New sharks and other chondrichthians from the latest Maastrichtian (Late Cretaceous) of North America

Terry A. Gates, 1,2 Eric Gorscak, 2 and Peter J. Makovicky 2

1Dept of Biological Sciences, North Carolina State University, Raleigh, NC, 27695, USA <tagates@ncsu.edu>
2Dept of Geology, Field Museum of Natural History, Chicago, IL, 60605, USA <egorscak@fieldmuseum.org>, <pmakovicky@fieldmuseum.org>

Abstract.—Cretaceous aquatic ecosystems were amazingly diverse, containing most clades of extant aquatic vertebrates as well as an array of sharks and rays not present today. Here we report on the chondrichthyan fauna from the latest Maastrichtian site that yielded the Tyrannosaurus rex skeleton FMNH PF 2081 (“SUE”). Significant among the recovered fauna is an unidentified species of carcharhinid shark that adds to the fossil record of this family in the Cretaceous, aligning with estimates from molecular evidence of clade origins. Additionally, a new orectolobiform shark, here named Galagadon nordquistae n. gen. n. sp., is diagnosed on the basis on several autapomorphies from over two-dozen teeth. Common chondrichthyan species found at the “SUE” locality include Lonchidion selachos and Myledaphus pustulosus. Two phylogenetic analyses (Maximum Parsimony and Bayesian Inference) based on twelve original dental character traits combined with 136 morphological traits from a prior study of 28 fossil and extant taxa, posited Galagadon n. gen. in two distinct positions: as part of a clade inclusive of the fossil species Cretorectolobus olsoni and Cederstroemia triangulata plus extant orectolobids from the Maximum Parsimony analysis; and as the sister taxon to all extant hemiscyllids from the Bayesian Inference. Model-based biogeographical reconstructions based on both optimal trees suggest rapid island hopping-style dispersal from the Western Pacific to the Western Interior Seaway of North America where Galagadon n. gen. lived. Alternatively, the next preferred model posits a broader, near-global distribution of Orectolobiformes with Galagadon n. gen. dispersing into its geographic position from this large ancestral range.

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Introduction

Chondrichthyan species are rare in modern freshwater ecosystems (Compagno et al., 2005). However, ancient aquatic systems abounded in sharks and rays, especially those with linkages to marine habitats (e.g., Kirkland et al., 2013). Throughout the latter half of the Cretaceous an intercontinental seaway flooded part of the North American continent, providing ideal conditions for the invasion of freshwater ecosystems by sharks and rays as well as the subsequent preservation of those ecosystems in the fossil record.

The latest Cretaceous Hell Creek Formation is a set of siltstone-/sandstone-dominated rocks named for a section exposed in the Hell Creek valley of Montana (Hartman et al., 2014). In South Dakota, where fossils from the “SUE” locality are located, the age of the Hell Creek Formation lies between 67.4 Ma and 65.5 Ma, according to LeCain et al. (2014). These sediments were deposited at a time when the intercontinental seaway was retreating from North America due in part to uplift of the Laramide orogeny (Lawton, 2008). This created a wide coastal plain where abundant aquatic communities containing crocodylians (Carpenter and Lindsey, 1980), choristoderans (Matsumoto and Evans, 2010), turtles (Holroyd and Hutchison, 2002), amphibians (Demar, 2013), osteichthyan fish (Brinkman et al., 2014), and chondrichthians (Cook et al., 2014) flourished. Estes et al. (1969) were the first to examine the Hell Creek Formation freshwater fauna comprehensively. Subsequent researchers have used similarly rich microvertebrate localities to identify new species from the aquatic ecosystems (e.g., Bryant, 1989; Demar, 2013; Cook et al., 2014) and to elucidate preservational biases among fossil sites (e.g., Wilson, 2008; Peterson et al., 2011).

Here we present the chondrichthyan fauna from the Field Museum of Natural History “SUE” locality (Fig. 1), a highly productive microsite locality that also contained the nearly complete Tyrannosaurus rex (Osborn, 1905) specimen “SUE” (FMNH PF 2081) and a diverse microvertebrate fauna that included small theropod dinosaurs (Gates et al., 2013), actinopterygian and sarcopterygian fish, lissamphibians, and reptiles (Lyson and Joyce, 2009; Gates et al., 2010b), all collected either during the excavation of the dinosaur specimen or subsequently during screen washing of the associated matrix in this study. The taxa identified in our fauna contain several observed in other localities as well as two new species. The results of this study illuminate the potential for new fossil taxon discoveries in the Hell Creek Formation and promote the utility of using microsite data from macrofossil assemblages.
Materials and methods

Geologic setting.—The “locality is situated in the lower portion of the Hell Creek Formation, ∼4.8 m above the contact with the Fox Hills Formation (Larson, 2008). It contains distinct layers of siltstone, fine-grained sandstone, and mudstone, with interbedded plant fossils. This association was possibly produced on a point bar deposit within a meandering river because no geologic evidence exists of a channel (e.g., rip-up clasts, coarser-grained sediment from a thalweg, or fall bocks from a cut-bank).

Matrix preparation.—Approximately two tons of matrix ranging in grain size from sandy siltstone to mudstone were obtained from fossil jackets containing the bones of Tyrannosaurus rex, FMNH PF 2081. No additional sediment was sampled from the site. Precise locality information can be obtained from the Field Museum of Natural History. The site likely represents a point bar deposit within a meandering river channel, as evinced by alternating layers of mud-rich sediment containing leaf fossils and sandy siltstone.

Sediment was washed in nested screens of sieve openings 4.6 mm (mesh #4), 0.841 mm (mesh #20), and 0.595 mm (mesh #30) utilizing an overhead sprinkler system separated from the matrix by a 0.841 mm (mesh #20) screen in order to decrease damage to the fossils. The fossil concentrate was manually picked under a stereo-microscope. All orectolobiform teeth were found within the 0.595 mm size fraction, whereas the carcharhiniform tooth was found in the 0.841 mm mesh concentrate. The teeth were photographed on a Keyence VHX-5000 microscope. Scanning electron microscope imaging of FMNH PF 15982 was performed on a Carl Zeiss NTS EVO60 XVP scanning electron microscope in both high and variable pressure vacuum modes, using VPSE detectors without extraneous coating. All fossils are accessioned at the Field Museum of Natural History.

Phylogenetic analyses.—In order to better determine the phylogenetic affinities of Galagadon n. gen. and other possible Cretaceous orectolobids, we identified a dozen traits related to dental anatomy (Table 1), following the pioneering work of Adnet and Cappetta (2001) and Shimada (2005) demonstrating that dental traits can provide phylogenetically useful information. We combined these twelve traits with a larger morphological analysis for orectolobiformes by Goto (2001) that includes 136 morphological traits arrayed across the body (Step II in his paper). Of those original traits, however, only two are related to teeth (characters 36 and 37), and of these, only character 37 involves the morphology of the teeth (apron breadth). The modified dataset consists of 148 morphological characters (ordered as in the original publication) and 28 ingroup taxa (Table 2), including several fossil forms.

Coding for the ingroup extant species was performed using Herman et al. (1992) and Compagno et al. (2005). Goto (2001) used a composite outgroup, which was coded originally as an amalgamation of Lamniformes and Carcharhiniformes. Our coding of tooth characters was based on the lamniform Otodus obliquus (Agassiz, 1843) and the carcharhiniform Carcharhinus leucas (Müller and Henle, 1839), which were combined into a composite scoring for the outgroup taxon following Goto’s (2001) protocol, resulting in numerous polymorphic characters.

Parsimony analysis.—We conducted several parsimony-based phylogenetic analyses (one of which was backed by molecular data), including: 1) a Maximum Parsimony (MP) analysis on the full data with characters ordered following Goto (2001); 2) an MP analysis in which the three Cretaceous fossil taxa Cretorectolobus olsoni, Cederstroemia triangulata, and Galagadon nordquistae n. gen. n. sp. are constrained as a monophyletic cluster endemic to the Western Interior Seaway (WIS); and 3) an MP analysis in which extant orectolobid species were arranged according to a backbone constraint following analysis of molecular data by Corrigan and Beherenegray (2009), but positions of fossils were left unconstrained. Importantly, this
Table 1. Phylogenetic characters created in this study. This list is amended to Goto (2001) and found in the complete NEXUS file included in the Supplementary Information. Numbering of characters follows that of the character matrix.

Crown

137) Labial surface of crown ornamented with folds and plications: present (0); absent (1).
138) Cusplets: present (0); absent (1).
139) Cusplets on crown: absent (0); one cusplet present on at least a single side of the tooth, but no more than one seen on both sides (1); multiple cusplets present on at least one side of crown (2).
140) Heels: (0) present; absent (1).
141) Serrated cutting edge present on central cusp or shoulders: present (0); absent (1).
142) Apron: absent (0); present (1).
143) Apron, when viewed labially, the depth to width ratio: less than or equal to 0.5 (0); greater than 0.5 (1); absent (2).
144) Labial face generally: flat (0); convex (1).

Root

145) Root lobes, when viewed basally the lateral edges of root lobe: straight (0); has a slight curvature (1); has a dramatic curvature making the labial half bulge into distinct lobes (2).
146) Root lobes, when viewed basally root lobes: are labiolingually restricted, forming narrow columns or struts that are oriented basally (0); form a single strut that runs medio-distally across the tooth (1); labiolingually expanded struts oriented diagonally from the labial midline to a lingual lateral position (2); labiolingually expanded struts oriented nearly labio-lingually (3).
147) Root development: hemiholaulcorhize (0); anaulacorhize (1); holaulacorhize (2).
148) Labiolingual fossa along midline of root open in at least some specimens: present (0); absent (1).

Table 2. List of the 28 ingroup taxa included in the phylogenetic analysis.

<table>
<thead>
<tr>
<th>Taxon Name</th>
<th>Author and Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>C. variolatum</td>
<td>Duméril, 1853</td>
</tr>
<tr>
<td>C. collare</td>
<td>Ramsey and Ogilby, 1888</td>
</tr>
<tr>
<td>C. ferrugineum</td>
<td>McCulloch, 1911</td>
</tr>
<tr>
<td>C. sp. (Gill, 1862)</td>
<td></td>
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<tr>
<td>C. japonicum</td>
<td>Regan, 1906</td>
</tr>
<tr>
<td>C. ornatum</td>
<td>De Vis, 1883</td>
</tr>
<tr>
<td>C. maculatus</td>
<td>Bonnaterre, 1788</td>
</tr>
<tr>
<td>C. waddi</td>
<td>Whitley, 1939</td>
</tr>
<tr>
<td>E. dasypogon</td>
<td>Bleeker, 1867</td>
</tr>
<tr>
<td>B. waddi</td>
<td>Bloch and Schneider, 1801</td>
</tr>
<tr>
<td>C. plagiosum</td>
<td>Bennett, 1830</td>
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<tr>
<td>C. hasselli</td>
<td>Bleeker, 1852</td>
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<td>C. punctatum</td>
<td>Müller and Henle, 1838</td>
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<tr>
<td>C. indicum</td>
<td>Gmelin, 1789</td>
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<tr>
<td>H. ocellatum</td>
<td>Bonnaterre, 1788</td>
</tr>
<tr>
<td>H. trifasciata</td>
<td>Richardson, 1843</td>
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<tr>
<td>H. freycineti</td>
<td>Quoy and Gaimard, 1824</td>
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<td>G. crassilobata</td>
<td>Bonnaterre, 1788</td>
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<td>S. fasciata</td>
<td>Hermann, 1783</td>
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<tr>
<td>R. typus</td>
<td>Smith, 1828</td>
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<tr>
<td>S. tentaculata</td>
<td>Peters, 1864</td>
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<tr>
<td>G. nordquisti</td>
<td>Smith and Radcliffe, 1913</td>
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<td>P. vespertinus</td>
<td>Casier, 1946</td>
</tr>
<tr>
<td>O. freeman</td>
<td>Underwood and Ward, 2004</td>
</tr>
<tr>
<td>E. numidica</td>
<td>Arambourg, 1952</td>
</tr>
<tr>
<td>C. triangulata</td>
<td>Siverson, 1995</td>
</tr>
<tr>
<td>C. olsoni</td>
<td>Case, 1978</td>
</tr>
</tbody>
</table>

Biogeographic analysis.—Select results of our phylogenetic analyses were subjected to biogeographic analysis using the BioGeoBEARS package in R (Matzke, 2013), which allows multiple models and great flexibility in user-applied geographic constraints and time slicing protocols. Six biogeographic areas were defined for these analyses: Western Pacific, Indian Ocean, Eastern Pacific, and Atlantic for living taxa, and Tethys/Paratethys (combined as one area for the analyses) and the North American Western Interior Seaway (WIS) for fossil taxa. Distributions for the living species were based on FAO Marine Fishing Areas, as reported by the IUCN Red List (http://www.iucnredlist.org/) and on locality data reported in Fossilworks.org (Uhen et al., 2017) for fossils. The best fitting MCC BI tree and a randomly chosen MPT from the scaffold MP analysis were employed for biogeographic evaluation because the analysis requires a fully resolved bifurcated tree. Two different branch scaling methods were examined for each tree: the first applied an arbitrarily chosen minimum branch length (here 2 MYA) to every zero length branch (Laurin, 2004), the second used the method of dividing a root time.
increment over those branches that have no length due to them being the oldest member of a lineage (Brusatte et al., 2008). Tree scaling was conducted using the STRAP routine (Bell and Lloyd, 2015) in the R environment. Living taxa were given ages either according to molecular estimates produced by Corrigan and Beheregaray (2009), or the default value of 2MYA, whereas fossils were calibrated according to their data provided in Fossilworks.org (Uhen et al., 2017). The Minimum Branch Scaling method provided calibrated trees that were more congruent with the molecular divergence age estimates recovered by Corrigan and Beheregaray (2009), whereas the second method pushed many orectolobid divergences between extant species into the Paleogene or even Cretaceous. We, therefore, will present results of our biogeographic analyses on the best fitting BI tree, equal/variable rates model, and MP scaffold tree scaled using the first method.

Each input tree was analyzed using three different base models: BAYAREALIKE (likelihood interpretation of BayArea; Landis et al., 2013; Matzke, 2013), DEC (Rex and Smith, 2008), and DIVALIKE (likelihood interpretation of DIVA; Ronquist, 1997) and three alternative models that include a parameter governing founder-event speciation, j, for each of the base models (Matzke, 2013). Seven dispersal constraint matrices were constructed to reflect different continental configurations and oceanic connections over the last 255 Ma of Earth history. These matrices and the time slices for which each applies are provided in the Supplemental Data. Results from different model applications were compared using both the log-likelihood and AIC score of each respective model. Analyses of both trees recovered BAYAREALIKE + j as the best-fitting model, as measured by model log likelihood and AIC scores regardless of optimality criterion and branch scaling method. Alternatively, if the recently employed + j parameter is excluded, then the DEC model is the best fitting model for the MP topology and BAYAREALIKE model for the BI topology.

Repositories and institutional abbreviations.—FMNH: Field Museum of Natural History, Chicago, IL, USA; ROM: Royal Ontario Museum, Toronto, ON, CAN; UCMP: University of California Museum of Paleontology, Berkeley, CA, USA.

Systematic paleontology

Class Chondrichthytes Huxley, 1880
Subclass Elasmobranchii Bonaparte, 1838
Superfamily Hybodontoidea Zangerl, 1981
Family Lonchidiidae Herman, 1977
Genus Lonchidion Estes, 1964

Type species.—Lonchidion selachos, UCMP 53897, isolated tooth.

Lonchidion selachos Estes, 1964

Figure 2.1–2.4

Type specimen.—UCMP 52897, isolated tooth.

Occurrence.—Field Museum “SUE” locality; Hell Creek Formation; Faith, South Dakota, USA; Upper Maastrichtian.

Description.—The two Lonchidion specimens recovered from the “SUE” locality (Fig. 2.1–2.4) are identical to those described in other studies (Estes, 1964; Estes et al., 1969; Bryant, 1989; Cook et al., 2014). Their occlusal surfaces are marked by a single moderately sharp ridge that runs the entire mesiodistal length of the central cusp. Other than the lingual protuberance, the central cusps are unornamented.

Materials.—FMNH PF 15756.

Remarks.—When compared to other larger specimens of this species (e.g., Estes, 1964; Estes et al., 1969; Bryant, 1989; Cook et al., 2014), the teeth recovered in this study are identical to other specimens. Therefore, it appears that the teeth of Lonchidion selachos do not undergo ontogenetic shape change.

Class Chondrichthytes Huxley, 1880
Subclass Elasmobranchii Bonaparte, 1838
Superorder Galeomorphii Compagno, 1973
Order Carcharhiniformes Compagno, 1973
Family Carcharhinidae Jordan and Evermann, 1896
Genus gen. indet. sp. indet.

Figure 2.5–2.8

Occurrence.—Field Museum “SUE” locality; Hell Creek Formation; Faith, South Dakota, USA; Upper Maastrichtian.

Description.—Anatomical terminology follows Shimada (2002). A single rounded tooth attributable to the Carcharhinidae (Fig. 2.5, 2.6; FMNH PF 15982) was recovered during this study. General morphology of this specimen suggests a member of the genus Carcharhinus, although this is uncertain. This tooth clearly suffered taphonomic alteration, but many important characteristics are still visible. The tooth is labio-lingually compressed with a central cusp that is broadly triangular with no obvious evidence of a serrated cutting edge, although the pre-burial rounding of the specimen may have eroded a small serrated cutting edge. There are no lateral cusplets or heels. Diagnostic traits of the crown are few, but the root possesses many traits that aid in assignment to the Carcharhinidae. A pronounced nutrient groove runs labio-lingually in the midline of the root, terminating just after reaching the lingual face. There appears to be a weak lingual protuberance, although not like the prominent protuberance easily identified on triakid or hemigaleid sharks. Root lobes are expanded with the basal margin flat and slightly widened, giving the tooth and overall plateaued appearance. Finally, the labial root face possesses a shallow ovoid impression presumably to house the weak lingual protuberance of the preceding tooth within the tooth family.

Materials.—FMNH PF 15982, isolated tooth.

Remarks.—Scanning electron microscopy of FMNH PF 15982 reveals no pitting on the crown enameloid as would be expected of a tooth that has undergone prolonged digestion. However, microabrasions are abundant across the enameloid surface (Fig. 2.7, 2.8) and large pits are present across the root, of which
the latter maybe natural. Possibilities for the relocation from marine to terrestrial sediments include: (1) reworking into the terrestrial “SUE” Locality sequence from another stratigraphic interval (possibly the underlying marine Fox Hills Formation); or (2) loss of the tooth by a carcharhinid shark traveling in freshwater during deposition of the Hell Creek Formation.

Carcharhinid species have been recorded from sediments prior to the Paleocene (Cappetta, 2012); however, Underwood and Ward (2008) described teeth they considered carcharhinid from Santonian rocks of Britain, which Guinot et al. (2013) challenged, instead placing the teeth in the Triakidae as opposed to the Carcharhinidae. Later, Guinot et al. (2014) named *Archaeogaleus lengadocensis*, a new carcharhinid taxon from the Valanginian of France, making that the earliest fossil occurrence of the species. Molecular phylogenies suggest that carcharhinid sharks evolved in the Cretaceous, or possibly the Jurassic, according to the calibrations used in Sorenson et al. (2014). Therefore, the discovery of teeth from this family in uppermost Cretaceous rocks should not come as a surprise. If the tooth should be found to represent the genus *Carcharhinus*, it means that FMNH PF 15982 pushes the oldest known occurrence of this genus into the upper Maastrichtian. At least two published instances document the discovery of marine shark teeth in Maastrichtian fluvial environments of western North America (Wroblewski, 2004; Wynd et al., 2018).

**Figure 2.** *Lonchidion selachos* (FMNH PF 15756) and Carcharhinidae indet. (FMNH PF 15982) teeth from the FMNH “SUE” locality. FMNH PF 15756 in (1) lingual, (2) labial, (3) basal, and (4) occlusal views. FMNH PF 15982 in (5) lingual and (6) labial views. Scanning electron microscopic image of Carcharhinidae indet. FMNH PF 15982 (7) and (8). Scale bar (1–4) equals 1 mm. Scale bar (5, 6) equals 2 mm. Scale bar in (7) equals 10 μm. Scale bar in (8) equals 100 μm.

Class Chondrichthyes Huxley, 1880
Subclass Elasmobranchii Bonaparte, 1838

Order Orectolobiformes Applegate, 1972
Family incertae sedis
*Galagadon* new genus

*Type species.*—*Galagadon nordquistae* n. gen. n. sp. by monotypy.

*Diagnosis.*—Same as for type species by monotypy.

*Etymology.*—*Galaga*, named for the shape of the teeth, which when seen in different views resemble the spaceships in the arcade game “Galaga” developed and copyrighted by the company Namco; and *don-* Greek for “tooth.”

*Remarks.*—Designation of this new genus is based on distinct traits seen across the entire series of teeth of the current hypodigm, which is consistent with variation found in other Orectolobiformes genera. All teeth come from a single locality, which is lower in section than other sites (e.g., Bug Creek Anthills). Other localities at this stratigraphic level may reveal more specimens of this genus.

*Galagadon nordquistae* new species

*Figures 3–7*

*Holotype.*—FMNH PF 15769, an isolated tooth, possibly from an anterolateral position.
Diagnosis.—Orectolobiform shark possessing the following autapomorphies: central cusp with distinctly swollen lingual face forming a clearly demarcated constriction, or neck, between the cusp and the root; labial surface of central cusp ornamented with a raised ridge or closely arrayed plications, which in most cases are distributed in such a pattern as to follow the slope of the heels; convex heels, well developed both mesial and distal to the central cusp with a distinct convex angle mid-distance along slope seen in anterior teeth and some lateral teeth. This taxon is further differentiated by the following suite of shared characters: anterior teeth possess distinct, high central cusp; heels on lateral teeth slightly serrated; heels of anterior teeth and some lateral teeth do not slope gradually toward root lobes, but are instead squared-off or rounded at their terminus; one or two rounded diminutive cusplets may be present although this trait varies among individual teeth; apron is generally broadly rounded and shows a bifid habit on some but not all specimens; root lobes enlarged on lingual side of tooth compared to more constricted structure on labial side, bestowing an exaggerated heart-shape in basal view; a central foramen pierces between the root lobes with the foramen divided by thin struts in some teeth.

Occurrence.—Field Museum “SUE” locality; Hell Creek Formation; Faith, South Dakota, USA; Upper Maastrichtian.

Description.—The tooth series was reconstructed based on a combination of tooth central cusp size and distal recurvature.

Figure 3. Anterior teeth of Galagodon nordquistae n. gen. n. sp. from the FMNH “SUE” locality. FMNH 15759 (1, 5, 9, 13, 17); FMNH 15761 (2, 6, 10, 14, 18); FMNH 15765 (3, 7, 11, 15, 19); FMNH PF 15766 (4, 8, 12, 16, 20). Specimens in lingual view (1–4), labial view (5–8), lateral view (9–12), basal view (13–16), and occlusal view (17–20). Scale bars equal 1 mm.
of the central cusp. Teeth with less distal recurvature are considered anterior teeth, slight distal recurvature typifies lateral teeth, whereas shorter central cusps represent posterior teeth.

The root of each *Galagadon nordquistae* n. gen. n. sp. tooth is heart-shaped and more similar to *Eometlaonia* spp. (Noubhani and Cappetta, 2002), *Restesia americana* (Estes, 1964) (Cook et al., 2014), and *Chiloscyllium* spp. (Cook et al., 2014) than to other species, yet remains distinct (Figs. 3, 4, 5, 7, 8, 10, 11, 13, 14, 16). The lingual extensions of the root lobes are narrow, beginning sub-parallel in anterior teeth and widening into a broad “V”-shape in more posterior teeth (e.g., Figs. 3, 6, 13–6, 16); note that we define posterior teeth by their relatively shorter crowns compared to other teeth in the hypodigm). Similar to the fashion in which the lingual root lobes vary anteriorly to posteriorly, the labial root lobes expand into the upper curved portion of the heart-shape with different degrees of exaggeration. In this case, anterior teeth (Figs. 3, 4, 13–4, 16) tend to possess an abrupt contact between the mesial and distal halves of the root, which produces a mouse-eared look to the entire root lobe; however, the lateral files show a much more gradual progradation from narrow to expanded root lobes (Figs. 5, 3, 5, 4, 6, 13–6, 16). As such, the sharpness of the contact between the lingual and labial regions of the root appears less abrupt than in more anterior teeth. Even the roots of *Restesia*, a taxon also present in the Hell Creek Formation (Cook et al., 2014), are much more subdued...
Figure 5. Anterolateral and lateral teeth of Galagadon nordquistae n. gen. n. sp. from the FMNH “SUE” locality. FMNH PF 15768 (1, 3, 5, 7, 9) and 15771 (2, 4, 6, 8, 10) in (1, 2) lingual view; (3, 4) labial view; (5, 6) lateral view; (7, 8) basal view; (9, 10) occlusal view. Scale bars equal 1 mm.

distally. The central cusp is swollen compared to the root, which produces a distinct constriction of neck between the tooth regions. Mesial and distal heels slope basally from the central cusp. In anterior teeth (Figs. 3, 4), the heels originate near the tip of the central cusp, whereas on lateral teeth the heels originate lower on the central cusp (Figs. 5, 6). The general shape of the heel changes posteriorly—the anterior teeth have mesio-distally narrower heels than lateral teeth, as a result of the narrower crown base compared to the laterals, and they show a distinct convex angle midway towards the tips (Figs. 3, 4, 5.1, 5.2, 6.1–6.4). Sharpness of this angle varies between specimens. Laterally, the heels largely lose the convex angle and become less steep. Some of the heels in the lateral series have slightly serrate apical surfaces. A small, diminutive cusplet is present on some teeth (Fig. 6.1). Development of this feature does not seem to follow a pattern through the tooth series. On both mesial and distal margins of the tooth, the central cusp-root intersection is positioned at the mid-point of a deep embayment formed by the heel and the root lobes. Such a feature is common in most orectolobiform teeth in which the root is as wide or wider than the crown. Depth of the constriction between the tooth central cusp and root similar to that of Galagadon n. gen. (well developed on the type specimen FMNH PF 15769) is visible on teeth of Cantioscyllium markaguntensis Kirkland, Eaton, and Brinkman, 2013 (Kirkland et al., 2013, fig. 9.9) and Palaebraeuchaelurus spp. (Cappetta, 2012, fig. 154).

The lingual bulge is prominent. When a tooth is viewed laterally, the lingual bulge can be seen to extend lingually equal to the lingual extent of the central cusp. In other words, a vertical line drawn from the tip of the central cusp to the base would form a tangent to the lingual bulge. A single, large lateral foramen is present on each side of the lingual protuberance. There is much variation in the size of this lateral foramen among orectolobiform species (Cappetta, 2012). That seen on Galagadon n. gen. is one of the larger lateral foramina, similar to Eometlaouia spp., Hemiscyllium ocellatum, and many members of the Ginglymostomidae.

Another comparison between Galagadon n. gen. and other species is the apical angle between the crown and the lingual protuberance. Variation in this angle seems wide-ranging, even among supposedly closely related species. For instance, Galagadon n. gen. has an arch generally similar in aspect to some specimens of Chiloscyllium missouriense (Case, 1979) (Kirkland et al., 2013, fig. 9.10CC), Restesia americana (Cook et al., 2014, fig. 3), or Eometlaouia numidica (Cappetta, 2012, fig. 145D). This is in contrast to the wider arch of Braeuchaelurus waddi or the narrower arch of the brachaelurid Eosteogostoma spp. (Cappetta, 2012, fig. 153C, G). Given the variation seen in this trait, it seems unlikely that it will be useful for morphological phylogenetic analysis.

The labial face of Galagadon n. gen. teeth is more or less triangular. An absence of cusplets creates an uninterrupted edge from the apex to the lateral corners of the crown. The labial face of each Galagadon n. gen. central cusp is flat to slightly convex (Figs. 3–6). The plications and raised ridges of the labial face can also be seen. In some teeth, the labial face has a large section raised to form a slight plateau-like feature that can occupy nearly the entire central portion of the central cusp. Along the base of the labial face, the apron extends basally for...
only a short distance, but spreads widely laterally. There is a
difference between the anterior and lateral teeth in that the
apron is slightly more extended in anterior teeth (Figs. 3, 4,
5.3, 5.4, 6.5–6.8). The basal aspect of the apron ranges in
shape from rounded to flat to faintly bifid and shows no particu-
lar pattern among tooth positions. The apico-basal shortness
of the apron differs from the elongated aspect observed in brachae-
lurids, and is much more similar to species of ginglymostomids
and some orectolobids (although there is much more variation in
apron morphology among this clade). Across the labial face of
the crown are numerous enameloid folds that form bumps,
short ridges, and elongate ridges, all coalescing into a chevron
with the terminal ends occupying the basal mesial and distal cor-
ners of the labial face, and the apex positioned along the midline
at various places between the central cusp apex and base of the
apron, depending on the individual tooth (Figs. 3, 4, 5.3, 5.4,
6.5–6.8). Some teeth, such as FMNH PF 15773 (Fig. 4.2)
show a well-defined triangular ridge with an accessory ridge
spanning from the apex of the triangular ridge to the apex
of the central cusp. Most other teeth exhibit a lesser degree
of enameloid folding than is seen on the aforementioned specimen.
Nonetheless, all specimens have ornate labial central cusps and
the plications form the same basic shape of an arc or triangle in
the center of the crown (except FMNH PF 15770 whose labial
face has a single apically oriented ridge).

A labial crown face adorned with bumps, ridges, and plica-
tions is a common morphological feature of species within the
Ginglymostomidae and other orectolobiform species (Cappetta,
2012). Protoginglymostoma, as illustrated in Cappetta (2012,
fig. 163A) exhibits small folds that look similar to those
observed on several specimens of Galagodon n. gen., although
these folds are too few in number and in a different position, with
exception of the Protoginglymostoma (Cantioscyllium) estesi
specimen UALVP 53531 (Cook et al., 2014), which shows no
ornamentation. Other ginglymostomid species that show orna-
mentation similar to Galagodon n. gen. include Plicatoscyllium,

Figure 6. Lateral teeth of Galagodon nordquistae n. gen. n. sp. from the FMNH “SUE” locality. FMNH 15762 (1, 5, 9, 13, 17); FMNH 15763 (2, 6, 10, 14, 18); FMNH 15766 (3, 7, 11, 15, 19); FMNH PF 15983 (4, 8, 12, 16, 20). Specimens in lingual view (1–4), labial view (5–8), lateral view (9–12), basal view (13–16), and occlusal view (17–20). Scale bars equal 1 mm.
which possesses ridges in a general triangular pattern, and *Can-
tioscyllium* spp., which possesses apico-basally oriented ridges, especially on *C. markaguntensis* (Kirkland et al., 2013). Outside of the Ginglymostomidae, *Orectoloboides* has irregularly apico-basally oriented ridges (Cappetta, 1977). Guinot et al. (2014) illustrated a specimen of *Ornatoscyllium* that exhibits a raised ridge following the heels in a similar fashion to *Galagadon* n. gen., and specimens figured in Underwood and Ward (2004) have less ornamentation. Also, Neuman and Brinkman (2005, fig. 9.1) figured an orectolobiform tooth (TMP 93.116.2) that has ample ornamentation.

Four teeth that may belong to *Galagadon* n. gen. (Fig. 7) are less certain with respect to their affinities. They could represent upper teeth of a heterodont dentition, ontogenetic or sex differences, or simply individual variation. The heel ridges on these teeth are more distinct—forming incipient cusplets—and the ornamentation is varied more than in the other teeth, albeit still similar in overall appearance. FMNH PF 15777 (Fig. 7.1, 7.5, 7.13, 7.17) has the most divergent labial ornamentation in that they are small bumps arrayed in an arc.

**Etymology.**—The species name honors Ms. Karen Nordquist for her decades of support and volunteering at the FMNH, including as a matrix sorter, during which she discovered the teeth described here.
Materials.—FMNH PF 15758, isolated tooth; FMNH PF 15759, isolated tooth; FMNH PF 15760, isolated tooth; FMNH PF 15761, isolated tooth; FMNH PF 15762, isolated tooth; FMNH PF 15763, isolated tooth; FMNH PF 15764, isolated tooth; FMNH PF 15765, isolated tooth; FMNH PF 15766, isolated tooth; FMNH PF 15767, isolated tooth; FMNH PF 15768, isolated tooth; FMNH PF 15770, isolated tooth; FMNH PF 15771, isolated tooth; FMNH PF 15772, isolated tooth; FMNH PF 15773, isolated tooth; FMNH PF 15774, isolated tooth; FMNH PF 15775, isolated tooth; FMNH PF 15776, isolated tooth; FMNH PF 15777, isolated tooth; FMNH PF 15778, isolated tooth; FMNH PF 15780, isolated tooth; FMNH PF 15781, isolated tooth; FMNH PF 15783, isolated tooth; FMNH PF 15784, isolated tooth.

Remarks.—The Galagadon n. gen. teeth from the “SUE” site are some of the lowest orectolobiform teeth yet discovered in the Hell Creek Formation. More discoveries of carpet sharks throughout the formation will yield information on the dynamics of the Cretaceous Western Interior Seaway and its influence of the preserved chondrichthyan fauna.

Class Chondrichthyes Huxley, 1880
Subclass Elasmobranchii Bonaparte, 1838
Order Rajiformes Berg, 1940
Suborder Rhinobatoidei Fowler, 1941
Family Rhinobatidae incertae sedis sensu Cappetta, 2012
Genus Myledaphus Cope, 1876

Type species.—Myledaphus bipartitus Cope, 1876.

Myledaphus sp.  
Figure 8.13–8.18

Materials.—FMNH PF 15989, isolated denticle; FMNH PF 15990, isolated denticle.

Remarks.—Included within the rhinobatid remains from this site are two dermal denticles (FMNH PF 15989, FMNH PF 15990; Fig. 8.13–8.18). These dermal denticles are virtually identical to those described by Neuman and Brinkman (2005), but no comparative material exists that is definitely attributable to Myledaphus pustulosus in order to define dermal denticate morphology beyond Myledaphus sp. (sensu Cook et al., 2014).

Myledaphus pustulosus Cook et al., 2014  
Figure 8.1–8.12

Type specimen.—UCMP 197869, isolated tooth.

Description.—Ray teeth attributed to Myledaphus pustulosus dominate the selachian fauna at the “SUE” locality. This species is diagnosed on the basis of tubercles on the labial side of the tooth occlusal surface. Cook et al. (2014) further described M. pustulosus as bearing transverse ridges that are discontinuous toward the lingual edge of the tooth. Figure 8 shows teeth from the “SUE” locality that possess each of the autapomorphic traits of M. pustulosus.

Materials.—FMNH PF 15313, isolated tooth; FMNH PF 15314, isolated tooth; FMNH PF 15786, isolated tooth; FMNH PF 15787, isolated tooth; FMNH PF 15788, isolated tooth; FMNH PF 17991, 44 isolated teeth.

Remarks.—Myledaphus pustulosus is the taxon with the most numerous remains from this locality, to date.

Results

Phylogenetic analysis.—Many extant shark families and subfamilies are thought to have arisen in the Mesozoic based on the fossil record (Frickhinger, 1999; Cappetta, 2012); however, there has been disagreement regarding the phylogenetic affinities of Mesozoic Orectolobidae. Cappetta (2012) classified the Cretaceous taxa Cretorectolobus olsoni and Cederstroemia triangulata as orectolobids, but other authors (e.g., Siverson, 1995; Underwood and Mitchell, 1999; Underwood et al., 1999) have treated these as Orectolobiformes incertae sedis. The earliest record of orectolobids within their present day biogeographic distribution appears to be Miocene in age (Kemp, 1991), and molecular analysis suggests that the extant taxa are members of a comparatively recent and rapid radiation, likely starting in the Miocene with many species originating as recently as the Pleistocene (Corrigan and Beheregaray, 2009).

Parsimony analysis.—The first analysis resulted in 468 Most Parsimonious Trees (MPTs) with a length of 263, Consistency Index of 0.692, Rescaled Consistency Index of 0.617, and Retention Index of 0.891 (the strict consensus trees are shown in Figure 9.1). Galagadon n. gen., as well as the other WIS fossil taxa Cretorectolobus olsoni and Cederstroemia triangulata, are nested deep within the Orectolobidae, thus supporting the placement originally proposed by (Cappetta, 2012), despite prior publications placing these genera as Family incertae sedis (e.g., Siverson, 1995; Underwood and Mitchell, 1999; Underwood et al., 1999). Our results agree furthermore with those of Goto (2001) and Corrigan and Beheregaray (2009) in finding that the genus Orectolobus is polyphyletic. Surprisingly, our analysis recovers the three WIS taxa as successive sister taxa to a more exclusive clade that includes some, but not all, living wobbegong species rather than forming an endemic clade, although this result is not robust and is not recovered in Bayesian analyses (see below). Another relevant result of our phylogenetic analysis is that the two fossil taxa attributed to Ginglymostomidae, Protoginglymostoma ypresiensis, and Ornatoscyllium freemani, are recovered within a clade with other nurse sharks, albeit in a polytomy (Fig. 9.1).

Our second analysis, which enforced monophyly of the three fossil species from the WIS, is only one step longer than the MPTs recovered from the first analysis. The constrained monophyletic clade still occupies the same position relative to living species where the three fossil species were recovered in unconstrained analyses; that is, sister to the remaining species of Orectolobus and Eocrossorhinus and the Orectolobus wardi and Sutorectus tentaculatus clade serving as the earliest diverging sister group to other orectolobids.
Figure 8. Examples of ray teeth and dermal denticles from the FMNH “SUE” locality. FMNH PF 15787 (1–3); FMNH PF 15313 (4–6); FMNH PF 15786 (7–9); FMNH PF 15788 (10–12) in lingual view (1, 4, 7, 10), basal view (2, 5, 8, 11), and occlusal view (3, 6, 9, 12). FMNH PF 15990 (13–15); and FMNH PF 15989 (16–18) in lateral view (13, 16), caudal view (14, 17), and apical view (15, 18). Scale bars equal 1 mm.
With a molecular scaffold constraint imposed on the data, resolution among orectolobid species is reduced in the strict consensus of the resulting 6318 MPTs. Only two nodes within Orectolobidae are recovered across all MPTs: (1) the family is recovered as monophyletic and sister to *Brachaelurus*; and (2) the late diverging extant species *Orectolobus japonicus*, *O. maculatus*, *O. ornatus*, *Sutorectus tentaculatus*, and *Eocrocosrhinos dasypogon* form a clade to the exclusion of *O. wardi* and the three fossil orectolobids, which still fall within the crown group.

Bayesian analysis.—In the favored single partition model (Table 3), all major clades are generally retained as reconstructed in MP analyses. Several differences, however, are found within the major clades because the hierarchical ordering of certain taxa differs from the MP analyses. For example, *O. wardi* is recovered as the sister taxon to *G. nordquisti* n. gen. n. sp. and the rest of the orectolobids in the MP topology, but in the BI topology, *O. wardi* is nested within orectolobids and as the sister taxon to *S. tentaculatus* with a posterior probability of 95%.

Based on comparisons of Bayes Factors across analyses, results of the analysis with separate partitions for non-dental and dental anatomical traits, with equal/variable rates model for the former partition, is preferred (Table 3). The maximum clade credibility (MCC) tree (Fig. 9.2) from that analysis places the fossil taxa *Cederstroemia* and *Cretorectolobus* as stem orectolobids, with all extant species forming a monophyletic clade, in contrast to the MP results. This is potentially more consistent with respect to the timing of the radiation of extant wobbegongs, which has been calibrated to be Miocene and younger based on analyses of mitochondrial DNA (Corrigan and Beheregaray, 2009), than the MP results that nest these Cretaceous fossils within the crown and thus dictate a Mesozoic age for the earliest divergences between extant lineages.

Figure 9. Phylogenetic trees positing position of *Galagadon nordquisti* n. gen. n. sp. among Orectolobiformes. (1) Majority Rules consensus tree for maximum parsimony analysis. (2) Bayesian Inference maximum clade credibility tree. For Figure 9.1, regular font numbers are frequencies of clades in the consensus tree; numbers in italics are the bootstrap values calculated in PAUP with the same settings described in body text for 100 replicates, and numbers in bold are the decay/Bremer values calculated in TNT 1.1 (Goloboff et al., 2008).
of which combination of analytical parameters is employed. However, the posterior probability is relatively low for its current placement (∼30%) and adjacent nodes suggest some ambiguity for this new taxon near the orectolobid (wobbegong) and hemiscyllid (bamboo) shark split (posterior probability of ∼28% at the node uniting both clades). In order to assess whether this difference could be due to potential biases in the placement of taxa with large amounts of missing data in Bayesian Inference analyses (Goloboff and Pol, 2005), we ran a series of tests in which fossil taxa were serially excluded, but no change in tree topology was noted. Taxon exclusion experiments conducted in MP suggest that the nested position of Galagadon n. gen. within Orectolobidae is dependent on inclusion of at least one of the other Cretaceous WIS species. When both Cederstroemia and Cretorectolobus are excluded, the strict consensus is unresolved to a large degree with neither Orectolobidae nor Hemiscyllidae recovered as monophyletic groups across all MPTs, indicating that Galagadon n. gen. can group with either of these shark families in the absence of the two other WIS fossil taxa. The discrepancy between BI and MP results is thus best explained as the result of different optimality criteria operating on a taxon characterized by homoplasic characters that can support at least two distinct phylogenetic positions.

Biogeographic Analysis.—Results of the biogeographic analysis (Table 4) of the BAYAREALIKE + j model find the founder-effect speciation (i.e., the ’j’ parameter [Matzke, 2013]) serves as a significant and more frequent mechanism to explain the observed distributions than dispersal, extinction, or vicariance. Founder-effect speciation is often invoked in instances of island biogeography (e.g., Matzke, 2013), especially in regards to oceanic islands, for which there is scant evidence of historic range expansions or connections to other areas. As such, this model parameter may be ill suited to explain distributions of marine organisms distributed in oceans and epeiric seaways that were never isolated from each other, and which, although ephemeral in some instances, persisted for tens of millions of years or more.

In light of this concern, we also considered the best fitting alternative model that did not employ the j parameter to examine potential biogeographic histories for orectolobiform sharks: the DEC model (Ree and Smith, 2008) for the MP scaffold tree and the BAYAREALIKE model for the BI tree. For the MP topology, ancestors to clades that encompass the Cretaceous WIS taxa as well as extant orectolobids, are reconstructed as being widespread and occurring in both the western Pacific and WIS, and including the (Para)Tethys for the node from which Eometlaouia -c and Indian Oceans. Furthermore, for the BI topology, the clades that include the WIS taxa and their respective extant carpet- and bamboo-shark relatives, the ancestral areas are also reconstructed to be more widespread and include the WIS and western Pacific, implying a wider geographic range than the current distribution suggests. These reconstructions then explain the present-day distributions of carpet and bamboo sharks as resulting from local extirpation of WIS taxa and secondary range expansion into the Indian Ocean as it achieved its current geography following the docking of India and extrusion of Indochina.

Discussion

Our study of the chondrichthyan fauna of the “SUE” quarry yielded both common Late Cretaceous taxa such as the batoid *M. pustulosus* and the hybodont shark *Lonchiodon*, but also

### Table 3. Model scores for the various Bayesian phylogenetic inferences produced for this study. The merged cells represent a single data partition. Abbreviations: LnL, harmonic mean log likelihood values.

<table>
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<tr>
<th>Body Partition</th>
<th>Dental Partition</th>
<th>LnL</th>
<th>Bayes Factor</th>
</tr>
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<td>−</td>
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<tr>
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<td>Variable</td>
<td>−1034.599</td>
<td>55.786</td>
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</tbody>
</table>

### Table 4. Model scores for the Maximum Parsimony and Bayesian Inference trees used in the biogeographic analysis. Abbreviations: AIC, Akaike Information Criterion; LnL, Log likelihoods.

<table>
<thead>
<tr>
<th>Model</th>
<th>LnL</th>
<th>AIC</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>DEC</td>
<td>−87.38</td>
<td>178.8</td>
<td>0.0002</td>
</tr>
<tr>
<td>DEC + j</td>
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<td>167.3</td>
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<td>BAYAREA + j</td>
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<td>145.1</td>
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### Bayesian Inference

<table>
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<th>AIC</th>
<th>p-value</th>
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</tbody>
</table>

However, specific routes for these independent dispersal events into the WIS are unclear in the best-fitting analyses that rely on the j founder-event parameter to explain the distributions of the endemic WIS Cretaceous taxa, given that there is a lack of close relatives spanning the areas between the western Pacific Ocean and the WIS (see Discussion). Subsequent dispersals from the western Pacific to the Indian Ocean are reconstructed in the Neogene to explain the current distributions of carpet and bamboo sharks in tropical to temperate parts of the western Pacific and Indian Oceans.

Biogeographic reconstructions on the MP scaffold tree differ in reconstructing the WIS as the ancestral area to the orectolobid crown and Cretaceous stem taxa (Fig. 10.2). Two separate dispersal events to the western Pacific in the Late Cretaceous are invoked to account for the present-day distribution of *O. wardi* and the *Sutorectus-Eucrossorhinus-Orectolobus* (pars) clade. As in the results of the BI analysis, this could also indicate a more widespread ancestral area for the orectolobids given that the reconstructed ancestral areas include two disconnected areas as defined in the current analysis.

Both the BI and MP analyses using the BAYAREALIKE + j model find the founder-effect speciation (i.e., the ‘j’ parameter [Matzke, 2013]) serves as a significant and more frequent mechanism to explain the observed distributions than dispersal, extinction, or vicariance. Founder-effect speciation is often invoked in instances of island biogeography (e.g., Matzke, 2013), especially in regards to oceanic islands, for which there is scant evidence of historic range expansions or connections to other areas. As such, this model parameter may be ill suited to explain distributions of marine organisms distributed in oceans and epeiric seaways that were never isolated from each other, and which, although ephemeral in some instances, persisted for tens of millions of years or more.
an unexpected diversity of shark teeth. This includes a specimen that might mark a First Appearance Datum (FAD) for the carcharhiniform genus *Carcharhinus*, which had fossil records restricted to the Cenozoic (Maisey, 2012; Fig. 3). A recent molecular analysis of shark relationships posited the last common ancestor of hemigaleids and carcharhinids as sister to the triakid sharks (Veléz-Zuazo and Agnarsson, 2011), which have a fossil record extending into the Late Cretaceous (see Becker et al., 2006; Maisey, 2012; Sorenson et al., 2014).

Recovery of the carcharhinid from the "SUE" Locality is surprising given the freshwater environment from which the fossils derive. This specimen shows clear evidence of reworking (rounded edges and a shiny patina), suggesting that it was not originally deposited at the same time as the *Tyranosaurus rex* fossil. That said, there are three modern species of carcharhinid shark that either regularly invade euryhaline habitats or are restricted to freshwater ecosystems: *Carcharhinus leucus* (Bull shark) (Müller and Henle, 1839) and two species of *Glyphis* (including the Ganges shark) (Compagno and Cook, 1995; Martin, 2005). The combination of evidence for taphonomic reworking with the known occurrence of some modern relatives invading freshwater habitats makes it difficult to infer life habits of this shark.

The second novel discovery is *Galagadon nordquistae* n. gen. n. sp. As mentioned above, there are currently three species of orectolobiform chondrichthyan known from the Hell Creek Formation: *Protoginglymostoma estesi* (Herman, 1977) (Cappetta, 2006), *Restesia americana* (Cook et al., 2014), and *Chiloscyllium* sp. (Cook et al., 2014). *Galagadon* n. gen. is the only Hell Creek Formation orectolobiform currently known that possesses abundant folds and plications on the labial tooth surface. With the inclusion of *Galagadon* n. gen., a fourth species can now be included within the freshwater ecosystems. Increasing the species-richness of aquatic ecosystems is not surprising when one considers the complexity and heterogeneity of habitats available to orectolobiformes during the Late Cretaceous. For over 30 million years prior to the deposition of the Hell Creek Formation, the WIS inundated the central portion of North America, generating marine and estuarine ecosystems for orectolobiforms to occupy. Despite all extant orectolobiforms living strictly in marine ecosystems (Martin, 2005),
tolerance to varying salinity levels within this group is documented by occurrences of orectolobiform fossils discovered across salinity gradients from the marine into upland river systems (e.g., Kirkland et al., 2013), and the “SUE” locality is considered a freshwater environment based on both its distance to the paleoshoreline and the occurrence of amphibian and bony turtle remains. Terrestrial landscape heterogeneity due to topographic and climatic microcosms during the Late Cretaceous in Laramidia is posited to have caused an increase in vertebrate biodiversity (Gates et al., 2010a, 2012), which is similar to biocenosis identified by Nicholls and Russell (1990) for the seaway itself. As such, it seems reasonable that a number of small-bodied sharks could live together in such diverse habitats, or perhaps they are occurring in short, successive stratigraphic intervals. Many extant carpet-and bamboo shark species also exhibit overlapping or peripatric distributions (Corrigan and Beheregaray, 2009; Compagno et al., 2005).

Past taxonomic practice often has been to assign fossil shark teeth to modern families based on morphological similarity, but the phylogenetic accuracy of such referrals has been questioned, especially for very old records including fossil Orectolobiformes (Glikman, 1967; Maisey, 2012). Here we have attempted to move beyond this concern by providing the first quantitative phylogenetic analysis that combines fossil and living Orectolobiformes. Although we have been unable to sample all Mesozoic fossils that have historically been assigned to Orectolobiformes, our analyses provide support for the phylogenetic placement of several contentious Mesozoic shark taxa within Orectolobiformes and its subclades. For example, both our Bayesian and MP analyses recovered *Cretorectolobus* and *Cederstroemia* as orectolobid relatives, which were originally designated by Siverson (1995) as simply Family Orectolobiformes incertae sedis. We also found support for grouping fossil and extant ginglymostomatids (nurse sharks).

Concerns related to phylogenetic assignments based on similarity alone are highlighted by our case study of *Galagodon* n. gen. All our MP analyses, whether constrained or not, posit *Galagodon* as an orectolobid closely related to the other WIS taxa and most crown clade species, whereas our BI analyses consistently recovered it as a stem hemiscyllid (however, relatively low posterior probability at the nodes uniting and nested within carpet and bamboo sharks indicates its phylogenetic placement with less precision). Taxon exclusion experiments suggest that the differences in topological results are not a result of methodological bias, but instead reflect how different models handle taxa with conflicting character data. Such insights regarding possible phylogenetic placement of fossils can only be gained by applying a range of quantitative phylogenetic analyses.

The placement of fossil species relative to modern ones differed notably between BI topologies and MP ones. In the latter, Cretaceous fossil species were recovered inside the orectolobid crown group forcing a Cretaceous minimum date on its earliest divergences. The BI results, on the other hand, place all WIS fossil species as stem taxa to the extant diversity of carpet and bamboo sharks, thus allowing all divergences among extant species to be much younger, in better agreement with the molecular divergence estimates reported by Corrigan and Beheregaray (2009). While the latter study was not calibrated against ingroup fossils, its estimates of very young (Plio-Pleistocene) divergences for most orectolobid species is consistent with the geological history and oceanography of the Indopacific region.

Our combined fossil and extant phylogenies allow for an examination of the biogeographic history of parts of the orectolobiform radiation. Results of the best-supported models suggest that a series of long distance dispersals or founder events account for most of the distribution of orectolobiformes over geological history, with only minor roles for dispersal through range expansion and vicariance. This result runs counter to most biogeographic theory in which the latter processes are thought to account for the bulk of distribution patterns among organisms (e.g., Ree and Smith, 2008), with founder events playing a rare though important role, especially for island systems (Matzke, 2013). When considering other models that omit the stochastic founder event parameter, the best-fit models indicate that present-day orectolobid and hemiscyllid distributions are relics of wider ancestral ranges with a secondary, but more limited radiation within the Indopacific region. This result is more consistent with traditional interpretations of orectolobiform history based on the fossil record, which comprises many named fossil species from shallow marine deposits in Europe, North Africa, the Middle East, and even Antarctica, as well as from the WIS (Compagno et al., 2005; Maisey, 2012; Engelbrecht et al., 2017). We hypothesize that our initial results are biased by the patchy nature of the orectolobiform record (Maisey, 2012), which is further exacerbated by our sampling of only those fossil species for which descriptions allow for character coding. The oldest fossils are Tethyan, whereas all Cretaceous fossils come exclusively from the WIS. Because these seaways disappear in the Mesozoic and thus cannot be shared with taxa not be invoked in the dispersal of extant crown clades, the analysis forces founder events to explain past and present-day distributions. This problem may be remedied by better sampling of the fossil record and we hope that future work will expand our preliminary phylogenetic analysis to include more critical fossils and in turn allow for a more comprehensive analysis and understanding of shark biogeography.

**Conclusions**

Orectolobiform sharks held a rich ancient biodiversity within both strictly marine ecosystems through freshwater riverine habitats. The freshwater “SUE” locality from the latest Cretaceous Hell Creek Formation of South Dakota preserves a new orectolobiform shark, *Galagodon nordquistae* n. gen. n. sp., diagnosed based on a series of unique and shared traits, such as a constriction between the central cusp and root and a convex angle on the heles. In addition to *Galagodon* n. gen., other chondrichthyes species found at the site include *Lonchodion selaches*, *Mylephalus pustulosus*, and an unidentified tooth from the family Carcharhinidae. The latter species marks a new occurrence for the respective family, firmly pushing the fossil record for their origination into the Maastrichtian. Taken together, these finds add to an already considerable diversity of WIS chondrichthyes and corroborate that Mesozoic epeiric seaways played an important role in the diversification of modern sharks and rays, as suggested by Maisey (2012).
We provide the first analytically supported phylogenetic positions for several Mesozoic orectolobiform taxa by incorporating both modern and ancient taxa through combined evidence data matrices, allowing for preliminary exploration of the evolution and biogeography of the clade. This study adds to others (e.g., Adnet and Capetta, 2001; Peart et al., 2015) that combine extant and fossil species to explore the history of shark evolution.

Accessibility of supplemental data

Data available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.m276f51

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