Scolex morphology of monozoic tapeworms (Caryophyllidea) from the Nearctic Region: taxonomic and evolutionary implications

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Abstract: A comparative study of the scolecites of monozoic tapeworms (Cestoda: Caryophyllidea), parasites of catostomid and cyprinid fishes (Teleostei: Cypriniformes) in the Nearctic Region, was carried out using light and scanning electron microscopy. Scolecites of 22 genera of North American caryophyllideans were characterised and their importance for taxonomy, classification and phylogenetic studies was critically reviewed. Nearctic genera exhibit a much higher variation in the shape and form of scolecites compared with taxa in other biogeographical regions. The following basic scolex types can be recognised in Nearctic caryophyllideans: monobothriate (Promonobothrium Mackiewicz, 1968), loculotruncate (Promonobothrium, Dieffluvium Williams, 1978), bothrioloculodiscate (Archigetes Leuckart, 1878, Janiszewska et al., 2017), monobothrium (Caryophyllaeides Mackiewicz et Deutsch, 1976, Penarchigetes Mackiewicz, 1969, Pseudoglaridacris Oros, Uhrovčić et al., 2018), fixomegabothriate (Capingens Hunter, 1927), bulbate and bulbocacuminate (Atractolytocestus Anthony, 1958), cuneiform (Hypocaryophyllaeus Hunter, 1927, Rowardleus Mackiewicz et Deutsch, 1976, Spartoidea Hunter, 1929), biacetabulate, bulboloculate, bothrioloculodiscate (Biacetabulum Hunter, 1927), hololete (Hunterekia Mackiewicz et McCrae, 1962), cuneiform (Khavia Hsi, 1935), cuneiform (Calentinella Mackiewicz, 1974, Caryophyllaeides Nybelin, 1922, Edlintonia Mackiewicz, 1970), hastate (Pseudolytocestus Hunter, 1929), loculohololete (Bialovarium Fischthal, 1953, Pliovitellaria Fischthal, 1951), and cuneiformo-loculoculate (Glariacris Cooper, 1920, Isoglaridacris Mackiewicz, 1965). The same type of scolex may be shared by species of different genera or families and species of the same genus can have a scolex of conspicuously different morphology, e.g. in Promonobothrium. Scolex morphology may be therefore of limited use in generic designation.

Key words: Cestoda, scolex terminology, fish, Catostomidae, comparative morphology, scanning electron microscopy, identification, North America

The scolex of tapeworms plays a key role in their attachment in the intestinal lumen of the definitive host and is the site of the most intimate interface between the parasite and its host (Smyth and McManus 1988). Scolex morphology is thus a result of long-term adaptation of the parasite to its host. Because of conspicuous differences between tape worms from different groups of vertebrate hosts, scolex morphology has been used as one of the key characters for a higher-level (ordinal) classification of tape worms (Schmidt 1986, Khalil et al. 1994) and it is also widely used for taxonomy at the generic level (Schmidt 1986, Caira and Jensen 2017).

In caryophyllidean tapeworms, which represent one of the dominant groups of intestinal parasites of cypriniform and siluriform fishes (Scholz and Kuchta 2017), scolex morphology has been used to characterise individual genera (Mackiewicz 1994), but also to distinguish congeneric species (Ash et al. 2011a). No single scolex type characterises caryophyllideans (Mackiewicz 1970). Oros et al. (2010) studied scolex morphology of Palaeartic species and found it to represent a useful tool for species identification. However, studies on the species of Caryophyllaeus Gimelin, 1790 in the Palaeartic Region have revealed morphological plasticity of scolex shape in tapeworms from different hosts, casting doubts upon general suitability of scolex morphology in caryophyllidean taxonomy (Baráč et al. 2014, 2017, Hanzelová et al. 2015).

In North America, caryophyllideans are the dominant group of cestodes in catostomid and, to a much lesser extent, cyprinid fishes, being represented by 55 species in 22 genera (Mackiewicz 1972, Hoffman 1999, Scholz and Oros 2017). They represent 56% of all species of adult cestodes...
in freshwater fishes of the Nearctic Region (Scholz and Kuchta 2017). Except for three genera, namely Biacetabulum Hunter, 1927, Isoglaridacris Mackiewicz, 1965 (both genera with 10 spp. each) and Promonobothrium Mackiewicz, 1968 (9 spp.), Nearctic genera are species-poor and 12 genera are monotypic (Scholz and Oros 2017).

Most of these genera are distinguished from each other by characteristics of the reproductive system (see the keys in Mackiewicz 1994), but scolex morphology was proposed as a useful tool for classification and identification of North American species by numerous authors (e.g., Hunter 1930, McCrae 1962, Mackiewicz 1970, 1972, 1994, Mackiewicz and McCrae 1965, Williams 1977). Hayunga and Mackiewicz (1988) studied histology of the scolex and neck region of Glaridacris laruei (Lamont, 1921) (= Pseudoglaridacris laruei) and Glaridacris catostomi Cooper, 1920, and Hayunga (1979) studied the intestinal pathology caused by three species of North American caryophyllideans to their host, the white sucker, Catostomus commersonii (Lacépède). However, a detailed comparative study on the scolex morphology of Nearctic caryophyllideans based on both light and scanning electron microscopy had not been carried out.

As part of a collaborative effort to better characterise the global diversity of cestodes (Caira and Jensen 2017, Scholz and Kuchta 2017), scolex morphology of caryophyllideans parasitising North American freshwater fishes was critically reviewed in this study using light and scanning electron microscopy. Whenever possible, material newly collected by the present authors was processed using the same method (heat-fixation) to ensure comparability of morphological and biometrical data.

MATERIALS AND METHODS

Newly collected specimens of 31 species (see below for the list of species) were processed for light (LM) and scanning electron microscopical (SEM) observations. Methods of processing were the same as previously reported in detail (Oros et al. 2010, 2016, Scholz et al. 2015). Briefly, worms were removed from the intestine of freshly killed fish host, rinsed in saline, fixed with hot (almost boiling) 4% neutral buffered formaldehyde solution (= 10% buffered formalin). For light microscopical examination, specimens were stained with carmine, dehydrated in a graded ethanol series, cleared in clove oil, and mounted in Canada balsam. Drawings were made using a drawing attachment on a Leica DM 5000B (Leica Microsystems, Wetzlar, Germany) light microscope. For histological study specimens were embedded in paraffin wax, sectioned at 12–15 µm thickness, and stained with Weigert’s haematoxylin.

For SEM studies, formalin-fixed specimens were dehydrated through a graded ethanol series, followed by a graded amylacetate series, dried by HMDS (hexamethyldisilazane), sputter-coated with 20–25 nm of gold and examined with JEOL JSEM 6510LV microscope (JEOL Ltd., Akishima, Tokyo, Japan). Scientific and common names of fish hosts follow FishBase (Froese and Pauly 2019). Terminology of microtriches follows Chervy (2009), that of scolex types Mackiewicz (1994; see his figs. 5.1–5.21).

Despite the considerable efforts of the present authors, it was not possible to collect representatives of all Nearctic genera. Therefore, material from museum collections was also studied. Voucher specimens of newly collected material and museum material are deposited in the Helminthological Collection of the Institute of Parasitology, Biology Centre of the Czech Academy of Sciences, České Budějovice, Czech Republic (IPCAS) and in the Smithsonian National Museum of Natural History, Washington, DC, USA (USNM; now hosting the previous U.S. National Parasite Collection formerly in Beltsville, Maryland – USNPC).

A list of specimens studied follows below. Species for which material was not fixed with hot fixative are marked with an asterisk (*).
Family Capingentidae Hunter, 1930
1. Capingens singularis Hunter, 1927 from notchlip redhorse, Moxostoma collapsum (Cope) (host code US 274a/PBI [Planetary Biodiversity Inventory code of molecular voucher] 470), Congaree River at Columbia, South Carolina, USA.
2. Edlintonia ptychocheila Mackiewicz, 1970 from northern pikeminnow, Ptychocheilus oregonensis (Richardson), Bonner Co., Idaho, USA (IPCAS C-553/1); from peamouth, Mylocheilus caurinus (Richardson), Middle Fork Willamette River, Oregon, USA.
3. "Pseudolytocestus differtus" Hunter, 1929 from smallmouth buffalo, Ictiobus bubalus (Rafinesque), Tallahatchie River, Mississippi, USA (holotype; USNM 1350001).
4. "Spartoides wardi" Hunter, 1929 from river carpsucker, Carpiodes carpio (Rafinesque), North Platte River, Nebraska, USA (IPCAS C-545/1).

Family Caryophyllaeidae Leuckart, 1878
5. "Archigetes iowensis" Calentine, 1962 from red worm, Limnodrilus hoffmeisteri Claparède USA (IPCAS C-588).
6. Biacetabulum carpodi Mackiewicz, 1969 from black buffalo, Ictiobus niger (Rafinesque) (US 244b/PBI 417), Chotard Lake, Vicksburg, Mississippi, USA.
7. Biacetabulum infrequens Hunter, 1927 from spotted sucker, Minntrena melanops (Rafinesque) (US 217b/PBI 415), Fish Lake near Pascagoula River at Benndale, Mississippi, USA.
8. Calentinella etnieri Mackiewicz, 1974 from western creek chubsucker, Erimyzon claviformis (Girard) (US 668), Sugar Creek, Cross Co., Arkansas, USA (IPCAS C-550/2).
9. "Dieffluvium unipapillatum" Williams, 1978 from river redhorse, Moxostoma carinatum (Cope), Cahaba River, Dallas Co., Alabama, USA (holotype; USNM 1369056).
10. Glaridacris catostomi Cooper, 1920 from white sucker, Catostomus commersonii, Assiniboine River, Manitoba, Canada (IPCAS C-5/1).
11. Hunterella nodulosa Mackiewicz et McCrae, 1962 from northern hog sucker, Hypentelium nigricans (Le Sueur), Waupaca River, Wisconsin, USA (IPCAS C-3); from Catostomus commersonii (DP 134), Duck Creek, Ashwaubenon and De Pere, Wisconsin, USA.

Fig. 2. Line drawings of the anterior parts of the Nearctic species of the Caryophyllidea with the anteriormost vitelline follicles (black) and testes (stippled). A – Calentinella etnieri Mackiewicz, 1974 ex Erimyzon claviformis (Girard); B – Caryophyllaeides fennica (Schneider, 1902) ex Acrocheilus alataucus Agassiz et Pickering; C – Edlintonia ptychocheila Mackiewicz, 1970 ex Ptychocheilus oregonensis (Richardson); D – Hunterella nodulosa Mackiewicz et McCrae, 1962 ex Hypentelium nigricans (Le Sueur); E – Promonobothrium hunteri (Mackiewicz, 1963) ex Hypentelium nigricans; F – Pseudolytocestus differtus Hunter, 1929 ex Ictiobus bubalus (Rafinesque); G – Dieffluvium unipapillatum Williams, 1978 ex Moxostoma carinatum (Cope); H – Pliovitellaria wisconsinensis Fischthal, 1951 ex Notemigonus crysoleucas (Mitchell).
12. *Hypocaryophyllaeus gilae* Fischthal, 1953 from Utah chub, *Gila atraria* (Girard), Emma Matilda River, Teton Co., Wyoming, USA (USNM 1347575).

13. *Isoclaridacris wisconsinensis* Williams, 1977 from shorthead redhorse, *Moxostoma macrocephalum* (Le Sueur) (US 282a), Congaree River at Columbia, South Carolina, USA (PCAS C-652/1); from *H. nigricans* (DP 148/09b/PBI 413), Waupaca River, Wisconsin, USA.

14. *Janiszewskella fortobothria* Mackiewicz et Deutsch, 1976 from *Carpiodes cyprinus* (Le Sueur), Susquehanna River, Pennsylvania, USA (IPCAS C-529/1).

15. *Penarchigetes oklensis* Mackiewicz, 1969 from *Ictiobus bubalus*, North Eastern Outing Club Lake, Cherokee Co., Oklahoma, USA (IPCAS C-555/1) and from *M. melanops*, Oxbow Lake of Illinois River, Cherokee Co., Oklahoma, USA (holotype; USNM 1366840).

16. *Pliovitellaria wisconsinensis* Fischthal, 1951 from golden shiner, *Notemigonus crysoleucas* (Mitchill), Westhampton, Massachusetts, USA (IPCAS C-533/1); from northern redbelly dace, *Chrosomus eos* Cope, Bolger Bog, Upper Peninsula of Michigan, Michigan, USA.

17. *Promonobothrium currani* Oros, Brabec, Kuchta, Choudhury et Scholz, 2016 from *I. niger* (US 244c/PBI 460), Chotard Lake, Mississippi, USA (IPCAS C-696/2).

18. *Promonobothrium hunteri* (Mackiewicz, 1963) from *H. nigricans*, Grindstone Creek, New York, USA (IPCAS C-505/3).

19. *Promonobothrium rogersi* (Williams, 1980) from highfin carp sucker, *Carpiodes velifer* (Rafinesque) (US 216a/PBI 472), Pascagoula River, Mississippi, USA (IPCAS C-698).

20. *Pseudoglaridacris confusa* (Hunter, 1929) from *Ictiobus cyprinellus* (Valenciennes) (US 246a/PBI 463), Chotard Lake, Mississippi, USA (IPCAS C-528).

21. *Pseudoglaridacris laruei* (Lamont, 1921) from *Moxostoma anisurum* (Rafinesque) (DP 93/09a, 111/09a), Wolf River, Wisconsin, USA (IPCAS C-322/3); from *Catostomus commersonii* (US 66a), Lake Winnebago, Wisconsin, USA (IPCAS C-322/2).

22. *Rowardleus pennensis* Mackiewicz et Deutsch, 1976 from *C. cyprinus* (PBI 405), Lake Winnebago, Wisconsin, USA (IPCAS C-545).

**Family Lytocestidae Hunter, 1927**

23. *Atractolytocestus huronensis* Anthony, 1958 from common carp, *Cyprinus carpio* Linnaeus, Reelfoot Lake, Tennessee, USA (IPCAS C-472).

24. *Caryophyllaeides femica* (Schneider, 1902) from chiselmouth, *Acrocheilus altanavus* Agassiz et Pickering, Trout Creek, Oregon, USA (IPCAS C-2/20).

25. *Khawia japonensis* (Yamaguti, 1934) (syn. *K. iowensis* Calentine et Ulmer, 1961) from *C. carpio*, Mohawk River, Albany, New York, USA and Lake Texoma, Oklahoma, USA (IPCAS C-528).

No specimens of species of *Bialovarium* Fischthal, 1953 and *Homeomorpha* Dutton et Barger, 2014 were available. For characterisation and line drawings of their scoleces, the original descriptions were used. In addition, the two invasive species *A. huronensis* and *K. japonensis*, previously reported in detail (see Oros et al. 2010, 2015, Scholz et al. 2011), were also included in the present study.

**RESULTS**

Drawings of the scoleces (and the anterior extent of the testes and vitelline follicles in some of them) of 23 representative species of 20 genera are provided in Figs. 1–4. Scanning electron micrographs (SEM) of the scoleces of 14 species of 11 genera of caryophyllidean cestodes of all three families from North American catostomid and cyprinid fishes are presented in Figs. 5, 6. The following types of the scoleces can be recognised:

1. **Afossate**
   1.1. bulboacuminate (*Atractolytocestus* – Fig. 1A);
Fig. 4. Line drawings of the anterior parts of the Nearctic species of the Caryophyllidea with the anteriormost vitelline follicles (black) and testes (stippled). A – Glaridacris catosomi Cooper, 1920 ex Catostomus commersonii (Lacépède); B – Isoglaridacris wisconsinensis Williams, 1977 ex Moxostoma macrolepidotum (Le Sueur); C – Rowardleus pennisensis Mackiewicz et Deutsch, 1976 ex Carpiodes cyprinus (Le Sueur); D – Spartooides wardi Hunter, 1929 ex Carpiodes carpio (Rafinesque); E – Pseudoglaridacris laruei (Lamont, 1921) ex Catostomus commersonii; F – Pseudoglaridacris confusa (Hunter, 1929) ex Ictiobus cyprinellus (Valenciennes); G – Pronomonobothrium carrani Oros, Brabec, Kuchta, Choudhury et Scholz, 2016 ex Ictiobus niger (Rafinesque); H – Pronomonobothrium rogersi (Williams, 1980) ex Carpiodes velifer (Rafinesque); I – Hypocaryophyllaeus gilae Fischthal, 1953 ex Gila atraria (Girard); J – Janiszewskella foriobothria Mackiewicz et Deutsch, 1976 ex Carpiodes cyprinus; K – Penarchigetes oklensis Mackiewicz, 1969 ex Minytrema melanops (Rafinesque); L – Capingens singularis Hunter, 1927 ex Moxostoma collapsum (Cope).

1.2. cuneiformbrite (Khawia Hsü, 1935 – Fig. 1D); 1.3. cuneiform (Calentinella Mackiewicz, 1974, Caryophyllaelides Nybelin, 1922, Edlintonia Mackiewicz, 1970 – Figs. 2A–C, 5A,C); 1.4. hastate (Pseudolytocestus Hunter, 1929 – Fig. 2F); 1.5. tholate (Hunterella Mackiewicz et McCrae, 1962 – Figs. 2D, 5B).
2. Fossate

2.1. biacetabulate (Biacetabulum – Figs. 3C, 6D);

2.2. bothrioloculodiscate (Archigetes Leuckart, 1878, Biacetabulum, Janiszewskella Mackiewicz et Deutsch, 1976, Penarchigetes Mackiewicz, 1969, Pseudoglaridacris Oros, Uhrovč et Scholz, 2018 – Figs. 3A, 4E,F,J,K, 6F,G);

2.3. bulboloculate (Biacetabulum, Janiszewskella – Figs. 3B, 4J, 6H);

2.4. cuneiformoloculate (Glaridacris Cooper, 1920 – Figs. 4A, 6C);

2.5. cuneiloculate (Homeomorpha, Hypocaryophyllaeus Hunter, 1927, Isoglaridacris, Rowardeus Mackiewicz et Deutsch, 1976, Spartoides Hunter, 1929 – Figs. 1C, 4B–D,I, 5E, 6B);

2.6. digitiform-loculate (Promonobothrium – Figs. 2E, 6A);

2.7. fixomegabothriate (Cepingens Hunter, 1927– Figs. 4L, 5F);

2.8. loculotholate (Bialovarium, Pliovitellaria Fischthal, 1951 – Figs. 1B, 2H, 5D);

2.9. loculotruncate (Diefflavium – Fig. 2G, Promonobothrium – Fig. 4H);

2.10. loculomonobothriate (Promonobothrium – see fig. 1B in Oros et al. 2016)

2.11. monobothriate (Promonobothrium – Fig. 2E, 6A)

Characterisation of the scoleces of caryophyllidean genera from the Nearctic Region

Family Capingentidae Hunter, 1930 (4 genera)

Capingens Hunter, 1927  Figs. 4L, 5F
Scolex fixomegabothriate – massive and robust, about 1/5 to 1/4 of total body length, with large, well-developed ventral and dorsal bothria; first testes begin at base of scolex.

Edlintonia Mackiewicz, 1970  Figs. 2C, 5C
Scolex cuneiform – simple, aculate, chisel-shaped, sagittate in lateral view; neck absent; first vitelline follicles begin posterior to first testes.

Pseudolytocestus Hunter, 1929  Fig. 2F
Scolex weakly hastate, aculate – slightly sagittate, partially narrowed in neck; first testes begin posterior to first vitelline follicles.

Spartoides Hunter, 1929  Figs. 4D, 5E
Scolex cuneiloculate – distinct, with three pairs of shallow loculi, sagittate in lateral view; neck long, narrower than scolex; first testes begin posterior to first vitelline follicles.
Family Caryophyllaeidae Leuckart, 1878 (16 genera)

Archigetes Leuckart, 1878
Scolex bothrioloculodicate – spherical with two pairs of deep loculi and a pair of deep central depressions, terminal disc present; neck absent; first testes begin at base of scolex, first vitelline follicles begin posterior to first testes.

Biacetabulum Hunter, 1927
Scolex bulboloculate, bothrioloculodicate, biacetabulate – relatively large scolex slightly wider than body, with or without one pair of deep central acetabulum-like structure and two pairs of less pronounced lateral loculi or just one pair of weak loculi present on scolex; variability of shape and type of scoleces very broad.

Bialovarium Fischthal, 1953
Scolex loculotholate – poorly defined, blunt, bearing a pair of shallow loculi, dorsoventrally compressed; neck absent; first testes begin posterior to first vitelline follicles.

Calentinella Mackiewicz, 1974
Scolex cuneiform – blunt-ended, slightly dorsoventrally compressed, wider than neck; neck present, long; first testes begin posterior to first vitelline follicles.

Dieffluvium Williams, 1978
Scolex loculotruncate – expanded posteriorly within the pair of loculi in expanded portion; neck present, long; first testes begin posterior to first vitelline follicles.

Glaridacris Cooper, 1920
Scolex cuneiformoloculate – wedge-shaped, with three shallow loculi on dorsal and ventral side of scolex; neck present, short; first testes begin posterior to first vitelline follicles.

Homeomorpha Dutton et Barger, 2014
Scolex cuneiloculate – shallowly ovoid, with three pairs of shallow loculi; neck present, long; first testes begin posterior to first vitelline follicles.

Hunterella Mackiewicz et McCrae, 1962
Scolex tholate – simple, non-specialised with round to conical shape; neck absent; first testes begin immediately posterior to scolex, vitelline follicles begin posterior to first testes.

Hypocaryophyllaeus Hunter, 1927
Scolex cuneiloculate – indistinctly defined from neck, bearing a flattened terminal disc and three pairs of distinct loculi; neck absent; first vitelline follicles begin anterior to first testes.
Isoglaridacris Mackiewicz, 1965  Figs. 4B, 6B
Scolex cunei locate – fan-shaped, with three pairs of shallow loculi; neck present; first testes begin posterior to first vitelline follicles.

Janiszewska Mackiewicz et Deutsch, 1976  Fig. 4J
Scolex bothrio loculodiscate – slightly spherical with two pairs of deep loculi and pair of deep central bothria; neck short; first testes begin posterior to first vitelline follicles.

Penarchigetes Mackiewicz, 1969  Fig. 4K
Scolex bothrio loculodiscate – slightly cylindrical or conical, wider than the body, with one pair of median loculus and two pairs of lateral loculi, apical disc small; neck indistinct or distinct; vitelline follicles begin slightly anterior or posterior to first testes, to posterior end of scolex or to neck region only.

Pliovitellaria Fischthal, 1951  Figs. 2H, 5D
Scolex loculotholate – rounded, dorsoventrally compressed, with one central shallow depression, wider than the body; neck absent; first testes begin posterior to first vitelline follicles.

Promonobothrium Mackiewicz, 1968  Figs. 4G,H, 6E
Scolex digitiform, digitiform-loculate or loculotruncate – with a terminal introvert on apex and one or three pair(s) of shallow dorsoventral depressions.; neck short to moderately long; variability of shape of scoleces and arrangement of vitelline follicles and testes very broad.

Pseudoglaridacris Oros, Uhrovič et Scholz, 2018  Figs. 4E,F, 6F,G
Scolex bothrio loculodiscate – forming definite terminal (apical) disc and three pairs of loculi; relatively short neck; variability of anterior position of vitelline follicles and testes broad.

Rowardleus Mackiewicz et Deutsch, 1976  Fig. 4C
Scolex cunei locate – round to ellipsoidal, with three pairs of featureless loculi in one pair shallow depression; neck region long; first testes begin posterior to first vitelline follicles.

Family Lytocestidae Hunter, 1927  (3 genera)

Atractolytocestus Anthony, 1958  Fig. 1A
Scolex bulbocaminate – arrowhead-shaped (tapered) or bluntly ended, with smooth anterior margin, scolex wider than distinct neck; first testes begin posterior to vitelline follicles.

Caryophyllaeides Nybelin, 1922  Figs. 2B, 5A
Scolex cuneiform – blunt-ended, slightly dorsoventral compressed, aloculate; neck absent; first testes begin posterior to first vitelline follicles.

Khavia Hsü, 1935  Fig. 1D
Scolex cuneiformiate – lacking apical structures and lateral loculi, with frilled or deeply folded anterior margin, scolex wider than neck region; first testes and first vitelline follicles start at approximately same level, or first testes begin posterior to first vitelline follicles.

DISCUSSION

The present study has confirmed high diversity of scolex types in the Nearctic species of caryophyllidean cestodes. In contrast, species from other zoogeographical regions are rather uniform in scolex morphology (see Oros et al. 2010 for Palaearctic species, Ash et al. 2011a,b for Indomalayan taxa). As many as 16 types of scoleces can be recognised among Nearctic taxa, but intrageneric variation may differ considerably between individual genera. Species of most genera have rather uniform shape (31 species studied), but the genus Promonobothrium as recently re-defined by Oros et al. (2016) includes taxa with conspicuously different scolex types, i.e. monobothriate, loculotruncate, biacetabulate and digitiform-loculate (Figs. 2E, 4G,H, 6E). As new data from further collections and molecular analysis become available, changes will be necessary for terminology of scolex types.

The scolex of caryophyllidean tapeworms is highly mobile and variable, with no single type being characteristic of this group (Mackiewicz 1970, 1972, 1994). Mackiewicz (2003) distinguished two basic scolex types among caryophyllideans of the world and indicated that 24 genera (59%) of all caryophyllideans recognised at that time possess afoossate scoleces and 17 genera (41%) possess fossate scoleces. As obvious from the present data, the number of afoossate genera among only Nearctic caryophyllideans is much lower (only 7 genera out of 23, i.e. 30%).

The scolex has been suggested as a useful structure for generic and/or species identification (Kulakovskaya 1961, Chubb et al. 1987, Dubinina 1987, Scholz 1989, Protasova 1990, Oros et al. 2010), but the use of scolex morphology can be limited in some taxa. A detailed study of two species of the Palaearctic genus Caryophyllaeus, C. brachycollis Janiszewska, 1953 and C. laticeps (Pallas, 1781), has revealed high plasticity in their scolex morphology (Barčák et al. 2014, Hanzelová et al. 2015). This inaspecific, possibly host-related morphological variability represents a serious obstacle for reliable identification of species of this genus (Barčák et al. 2017).

The present study of Nearctic species has shown that form and shape of the scolex may help in differentiating most Nearctic genera, especially those with a low number of taxa that have a unique form of scolex (e.g., Capingens, Hunterella, Pliovitellaria and Pseudolytocestus). However, it should be mentioned that different methods of fixation may greatly influence scolex morphology (Oros et al. 2010). Live material should also be studied to observe the natural extension and variation of the scolex during normal contractions (Mackiewicz 1972). For example, species of Pseudoglaridacris are characterised by a bothrio loculodiscate scolex, i.e. the scolex forming a definite terminal (apical) disc and three pairs of loculi. However, the normal flat terminal disc may naturally push forward into a dome configuration and the rest of the scolex may have knobs formed by the expanded muscles that form the disc (Oros...
et al. 2018). This abnormal extent can be readily seen, for example, in species of *Pseudoglaridacris* (Fig. 4E,F) and *Biacetabulum* (Fig. 3B,C) of the present study.

Examination of the holotype of *Diefflevium unipapillatum* (USNM 1369056) and two other gravid individuals determined to be that species indicates that the scolex of the holotype has been compressed and misshapen. There is no papilla-like structure as described and incorrectly illustrated by Williams (1978). Based on non-compression, we interpret the scolex to be most similar to the loculotruncate type. As a result the previous designation of loculopapillate is considered inappropriate and inaccurate and has been based on an abnormal, compressed scolex. As a consequence of this mistake in terminology, the figures of Williams (1978: figs 1 and 5) as loculopapillate and subsequently utilised by Mackiewicz (1994) and Hoffman (1999) should no longer be used. A similar observation has been made independently by Herzog and Barger (2019).

A robust phylogenetic hypothesis of relationships of caryophyllidean cestodes is not yet available (see Scholz and Oros 2017 for preliminary data on interrelationships of these cestodes). Brabec et al. (2012) included in their phylogenetic analyses only three Nearctic taxa, namely *Glaridacris catostomi*, *Hunterella nodulosa* and *Promonobothrium hunteri* (as *Monobothrium hunteri* Mackiewicz, 1963), all of the family Caryophyllaeidae. Scholz et al. (2015) and Oros et al. (2016) enlarged the number of species of *Promonobothrium*, but no representatives of other Nearctic genera were analysed. Therefore, it is not possible to reliably assess the homology of individual types of scoleces of Nearctic caryophyllideans, which possess structures absent in species from other zoogeographical regions (Mackiewicz 1972, Oros et al. 2010). Preliminary molecular phylogenetic data generated by A. Waeschenbach (see Scholz and Oros 2017) indicate that Nearctic taxa form a monophyletic, more recently diverging group distant from the remaining caryophyllideans from other zoogeographical regions.

The presence of the same scolex type in genera of different families (cuneiform in the capingentid *Edlintonia*, caryophyllaeid *Calentinella* and lytocestid *Caryophyllaeides*, and cuneilocate in the capingentid *Spartoides* and caryophyllaeids *Hypocaryophyllaeus* and *Rowardleus*) indicates that scolex morphology does not necessarily reflect the evolutionary history and relatedness of these taxa. Indeed, the current system of families is undoubtedly artificial as evidenced by several studies (Oros et al. 2008, 2016, Brabec et al. 2012, Scholz and Oros 2017). Similarly as in some other cestode groups (e.g., Scholz et al. 2013), the scolex morphology can be a result of adaptation to a particular host physiology, intestinal region or intestinal surface architecture rather than it witnesses co-evolutionary history of taxa with a similar type of scolex. Interestingly, North American caryophyllideans from cyprinids, i.e. species of *Atractolytocestus*, *Caryophyllaeides*, *Edlintonia*, *Khawia* and *Pliovitellaria*, have simpler scoleces compared to those of caryophyllideans from catostomids.

Scoleces of some species of *Biacetabulum* invariably possess a large, deep central structure on the dorsal and ventral sides with shallower, lateral loculi, but the shape and depth of loculi and central structure may differ between species and individual specimens (Figs. 3B,C, 5D,H). According to Hunter (1930), McCrae (1962), Mackiewicz (1972, 1994), Williams (1977), Mackiewicz et al. (1972), the central structure represents a pair of true acetabula that function like suckers, i.e. acetabulate attachment organs (small suckers) of more recently diverged ‘tetraphyllid-like’ cestodes of Olson et al. (2001) and Cairns and Jensen (2017) (see Mackiewicz et al. 1972: fig. 18 for photo of *in situ* attachment of the ‘acetabulum’). However, the anatomy of this pair of large structures, which appears to be well demarcated with a basement membrane only in its posterior part, does not seem to be homologous with acetabulate structures (four small suckers) of oncoproteinophylideans, cyclophyllideans and other tapeworms (see Olson et al. 2001, Cairns et al. 2014, Cairns and Jensen 2017). Most likely, the ‘acetabulum’ in species of *Biacetabulum* and *Promonobothrium* represents an example of convergent evolution of similar but larger pair of sucker-like attachment (acetabulate) structures in cestodes.

In addition to documenting extraordinary shape diversity of the scoleces of North American taxa using light and scanning electron microscopy, the present study has also revealed that the diversity of this cestode group is not fully known. Some of the specimens studied may represent new, undescribed taxa whose formal descriptions are pending. For a more conclusive study, examination of missing taxa, in particular those from endemic catostomids and cyprinids in the western part of North America, is also highly needed to generate a robust phylogenetic hypothesis of the group. Such new data would enable us to test homology of scolex forms in North American caryophyllideans and their importance for classification of these cestodes.

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