jaw excavated from Beremend 39, another karst filling of
the Beremend karst complex, which apparently is coeval to
Beremend 26, at least judging from the accompanying fauna
(Pongrácz in litt. ; Čermák & Wagner 2013), do not show
substantial differences in morphology and measures compared
to P . bilobus (cf. Tables 1; 3). This taxonomic incongruence
between materials from coeval and neighbouring fissure fillings
does not find an explanation for the moment and, though in
our opinion Prolagus from Beremend 26 represents a distinct
species, we provisionally keep all the available material from
Beremend in open nomenclature as Prolagus sp.

It is really tempting to suppose that the p3 and only Prolagus
specimen from Kisláng (Kretzoi 1954) may be related to Prola-
gus sp. from Beremend 26. Indeed, the geographical proximity
of the fossil sites, and the age range of the mixed assemblage
of Kisláng (MN15-MQ1; Mayhew 2012) are compatible with
this hypothesis. However, the morpho-dimensional features of
the p3 remain from Kisláng described in literature are too
unclear to make any taxonomic speculation. Kretzoi (1954: 247)
provided only indications about the presence of a crochet, and
hypothesized a taxonomic proximity of the Kisláng specimen
to Prolagus from Gundersheim. This is in contrast with our
observation about the lower series from Beremend 26 (totally
different from Gundersheim). We envisage the retrieval of the
Prolagus materials from Kisláng, in order to solve this open issue.

Table 3. — Cranial and dental measurements of Prolagus sp. from the early Pliocene (MN15b) localities Beremend 26 and 39. See Material and methods for abbreviations.

| Mandible and maxillae | Lower teeth | Upper teeth |
|-----------------------|-------------|-------------|
| Total height of mandible | 21.37 | P3 | L 1.89 |
| Lingual height of mandible at p3 | 6.46 | P3 | L 2.04 |
| Lingual height of mandible at p4 | 8.18 | W 1.91 |
| Lingual height of mandible at m2 | 7.92 | P4 | L 1.76 |
| Length of diastema | 8.18 | Ltrig 1.05 |
| Coronar length of p4-m2 | 6.08 | Wtrig 1.93 |
| Alveolar length of p3-m2 | 9.16 | W 2.68 |
| Mandible width at m2 | 3.20 | m1 | L 1.79 |
| Mandible width at p4 | 4.34 | Ltrig 1.05 |
| Width of articular head | 3.95 | Wtrig 2.01 |
| Alveolar length of P2-M2 | 9.54 | W 2.85 |
| Length of P2 alveolus | 1.95 | W 2.54 |
| Width of P2 alveolus | 2.86 | PH 2.16 |

M1 (Fig. 7D [in part], E [in part])
The mesial hypoflexus is rather thin, longer than the distal
one. The hypoflexus is rather long (c. 78% of W, c. 88%
of AA; N = 2), posteriorly curved; in 1 of 2 individuals, a
round, small fossette is present posteriorly to the labial tip
of the hypoflexus.

M2 (Fig. 7D [in part], E [in part])
The general features are the same as M1, with a relatively
longer hypoflexus (c. 82% of W, c. 92% of AA; N = 1) and
no fossette.

Upper jaw (Fig. 7A-C)
The hard palate is of average width. The posterior edge of the
incisive foramen reaches the posterior edge of P2. The ante-
rior edge of the choanae reaches the half of P4. The premolar
foramen is large, oval, positioned medially to P4. Next to
the maxillopalatine suture, there is a large palatine foramen,
posteriorly accompanied by several smaller foramina. The
masseteric spine is short and triangular in shape. The infraor-
bital foramen is oval, prolonged only slightly in dorsoventral
direction. The infraorbital canal is short; in relation to the
wall of P2 alveolus, placed rather dorsally.

Taxonomic comparison and remarks

The sole Prolagus specimen from Beremend 26, represented by
an almost complete mandible, is extremely peculiar. Evident
differences can be observed with some late Miocene
congeneric species (P. sorbini i P. latiuncinatus), namely
in its very large triangular anteroconid of p3. Prolagus sp.
from Beremend differs from the Pliocene-Holocene species
P. caucasicus, P. michaetxi, P. ibericus, P. depereti, P. figaro,
P. bilobus, P. savagei, P. italicus, and P. sardus, as its anteroconid
is not displaced towards the lingual side. Moreover, the
p3 of Prolagus sp. from Beremend 26 differs from that in:
1) P. pannonicus, P. latiuncinatus, P. caucasicus, and P. ibericus
in having a larger size; (Fig. 5); 2) P. depereti, P. savagei, and
P. bilobus in having a smaller size (Fig. 5); 3) P. pannonicus and
P. caucasicus in having a larger and wider p3 anteroconid; and
4) P. pannonicus, P. latiuncinatus, and P. bilobus in having a
significantly less developed crochet.
RESULTS AND DISCUSSION

NOTES ON THE ONTOGENY OF *P. bilobus*

Mandibles

The available jaws of *P. bilobus* cover various ontogenetic stages (Fig. 4) and show a noticeable size range (Table 2), typical of lagomorphs. Several allometric changes distinguish juvenile and adult individuals (see Fig. 4):

- the diastema is relatively shorter (compared to the alveolar row length) in the juvenile mandible (SMF1985/165) than in the unbroken aged specimen (ZPAL M.10). The values of MdL are 138% vs 108%, respectively.
- the mandibular body is relatively lower (compared to alveolar row length) in the juvenile mandible (SMF1985/165) in contrast to that in the adult (SMF 1996/156) and aged (ZPAL M.10) ones. The values of MH are 70 vs 84 and 89%, respectively (to exclude the influence of ventral curvature of the mandibular body, the height was calculated as the mean value of measurements at p3 and m2).
- a relative height increase of the mandibular body is associated with the curvature increase of its ventral margin; this phenomenon was also observed in the well-known and documented species *Prolagus sardus* (Tobien 1935: Fig. 2, p. 272).

The above described ontogenetic variations observed in the jaws of *P. bilobus* are quite standard for the genus. In *Prolagus*, though, also the position of the jaw posterior foramen slightly varies during the ontogeny, but in *P. bilobus* this feature remains unchanged in all the available specimens.

**Teeth**

In the genus *Prolagus*, the morphology of juvenile permanent teeth is quite different from the adults of the same species. This does not just depend by the fact that the cusps may be unworn: in fact, especially in p3, the relative size of the cusps, their connection, their position, and also their presence, are prone to noticeable variations during the ontogeny. Due to the progressive inclination of the wear surface (W) and the conical shape of the juvenile tooth shaft (L and W), also the size ranges of the teeth show a wide variation if juveniles are included in the measurements. It has even been observed, since the early studies about this taxon, that in some tooth positions the measurements cluster in two very distinct clouds, one for permanent juvenile teeth and one for adults (López Martínez 1974).

Also in the case of *P. bilobus*, it is possible to notice the remarkable ontogenetic changes in p3. Moreover, in this case, thanks to the stable occlusal surface morphology which characterizes this species (see “Taxonomic comparison and remarks” of *P. bilobus*) and thanks to a sufficient number of specimens for both ontogenetic categories (6 conical p3 in the initial stage of abrasion = permanent juvenile teeth; and 14 p3 with prismatic shafts = permanent adult teeth), we were able to observe in the occlusal and “radical” parts of the teeth the following pattern (young to old):

- the enlargement, deformation, and tilting of the anteroconid;
- the deformation of the metaconid;
– the noticeable thickening of the protoconulid;
– the thickening of the entoconid;
– the appearance or the enlargement of the crochet.

The ontogenetic changes observed in *P. bilobus* can not represent a pattern perfectly applicable to other congeneric species. However, it would be worth to verify if the patterns of ontogenetic trajectories are similar in phylogenetically close species.

It is very interesting to notice the extreme ontogenetic dimensional change between the deciduous and the permanent teeth of *P. bilobus* (Fig. 8). Indeed, López Martínez (1974) noticed that the relative proportion of the length of d3 and p3 varies through time in the species of the genus *Prolagus*, but this observation had no further development in literature. Thirteen selected fossil localities ranging in age from the early Miocene (MN2b) to the Late Pleistocene with sufficient materials of d3/p3 were selected to verify and further investigate López Martínez’s (1974) remark. Materials from different ages, but also from different geographical areas were selected, in order to exclude possible biases. It appears (Fig. 8) that in early Miocene to early middle Miocene species, Ld3 is larger than Lp3 (mean value of the $K_{dp}$: 110–123%), as occurs in *Prolagus*’ ancestor *Piezodus* Viret, 1929 (see López Martínez 1974). Since the late middle Miocene, the size ratio is reversed, i.e., Lp3 is larger than Ld3. Moreover, it is remarkable that after the reversal of size proportions, the size of d3 not only remains small, but also relatively stable. The size of p3, on the contrary, though showing a general increase, is variable, consequently the size differences between d3 and p3 are also quite variable. The mean value of $K_{dp}$ ranges from 98% in *P. tobieni* from the MN7/8 of Escobosa to 55% in *P. bilobus* from the MN15b of Gundersheim-Findling, which is the species with the largest d3/p3 size difference among the studied taxa (Fig. 8). The percentage of size difference between d3 and p3 in the analyzed taxa does not evidence a gradual trend depending on the geologic age, as hypothesized by López Martínez (1974: 142). Of particular interest in this sense, is to observe that two distinct species, *P. tobieni* and *P. oeningensis*, recorded from the same locality (Escobosa, Spain), show a quite different $K_{dp}$. It can not be excluded that, within a mosaic evolutionary pattern, which is commonly observed in *Prolagus*, the size differences between d3 and p3 may carry a signal of phylogenetic closeness of species within distinct clades. At any rate, this parameter may be useful for species discrimination (cf. Álvarez-Sierra et al. 1990: 16), and it may have a biochronological significance, being a real turning point in the evolution of *Prolagus* similarly to:

– the appearance of the centroflexid. The presence of a long centroflexid became a stable feature of the genus after the *P. vasconiensis/oeningensis* transition, near the early/middle Miocene boundary. The only exception is a conservative clade endemic to Spain, whose last representant was *P. tobieni* (still possessing a very incipient centroflexid), a species which survived until the early late Miocene (according to Hordijk & van der Meulen 2010).

– the appearance of the mesial hyperloph in P2 coupled with the sudden increase of Lp3. These features make a pan-European, “simultaneous” appearance between MN12 and MN13 (Angelone 2007; Angelone & Čermák 2015 with references). The only known exception is the MN15 *P. ibericus*, an extremely peculiar species whose phylogenetic history is quite mysterious and characterized by several “incongruent” features compared to coeval ones (first of all the very small size), known only from the type locality Layna (NE Spain; López Martínez 1989). It must be remarked, though, that in spite of all its peculiarities, the $K_{dp}$ of *P. ibericus* is congruent with that of coeval species.

**EVOLUTIONARY TRENDS IN *PROLAGUS* P3**: CENTRAL VS WESTERN EUROPEAN PLEISTOCENE SPECIES

**Size**

The dental size (p3) of European prolagids shows a general increase trough time, with a few countertrending episodes (e.g. *P. oeningensis* → *P. crusafontii*; López Martínez 1989) and an interesting sudden enlargement in correspondence of the MN15 (Angelone 2007). Until Miocene, though, the size range of the p3 of coeval *Prolagus* species is comparable (with punctual exception as *P. major* in the MN6 of Spain; López Martínez 1989). During the Pliocene, instead, *Prolagus* shows a wide range of size (Fig. 5): from quite small (e.g. *P. ibericus*; Layna, MN15) to very large (e.g. *P. bilobus*; Gundersheim-Findling, Raciszyn 1, both MN15b, which is the largest continental species of the genus).

**Enamel folding and entoconid thickness**

Morphological patterns corresponding to temporal trends were individuated in the p3 of western European Plio-Pleistocene species, namely the simplification of the enamel folding pattern (particularly the loss of the crochet and the smoothing of anteroconid and metaconid folds) and the reduction of the entoconid (Angelone 2008a).

The detailed study of *P. bilobus* and *Prolagus* sp. from Beremend 26 allowed to verify if those trends are shared also by Pliocene *Prolagus* of central Europe. The answer is negative. *Prolagus bilobus* and *Prolagus* sp. from Beremend 26 do not follow the above listed morphological trends, possessing a folded enamel and a thicker entoconid (this feature is particularly evident in the former species). In this sense, they are similar to *P. italicus*, whose peculiarities were explained by Angelone (2008a) as a consequence of its status of peninsular Italian endemic species. Hypothesizing that the morphological analogies between *P. italicus*, *P. bilobus*, and *Prolagus* sp. from Beremend 26 are common to continental isolated species, however, is quite simplistic in our view. In fact, the conditions of isolation (geographical extent, duration, dynamics, and climate) of these species are too different to justify a shared common morphological trend in the teeth occlusal surface. Possibly, some of the similarities shared by *P. bilobus*, *P. italicus*, and *Prolagus* sp. from Beremend 26 and, consequently, their divergence from Pliocene western European species, may derive from a common origin from an eastern European stock. In fact, *P. italicus*, though known only from the MN17 of central Italy, stemmed from the eastern European species *P. torbinii* just after its dispersal into Italy, an event occurred in the Messinian (see cladistics
analysis in Angelone et al. 2015; palaeobiogeographical and taxonomic details in Angelone et al. 2019, 2020). There is no available analysis at present to support the origin of *P. bilobus* and *Prolagus* sp. from Beremend 26 from eastern European species, though it can be a palaeobiogeographically sensible hypothesis. A definitive answer can be obtained only after the revision of *Pliocene Prolagus* species from eastern Europe and Anatolia.

**Entoconid enamel hiatus**

As stated above, an eastern origin for *P. bilobus* and *Prolagus* sp. from Beremend 26 can only be hypothesized, but an encouraging hint in that sense (waiting for a phylogenetic analysis) can be the presence of an enamel hiatus in the entoconid of p3. Angelone (2008a) hypothesized a delayed acquisition of the enamel hiatus in the Italian *P. sorbinii-italicus* stock (in *P. sorbinii* the hiatus is absent, whereas in *P. italicus* only a partial hiatus may be present) in contrast to "the European lineage" i.e., western European species, which indeed represent a separate clade; see Angelone et al. 2015, and in which the presence of the enamel hiatus can be observed since the latest Miocene (MN13, Messinian). However, other studies (Čermák & Angelone 2013; Angelone & Čermák 2015; this study) documented the absence of entoconid enamel hiatus also in the central European taxa *P. latiuscinatus* (Polgárdi 2, MN13), *P. bilobus* (Gundersheim-Findling, Raciszyn 1, both dated to MN15b), and *Prolagus* sp. (Beremend 26, MN15b). This evidence reinforces the hypothesis of a south-eastern European phylogenetic affinity of *P. bilobus* and *Prolagus* sp. from Beremend 26.

**EXPLAINING THE CENTRAL EUROPEAN DISTRIBUTION OF PROLAGUS DURING THE PLIOCENE**

The Pliocene remains of *Prolagus* recorded at the northern border of its geographical range were generally considered to be isolated endemic populations, relics left behind after the reduction of the formerly homogenous Miocene distribution of the genus, a phenomenon that became quite evident since the Pliocene (López Martínez 2001). Nevertheless, in our view, there may be an alternative explanation for the *Prolagus* occurrences in the Pliocene of central Europe.

**Palaeoclimatical background**

The genus *Prolagus* appeared during the early Miocene (MN2b), i.e., towards the end of a period of stable climatic conditions (late Oligocene-early Miocene, MP28-MN3, c. 27-17 Ma; Maridet et al. 2007). In this period, Europe was characterized by a homogeneous mammal fauna with high interregional taxonomic affinities (sensu Raup & Crick 1979). The situation became increasingly heterogeneous since the end of the early Miocene (MN4, c. 17-16.5 Ma). At a larger scale, we assist at a general global cooling trend started about 14 Ma (Zachos et al. 2001). In Europe, an abrupt end of the warm period has been recorded at 14-13.5 Ma (Böhme 2003), and since the middle Miocene, several cooling and warming pulses, as well as several fluctuations in the precipitation rate occurred (Böhme et al. 2008 with references). In particular, the transition to a drier climate and a concomitant cooling episode (c. 9.7-9.5 Ma) determined the biotic crisis known as the Vallesian crisis (but see Casanovas-Vilar et al. 2014). The climatic trend induced a strong latitudinal differentiation in the European climates. In the precipitation regime, a quite evident latitudinal differentiation between northern and southern Europe occurred at c. 8 Ma (Böhme et al. 2008). The study of European faunal assemblages evidences the strong isolation of some regions during the late Miocene. Since the latest Miocene and until the first part of the early Pliocene (MN13-MN14, c. 6.5-5 Ma), a renewed faunal homogeneity shortly occurred in the peri-Mediterranean area of Europe (Maridet et al. 2007).
The early Pliocene is considered to be a period of globally warm climate (De Schepper et al. 2014 with references), though interrupted by short-lived glacial events (c. 4.9/4.8 Ma, and c. 4 Ma). Since the early/late Pliocene transition (3.6 Ma), the relatively stable humid conditions which characterized Europe during the early Pliocene came to an end. The increase in coolness and dryness culminated with a global glaciation at c. 3.3 Ma (MIS M2; see De Schepper et al. 2009 and Tan et al. 2017, both with references). The glaciation, though intense (comparable to early Quaternary glaciations) was short-lived (50 ky) and followed by the mid-Piacenzian Warm Period (c. 3.3-3.0 Ma) (Tan et al. 2017). Though characterized by a climate deterioration if compared to the early Pliocene, the interval 3.6- c. 3.0 Ma was "the last sustained interval in Earth's history when global climate was warmer than today" (De Schepper et al. 2009). Between c. 3.0-2.7 Ma is recorded the onset of the Northern Hemisphere glaciation. Such event, contrarily to the quite ephemeral late Eocene to early Pliocene glaciations (Lunt et al. 2008), is the forerunner of the glacial-interglacial cycles of the Quaternary (Ganopoulos & Calov 2011).

The Pliocene environmental changes impacted on mammalian evolution and dynamics, driving extensive rearrangements of community and faunal structures, including large scale migrations and rearrangements of the selection pressure upon the phenotypic design of particular taxa. A short period near the early/late Pliocene boundary c. MN15-16) represents one of the most dramatic global turning points of the late Cenozoic faunal history. Nearly a half of the Neogene mammalian taxa disappeared at that time, and almost all the elements constituting the modern Quaternary communities appeared in that stage as well (Bernor et al. 1996; Kössner & Heissig 1999; Reumer & Wessels 2003). The aridisation and spread of open ground habitats grew particularly pronounced at the MN16/17 boundary (c. 2.5 Ma), as attested by the loess deposits recognized throughout the northern hemisphere (Rabeder 1981; Rabeder & Verginis 1987; Tedford et al. 1991; Shi 1994; Xue et al. 2006).

A focus on Pliocene palaeoecology of central Europe with emphasis on the distribution of lagomorphs. In the Pliocene of Europe, the climatic changes resulted in a decoupling of the climatic regime of northern and southern Europe. Relatively warmer and arid conditions characterized the Mediterranean area, in contrast to the onset of cooler and humid conditions in the northern parts of central Europe (Popov 2004). The plant record indicates that, in the early Pliocene (before c. 4 Ma), mixed mesophytic forests and broad-leaved deciduous forest prevailed in the studied area (Kovar-Eder et al. 2008). Indeed, Pliocene faunas of central Europe (documented particularly from the localities Gundersheim-Findling, Wèze 1, and Csarnóta 2) contain a large proportion of mesophilous lypoathyphlan and rodent species (>70%; ecogeographical group 1 sensu Popov 2004). This is indicative for an even and humid climate. The other 30% of the assemblages is represented by species with relatively wide tolerances, adapted to a wide range of biotopes under more continental and arid climate (ecogeographical group 2 sensu Popov 2004), inhabiting edge habitats such as dry open forests, bushes, dry meadows, etc. In the Mediterranean area, the genera of group 1 are absent or are very poorly represented, whereas those of group 2 are predominant, together with genera characteristic of dry open habitats (mostly dry savanna dwellers ecogeographical group 3 sensu Popov 2004). See Kretzoi (1956, 1962), Sulimski (1959, 1962, 1964), Kowalski (1960), Repenning (1967), Michaux (1971), Jánossy (1972, 1986), Van de Weerd & Daams (1978), Van de Weerd (1979), Skočzen (1980), Van de Weerd et al. (1982), Reumer (1984, 1989), Van der Meulen & Van Kolfschoten (1986), Fejfar & Storch (1990), Terzea (1997), Šen (1998), Šen et al. (1998), Šen & de Brujin (1998), Popov (2001, 2003, 2004), Fejfar et al. (2006) for details.

The lagomorph diversity in the late Ruscinian localities (MN15b, i.e., coeval with the Prolagus samples under study) of central and south-eastern Europe is quite high (see Kretzoi 1962; Syč 1965; Daxner & Fejfar 1967; Flederer & Reiner 1996; Terzea 1997; Dahlmann 2001; Popov 2004; Čermák 2007, 2009; Čermák & Angelone 2013; Čermák & Wagner 2013). Five species of lagomorphs pertaining to five distinct genera (Ochotonoma Šen, 1998, Prolagus, Hypolagus, Dice, 1917, Pliopentalagus Gureev & Konkova in Gureev, 1964, Trischizolagus Radulessco & Samson, 1967) are recorded. The palaeobiogeographical trajectory and the biochronological importance as MN15b markers of some of them is well documented (Angelone 2008b; Čermák & Wagner 2013). The most common lagomorph was H. petenyii, which occurred in almost all central and south-eastern MN15b European localities. This species is usually regarded as a forest dweller (Flederer & Reiner 1996). Trischizolagus dumitrescuae Radulessco & Samson, 1967 and O. csarnotana generally occur together in the Pliocene of Europe. Their presence indicates a rather dry and hot climate, and the presence of vast areas of savanna-steppes (Popov 2004). Most of the available remains of T. dumitrescuae are known from the northern part of the peri-Paratethyan area, in fossil sites of early Ruscinian age (MN14). Western occurrences of T. dumitrescuae are rather rare and exclusively limited to the late Ruscinian (MN15b). Ochotonoma csarnotana from Beremend 26 (Čermák & Wagner 2013) so far represents the westernmost proven record of the species. European occurrences of O. csarnotana are limited to south-eastern regions and are closely related to the distribution of Ochotonoma in Anatolia (see Čermák 2007 for details). The distribution range of T. dumitrescuae and O. csarnotana in central Europe does not exceed latitude N46° (occurrences in Beremend 26/39 and Csarnóta 2). On the contrary, the distribution range of Pliopentalagus dietrichi clearly exceeds this latitude and reaches the Rhine Graben (western Germany), together with P. bilobus (co-occurrences in Gundersheim-Findling and probably in Wölfersheim). Pliopentalagus dietrichi may be considered, by analogy with its extant relative Pentalagus furnessi (Stone, 1900), a forest dweller inhabiting dense subtropical forests characterized by a low cursorial ability. In this framework, it is difficult to guess the environmental requirements of P. bilobus and Prolagus sp.

\[ \text{Čermák S. et al.} \]
from Beremend. Possibly the latter was more related to a dry and open habitat, given the ecological requirements of the other lagomorphs found in the same assemblage, but the former may have pertained to either ecogeographical group 1 or 2 (sensu Popov 2004).

It is evident that in MN15b a conspicuous number of the lagomorph genera reached central Europe, most of them from eastern or south-eastern regions. Such appearance of lagomorph taxa with very different ecological requirements, all together in a short time span, suggests that the onset of favourable climatic conditions was not the only and/or primary driving factor of their coeval dispersal.

**The post-Miocene fate of Prolagus: climate change, palaeogeography, competitive pressure**

It is clear from the climatic and ecological frameworks above illustrated that, except for the earliest stages (MN2b-MN3), *Prolagus* developed and passed unscathed through periods of profound climatic/environmental changes for at least 14 of the c. 20 my of its evolutionary history. Actually, *Prolagus* species had a wide range of environmental adaptations. Several species lived in subtropical forest or wetland lacustrine habitats, nevertheless other species, i.e., *P. ibericus* from Layna (MN15) or *Prolagus* sp. from the Gargano palaeoarchipelago, inhabited arid environments (López Martínez 2001). It is also possible that several of those species did not have strict ecological requirements. Anyway, *Prolagus* was quite adaptable to different and variable climatic conditions. Thus, it is reasonable to suggest that climate change sensu stricto was not the primary reason for the limited distribution of *Prolagus* in the Pliocene of central Europe.

A different perspective on the distribution of the genus *Prolagus* in Europe comes from the palaeogeographical approach. Until the middle Miocene, the geographic distribution of *Prolagus* was arranged in a roughly latitudinal band (as far as the distribution of emerged lands permitted; cf. López Martínez 2001 for *Prolagus* distribution data, and Popov et al. 2004 for updated palaeogeographical setting). During the late Miocene, the area in which *Prolagus* was distributed was deformed by the Alpine arc in its central part (the radial axis of the arc, coinciding with the Rhine graben, was “pushed” northwards; see Sue et al. 2007). The bending movement may have had an effect on isolation (the question is how much) between western and eastern *Prolagus* populations, with the Alpine arc and the Bohemian Massif acting as physiographic barriers. Indeed, significantly increasing differences between western European *Prolagus* species and central-eastern European ones start to be noticed at least since MN10 (Angelone & Veitschegger 2015).

The decline in *Prolagus* abundance in northern central Europe appears evident already since the late Miocene (López Martínez 2001). Although the mammalian fossil record around the Miocene/Pliocene boundary (late MN13 to MN14) is extremely rare north of the Alps, *Prolagus* had probably disappeared there since the latest Miocene. Further south, in the Pannonian Basin, the genus was still present in the MN13 (Angelone & Čermák 2015). For sure, *Prolagus* is absent in the Pliocene of central Europe except for MN15b: c. 30 vertebrate-bearing fossiliferous localities are known between MN14-16 (Wagner et al. 2009) and the occurrences of *Prolagus* are exclusively limited to the 6 localities (out of a total of the 11 available for MN15b). Such chronologically limited occurrences are indicative of a dispersal, rather than of a progressive Pliocene shrinking of the Miocene distribution area that left isolated endemic populations. This dispersal most probably occurred from south-eastern regions of Europe (e.g. the NW peri-Black Sea area), as the main morphological characters of central European Pliocene *Prolagus* here analyzed is quite different from those of western European ones. Moreover, a dispersal wave from eastern regions fits quite well in the framework of the above discussed extensive faunal rearrangements occurred during the early Pliocene.

The subsequent, relatively sudden, disappearance of *Prolagus* from central Europe after MN15b can be explained by the increase of the competitive pressure related to the continuous global climatic degradation during the late Pliocene. In this sense, *Prolagus* met the same fate of *Pliopentatagus dietrichi* in Europe (see Čermák & Wagner 2013). In fact, the Pliocene climatic deterioration caused an increase of the aridisation and the spreading of open ground habitats in central Europe. This opened the way for *Ochotona* Link, 1795 and *Lepus* Linnaeus, 1758, very well adapted for those landscape settings, and the *Prolagus* “story” in central Europe was over and was never repeated again.

**CONCLUSIONS**

This paper fills a gap in the study of Pliocene lagomorphs, until now mainly centered on western European species, performing a detailed analysis of all available published and unpublished Pliocene materials of *Prolagus* from central Europe.

Most of the record of central European Pliocene *Prolagus* pertains to *Prolagus bilobus*. This species is reported here from the MN15b localities of Gundersheim and Raciszyn 1. Some materials of *P. bilobus* had already been revised and its diagnosis emended (Čermák & Angelone 2013), however the availability of relatively rich materials allowed to define here some additional peculiarities in tooth positions other than p3. The extremely low intraspecific variability of the specimens of *P. bilobus* is striking. This characteristic allowed to easily describe the ontogenetic trajectory of the p3 of *P. bilobus*. From juvenile to adult, the p3 undergoes the enlargement, deformation, and tilting of the anteroconid, the deformation of the metaconid, a noticeable thickening of the protoconulid, the thickening of the entoconid, and the enlargement of the crochet. In the mandibles of adult specimens, compared to the juveniles, it is possible to observe an enlargement of the diastema in relation to the alveolar row length, the heightening of the mandibular body, and an increase of the curvature in the outline of the ventral margin.

The study of the ontogenetic development of *P. bilobus* was extended to the deciduous teeth (d3), and the dimensional results were compared with *Prolagus* species from several fossil
localities ranging in age from the early Miocene (MN2b) to the Middle-Late Pleistocene. In early Miocene-early middle Miocene taxa, the d3 is larger than p3, as occurs in *Piezodus* (López Martínez 1974). However, since late middle Miocene, the size relationship appears reversed, and the values of d3 remain quite stable. The reversal is not gradual: the d3/p3 relative length proportion is one of the characters that, in the genus *Prolagus*, change their state at a certain geological moment and in a trans-specific way, and, in this sense, it may have a biochronological value. At any rate, the size difference between d3/p3 in *P. bilobus* is clearly the largest among the studied taxa.

New materials from Beremend 26 and 39 (MN15b) revealed the presence of a possible new species of *Prolagus*. A lower jaw of an adult individual from Beremend 26 appears clearly distinct from other Pliocene-Holocene congenic species. In particular, the differences with the coeval *P. bilobus* are striking. Teeth and mandible dimensions fall below the lower range of *P. bilobus*, and also the morphology of p3 is incompatible with *P. bilobus*. Nevertheless, an upper jaw excavated from the neighbouring fissure filling Beremend 39 does not show substantial differences in morphology and measures with *P. bilobus*. The two fillings seem coeval, basing on the biochronological indications given by the faunal content. These contradictory taxonomic evidences can be solved only by the retrieval of additional material, and the prolagids from Beremend 26 and 39 are provisionally left in open nomenclature as *Prolagus* sp.

Ukrainian and Moldavian MN15-216 materials attributed by Tesakov & Averianov (2002) to *P. bilobus* do not match the diagnosis of the species and, in our opinion, should be provisionally referred to *P. aff. sorbinii*. The material from Kislâng (mentioned by Kretzoi 1954) could have shed light on the subject. Unfortunately, the material is currently unavailable. The same fate was suffered by the *Prolagus* remains from Wölfersheim (faunal list in Tobien 1977; eventually reported also by Dahlmann 2001).

Contrarily to the “traditional” view (López Martínez 2001), we hypothesize that central European *Prolagus* are not relic species “left behind” after a progressive reduction of the geographic distribution of the genus due to climate deterioration s.s. occurred during the Pliocene. *Prolagus* disappeared from northern central Europe since the latest Miocene to reappear briefly in the late early Pliocene (MN15b). This evidence rather indicates a sudden dispersal, most probably from south-eastern regions of Europe, as the morphology of *P. bilobus* and *Prolagus* sp. studied herein exclude close phylogenetic affinities with western European species. *Prolagus* is not the only lagomorph to appear in central Europe in MN15b: others are e.g. *Plipopenatalagus* and *Ochotonoma*, two taxa of clear eastern origin.

The MN15b central and south-eastern Europe lagomorph palaeobiogeographical dynamics represent only a tile of the extensive faunal rearrangements that characterize the entire Europe during the Pliocene. The post-Miocene fate of *Prolagus* in central Europe was determined by a combination of palaeogeographic and climatic changes which caused extensive rearrangements of community and faunal structures that increased the competitive pressure on the genus.

Acknowledgements

We would like to express our thanks to László Pongrácz (Györ), Katrin Krohmann (SMF), and Adam T. Halanski (ZPAL) and M. Borsuk-Bialynicka (ZPAL) for providing us the fossil material in their care. We are grateful to two anonymous reviewers for their valuable comments and critical remarks. The study was supported by institutional support RVO67985831 of the Institute of Geology of the Czech Academy of Sciences. CA was supported by: Visiting Professor grant of the President’s International Fellowship Initiative of the Chinese Academy of Science; Spanish Agencia Estatal de Investigación and the European Regional Development Fund of the European Union (CGL2016-76431-P); CERCA Program, Generalitat de Catalunya; Grant to Department of Science, Roma Tre University (MIUR-Italy Dipartimenti di Eccellenza, ART. 1, C. 314-337 L. 232/2016). BMS was supported by Xunta de Galicia (ED481B 2018/046, Axudas á etapa postdoutoral da Xunta de Galicia 2018-Modalidade A).

REFERENCES

AGADJANIAN A. K. & ERBAJEVA M. A. 1983. — [Late Cenozoic rodents and lagomorphs of the USSR]. Nauka, Moscow, 189 p. (in Russian).

ÁLVAREZ-SIERRA M. A., DAAMS R., LACOMBA J. I., LÓPEZ-MARTÍNEZ N., VAN DER MUELEN A. J., SÉSE C. & DE VISSER J. 1990. — Palaeontologie and biostratigraphy (micromammals) of the continental Oligocene-Miocene deposits of the North-Central Ebro basin (Huesca, Spain). *Scripta Geologica* 94: 1-77.

ANGELONE C. 2007. — Messinian *Prolagus* (Ochotonidae, Lagomorpha, Mammalia) of Italy. *Geobios* 40: 407-421. [https://doi.org/10.1016/j.geobios.2006.04.004](https://doi.org/10.1016/j.geobios.2006.04.004)

ANGELONE C. 2008a. — *Prolagus italicus* n. sp. (Ochotonidae, Lagomorpha, Mammalia) a new Pliocene species of peninsular Italy. *Geobios* 41: 445-453. [https://doi.org/10.1016/j.geobios.2007.12.001](https://doi.org/10.1016/j.geobios.2007.12.001)

ANGELONE C. 2008b. — Family Ochotonidae (Lagomorpha) and its application in biochronology: some case studies from the Plio-Quaternary of Eurasia. *Quaternary International* 179: 5-8. [https://doi.org/10.1016/j.quaint.2007.08.019](https://doi.org/10.1016/j.quaint.2007.08.019)

ANGELONE C. & ČERMÁK S. 2015. — Two new species of *Prolagus* (Lagomorpha, Mammalia) from the Late Miocene of Hungary: taxonomy, biochronology, and palaeobiogeography. *Paläontologische Zeitschrift* 89: 1025-1038. [https://doi.org/10.1007/s12542-014-0247-z](https://doi.org/10.1007/s12542-014-0247-z)

ANGELONE C. & ROOK L. 2012. — Late Neogene and Quaternary lagomorphs from Tuscany: a revision based on specimens in Basel Naturhistorisches Museum and Florence University collections. *Swiss Journal of Geosciences* 131: 127-145. [https://doi.org/10.1007/s13358-011-0035-2](https://doi.org/10.1007/s13358-011-0035-2)

ANGELONE C. & SÉSE C. 2009. — New characters for species discrimination within the genus *Prolagus* (Ochotonidae, Lagomorpha, Mammalia). *Journal of Paleontology* 83: 80-88. [https://doi.org/10.1666/07-067R2.1](https://doi.org/10.1666/07-067R2.1)

ANGELONE C. & VETSCHEGGER K. 2015. — MN10 *Prolagus* (Ochotonidae, Lagomorpha) from Austria: a new tile for the central European palaeogeography of the genus. *Neues Jahrbuch für Geologie und Paläontologie* 275: 1-10. [https://doi.org/10.1127/njgp/2015/0444](https://doi.org/10.1127/njgp/2015/0444)

ANGELONE C., ČERMÁK S. & KOTSARKIS T. 2015. — The most ancient lagomorphs of Sardinia: an overview. *GeoBiologia* 48: 287-296. [https://doi.org/10.1016/j.geobios.2015.06.002](https://doi.org/10.1016/j.geobios.2015.06.002)

ANGELONE C., MONCUNILL-SOLE B. & KOTSARKIS T. 2019. — Contribution of fossil Lagomorpha (Mammalia) to the refinement of the late Miocene-Quaternary palaeobiogeographical setting
of Italy. *Comptes Rendus Palevol* 18: 1025-1040. https://doi.org/10.1016/j.crpv.2019.10.002

ANGELONE C., MONCUNILL-SOLE B. & KOTSARI T. 2020. — Fossil Lagomorpha (Mammalia) of Italy: systematics and biochronology. *Rivista Italiana di Paleontologia e Stratigrafia* 126: 157-187. https://doi.org/10.1340/2039-4942/130/14

AYERIANOV A. O. & TESAKOV A. S. 1998. — Lagomorphs (Mammalia, Lagomorpha) from early Pliocene locality of Kosyakino of the Northern Caucus. *Paleontological Journal* 32: 305-309.

BERNOR R. L., KOFOUS G. G., WOODBURN M. & FORTELIUS M. 1996. — The evolutionary history and biochronology of europeen and southeastern asian late Miocene and Pliocene hippocarnion horses, in BERNOR R. L., FAHLSCHU & MÜTTMANN H. W. (eds), *The evolution of Western Eurasian Neogene mammal faunas*. Columbia University Press, New York: 307-338.

BERZI A. 1967. — Lagomorpha from the type Villafranchian of Villafranca d’Asti (Italy). Preliminary note. *Giornale di Geologia* 35: 137-150.

BOHME M. 2003. — The Miocene Climatic Optimum: evidence from ectothermic vertebrates of Central Europe. *Palaeogeography, Palaeoclimatology, Palaeoecology* 199: 389-401. https://doi.org/10.1016/S0031-0182(03)00367-5

BOHME M., ILG A. & WINKLOHER M. 2008. — Late Miocene “washhouse” climate in Europe. *Earth and Planetary Science Letters* 275: 393-401. https://doi.org/10.1016/j.epsl.2008.09.011

BRANDT J. F. 1855. — Beiträge zur näheren Kenntniss der Säugertiere Russlands. *Mémoires de l’Académie impériale des sciences de St.-Pétersbourg* 9: 1-365.

CASSANOVAR-VILAR I., VAN DEN HOEK OSTENDE L. W., FURIO M. & MADERNO A. P. 2014. — The range and extent of the Vallesian Crisis (Late Miocene): new prospects based on the micromammal record from the Vallés-Penedés basin (Catalonia, Spain). *Journal of Iberian Geology* 40: 29-48. https://doi.org/10.5209/rev_JIGE.2014.v40.n1.44086

ČERMÁK S. 2007. — New finds of *Ochotonoma carnotana* (Lagomorpha, Ochotonidae) from the Pliocene of Hungary: a new look on the species. *Neues Jahrbuch für Geologie und Paläontologie* 246: 247-256. https://doi.org/10.1127/0077-7749/2007/0246-0247

ČERMÁK S. 2009. — The Plio-Pleistocene record of *Hypolagus* (Lagomorpha, Leporidae) from the Czech and Slovak Republics with comments on systematics and classification of the genus. *Bulletin of Geosciences* 84: 497-524. https://doi.org/10.3140/bull.geosci.1104

ČERMÁK S. & ANGELONE C. 2013. — Revision of the type material of the Pliocene species *Prolagus bilobus* Heller, 1936 (Mammalia, Lagomorpha), with comments on the taxonomic validity of *P. osmolskae* Fostowicz-Frelik, 2010. *Bulletin of Geosciences* 88: 45-50. https://doi.org/10.3140/bull.geosci.1369

ČERMÁK S. & WAGNER J. 2013. — The Pliocene record of *Trischizolagus* and *Pliopentalagus* (Leporidae, Lagomorpha, Mammalia) in Central Europe, with comments on taxonomy and evolutionary history of Leporinae. *Neues Jahrbuch für Geologie und Paläontologie* 268: 97-111. https://doi.org/10.1127/0077-7749/2015/0321

CZASZÁR G. & KORDOS L. 2004. — Bere mend, Kőfejtő. *Pragnum, eliadádiskvonatok, kirándulóvezetékek*, 7: *Magyar Öslényi Vágtorlók*. 2004: 51-57.

DAHLMANN T. 2001. — Die Kleinsäuger der unter-pliozänen Fund stelle Wolfsheim in der Wetterau (Mammalia: Lipotyphla, Chiroptera, Rodentia). *Kurator Forschungsinstitut Senckenberg* 227: 1-129.

DAHLMANN T. & STORCH G. 1996. — Eine pliozoische (ober-ruschnische) Kleinsäugerfauna aus Gundersheim, Rheinhessen: 1. Nagetiere: Mammalia, Rodentia. *Senckenbergiana lethaea* 71: 139-184.

FEOJAR O., HEINRICI W.-D. 1983. — Arvicoliden-Sukzession und Biostatigraphie des Oberpliozän und Quartärs in Europa. *Schriftenreihe für geologische Wissenschaften* 19: 20-61-109.

FEOJAR O. & STORCH G. 1990. — Eine pliozoische (ober-ruschnische) Kleinsäugerfauna aus Gundersheim, Rheinhessen: 1. Nagetiere: Mammalia, Rodentia. *Senckenbergiana lethaea* 71: 139-184.

FEOJAR O., HEINRICI W.-D. & LINDAY E. H. 1998. — Updating the Neogene Rodent biochronology in Europe. *Mededelingen Nederlands Instituut voor Toegestane Geowetschappen TNO* 60: 533-554.

FEOJAR O., STORCH G. & TOBIEN, H. 2006. — Gundersheim 4, ein dritter Rüsclies-Chiroptera, Rodentia). *Palaeontographica A* 278: 97-111. https://doi.org/10.1127/pala/278/2006/97

FLADERER F. A. & REINER G. 1996. — Evolutionary shifts in the first premolar pattern of *Hypolagus beremenensis* (Petényí, 1864) (Lagomorpha, Mammalia) in the Plio-Pleistocene of Central Europe. *Acta Zoologica Croatica* 39: 147-160.

FOSTORCZ-FRELIK Ł. 2010. — A new species of Pliocene *Prolagus* (Lagomorpha: Ochotonidae) from Poland in the northermost record of the genus. *Journal of Vertebrate Paleontology* 30: 609-612. https://doi.org/10.1080/07224631003621789

GANOPOLSKI A. & CALOV R. 2011. — The role of orbital forcing, carbon dioxide and regolith in 100 kyr glacial cycles. *Climate of the Past* 7: 1415-1425. https://doi.org/10.5194/cp-7-1415-2011

GUREEV A. A. 1960. — [Lagomorphs (Lagomorpha) from the Oligocene of Mongolia and Kazakhstan]. *Trudy Paleontologicheskogo Instituta Akademii Nauk SSSR* 77: 5-34. (in Russian).

GUREEV A. A. 1964. — [Fauna of the USSR (Lagomorpha), Vol. 3 (10)]. Nauka, Moscow & Leningrad, 276 p. (in Russian).

HALLER F. 1936. — Eine oberpliozoische Wirbeltierfauna aus Rhein hessen. *Neues Jahrbuch für Mineralogie, Geologie und Paläontologie, Abteilung B* 76: 99-160.

HORDIK J. & VAN DER MEULEN A. J. 2010. — Systematics of resident species of *Lagopus* and *Prolagus* (Ochotonidae, Lagomorpha, Mammalia) from the late early and middle Miocene of northeastern Central Spain. *Geologia Ultraceincta* 533: 17-85.

JANOSZ D. 1972. — Middle Pliocene microvertebrate fauna from the Ostromats Loc. 1. (Northern Hungary). *Annales historico-naturales Musei nationalis hungarici* 64: 27-50.

JANOSZ D. 1986. — Pleistocene vertebrate faunas of Hungary. *Aka démiai Kiadó, Budapest*, 208 p.

KOENIGSWALD W. VON & TOBIEN H. 1990. — Important arvicolid faunas from the Late Pliocene to Early Holocene in Western Germany (FRG). In FEOJAR O. & HEINRICI W.-D. (eds), *International symposium evolution, phylogeny and biostatigraphy of arvicolids (Rodentia, Mammalia)*. Czech Geological Survey, Prague: 231-254.

KÖNIG C. D. E. 1825. — *Icones Fossilium Sectiles, Centuria Prima*. G.B. Sowerby, London, 4 p.
Prolagus in the framework of the Pliocene faunal rearrangements in central Europe

Trobie 1935. — Über die pleistozenischen und postpleistozenischen Prolagusformen Korsikas und Sardiniens. Berichte der naturforschenden Gesellschaft zu Freiburg im Breisgau 34: 253-344.

Trobie H. 1977. — 7. Fauna, in Boenigk W., Brulé G., Brunacker K., Kempf E. K., Kočí A., Schirmer W., Stadler G., Streit R. & Trobie H. (eds), Jungtertiär und Quartär im Horloff-Graben, Vogelsberg, Geologische Abhandlungen Hessen 75: 65-68.

Trobie H. 1980. — Säugeraffen von der Grenze Pliozän/Pleistozen in Rheinhessen. 1. Die Spaltfüllungen von Gundersheim bei Worms. Mainzer geowissenschaftliche Mitteilungen 8: 209-218.

Topacheyevsky V. O. 1962. — [On the Geological Age of Early Alluvial deposits of the Lower Dniester Valley]. Geologicheskii zhurnal 22: 106-109 (in Ukrainian).

Únay E. & de Bruijn H. 1998. — Plio-Pleistocene rodents and lagomorphs from Anatolia. Mededelingen Nederlands Instituur voor Toegepaste Geowetenschappen TNO 60: 431-466.

Van der Meulen A. & van Kolschot T. 1986. — Review of the Late Turolian to Early Biharian mammal faunas from Greece and Turkey. Memoria Società Geologia Italiana 31: 201-211.

Van de Weerd A. 1979. — Early Ruscinian rodents and lagomorphs (Mammalia) from the lignites near Pollemais (Macedonia, Greece). Proceedings of the Koninklijke Nederlandse Akademie van Wetenschappen, Series B Physical Sciences 82: 127-168.

Van de Weerd A. & Daams R. 1978. — Quantitative composition of rodent faunas in the Spanish Neogene and palaeoecological implications. Proceedings of the Koninklijke Nederlandse Akademie van Wetenschappen, Series B Physical Sciences 81: 448-473.

Van de Weerd A. & de Vos J. 1982. — Pliocene mammals from Apolakkia Formation (Rhodes, Greece). Proceedings of the Koninklijke Nederlandse Akademie van Wetenschappen, Series B Physical Sciences 85: 89-112.

Viret J. 1929. — Les faunes de mammifères de l’Oligocène supérieur de la Limagne bouronnaise. Annales de l’Université de Lyon, nouvelle série 1 47: 1-327.

Wagner R. 1829. — Über die Knochenbrekzie in Sardinien und die darin gefundenen Thiere, so wie über einige andere hierher gehörige Erscheinungen. Archiv für die gesammte Naturlehre 15: 10-31.

Wagner J., Čermák S., Horáček I., Fejfar O. & Mihelčič A. 2009. — New mammalian fossil records refining a view on Early/Late Pliocene faunal turnover in Central Europe. Journal of Vertebrate Paleontology 29 (Supl. 3): 197A.

Wible J. R. 2007. — On the Cranial Ostegology of the Lagomorphs. Bulletin of Carnegie Museum of Natural History 39: 213-234. https://doi.org/10.2992/0145-9058(2007)39[213:OTOCO]2.0.CO;2

Xue X., Zhang Y. & Yue L. 2006. — Paleoenvironments indicated by the fossil mammalian assemblages from red clay-loess sequence in the Chinese Loess Plateau since 8.0 Ma. B.P. Science in China Series D 49: 518-530. https://doi.org/10.1007/s11430-006-0518-y

Zachos J. C., Pagani M., Sloan L. C., Thomas E. & Billups K. 2001. — Trends, rhythms, and aberrations in global climate 65 Ma to present. Science 292: 686-693. https://doi.org/10.1126/science.1059412