The Late Cretaceous belemnite family Belemnitellidae: Taxonomy and evolutionary history

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The Late Cretaceous belemnite family Belemnitellidae Pavlov, 1914 occurs only in the Northern Hemisphere and includes nine genera and two subgenera: Praeactinocamax Naidin, 1964b, Actinocamax Miller, 1823, Belemnocamax Crick, 1910, Goniateuthis Bayle, 1878, Belemnellococamax Naidin, 1964b, Gonocamax Naidin, 1964b, Belemnitella d’Orbigny, 1840, Belemnella (Belemnella) Nowak, 1913, Belemnella (Pachybelemnella) Schulz, 1979, Belemnella (Neobelemnella) Naidin, 1975 and Fusiteuthis Kongiel, 1962. The latter is most likely a nomen dubium. Diagnoses of the genera and subgenera are provided.

The origin of the family is poorly known. The number of genera and subgenera fluctuated during the Late Cretaceous. It was one to two in the Cenomanian, increased gradually to a maximum of six in the Early Santonian, decreased gradually to one during most of the Late Campanian and increased to two or possibly three in the Maastrichtian. The belemnitellids occur in the North European and North American palaeobiogeographical Provinces of the North Temperate Realm, in addition to the northern European margin of the Tethyan Realm. The belemnitellids occur in the North European and North American palaeobiogeographical Provinces of the North Temperate Realm, in addition to the northern European margin of the Tethyan Realm. The centre of evolution lay in the North European Province and all known genera and subgenera occur there. Species of five genera and two subgenera occur on the northern European margin of the Tethyan Realm and the majority of these are conspecific with species from the North European Province. Species of essentially two genera occur in the North American Province and these are endemic, with a few exceptions.

Key words: Belemnitellidae, taxonomy, evolutionary history, Northern Hemisphere, Late Cretaceous.

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During the Late Cretaceous, except in the Cenomanian, belemnites had a bipolar distribution. The Late Cretaceous family Belemnitellidae Pavlov, 1914 inhabited the North Temperate Realm (= Boreal Realm of authors) and the late Early and Late Cretaceous (Aptian to Maastrichtian) family Dimitobelidae Whitehouse, 1924 inhabited the South Temperate Realm (= Austral Realm). The last genera of the family Belenopseidae Naef, 1922, the mid-Cretaceous (Aptian to Cenomanian) Neohibolites Stolley, 1911 and Parahibolites Stolley, 1919 inhabited the Tethyan Realm, but occurred also in the North Temperate and South Temperate Realms, where they may be locally abundant. They became extinct in the Early and Middle Cenomanian (Combémorel, Christensen, Naidin & Spaeth 1981). Therefore, a Tethyan Realm cannot be defined on the basis of belemnites alone after the Cenomanian, but it can be recognized on the basis of other fossil groups, such as rudists, ammonites, echinoids, actaeonellid gastropods and larger foraminifera.


The belemnitellids were distributed in the North American and North European palaeobiogeographical Provinces of the North Temperate Realm, in addition to the northern part of the Tethyan Realm in Europe (Christensen 1975a, 1976, 1988, 1993b). The centre of evolution of the belemnitellids lay in the North
European Province, and they invaded the North American Province and Tethyan Realm at various times during the Late Cretaceous. The belemnitellids are common in the North American Province and all known genera and subgenera are recorded. They are generally rare in the North American Province, and essentially two genera are recorded. The belemnitellids occur rarely in the Tethyan Realm in Europe and five genera and two subgenera are reported.

The belemnitellids were stenothermal shelf dwellers, but it appears that the breeding, spawning, hatching, and, possibly for the females at least, dying grounds, were inner neritic, shallow water environments.

The belemnitellids are of fundamental importance in biostratigraphy and correlation in Europe, because they are common, widely distributed and their fossilization potential is high (Christensen 1990b).

The aim of this contribution is to discuss the systematic palaeontology and evolutionary history of the belemnitellids. Representative species of the genera are figured on Plates 1–2.

Classification of the Belemnitellidae

Up to 1964, five genera within the Belemnitellidae had received general recognition by most authors, including Wright & Wright (1951): Actinocamax Miller, 1823, Goniotethis Bayle, 1878, Belemnocamax Crick, 1910, Belemnitella d'Orbigny, 1840 and Belemnella Nowak, 1913. In contrast, Kongiel (1962) did not recognize Belemnella.

Naidin (1964b) proposed a new classification of the belemnitellids without a true alveolus, that is species with a convexly conical alveolar fracture, a flat anterior end or a pseudoalveolus (see below). He erected two new subgenera of Actinocamax, A. (Praeactinocamax) and A. (Paractinocamax); one new genus, Belemnelloccamax; and one new subgenus of Goniotethis, G. (Goniocamax). He placed two small species, A. verus Miller, 1823 and A. laevigatus Arkhangelsky, 1912, in Actinocamax (Actinocamax); two large species, A. primus Arkhangelsky, 1912 and A. plenus (Blainville, 1825–1827), in Actinocamax (Praeactinocamax); one species, A. grossouvreii Janet, 1891, in Actinocamax (Paractinocamax); and one species, A. mammillatus (Nilsson, 1826), in Belemnelloccamax.

The following species from the North European Subprovince were placed in Goniotethis (Goniocamax): A. lundgreni Stolley, 1897, the type species, A. westfalicus (Schlüter, 1874) and A. intermedius Arkhangelsky, 1912, as well as two new species, matesovae Naidin, 1964b and medwedieicus Naidin, 1964b, A. strehensis (Fritsch and Schloenbach, 1872), A. bohemicus Stolley, 1916 and A. paderbornensis Schüter, 1894 were placed in the subgenus with a query. Species from the North American Province comprise A. manitobensis (Whiteaves, 1889), A. aff. plenus and A. walkeri Jeletzky, 1961. Three granulated taxa from this province: A. groenlandicus Birklund, 1956, A. aff. groenlandicus and A. sternbergi Jeletzky, 1961 were placed in Goniotethis (Goniocamax) with a query.

The classification of Naidin was discussed by J. A. Jeletzky in his unpublished manuscript for the Treatise on Invertebrate Palaeontology (copy obtained by me in 1972; it is referred to below as unpublished MS 1972). He placed Goniotethis in synonymy with Actinocamax (Praeactinocamax) and considered Actinocamax (Paractinocamax) as a subgenus of Belemnelloccamax. Jeletzky regarded granulation to be of great taxonomic significance (see below), and in contrast to Naidin (1964b) he placed the granulated or supposedly granulated species paderbornensis, bohemicus, groenlandicus and sternbergi in Actinocamax (Actinocamax). He placed the species westfalicus, lundgreni, manitobensis and intermedius in Actinocamax (Praeactinocamax). It has been shown subsequently that paderbornensis is not granulated (Christensen 1982) and manitobensis may be granulated (Christensen & Hoch 1983).

In this paper I recognize nine genera and two subgenera within the Belemnitellidae: Praeactinocamax Naidin, 1964b, Actinocamax Miller, 1823, Belemnocamax Crick, 1910, Goniotethis Bayle, 1878, Belemnelloccamax Naidin, 1964b, Goniocamax Naidin, 1964b, Belemnitella d'Orbigny, 1840, Belemnella (Belemnella) Nowak, 1913, Belemnellina (Pachybelemnella) Schulz, 1979, Belemnella (Neobelemnella) Naidin, 1975 and Fusiteuthis Kongiel, 1962 (Fig. 1). Belemnocamax and Fusiteuthis are monotypic, but the latter is most likely a nomen dubium (see below). The classification of Naidin and his generic assignment of various species are discussed further below.

Origin and evolutionary history of the Belemnitellidae

The earliest species of the Belemnitellidae, Praeactinocamax primus, appears abruptly in the Early Cenomanian, some way above the base of the stage (Christensen 1990a). Jeletzky (1946, unpublished MS 1972) suggested that the belemnitellids are derived from Aptian belemnopseids, either Neohibolites ewaldi (Strombeck, 1861) or N. clava Stolley, 1911, because the shape of the guard and the structure of the alveolar end of these species and the earliest belemnitellids are closely similar. Doyle (1987a, 1988a, 1992) suggested that the belemnitellids may have evolved from a northern endemic stock and the dimitobelids from a southern endemic stock of the Tethyan belemnopseid Hibolites de Montfort, 1808 during the later part of the Early Cretaceous. The sug-
gestions by Jeletzky and Doyle suffer the disadvan-
tage, however, that the earliest belemnitellid is sepa-
rated from Neohipolites and Hibolithes, respectively, 
by a considerable time gap. N. ewaldi is middle Aptian 
and N. clava is early Late Aptian in age (Mutterlose, 
Schmid & Spaeth 1983). Hibolithes became extinct 
in the basal Aptian in the Boreal Realm, in the middle 
Barremian in the Mediterranean Province and at the 
Hauterivian-Barremian boundary in the Indo-Pacific 
Province (Mutterlose 1988). The time gap between 
the latest belemnopseids and the earliest belemnitellid 
is thus at least about 20 million years (Ma).

Naidin & Alekseev (1975) suggested that P. primus 
was derived from the early Middle Cenomanian Neohipo-
lites repentinus Naidin & Alekseev, 1975, which 
occurs in the Crimea. This suggestion was based on 
the similarity of the shape of the guard and the struc-
ture of the alveolar end, as well as other characters, in 
the two species. The suggestion by Naidin & Alekseev 
is not tenable, because P. primus appears earlier than 
N. repentinus. The similarity in various characters in 
P. primus and N. repentinus may be a case of synchro-

It can thus be concluded that the origin of the Belem-
nitellidae is poorly known.

Jeletzky (1946, unpublished MS 1972) further sug-
gested that all belemnitellid genera are derived from 
the Cenomanian Praeactinocamax rootstock, imply-

Fig. 1. Stratigraphical range and inferred phylogeny of belemnitellid genera and subgenera. Stage abbreviations after 
Fig. 2. Stratigraphical range diagram of Late Cretaceous belemnopseids and belemnitellids from the North European Province. 1, northwest Europe; 2, Russian Platform; 3, Bornholm, Denmark. Stage abbreviations after Harland et al. (1989). Ages in Ma after Obradovich (1994).

(Christensen 1993c). Moreover, it has been suggested by Jeletzky (unpublished MS, 1972) and Naidin (1964 b, Fig. 35) that Actinocamax, which appears in the Early Turonian (Naidin 1964b), is derived from Praeactinocamax. This suggestion is, however, very questionable, because the two genera differ in several critical characters, including the size of the guard, the length of the cone-shaped alveolar fracture, the size and shape of the juvenile guard and the growth relationship (see diagnoses). Christensen (1993b) and Christensen & Schulz (in press) showed that growth is isometric in Actinocamax, and, consequently, both juvenile and adult specimens are short and stout. Growth is allometric in Praeactinocamax (see below); juvenile specimens are long and slender (needle-shaped) and adult specimens are stouter. Thus, it seems not very likely that Actinocamax is derived from a pre-Turonian species of Praeactinocamax, that is P. primus or P. plenus. Oddly enough, Naidin (1964b: 171) also noted that Actinocamax may be derived from a small Albian-Cenomanian species of Neohibolites. In conclusion, the Belemnitellidae is probably a polyphyletic taxon.

Goniocamax enters at the base of the Coniacian (Fig. 1) (Christensen & Schulz, in press). Naidin (1964b) derived this genus from the Late Turonian species Goniothlystys (Goniocamax) medwedieicicus, which is here placed in Praeactinocamax (see below). Christensen (1988) suggested that the Belemnitella stock, including Goniocamax lundgreni, was derived from a Turonian-Coniacian species of Praeactinocamax from the Central Russian Subprovince. At that time, it was erroneously believed that G. lundgreni, the earliest species of the genus, was Late Coniacian in age. G.
lundgreni appears, however, at the base of the Coniacian. Christensen & Schulz (in press) noted that the origin of G. lundgreni is unknown.

Belemnitella appears at the base of the Santonian (Christensen & Schulz, in press). B. schmidtii Christensen & Schulz (in press), the earliest species of the genus, may have evolved from Goniocamax lundgreni by allopatric speciation.

Gonioteuthis enters in the Middle Coniacian (Ernst & Schulz 1974). Christensen & Schulz (in press) suggested that the earliest species, G. praevestfaliaea Ernst & Schulz, 1974, may have been derived from Goniocamax lundgreni.

Belemnellocamax appears probably at the base of the Santonian (Christensen 1986). Jeletzky (unpublished MS 1972) derived this genus from a Turonian species of Actinocamax (Praeactinocamax). Nadin (1964b) noted that the origin of the genus is unknown. Christensen (1988) suggested that Belemnellocamax may be derived from one of the large Cenomanian-Turonian species of Praeactinocamax from Europe due to the size of the guard, overall morphology and ontogeny. It is noteworthy, however, that the time gap between the two genera is at least 2.5 Ma.

Belemnella enters at the base of the Maastrichtian (Schulz 1979). It has been suggested that Belemnella may be derived from Belemnellocamax, because both genera have large guards, long and slender juvenile guards and a small Schatzky Distance. Furthermore, the vascular imprints branch off the dorso-lateral double furrows posteriorly at an angle exceeding 30 degrees in both genera (Nowak 1913; Jeletzky 1949a, 1951; Christensen 1975a). However, the latest species of Belemnellocamax, B. balsvikensis (Brotzen,
Fig. 3. Stratigraphical range diagram of Late Cretaceous belemnittellids of the North American Province. A. = Actinocamax. Stage abbreviations after Harland et al. (1989). Ages in Ma after Obradovich (1994).

The number increases to three to four in the Coniacian and reaches a maximum of six in the Early Santonian. Afterwards the number decreases to four from the Middle Santonian to basal Early Campanian and three in the late Early Campanian. Only one genus, Belemnita, occurs in the Late Campanian, except its basal part. The early Late Campanian Belemnelloccamax balsvikensis occurs commonly only in Scania and is virtually absent elsewhere (see discussion below). Three genera and subgenera occur in the Early Maas- trichtian and the number decreases to two or three in the Late Maastrichtian, depending the systematic status of Fusciteuthis.

Belemnitala is a very long-ranging genus, occurring from the Santonian to the Maastrichtian, c. 21 Ma longevity. The genera Actinocamax and Praeactinocamax are long-ranging, c. 11–12 Ma longevity. The longevity of the genera Belemnelloccamax and Gonio- teuthis is around 6–7 Ma. Gonio-ccamax and the sub- genera of Belemnella are short-ranging, around 2–4 Ma longevity.

The stratigraphical ranges of the belemnittellids of the North European and North American Provinces, in addition to the Tethyan Realm are shown in Figures 2–4.

Systematic Palaeontology

Class Cephalopoda Cuvier, 1795
Subclass Coleoidea Bather, 1881
Superorder Belemnoida Hyatt, 1884
Order Belemnitida Zittel, 1895
Suborder Belemnopseina Jeletzky, 1965
Family Belemnittellidae Pavlov, 1914

Diagnosis. - Belemnopseina with a conical depression, an alveolus, in anterior part of guard, which is connected through ventral fissure with surface; surface markings consist of dorso-lateral longitudinal depressions and double furrows, single lateral furrows, more or less distinct vascular imprints, which cover all or parts of guard, longitudinal striae, and granules, which may form corrugated transverse lines; pro- ostracum narrow and tongue-shaped, and phragmocone with a dorsal keel, which fits in a dorsal furrow in alveolus; guard prolonged ventrally around ventral fissure in a tongue-like extension; a shorter dorsal extension may also be present.

Remarks. - The guard is usually the only part of the skeleton preserved in belemnittellids, and its external and internal characters are used for taxonomic classification. These include 1) size of the guard, 2) shape of the guard, 3) structure of the adoral end, 4) surface markings, 5) internal characters, including the alveolar angle, fissure angle, Schatzky distance, and the
shape of the bottom of the ventral fissure, and 6) ontogeny. These were discussed in detail by Christensen (1986). The Riedel-Quotient of Ernst (1964) is the ratio of the length of the guard divided by the depth of the pseudoalveolus. The Riedel-Index of Ernst & Schulz (1974) is the depth of the pseudoalveolus as a percentage of the length of the guard. The Birkeland Index of Christensen (1995) is the length from the apex to the protoconch divided by the dorso-ventral diameter at the protoconch.

The guard is completely calcified only in the genera *Belemnitella* and *Belemnella*, and they therefore have a true alveolus adorally. However, complete guards are extremely rare, because the most anterior end is usually not preserved due to its fragility. However, Christensen (1972, 1975a) figured nearly complete guards of *Belemnitella mucronata* (Schlotheim, 1813), *B. praecursor* Stolley, 1897 and *B. junior* Nowak, 1913 and demonstrated that the guard is prolonged ventrally in a tongue-like extension. A shorter dorsal extension may also be present; this structure should not be confused with the pro-ostracum.

In the genera *Praeactinocamax*, *Actinocamax*, *Belemnocamax*, *Goniocamax*, *Gonioteuthis* and *Belemnellocamax* the adoral part of the guard was built up of aragonite, which was lost during fossilization (Bandel & Spaeth 1988). Therefore, the anterior end may be either conical or flat with a central pit (*Praeactinocamax* and *Actinocamax*) or developed as a pseudoalveolus (*Belemnocamax*, *Goniocamax*, *Gonioteuthis* and *Belemnellocamax*). Some species of *Praeactinocamax* and some specimens of *Actinocamax* may have a shallow pseudoalveolus.

Jeletzky (1950, 1961, unpublished MS 1972) stressed the taxonomic importance of granulation within the belemnitellids. Christensen (1982, 1993b), Christensen & Hoch (1983) and Christensen & Schulz (in press) did not follow this point of view and showed that too much importance has been attributed to this character. Granulation may be diagnostic at the species and generic level, but it may also occur extremely rarely in genera, which are not usually granulated (see later discussion).

Jeletzky (1950) stated that all Cenomanian and most Turonian and Coniacian species of *Praeactinocamax* are extremely variable with respect to the shape of the guard. I disagree, because biometric studies of the Middle Cenomanian *Praeactinocamax primus*, the Late Cenomanian *P. plenus* and the Middle Coniacian *P. cobbani* (Christensen, 1993b) have shown that these species do not exhibit an extreme variation (Christensen 1974, 1990a, 1993b). It was shown, however, that growth was allometric. Thus, adult specimens of these species are stouter and more lanceolate in ventral view than juvenile and adolescent specimens. It is consequently very important to study the relative growth of belemnitellid species by bivariate analysis. In case of isometric growth the ratio of two variates will remain constant, whereas in case of allometric growth the ratio will change with size. Ratios have been widely used in palaeontological studies, including belemnite studies, but ratios should be used only when the growth is isometric or when specimens of nearly the same size of different species are compared.

**Distribution.** – The belemnitellids occur in the North European and North American Provinces of the North Temperate Realm, and the northern margin of the Tethyan Realm in Europe.

**Genus Actinocamax** Miller, 1823

**Type species.** – By monotypy *Actinocamax verus* Miller, 1823, p. 63, Pl. 9: 17.

**Emended diagnosis.** – Small, stout belemnitellids

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![Graph](image.png)

Fig. 4. Stratigraphical range diagram of Late Cretaceous belemnitellids of the Tethyan Realm. *P.* = *Praeactinocamax*; *G.* = *Gonioteuthis*, *Blc.* = *Belemnellocamax*. Stage abbreviations after Harland et al. (1989). Ages in Ma after Obradovich (1994).
(guard up to 55 mm long) with isometric growth; generally long cone-shaped alveolar fracture; ventral fissure, ventral furrow, and ventral notch absent; single lateral furrows weakly developed or absent; granulation present in some species.

**Discussion.** — Naidin (1964b) recognized three subgenera of *Actinocamax*: A. *Actinocamax*, type species *A. verus*; A. *Praeactinocamax*, type species *A. minutus*; and A. *Laevigatus*, type species *A. grossouvrei*.

Christensen (1986) placed A. *Praeactinocamax* in synonymy with *Belemnellopax*, type species *Belemnites mammillatus* (see later discussion). The legitimacy of the subgenera *Actinocamax* and *Praeactinocamax* was initially questioned by Christensen (1982, 1986, 1991, 1993b, 1994), but they were recognized by Gale & Christensen (1996) and Christensen & Schulz (in press), because they differ in several critical characters (see discussion above). Subgenus *Praeactinocamax* is here elevated to full generic rank due to these differences.

*Actinocamax* has been the subject of excessive subdivision by Russian palaeontologists. Four species and nine subspecies have been established: *A. verus* (including six subspecies), *A. laevigatus* (including three subspecies), *A. minitus* Glazunova, 1972 and A. *quasi-verus* Naidin, 1953. These were discussed by Christensen & Schulz (in press), who recognized *A. v. verus*, *A. verus subfragilis* Naidin, 1964b, *A. verus antefragilis* Naidin, 1964b, *A. laevigatus* and *A. quasi-verus*.

**Distribution.** — *Actinocamax* occurs mainly in the North European Provinces and is recorded from the Early Turonian to the boundary between the lower and upper part of the Early Campanian (Fig. 1). It occurs extremely rarely in the North American Province. The stratigraphical ranges of the species are shown in Figures 2 and 4.

*A. v. verus* is widely distributed and occurs commonly in the North European Province. In northwest Europe it is recorded from the Santonian and early Early Campanian. On Bornholm it enters in the latest Early Santonian, slightly above the appearance of *Gonioteuthis westfalica westfalica* (Christensen & Schulz, in press). In offshore chalks it is most common in the late Late Santonian; it has an acme occurrence in the upper part of the early Late Santonian *Uintacrinus* Zone of southern England. On the Russian Platform it enters in the Coniacian.

*A. verus antefragilis* is recorded from the Early Turonian of the Russian Platform, and *A. verus* cf. *antefragilis* was reported recently from the late Early and Middle Coniacian of Bornholm (Christensen & Schulz, in press). *A. verus subfragilis* occurs in the late Coniacian of the Russian Platform.

*A. laevigatus* occurs on the Russian Platform in the so-called 'Pteria'-beds of earliest Early Campanian age; it possibly also occurs in beds of the same age in western Europe (Naidin 1964b). *A. quasiverus* occurs in the Late Santonian of Crimea.

**Genus Praeactinocamax** Naidin, 1964b

**Type species.** — *Belemnites plenus* Blainville, 1825–1827, p. 376, Pl. 11bis: 3, by original designation by Naidin (1964b: 34).

**Emended diagnosis.** — Medium-sized to large belemnitellids (guard up to 115 mm long) with allometric growth; adult specimens stouter and more lanceolate in ventral view than juvenile specimens; usually with short cone-shaped alveolar fracture, but some species with a very shallow pseudoalveolus; ventral fissure absent; ventral furrow and ventral notch sometimes present; juvenile guard long and slender (needle-shaped).

**Discussion.** — The genus is generally not granulated, but granulation may occur very rarely in some species and one species, *P. groenlandicus*, is granulated.

The earliest belemnitellid, *P. primus*, enters some way above the base of the Early Cenomanian in the North European Province and is followed upwards by the Late Cenomanian *P. plenus*. The two species are closely allied and form an evolutionary lineage (Christensen 1974, 1990a). These species are not granulated.

In addition to the two large Cenomanian species, I also include in this genus medium-sized to large species from the Turonian-Early Santonian of the North European and North American Provinces (Figs 2–3). Species from the Central European Subprovince comprise: *bohemicus*, *strehlensis* and *paderbornensis*. They were revised by Christensen (1982), who showed that *bohemicus* may be granulated, whereas the others are not granulated.

Species from the Central Russian Subprovince include: *planus* (Makhlin, 1965), *coronatus* (Makhlin, 1965), *intermedius*, *matesovae*, *medwedicicus*, *mujnakensis* (Naidin, 1964b) and *aralensis* (Arkhangelsky, 1912). These species are not granulated.

Species from the North American Province comprise: *manitobensis*, *cobbani*, *groenlandicus*, *walkeri*, *sternbergi*, aff. *primus* and aff. *groenlandicus* (Fig. 3). *P. groenlandicus* is granulated (Birkelund 1956) and *P. cobbani* is very rarely granulated (Christensen 1993b). *P. sternbergi* was established by Jeletzky (1961) on the basis of a single specimen, which differs from *P. manitobensis* only by being granulated. Jeletzky (1961) suggested that *P. aff. groenlandicus* from central West Greenland, which was similarly erected on the basis of a single, granulated specimen, may be either a geographical subspecies or a morphological variety of *P. sternbergi*. Christensen & Hoch (1983) suggested that *P. sternbergi* should be placed...
in synonymy with *P. manitobensis*, because granulation may occur very rarely in this species. The species from the North American Province were discussed by Christensen (1993b).

Birkelund (1956) referred one specimen of unknown age from central West Greenland to *P. aff. primus*. This determination is open to discussion, because the specimen is enclosed in matrix and only the side is exposed. It is so poorly preserved that it cannot be orientated with certainty. Moreover, *P. primus* occurs in the Lower and Middle Cenomanian in the North European Province, but Cenomanian belemnitellids are not recorded elsewhere in the North American Province.

*P. primus* usually has a short cone-shaped alveolar fracture. Specimens with a flat anterior end are rare and specimens with a shallow pseudoalveolus are extremely rare (Christensen 1990a). A specimen from Bornholm, MGUH 7834, figured by Birkelund (1957, Pl. 1: 2) and Christensen (1990a, Fig. 3C), has a Riedel-Quotient of c. 46 and a Riedel-Index of c. 2. Christensen (1974) analyzed a large sample of *P. plenus* from the Plenus Marls of the Betchworth Limeworks in southern England. He showed that most specimens have a low cone-shaped alveolar fracture, and that about 7% of the specimens have a shallow pseudoalveolus, with all intermediate forms. I have subsequently analyzed the variation of the Riedel-Quotient and Riedel-Index of the specimens with a shallow pseudoalveolus (Table 1).

*P. manitobensis* include morphological variants with either a low cone-shaped alveolar fracture, a flat anterior end or a shallow pseudoalveolus (Jeletzky 1950, 1961). According to Jeletzky (1961, Pl. 1: 2), specimen no. 7936-2 has a Riedel-Quotient between 17.5 and 19.4.

Five species from the Russian Platform have a shallow pseudoalveolus or a flat anterior end with a central pit. The Riedel-Quotient is 11 to 13 in the Late Coniacian *P. aralensis* and Late Turonian *P. mate- sovae*, 16 to 18 in the Late Turonian *P. coronatus*, 13 to 25 in the Late Coniacian *P. mujnakensis* and about 15 in the Late Turonian *P. medwedicicus*. In contrast to Naidin (1964b) and Makhlin (1965), who placed these species in *Goniotethys* (*Goniocamax*), they are here assigned to *Praeactinocamax*, because the pseudoalveolus is relatively shallow. However, these species may be transitional forms between the genera *Praeactinocamax* and *Goniocamax*.

**Distribution.** *Praeactinocamax* occurs in the North American and North European Provinces, in addition to the northern part of the Tethyan Realm in Europe. It appears in the Early, but not earliest, Cenomanian and continues into Early Santonian. The stratigraphical ranges of the species are shown in Figures 2-4.

**Genus Belemnocamax** Crick, 1910

**Type species.** By monotypy *Belemnocamax boweri* Crick, 1910, p. 364, Pl. 28: 1–2.

**Diagnosis.** Very small belemnitellid (guard up to 25 mm long), conical in lateral and ventral views, and with acute apex; anterior end with pseudoalveolus; Riedel-Quotient from about 4 to 7; walls of pseudo-alveolus sometimes with conelae; guard with long, broad, and deep ventral groove extending from the anterior end; length of groove 1/3 to 1/4 of length of guard; dorso-lateral longitudinal depressions distinct, delimiting the dorsal field; dorso-lateral depressions from anterior end and almost to apex; single lateral furrows present; guard with longitudinal striae, but otherwise smooth.

**Discussion.** Christensen (1993c) described *B. boweri* on the basis of a small sample, consisting of 18 specimens, from the limestone pit at Wunstorff, west of Hannover, Lower Saxony Basin. In addition, three specimens from Dörenthe, Münster Basin, and eight specimens from eastern England, including the holotype, were also studied.

*B. boweri* differs from all other belemnitellid genera by the size and shape of the guard, and by having a long, broad, and deep ventral groove.

The ancestry of *B. boweri* is unknown. J. A. Jeletzky (unpublished MS 1972) suggested that it probably represents an early independent offshoot of a still unknown pre-Cenomanian form between *Neohibolites* and *Actinocamax*, and that this aberrant genus died out without issue.

**Distribution.** *B. boweri* is very rare and has a very restricted distribution. It occurs in the early Middle Cenomanian of the Lower Saxony and Münsterland Basins in northwest Germany, the Cleveland Basin in eastern England and at Hunstanton, Norfolk.

**Genus Goniotethys** Bayle, 1878

**Type species.** *Belemnites quadratus* Blainville, 1827, p. 62, Pl. 1: 9, by original designation of Bayle (1878, caption to Pl. 22: 6–8).
Plate 1

Figs 1-3. *Praeactinocamax plenus* (Blainville, 1825-1827), Sedgwick Museum, B76709, Betchworth Limeworks, Surrey, England, Plenus Marls, middle Late Cenomanian; 1, dorsal view; 2, ventral view; 3, view of the anterior end showing concentric growth layers and radial ridges, x2. A large species of *Praeactinocamax*. Figured by Christensen (1974, Pl. 4: 4).


Figs 6-7. *Actinocamax verus* Miller, 1823, Geological Museum, Copenhagen, MGUH 23718, Bavndode Greensand Formation, Bornholm, Denmark, middle part of *Bavndode* zone, early, but not earliest, Late Campanian; 6, dorsal view; 7, lateral view. Figured by Christensen & Schulz (in press, Pl. 1: 2).

Figs 8-10. *Gonioteuthis quadraquadrata* quadraquadrata (Blainville, 1827), Geological Museum, Copenhagen, MGUH 17529; Vaals Formation, CPL quarry, Hallembeaye, Belgium, lower part of *Inoceramus lingua/Gonioteuthis quadraquadrata* Zone, early, but not earliest, Early Campanian; 8, dorsal view; 9, lateral view; 10, view of the anterior end, x1.5. A species with a deep pseudoalveolus and well-developed granulation. Figured by Christensen & Schmid (1987, Pl. 3: 8–11).

Figs 11-12. *Gonioteuthis westfalica* (Schlüter, 1874), Geological Institute, Lund, Sweden, LO 4859, Ringeleslätt, Scania, Sweden, early Santonian; 11, dorsal view; 12, view of the anterior end, x2. A specimen of *G. westfalica* which has a flat anterior end with a central pit. Figured by Christensen (1975a, Pl. 2: 4).


Figs 16-17. *Goniocamax lundgreni* (Stolley, 1897), Geological Museum, Copenhagen, Denmark, MMH 13078, Arnager Limestone Formation, west of Arnager, Bornholm, Early Coniacian; 16, dorsal view; 17, ventral view. Figured by Christensen & Schulz (in press, Pl. 2: 1).

Figs 18-19. *Belemnocamax boweri* Cric, 1910, Geological Museum, Copenhagen MOU 22068 (cast), Wunstorf, Lower Saxony, Germany, early Middle Cenomanian, *Acanthoceras rhomomagense* Zone, *Turrilites cosatus* Subzone; 18, ventral view, x3; 19, lateral view, x3. Figured by Christensen (1993c, Fig. 3F).

Figs 20-21. *Belemnittella mucronata* (Schlotheim, 1813), neotype, Niedersächsisches Landesamt für Bodenforschung, Hannover, Germany, kca 5/2, Germainia IV pit at Misburg near Hannover, middle part of the *basiplana/spiniger* Zone, early, but not earliest, Late Campanian; 20, ventral view; 21, lateral view. A large species of *Belemnittella* with a stout guard. Figured by Christensen et al. (1975, Pl. 1: 1).

All specimens are coated with ammonium chloride, and are natural size unless otherwise stated.

**Diagnosis.** — Medium-sized belemnitellids (up to 85 mm long), with a short cone-shaped alveolar fracture or pseudoalveolus of varying depth (Riedel-Quotient from 3 to 20); guard usually only slightly flattened or not flattened at all ventrally; generally subcylindrical or cylindrical in ventral view and cylindrical in lateral view; juvenile guard short and stout; surface markings, including dorso-lateral longitudinal depressions and double furrows, vascular imprints, striae, and granules, well developed; Schatzky distance small, usually 0.5 to 4.5 mm; bottom of ventral fissure commonly sinue-shaped forming a large angle with the wall of the pseudoalveolus; vascular imprints branch off dorso-lateral double furrows posteriorly at an angle less than 30 degrees; relationship between length of guard and dorso-ventral diameter at the alveolar end generally allometric.

**Discussion.** — Naidin (1964b) recognized two subgenera of *Gonioteuthis:* *G. (Gonioteuthis)*, type species *Belemnites quadratus* and *G. (Goniocamax)*, type species *Actinocamax lundgreni*. Christensen & Schulz (in press) elevated subgenus *Goniocamax* to full generic rank, following an earlier suggestion by Ernst & Schulz (1974) (see discussion below).

The Middle Coniacian to Early Campanian evolutionary lineage of *Gonioteuthis* includes, in ascending order, *G. praevestfalcica, G. westfalica, G. westfalica-granulata* (Stolley, 1897), *G. granulata* (Blainville, 1827), *G. granulataquadraquadrata* (Stolley, 1897), *G. q. quadraquadrata* and *G. quadraquadrata gracilis* (Stolley, 1892) (Fig. 2). It has been studied in great detail by German authors, including Stolley (1892, 1987, 1916, 1930), Ernst (1964, 1966, 1968), Ernst & Schulz (1974) and Ulbrich (1971), in addition to Christensen (1975a, 1975b, 1986, 1991, 1994), Christensen & Schmid (1987), Christensen & Schulz (in press), Jarvis (1980) and Jagt, Kennedy, Burnett, Christensen & Dhoodt (1995). In addition to the members of this evolutionary lineage the following species and subspecies are placed in *Gonioteuthis:* the Early Santonian *G. ernsti* Christensen & Schulz (in press), which occurs on Bornholm and in the Münster Basin and *G. quadraquadrata scaniensis* Christensen, 1975a from the latest Early Campanian of Scania (Christensen 1975a, 1986) (Fig. 2). According to Christensen (1975a, 1986) the latter taxon is a geographical subspecies, which is closely allied to *G. quadraquadrata gracilis*.

The *Gonioteuthis* lineage survived for around 7 Ma and has been considered as an example of phyletic gradualism, viz. gradual transformation of a suite of characters through time. The general trends in evolution are: (1) gradual calcification of the anterior end of the guard, (2) increasing size and stoutness of the guard and (3) gradual development of granulation (Ernst 1964, Ernst & Schulz 1974).

In the oldest member of the lineage, the late Middle Coniacian-early Santonian *G. praevestfalcica,
less than 10% of the specimens are granulated, whereas in the succeeding species, the late Early-early Middle Santonian *G. westfalica*, about the half of the specimens are granulated (Christensen & Schulz, in press). Granulation becomes a very prominent character in younger species. With respect to the depth of the pseudoalveolus, Christensen & Schulz (in press) showed that *G. praewestfalica* has a deeper pseudoalveolus than *G. westfalica* from the early Early Santonian, and the depth of the pseudoalveolus is virtually identical in *G. praewestfalica* and *G. westfalica* from the late Early Santonian. Thus, the depth of the pseudoalveolus first decreases from the late Middle Coniacian to the early Early Santonian and this trend is reversed in the late Early Santonian.

Ernst (1964) and Christensen (1991) noted that coeval populations of *G. q. quadrata* from the middle Early Campanian chalk of Lägerdorf and marl of Misburg/Höver in northwest Germany differ with respect to the depth of the pseudoalveolus. The specimens from Misburg/Höver have a slightly deeper pseudoalveolus. The specimens from Lägerdorf are thus phylogenetically retrograde compared to coeval specimens from Misburg/Höver. *G. quadrata scaniensis* from Scania is also phylogenetically retrograde in comparison with the coeval *G. quadrata gracilis* from northwest Germany. The mean Riedel-Quotient of *scaniensis* is 5.1 (Christensen 1975a), while it is ±4.5 in *gracilis* (Ernst 1964).

Mitchell (1994, 1995) showed that populations of *Gonioteuthis* from the Late Santonian *Marsupites testudinarius* and *Uintacrinus socialis* Zones of Yorkshire are phylogenetically retrograde in comparison with coeval populations from northwest Germany. He also demonstrated that populations of *Gonioteuthis* from the early Early Campanian of Yorkshire and northwest Germany have reached the same evolutionary development.

It can thus be concluded that the evolution of *Gonioteuthis* lineage with respect to the depth of the pseudoalveolus is not as straightforward as supposed earlier. The depth of the pseudoalveolus first decreases and then increases in its earliest members, and the evolutionary development of populations of the lineage may be delayed in some areas.

**Distribution.** — *Gonioteuthis* had its evolutionary centre in northwest Europe and is recorded almost exclusively from the Central European Subprovince. It occurs very rarely in the northern part of the European Tethyan Realm. The genus existed from the late Middle Coniacian to the boundary between the Early and Late Campanian (Fig. 1). The stratigraphical ranges of the species are shown in Figure 2.

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**Plate 2**

Figs 1-2. *Belemmitella najdini* Kongiel, 1962, British Geological Survey, Nottingham, BGS CTW 5478, Caistor, Norfolk, Beeston Chalk, between Flint 9-10, late Late Campanian; 1, ventral view; 2, view of the split anterior end, x1.5. A small, slender species of *Belemmitella*. Schatzky Distance small, 7.7 mm; fissure angle large, 120.5 degrees; Birkelund Index, 4.7. Figured by Christensen (1995, Pl. 4: 1-4).

Figs 3-5. *Belemmitella minor* Jeletzky, 1951 form II Christensen, 1995, holotype, Natural History Museum, London, BMNH C43553, Whittingham, Norfolk, Paramoudra; Chalk, late, but not latest, Late Campanian; 3, ventral view; 4, lateral view; 5, view of the split anterior end. A stout and very large species of *Belemmitella* with well-developed vascular markings. Schatzky Distance large, 12.0 mm; fissure angle small, 15.5 degrees; Birkelund Index, 3.4. Figured by Christensen (1995, Pl. 7: 3-6; Fig. 20D).

Figs 6-7. *Fusiteuthis polonica* Kongiel, 1962, holotype, Bochnotica, Poland, latest Maastrichtian; 6, ventral view; 7, ventral view of the anterior part of the guard, approximately x6. Figured by Kongiel (1962, Pl. 1: 1-3).

Figs 8-10. *Belemnella lanceolata* (Schlotheim, 1813), Geological Museum, Copenhagen, MMH 13115, Balsvik, Scania, Sweden, earliest Early Maastrichtian; 8, ventral view; 9, lateral view; 10, view of the split anterior end, x2. Figured by Christensen (1975a, Pl. 12: 4).

All specimens are coated with ammonium chloride, except Figs 6-7, and are natural size unless otherwise stated.
but a few granulated specimens of *B. ex gr. grossouvrei* and *B. mammillatus* have been recorded (Christensen 1986).

Naidin (1964b) placed species of the *grossouvrei* group in his new subgenus *Actinocamax* (*Paractinocamax*), type species *A. grossouvrei*, and *mammillatus* in his new genus *Belemnellolocamax*. He distinguished *A. (Paractinocamax)* from *Belemnellolocamax* on the basis of the size of the juvenile guard and claimed that *A. (Paractinocamax)* has a short juvenile guard, while *Belemnellolocamax* has a long and slender juvenile guard. Christensen (1986) noted, however, that specimens of the *grossouvrei* group from west Europe have a very elongated juvenile guard. He therefore placed the *grossouvrei* group in *Belemnellolocamax*, and *A. (Paractinocamax)* was considered a junior synonym of *Belemnellolocamax*. Christensen (1991) figured two juvenile specimens of *B. ex gr. grossouvrei* from southern England, both of which are very elongated.

The *grossouvrei* group is rare; about 60 specimens are known from west Europe, and about 60–70 specimens are recorded from the Russian Platform. The specimens from west Europe have been assigned to eight species and varieties: *grossouvrei, toucasi* (Janet, 1891), *alfridi* (Janet, 1891), *depressus* (Andreae, 1895), *depressus var. fusiformis* (Andreae, 1895), *mammillatus var. germanica* (Stolley, 1930), *mammillatus var. ornatus* (Moberg, 1885) and *blackmorei* (Crick, 1907).

Naidin (1964b) distinguished five subspecies of *B. grossouvrei*: *depressus, toucasi* and *alfridi* from west Europe, and two new subspecies, *pseudotoucasi* and *pseudolfridi*, from the Russian Platform. In addition, Nikitin (1958) established *B. toucasi var. seimensis* and Glazunova (1972) recorded *B. cf. toucasi* and *B. alfridi* from the Russian Platform.

To sum up, eleven taxa have been established within the *grossouvrei* group on the basis of relatively little material. Thus, the genus has been the subject of excessive subdivision by previous authors. The group is currently being revised by W. K. Christensen and M.-G. Schulz, Kiel on the basis of material from west Europe.

**Distribution.** — The genus appears probably at the base of the Santonian and continues into the early Late Campanian (Figs 1–2). The Santonian-Early Campanian *B. ex gr. grossouvrei* is widely distributed but rare in the North European Province; it occurs from Scania in southern Sweden in the north to the Corbières in the French Pyrénées in the south. The latest Early Campanian *B. mammillatus* is extremely common in Scania, but rare outside this area; a little over 100 specimens are recorded from northern Germany, Poland and the eastern part of the Russian Platform (Christensen 1975a). The early Late Campanian *B. balsvikensis* is also extremely common in Scania, but outside this area it is unknown except for two specimens from northwest Germany. Thus, the geographical distribution of the genus was gradually reduced during its stratigraphical range.

**Genus Goniocamax** Naidin, 1964b

**Type species.** — *Actinocamax lundgreni* Stolley, 1897, p. 285, Pl. 3: 16–20, non 15, by original designation of Naidin (1964b: 104).

**Emended diagnosis.** — Small to medium-sized (up to 80 mm long) non-granulated belemnitellids, with a shallow to relatively deep pseudoalveolus; guard markedly flattened ventrally, lanceolate in ventral view and subcylindrical in lateral view; juvenile guard short and stout; guard with dorso-lateral longitudinal depressions and double furrows, vascular imprints, and longitudinal striae; longitudinal striae usually more distinct than vascular markings; Schatzky distance small, 2 to 4 mm; bottom of ventral fissure commonly straight or slightly curved forming a medium-sized angle, about 30 to 50 degrees, with wall of pseudo-alveolus; vascular imprints branch off dorso-lateral double furrows posteriorly at an angle less than 30 degrees; allometric relationship of length of guard and dorso-ventral diameter at protoconch; adults stouter than juveniles.

**Discussion.** — Christensen & Schulz (in press) raised the subgenus *Gonioteuthis* (*Goniocamax*) to full generic rank and included the type species and its close allies in *Goniocamax*, that is *G. birkelundae* Christensen & Schulz (in press), *G. striatus* Christensen & Schulz (in press), *G. essenensis* (Christensen, 1982) and *G. mirabilis* (Arkhangelsky, 1912).

*Goniocamax* is closely similar to *Gonioteuthis*, but differs from that genus by being non-granulated, flattened ventrally, lanceolate in ventral view, and the bottom of the ventral fissure is usually straight or slightly curved forming a medium-sized angle with the wall of the pseudoalveolus.

**Distribution.** — *Goniocamax* occurs mainly in the Central Russian Subprovince and Balto-Scandia. It occurs very rarely in the Central European Subprovince. It appears at the base of the Coniacian and continues into the Early Santonian (Christensen & Schulz, in press) (Fig. 1). The stratigraphical ranges of most of the species are shown in Figure 2.

**Genus Belemnitta d’Orbigny, 1840**

[ICZN 1985, Opinion 1328; name no. 2269]

**Type species.** — *Belemnites mucronatus* Schlotheim, 1813, p. 111, by subsequent designation by Herrmannsen (1846: 105); ICZN Opinion 1328 (1985); name no. 2979.
Species of Belemnitella, NW Europe

**Fig. 5.** Stratigraphical range and inferred phylogeny of Santonian-Maastrichtian species of *Belemnitella*. Modified from Christensen (1995). Stage abbreviations after Harland et al. (1989). Ages in Ma after Obradovich (1994).

**Diagnosis.** Small to large belemnellids (length from apex to protoconch up to 80 mm) with a deep alveolus; anterior end of guard completely calcified and prolonged ventrally along ventral fissure in a tongue-like extension; well developed dorso-lateral longitudinal depressions and straight double furrows, in addition to single lateral furrows; longitudinal strie may be present; vascular imprints branch off double furrows posteriorly at an angle less than 30 degrees; juvenile guard short and stout; alveolar angle 17 to 24 degrees, Schatzky distance long, commonly larger than 4 mm; relationship of length from apex to protoconch and dorso-ventral diameter at protoconch generally isometric.

**Discussion.** Christensen (1995) introduced a classification of size ranges of species of *Belemnitella* based on the length from the apex to the protoconch. This is as follows: (1) guard small; length from apex to protoconch less than 55 mm; (2) guard large; length from apex to protoconch between 55 and 65 mm; (3) guard very large; length from apex to protoconch larger than 65 mm. He also introduced a classification of the relative length of species of *Belemnitella* based on the Birkelund Index. This is as follows: (1) guard stout; mean value of Birkelund Index less than 4; (2) guard slender; mean value of Birkelund Index 4 to 5; (3) guard very slender; mean value of Birkelund Index larger than 5. It should be stressed that it is acceptable to calculate the mean value of the Birkelund Index in species of *Belemnitella*, because the relationship of the length from the apex to the protoconch versus the dorso-ventral diameter is generally isometric (Christensen 1995).

More than two dozen species, subspecies, and varieties of *Belemnitella* from the Late Campanian and Early Maastrichtian have been established. Many of these are poorly understood, because they were commonly erected on the basis of relatively little material, and the variation of the critical characters was rarely studied (Christensen 1988, 1993a).

Christensen & Schulz (in press) established *Belemnitella schmidi*, which appears at the base of the Santonian. They considered this species as the earliest member of *Belemnitella*. It is a transitional form possessing characters in common with the genera *Goniocamax* and *Belemnitella*. It probably evolved from *Goniocamax lundgreni* by allopatric speciation. *B. propinqua* (Moberg, 1885) enters some way above the base of the Santonian and is probably the lineal descendent of *B. schmidi* (Christensen & Schulz, in press). *B. propinqua* was considered previously to be the earliest member of *Belemnitella* by Jeletzky (1949b, 1955), Naidin (1964a, 1964b, 1974) and Christensen (1971, 1973, 1991), although some authors, including Birkelund (1957) and Glazunova (1972), placed it in *Actinocamax*.

Christensen (1995) analyzed the evolutionary trends
of *Belemnitella* from the Early Santonian to the Late Maastrichtian on the basis of biometric analyses of 43 samples, representing 20 species and subspecies. He tentatively recognized two lineages of *Belemnitella* (Fig. 5):

1. A Santonian–Maastrichtian lineage, which is derived from *Goniocamax lundgreni*. It includes *B. schmidi*, *B. propinquus*, *B. praecursor*, *B.woodi* Christensen, 1995, *B. minor I* Jeletzky, 1951, *B. minor II* Christensen, 1995, *B. minor III* Christensen, 1995, *B. ex gr. junior sensu* Keutgen & van der Tuuk, 1990 and *B. junior*. These species are large to very large and stout to slender. In addition, the fissure angle is large to very large, the bottom of the ventral fissure is generally straight and the Schatzky distance is medium-sized to large.

2. A late Late Campanian–Early Maastrichtian lineage, which comprises the late Late Campanian *B. langei* Jeletzky, 1948 and *B. najdini* Kongiel, 1962, in addition to the Early Maastrichtian *B. pulchra* Schulz, 1982. These species are small and slender to very slender. Moreover, the fissure angle is large to very large, the bottom of the ventral fissure is irregular and the Schatzky distance is small. The origin of this lineage is unknown.

In addition to the species of the two lineages, other species are recorded from the Campanian and basal Maastrichtian. The early Early Campanian *B. alpha* Nadin, 1964a is probably an off-shoot derived from *B. praecursor* (see Christensen 1995). Species of unknown origin or poorly known species include the late Campanian *B. aff. langei* Jeletzky (1986, 1993a), *B. langei sensu* Schulz (1978), *B. cf. najdini* of Schulz (1978) and *B. pauli* Christensen, 1995, in addition to the basal Maastrichtian *B. minor sensu* Christensen (1975a) and *B. langei sensu* Birkelund (1957). The latter was placed in synonymy with *B. minor III* with a query by Christensen (1995), who also discussed the species mentioned above.

The little known late Campanian *Belemnitella hoeferi* (Schloenbach, 1867) from the Northern Calcareous Alps of Austria is a valid species, which belongs to the *B. mucronata* group of Christensen (1995). Christensen (submitted) redescribed this species, including biometric analysis, on the basis of material from the Gschliefgraben area in Austria. It can be distinguished from most species of the *B. mucronata* group on the basis of the slender guard (mean Birkelund Index about 4.5). *Belemnitella* sp. from the ‘Craie marnesse’ of Chartreuse in the Sub-Alpine Chain, Savoie, southeast France (Combémoré 1996) is closely similar to *B. hoeferi* with respect to size, shape and slenderness of the guard, in addition to surface markings. Thus, the two taxa may be conspecific, but a specific determination of the specimens from Chartreuse is not possible, because the internal characters are unknown.

**Distribution.** *Belemnitella* as interpreted here appears at the base of the Santonian and continues to the top of the Maastrichtian (Fig. 1). It is recorded from the North European and North American Provinces, in addition to the northern part of the Tethyan Realm. The stratigraphical ranges of the species are shown in Figures 2–5.

Genus *Belemnella* Nowak, 1913
[ICZN 1985, Opinion 1328; name no. 2270]

Type species. *Belemnites lanceolatus* Schlotheim, 1813, p. 111, by subsequent designation by von Bülow-Trümmel (1920: 195); ICZN Opinion 1328 (1985); name no. 2980.

**Diagnosis.** Large belemnitellids (length from apex to protoconch up to 110 mm) with a deep alveolus; anterior part of guard complete calcified and prolonged ventrally around ventral fissure in a tongue-like extension; guard markedly flattened ventrally; well-developed dorso-lateral longitudinal depressions and double furrows, which undulate posteriorly, in addition to single lateral furrows; vascular imprints weakly developed or not present in early forms, younger forms with distinct vascular imprints; vascular imprints branch off dorso-lateral double furrows posteriorly at an angle exceeding 30 degrees; juvenile guard long and slender (needle-shaped); alveolar angle small, 10 to 21 degrees; Schatzky distance short, 0 to 4.5 mm, commonly less than 4 mm; relationship of length from apex to protoconch and dorso-lateral diameter at protoconch allometric; adult specimens stouter than juvenile specimens.

**Discussion.** Three subgenera have been established, the non-nomotypical subgenus and *B. (Pachybelemnella)* Schulz, 1979, which are Early Maastrichtian (Schulz 1979), in addition to the Late Maastrichtian *B. (Neobelemnella)* Nadin, 1975. These subgenera include only large species in contrast to the genus *Belemnella* (see above).

Schulz (1979) studied the Early Maastrichtian subgena in great detail and stressed the importance of the shape of the guard in ventral view compared with the length from the apex to the protoconch. Since growth is allometric in *Belemnella* he introduced the derived variable ‘standardized length from the apex to the protoconch’ in order to compare specimens of different size. The calculation of this variable is rather laborious. The shape of the guard in ventral view is defined by an index, which includes three different measurements of the lateral diameter.

Schulz distinguished two Early Maastrichtian subgenera of *Belemnella*: the slender *B. (Belemnella)* and the stout *B. (Pachybelemnella)*.
Subgenus Belemnella (Belemnella) Nowak, 1913

**Diagnosis.** — *Belemnella* with a slender guard; species with a lanceolate guard very slender; species with cylindrical to cone-shaped guard slender.

**Discussion.** — Schulz (1979) recognized two, possibly three, evolutionary lineages within *Belemnella* (*Belemnella*). A lineage including *lanceolata* (oldest), *gracilis* (Arkhangelsky, 1912) and *fastigata* Schulz, 1979 (youngest) (Fig. 2). The general trends of this lineage are: (1) the decreasing length from the apex to the protoconch, (2) the shape of the guard in ventral view changes from lanceolate to cylindrical, (3) the increasing Schatzky Distance and (4) the increasing alveolar angle. Another lineage includes *lanceolata* and *longissima* Schulz, 1979 (Fig. 2). The latter taxon is characterized by its very large length from the apex to the protoconch and its very lanceolate shape in ventral view.

Schulz (1979) placed *B. praearkhangelskii* Naidin, 1964a in subgenus *Belemnella* with a query (Fig. 2). This species occurs only at a very restricted horizon at Hemmoor and Kronsmoor, that is in the middle part of the sumensis Zone, 35–38 m above the base of the Maastrichtian. He also recorded four unhorizoned specimens of this taxon from Møns Klint, Denmark, and Rügen, Germany, which may have come from the same zone. Keutgen in Jagt et al. (1995) has subsequently reported the species from the middle part of the sumensis Zone of northeast Belgium. Schulz (1979) suggested that *B. praearkhangelskii* probably belongs to another evolutionary lineage.

*B. (B.) lanceolata* has been widely used as a zonal index fossil of the early Early Maastrichtian (see discussion by Christensen 1996). Schulz' (1979) concept of this species is more restricted, however, than earlier authors, including Jeletzky (1951), Naidin (1952) and Birkelund (1957). He recorded *B. (B.) lanceolata* only from the earliest Early Maastrichtian.

**Distribution.** — This subgenus is distributed in the Early Maastrichtian (Fig. 1) and occurs in the North European Province and northern part of the Tethyan Realm. The stratigraphical ranges of the species are shown in Figure 2.

Subgenus Belemnella (Pachybelemnella) Schulz, 1979

**Type species.** — *B. (P.) obtusa* Schulz, 1979, p. 106, Pl. 9: 1–8, Pl. 12: 9, by original designation.

**Diagnosis.** — *Belemnella* with a stout guard; species with a very lanceolate guard slender; species with a lanceolate or cylindrical guard stout.

**Discussion.** — Schulz (1979) distinguished two lineages within this subgenus. A lineage including *inflata* (Arkhangelsky, 1912) (oldest), *pseudobtusa* Schulz, 1979, *obtusa, sumensis* Jeletzky, 1949a and *cimbrica* Birkelund, 1957 (youngest) (Fig. 2), which is derived from *B. (B.) lanceolata*. The general trends in this lineage are (1) the decreasing length from the apex to the protoconch, (2) the shape of the guard in ventral view changes from lanceolate to cylindrical, (3) the increasing Schatzky Distance and (4) the increasing alveolar angle. Another lineage includes *inflata* and *desnensis* (Jeletzky, 1941) (Fig. 2). The latter species has a guard which is strongly lanceolate in ventral view.

**Distribution.** — *Belemnella* (Pachybelemnella) is distributed in the Early Maastrichtian (Fig. 1) and occurs in the North European Province and northern part of the Tethyan Realm. The stratigraphical ranges of the species are shown in Figure 2.

Subgenus Belemnella (Neobelemnella) Naidin, 1975

**Type species.** — By monotypy *Belemnitella kazimirovensis* Skołosdrowna, 1932, p. 117.

**Diagnosis.** — *Belemnella* with a large alveolar angle, 20 to 23 degrees; large Schatzky Distance, 3 to 5 mm; well developed vascular markings; a short and cone-shaped juvenile guard; vascular imprints branch off dorso-lateral double furrows posteriorly at an angle less than 40 degrees.

**Discussion.** — Jeletzky (1951) spelled the species name *casimirovensis*, as did Birkelund (1957) and some later authors. According to the International Code of Zoological Nomenclature, Article 33c, 1985, this is, however, an incorrect subsequent spelling.

The subgenus *B. (Neobelemnella)* has several characters which otherwise are typical for the genus *Belemnitella*, namely a short and cone-shaped juvenile guard and well developed vascular imprints. Moreover, the Schatzky Distance is relatively large.


Naidin (1975) suggested a late Early Maastrichtian-Late Maastrichtian evolutionary lineage, which includes, in ascending order, *Belemnella sumensis sumensis* Naidin 1964a, *B. sumensis postsumensis* Naidin, 1964a, *B. sumensis praearkhangelskii* Naidin, 1964a, *B. sumensis kajnarensis* Naidin, 1964a and *Belemnella (Neobelemnella) kazimirovensis*. This lineage bridges the Early and Late Maastrichtian boundary in the eastern part of the Russian Platform. According to Schulz (1979), however, *B. sumensis* Naidin, 1964a is not conspecific with *B. sumensis* Jeletzky.
Central Russian Subprovince

Central European Subprovince

Boundary between North European Province and Tethyan Realm

Fig. 6. Map showing the Central European and Central Russian palaeobiogeographical Subprovinces within the North European Province as defined on belemnites. A third subprovince, the Balto-Scandian Subprovince, is recognized during the Early Coniacian and latest Early Campanian-early Late Campanian. It is characterized by Goniocamax lundgreni in the Early Coniacian and the genus Belemnellocamax in the latest Early Campanian and early Late Campanian. Late Cretaceous land and sea areas represent maximum inundation for all stages. The boundaries are not reliable in detail and the biogeographic units are typically gradational in character. After Christensen (1976).

1949a, and he placed B. s. sumensis and B. sumensis postsumensis in synonymy with B. (B.?) praearkhangel skii.

Distribution. — This subgenus is restricted to the Late Maastrichtian (Fig. 1) and occurs in the North European Province and northern part of the Tethyan Realm. B. (N.) kazimiroviensis occurs in the Late Maastrichtian in the eastern part of the Russian Platform, late Late Maastrichtian in Denmark and latest Late Maastrichtian in The Netherlands (Christensen 1996).

Genus Fusiteuthis Kongiel, 1962

Type species. — By monotypy F. polonica Kongiel, 1962, p. 28, Pl. 1: 1–3.

Diagnosis. — Medium-sized belemnitellids with a shallow alveolus; guard lanceolate in ventral and lateral views; dorso-lateral longitudinal depressions present, guard otherwise smooth.

Discussion. — F. polonica was established on the basis of a single specimen from the latest Maastrichtian of central Poland (Pl. 2: 6–7). Later, Naidin (1973, 1975) recorded two specimens of F. sp. from the Late Maastrichtian of Crimea.

J. A. Jeletzky (unpublished MS 1972) has suggested that the genus is dubious being based on a possibly pathological specimen. I agree.

Distribution. — This genus is restricted to the latest Maastrichtian and occurs in Poland and the Crimea.

Palaeobiogeography

The belemnitellids were distributed in the North American and North European Provinces of the North Temperate Realm, in addition to the northern European margin of the Tethyan Realm (Christensen 1976, 1988, 1993b).

North European Province

This province extends from Ireland in the west to the Ural Mountains and beyond in the east (Fig. 6). The centre of origin of the belemnitellids lay there for the following reasons. They are common and all known genera and subgenera occur there, the earliest species of the family, Praeactinocamax primus, appears in the Early Cenomanian, some way above the base the sub-stage, in this province and the earliest species of the...
Table 2. The occurrence of belemnitellid genera and subgenera in the North European and North American Provinces of the North Temperate Realm and the Tethyan Realm. *Fusiteuthis* is most likely a *nomen dubium*. The genera and subgenera are ranked with respect to their palaeobiogeographical distribution. Nine genera and two subgenera occur in the North European Province, five genera and two subgenera in the Tethyan Realm in Europe and essentially two genera in the North American Province. The numbers 1–12 refer to the first appearances of the genera and subgenera in the various palaeobiogeographical units. Detailed data are not available for the subgenera of *Belemnella*.  

<table>
<thead>
<tr>
<th>Genera and subgenera</th>
<th>North European Province</th>
<th>Tethyan Realm</th>
<th>North American Province</th>
</tr>
</thead>
<tbody>
<tr>
<td>Praeactinocamax</td>
<td>+¹</td>
<td>+²</td>
<td>+¹</td>
</tr>
<tr>
<td>Belemnella</td>
<td>+⁴</td>
<td>+⁵</td>
<td>+⁶</td>
</tr>
<tr>
<td>Belemnella (Belemnella)</td>
<td>+</td>
<td>+</td>
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</tr>
<tr>
<td>Belemnella (Pachybelemnella)</td>
<td>+</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Belemnella (Neobelemnella)</td>
<td>+</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Goniotethis</td>
<td>+⁷</td>
<td>+⁴</td>
<td></td>
</tr>
<tr>
<td>Belemnelloccamax</td>
<td>+⁹</td>
<td>+¹⁰</td>
<td></td>
</tr>
<tr>
<td>Actinocamax</td>
<td>+¹¹</td>
<td>(+)¹²</td>
<td></td>
</tr>
<tr>
<td>Goniocamax</td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Belemnocamax</td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fusiteuthis</td>
<td>+</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

other genera either occurs only there or appears earlier there than elsewhere (Table 2). The last belemnitellids became extinct at the Maastrichtian-Danian boundary. In addition to the belemnitellids, the genera *Neohibolites* and *Parahibolites* of the family Belemnopsidae also occur in this province.

The North European Province includes the Central European, Central Russian and Balto-Scandian Subprovinces (Fig. 6). The Central Russian and Central European Subprovinces are well-defined from the Middle Coniacian to the boundary between the Early and Late Campanian, a period of around 7 Ma. The two subprovinces are characterized by two independently evolving belemnite lineages: the *Goniotethis* stock inhabited the Central European Subprovince, and the *Goniocamax-Belemnella* stock inhabited the Central Russian Subprovince (Christensen 1975a, 1976, 1988, 1990b, Christensen & Schulz in press).

The belemnite faunas of Balto-Scandia show affinity to those of the Central Russian Subprovince at certain times and to those of the Central European Subprovince at other times. However, in the Early Coniacian and latest Early Campanian and early Late Campanian a third subprovince can be recognized, named here the Balto-Scandian Subprovince. It is characterized by *Goniocamax lundgreni* in the Early Coniacian, *Belemnelloccamax mammillatus* in the latest Early Campanian and *B. balsvikensis* in the early Late Campanian. *B. mammillatus* co-occurs with *Belemnittella mucronata*, *Gonioteuthis quadrata* and *Belemnelloccamax ex gr. grossouvrei* and comprises about 90–95% of the belemnite fauna (Christensen 1975a, Table 3). *B. balsvikensis* co-occurs with *Belem-
B. mucronata was recorded also from Austria, but this may be a misconception. B. hoeferi occurs in the Northern Calcareous Alps (Schloenbach 1867, Christensen, submitted; see above).

Species of the following genera and subgenera are recorded from the Tethyan Realm: Praeactinocamax, Gonioteuthis, Belemnelloccamx, Belemnitella, Belemnella (Belemnella), B. (Pachybelemnella) and B. (Neo-belemnella) (Fig. 4). The majority of the species occurring in the Tethyan Realm are conspecific with those from the North European Province and they thus provide a basis for correlation.


The middle Late Cenomanian Praeactinocamax plenus is the earliest belemnittellid recorded from the Tethyan Realm (Gale & Christensen 1996) and belemnittellids are not recorded from the Turonian, Coniacian and Early and Middle Santonian.

North American Province
This province includes Greenland, the Western Interior of North America, in addition to the Atlantic and Gulf coasts of the USA (Fig. 7). Belemnittellids are generally very rare and belemnopseids do not occur. The belemnite faunas are essentially restricted to species of Praeactinocamax (Turonian to Early Santonian) and Belemnitella (uppermost Santonian to Maastrichtian) (Fig. 3). Actinocamax is represented only by a single specimen of A. veras? from central East Greenland (Jeletzky in Donovan 1954) and two specimens of A. laevigatus from Kansas (Jeletzky 1961). However, Jeletzky (1961) suggested that the two specimens from Kansas may be juveniles of one of the large species of Praeactinocamax. The belemnite fauna of the middle Turonian seems to be rather diverse, but this may due to excessive subdivision (see above). The earliest belemnittellids from this province are Middle Turonian in age. The belemnittellids of the North American Province were discussed by Christensen (1993b).

Seibertz & Spaeth (1995) recorded three fragmentary belemnites from the Early Turonian of northern Mexico. The anterior end with the critical characters is missing in these specimens, but, nevertheless, Seibertz & Spaeth tentatively assigned them to Praeactinocamax cf. manitobensis. P. manitobensis occurs in the Western Interior of North America and has not been recorded previously south of Kansas (Cobban 1991). If the specimens from northern Mexico are correctly identified, which is open to discussion, then they are the most southerly belemnittellids recorded to date. Northern Mexico lay at palaeolatitude 15–20°N.

B. americana (Morton, 1830) occurs in the Mount Laurel Formation and basal part of the Navesink Formation of the Atlantic Coastal Plain (Owens & Sohl 1973, Fig. 4). In ammonite terms, the Mount Laurel Formation is late Late Campanian, but not latest Campanian in age (Kennedy & Cobban 1994), and the basal part of the Navesink Formation is latest Campanian and Early Maastrichtian in age (Kennedy, Johnson & Cobban 1995).

B. bulbosa occurs in the Fox Hills Formation of South Dakota, which is probably early Late Maastrichtian in age (Waage 1968, Landman & Waage 1993). A small species of Belemnitella, probably conspecific with B. bulbosa, occurs in the upper part of the underlying Pierre Shale, Baculites baculus and B. clinolobatus Zones. These zones are Early but not earliest Maastrichtian in age (Kennedy & Cobban 1993).

The majority of the belemnittellids of the North American Province are endemic, have a punctuated, strongly discontinuous stratigraphical distribution and are derived from species from the North European Province (Christensen 1993b). The endemic taxa include the Middle Turonian-Early Santonian species of Praeactinocamax and the Late Campanian-Maastrichtian species of Belemnitella. These taxa evolved from European species by allopatric speciation. Three European taxa, Actinocamax verus?, Belemnitella praecursor and B. ex gr. alpha/praecursor, occur in the uppermost Santonian-basal Early Campanian.
Since most of the belemnittellid species of the North American Province are endemic intercontinental correlation based on belemnites is not possible.

Mode of life of Late Cretaceous belemnites

The belemnittellids were neritic animals restricted to the shelf. They are common in nearshore sediments, such as biocalcarenites, greensands, marls and shallow-water chalks, and populations from these sedimentary deposits characteristically contain all growth stages. They are less common to virtually absent in deeper water chalks, and populations from these deposits consist mainly of adult specimens (Christensen 1976). W.J. Kennedy (personal communication, November 1996) noted that he has never seen a single specimen in cores from the Central Graben, of which he has examined several kilometres. It therefore appears that the breeding, spawning, hatching, and, possibly for the females at least, dying grounds, were inner neritic, shallow water environments. Adult specimens from offshore, deeper water chalks may be considered as strain fully-grown individuals that died outside their normal habitat (Christensen 1976, Surlyk & Birkelund 1977).

It is noteworthy that belemnittellids are absent or extremely rare at some horizons in the Late Campanian and Maastrichtian chalks of northwest Europe. At Kronsmoor in northwest Germany, the topmost five metres of the Late Campanian have not yielded belemnites, and *Belemnella* and *Belemnitella* are extremely rare in the basal three metres of the Maastrichtian (Schulz 1978, 1979, 1982). At Hemmoor in northwest Germany belemnites are virtually absent in the topmost five metres of the late Early Maastrichtian (only one specimen of *Belemnella fastigata* is recorded) and they are not recorded from the basal five metres of the early Late Maastrichtian (Schulz and Schmid 1983 b). They are virtually absent in the late Early Maastrichtian of Denmark (Surlyk 1970); only two specimens of *Belemnella cf. cimbrica* are recorded (Schulz and Schmid 1983b). The topmost six metres of the chalk of Hemmoor, which equates with the basal part of the late Late Maastrichtian *Belemnella kazimiroviensis Zone* of Denmark, have not yielded belemnites (Schulz and Schmid 1983b).

The belemnittellids were stenothermal animals adapted to life in warm-temperate waters according to the palaeotemperature curves by Jenkyns, Gale & Corfield (1994, Fig. 12; Cenomanian to Santonian of east Kent) and Schönfeld & Schulz (1996, Fig. 4; Late Campanian to Early Maastrichtian of northern Germany). These areas were situated at about palaeolatitude 40°N. According to Jenkyns et al. the palaeotemperature was about 21°C in the basal Cenomanian, increased gradually to a maximum of about 28°C at the Cenomanian-Turonian boundary and declined irregularly thereafter. It was about 22°C in the Coniacian and dropped to around 19°C in the Late Santonian. Following Schönfeld & Schulz the palaeotemperature was about 15°C in the early Late Campanian, increased to about 17°C in the earliest Maastrichtian and decreased to around 15°C in the middle Early Maastrichtian.

The South Temperate dimitobelids were also stenothermal shallow-water dwellers (Doyle & Howlett 1989), but they appear to be adapted to life in cooler waters than the belemnittellids. Pirrie & Marshall (1990) and Crame, Lomas, Pirrie & Luther (1996) suggested that the maximum surface water temperature was about 20°C in the Turonian-Coniacian and declined thereafter. It was about 14°C in the Santonian and Campanian and about 12°C in the Maastrichtian of James Ross Island, Antarctica (palaeolatitude 60–65°S).

Since both the belemnittellids and dimitobelids were shallow-water dwellers, oceans with deep water may have acted as physical barriers and precluded spread of these families. Warm tropical waters in the Tethyan Realm may also have acted as a physical barrier for these families.

The belemnopsoids *Neohibolites* and *Parahibolites* were widely distributed in the Albian-Cenomanian, from the Central European Subprovince of the North Temperate Province, across the Tethyan Realm to the South Temperate Province, and have been recorded from deep marine and continental slope deposits, as well as inner neritic sediments. The habitat of these genera may therefore have been surface oceanic, but they probably spent their breeding season inshore (Doyle & Howlett 1989). They were eurythermal, adapted to life in warm temperate as well as tropical waters.

Belemnittellid distribution

*Praeactinocamax* and *Belemnitella* are more widely distributed than the remaining genera and subgenera and occur in both the North European and North American Provinces, as well as in northern European part of the Tethyan Realm (Table 2). *Gonioteuthis, Belemnellocomax* and *Belemnella* are distributed in both the North European Province and the Tethyan Realm. *Actinocamax, Belemnocamax* and *Goniocamax* occurs only in the North European Province, except for a single specimen of *A. verus*? from East Greenland (see above), but they have different distribution patterns. *Actinocamax* is widely distributed in the North European Province, *Goniocamax* is mainly restricted to the Central Russian Subprovince and Balto-Scandia and *Belemnncamax* occurs only in northwest Germany and eastern England. Thus, the genera and subgenera can be ranked in the following way with respect to their
Fig. 8. Late Cretaceous belemnite stratigraphy of northwest Europe, Balto-Scandia and the Russian Platform. In order to ease reading of the diagram the following abbreviations are used for the genera.

- **P. = Praeactinocamax, A. = Actinocamax;**
- **Blt. = Belemnitella;**
- **Bln. = Belemnella;**
- **Blc. = Belemnellocamax;**
- **Gt. = Gonioteuthis;**
- **Gc. = Goniocamax; N. = Neohibolites. Vertical axis not to scale. Modified from Christensen (1986).**

<table>
<thead>
<tr>
<th>STAGES</th>
<th>BELEMNITE ZONES, NW EUROPE</th>
<th>ZONAL BELEMNITES, BALTO-SCANDIA</th>
<th>ZONAL BELEMNITES, RUSSIAN PLATFORM</th>
</tr>
</thead>
<tbody>
<tr>
<td>BELEMNITE ZONES, NW EUROPE</td>
<td><strong>Bln. kazimiroviensis</strong></td>
<td><strong>Bit. junior</strong></td>
<td><strong>Bln. kazimiroviensis</strong> (E)</td>
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<td><strong>UMBRIAN</strong></td>
<td><strong>Bln. fastigata</strong></td>
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<td></td>
</tr>
<tr>
<td><strong>NELSPRUIT</strong></td>
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</tr>
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<td><strong>Bln. obtusa</strong></td>
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<td><strong>Bln. lanceolata</strong></td>
<td><strong>B. lanceolata</strong></td>
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<td><strong>B. licharewi</strong></td>
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</tr>
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<td><strong>SUPERIAN</strong></td>
<td><strong>Bln. kazimiroviensis</strong></td>
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<td><strong>UPPER CAMPANIAN</strong></td>
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<td><strong>LOWER CAMPANIAN</strong></td>
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<tr>
<td><strong>CENOMANIAN</strong></td>
<td><strong>Bln. kazimiroviensis</strong></td>
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</tr>
</tbody>
</table>

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**Legend:**
- **P. = Praeactinocamax, A. = Actinocamax;**
- **Blt. = Belemnitella;**
- **Bln. = Belemnella;**
- **Blc. = Belemnellocamax;**
- **Gt. = Gonioteuthis;**
- **Gc. = Goniocamax; N. = Neohibolites. Vertical axis not to scale. Modified from Christensen (1986).**

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palaeogeographical distribution, from the largest to the smallest areal extent (Table 2):

(1) Praeactinocamax and Belemnitaella; (2) Gonioteuthis, Belemnellocamax and Belemnella; (3) Actinocamax; (4) Goniocamax; (5) Belemnocamax.

The palaeogeographical distribution of the belemnitellids was controlled by a number of factors, including sea-level changes, temperature, palaeocurrents, ecological tolerance and competition. Gale & Christensen (1996) suggested that the southwards migration of Praeactinocamax plenus into the Tethyan Realm in the Late Cenomanian was due to a significant fall of sea temperature, the so-called Plenus Cold Event, working in concert with the development of suitable shallow water habitats. In addition, lack of competition of the Tethyan belemnopseids, which disappeared in the Middle Cenomanian, may also have been of importance. Christensen (1976) suggested that the parallel evolution of the Gonioteuthis and Goniocamax-Belemnitaella stocks from the Middle Coniacian to the boundary between the Early and Late Campanian in the adjacent Central European and Central Russian Subprovinces, respectively, was due to mutual competition. The causes of the palaeogeographical distribution and migrations of the belemnitellids will be discussed in more detail in a forthcoming paper.

Late Cretaceous belemnite extinctions


The belemnitellids suffered three extinctions during the Late Cretaceous: (1) in the Middle Cenomanian-earliest Early Turonian, (2) around the Early and Late Campanian boundary and (3) at the Maastrichtian-Paleocene boundary. At the first extinction event, the belemnitellids Belemnocamax boweri and Praeactinocamax plenus of the P. primus-plenus lineage disappeared along with the belemnopseids. After this extinction the belemnitellids are extremely rare in the Turonian, which may be situated in the Central Russian Subprovince. The Central European Subprovince was repopulated by Gonioteuthis praewestfalica in the late Coniacian and belemnitellids did not invade the Tethyan Realm again until the Late Santonian.

The next extinction event occurred in the latest Early Campanian and early Late Campanian. Three genera disappeared, Actinocamax, Gonioteuthis and Belemnelloccax, and only one genus, Belemnitaella, survived. After this extinction, Belemnitaella expanded its area of distribution to cover the entire North European Province, where it is common. It also invaded the Tethyan Realm. However, only one species, B. mucronata, occurs in the lower part of the early Late Campanian (Fig. 2), that is a period of about 1.5 Ma.

The last extinction occurred at the Cretaceous-Paleocene boundary and no belemnites survived. Paleogene belemnites have been recorded earlier. However, the Eocene Bayanoteuthis rugifer Schloenbach, 1868, which occurs in southern Europe, has been reinterpreted as a pennatulacean coral (Riegraf 1991) and others are considered to be sepulids with a belemnite-like guard (Doyle, Donovan & Nixon 1994).

Late Cretaceous belemnite biostratigraphy of Europe

The belemnitellids are of fundamental importance in biostratigraphy and correlation of the Late Cretaceous of the North European Province, particularly during the Coniacian to Maastrichtian stages. They are common, widely distributed there and the fossilization potential is high (Christensen 1990b, 1996). 25 zones have been established in northwest Europe and a little less on the Russian Platform (Fig. 8).

The zonation of the west Europe and the Russian Platform is based upon the belemnopseid Neohibolites ultimus and species of Praeactinocamax for the Conenomanian, species of Belemnitaella for the Late Campanian, species of Belemnella for the Early Maastrichtian and species of Belemnitaella and Belemnella for the Late Maastrichtian. In the Middle Coniacian to Early Campanian the zonation of northwest Europe is based upon species of Gonioteuthis, while it is based upon species of Goniocamax and Belemnitaella on the Russian Platform (Fig. 8).

The Middle Coniacian to Early Campanian belemnite faunas of Balto-Scandia are of great importance, because they include species of both the Gonioteuthis and Goniocamax-Belemnitaella stocks, and thus provide a basis for correlation between the two subprovinces (Fig. 8).

Belemnitellids are extremely rare in the Turonian and early Coniacian of the Central European Subprovince and have a very restricted distribution in the Central Russian Subprovince (Christensen 1982). It appears that the belemnitellids retreated into refugia in the Turonian, which may be situated in the Central Russian Subprovince. The Central European Subprovince was repopulated by Gonioteuthis praewestfalica in the late Coniacian and belemnitellids did not
Some of the belemnite zones have been subdivided. For instance, Christensen (1995, 1996) subdivided the Late Campanian of Norfolk into four informal zones on the basis of large to very large species of Belemninitella (Fig. 9). These are in ascending order: the B. mucronata, B. woodi, B. minor I and B. minor II zones. The B. minor zones were subdivided into five informal subzones, mainly on the basis of small species of Belemninitella, B. langei, B. najdini and B. ex gr. langei/najdini. Further studies are necessary to see if these zones can be extended to other areas in Europe. Keutgen (1995) has shown subsequently that the zonation of Norfolk is applicable in northeast Belgium.

Christensen & Schulz (in press) subdivided the Coniacian and Lower Santonian of Bornholm, Denmark into three zones: the Coniacian Goniocamax lundgreni Zone, the early Early Santonian Gonioteuthis praewestfalica Zone and the late Early Santonian Gonioteuthis westfalica Zone (Fig. 10). These zones were subdivided into eight assemblage zones. However, it is very likely that the zones and assemblage zones are applicable only in Balto-Scandia for the following reasons. Some of the index species are very rare or not recorded from the Central European Subprovince (Goniocamax lundgreni, Belemninitella schmidi and B. propinqua), others are not recorded from the Central Russian Subprovince (Gonioteuthis praewestfalica, G. westfalica and G. ernsti), and Goniocamax birkelundae and G. striatus are not recorded outside Bornholm. Moreover, G. lundgreni occurs earlier on Bornholm (Early Coniacian) than in the Central Russian Subprovince (Middle Coniacian), and G. praewestfalica is late Coniacian in age in the Central European Subprovince, while it is early Early Santonian in age on Bornholm.

Acknowledgments
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Dansk sammendrag
Belemnit familien Belemnitellidae forekommer kun i Øvre Kridt på den nordlige halvkugle, fra Nedre Cenomanien til Øvre Maastrichtien. Den inkluderer følgende slægter og underslagter: Praeactinocamax, Actinocamax, Belemnocamax, Gonioteuthis, Belemnellophusus, Goniocamax, Belemninitella, Belemnella (Belemnella), Belemnella (Pachybelemnella), Belemnella (Neobelemnella) og den dubiose Fusiteuthis. Antallet af slægter og underslagter varierer gennem Øvre Kridt. Det er en til to i Cenomanien til stiger gradvist til seks i Nedre Santonian. Det aftager derefter gradvist til et nogenlig ni til tre i Maastrichtien.

Det evolutionære center for belemnitetilde er lå i
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**Fig. 10. Stratigraphical correlation diagram of the Coniacian and Santonian, showing faunal and Gonioteuthis zones of Lägerdorf and belemnite zones and assemblage zones of Bornholm.**

<table>
<thead>
<tr>
<th>STAGES</th>
<th>Lägerdorf, NW Germany</th>
<th>Bornholm, Denmark</th>
</tr>
</thead>
<tbody>
<tr>
<td>Faunal zones</td>
<td>Gonioteuthis Zones</td>
<td>Belemnite zones</td>
</tr>
<tr>
<td>Lower Santonian</td>
<td>westfalic granulata (RI: 11.0-12.5)</td>
<td>Gonioteuthis westfalica (RI &lt; 8.5)</td>
</tr>
<tr>
<td></td>
<td>westfalic</td>
<td>Gonioteuthis praewestfalica</td>
</tr>
<tr>
<td></td>
<td>(RI: 8.5-11.0)</td>
<td>Goniocamax lundgreni</td>
</tr>
<tr>
<td>Lower Cenomanian</td>
<td>westfalica</td>
<td>Goniocamax lundgreni</td>
</tr>
<tr>
<td></td>
<td>coranguinum/westfalica</td>
<td>Goniocamax lundgreni</td>
</tr>
<tr>
<td></td>
<td>westfalica</td>
<td>Goniocamax lundgreni</td>
</tr>
<tr>
<td></td>
<td>pachti/undulatoplicatus</td>
<td>Goniocamax lundgreni</td>
</tr>
<tr>
<td></td>
<td>bucailli/prae westfalica</td>
<td>Goniocamax lundgreni</td>
</tr>
<tr>
<td></td>
<td>involutus/bucailli</td>
<td>Goniocamax lundgreni</td>
</tr>
<tr>
<td></td>
<td>koeneni</td>
<td>Goniocamax lundgreni</td>
</tr>
<tr>
<td></td>
<td>not exposed</td>
<td></td>
</tr>
<tr>
<td>Coniacian</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
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</tbody>
</table>

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