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MORPHOLOGICAL EVOLUTION OF SILICA SCALES IN THE FRESHWATER GENUS *SYNURA* (STRAMENOPILES)1

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A high degree of morphological variability is expressed between the ornately sculptured siliceous scales formed by species in the chrysophycean genus, *Synura*. In this study, we aimed to uncover the general principles and trends underlying the evolution of scale morphology in this genus. We assessed the relationships among thirty extant *Synura* species using a robust molecular analysis that included six genes, coupled with morphological characterization of the species-specific scales. The analysis was further enriched with addition of morphological information from fossil specimens and by including the unique modern species, *Synura punctulosa*. We inferred the phylogenetic position of the morphologically unique *S. punctulosa*, to be an ancient *Synura* lineage related to *S. splendida* in the section *Curtispinae*. Some morphological traits, including development of a keel or a labyrinth ribbing pattern on the scale, appeared once in evolution, whereas other structures, such as a hexagonal meshwork pattern, originated independently several times over geologic time. We further uncovered numerous construction principles governing scale morphology and evolution, as follows: (i) scale roundness and pore diameter decreased during evolution; (ii) elongated scales became strengthened by a higher number of struts or ribs; (iii) as a consequence of scale biogenesis, scales with spines possessed smaller basal holes than scales with a keel and; and (iv) the keel area was proportional to scale area, indicating its potential value in strengthening the scale against breakage.

Key index words: chrysophytes; construction principles; microalgae; molecular; morphology; phylogeny; silica scales; *Synura punctulosa*; synurophytes; ultrastructure

Abbreviations: BIC, Bayesian information criterion; MCMC, Markov Chain Monte Carlo; mt, mitochondrial; NMDS, non-metric multidimensional scaling; nu, nuclear; psaA, photosystem I P700 chlorophyll a apoprotein A1; pt, plastid; SDV, silica deposition vesicle

Every day, we observe inorganic and living objects with a vast array of shapes, patterns, structures, and colors. Morphological variability expressed in living organisms is often explained by a random and passive tendency to increase complexity, constrained by physical and mechanical laws, and ultimately shaped by natural selection. Examples of such morphological diversification are those driven by sexual selection or social competition in plants and their pollinating insects (Castellanos et al. 2006, Whittall and Hodges 2007, Quesada-Aguilar et al. 2008), eccentrically colored birds (Endler and Day 2006, Rubenstein and Lovette 2009, Maia et al. 2013), and fishes such as cichlids (Elmer et al. 2010, Arbour and Lópéz-Fernández 2013, Felich 2016).

Compared to the macroscopic world, studies explaining morphological diversity among small, largely microscopic, euakaryotic organisms are few. Despite their small size, protists are extremely disparate with respect to their physiology and genetic makeup, and some groups display incredible morphological diversity. Among the most morphologically diversified protists are those organisms that construct structures using organic (e.g., glycoproteins and cellulose) or inorganic (e.g., silica, calcium, and strontium) materials.

*Synura* (Synurales, Chrysophyceae, and Stramenopiles) is a genus of freshwater algae consisting of species that form motile colonies, where each cell is covered with a highly organized series of overlapping siliceous scales (Starmach 1985, Leadbeater 1990, Siver 2015). The highly sculptured scales are positioned in spiral rows that can be traced from the flagellar end of the cell, around the body of the organism, to the posterior end. The great diversity
of scale ornamentation was initially revealed with transmission electron microscopy in the 1950s (Manton 1955, Petersen and Hansen 1956, Fott and Ludvik 1957) and later with scanning electron microscopy (Hartmann and Steinberg 1986, Siver 1987). Scales have a bilateral symmetry, range in length up to approximately 10 µm, and their designs are species-specific. All scales possess a basal plate perforated with pores, an upturned rim usually encircling half to two-thirds of the scale, and either a forward projecting spine or a raised elongated ridge positioned on the middle of the scale referred to as the median keel (Fig. 1). In addition to variability between species, environmental conditions and position on the cell body influence the morphology of *Synura* scales. *Synura* species are found under a wide range of ecological conditions (Kristiansen 1979, 2008), which, in turn, can result in subtle differences in scale morphology. Phenotypic plasticity of scale shape is also impacted by environmental stress (Sandgren et al. 1996, Saxby-Rouen et al. 1997). A number of studies have documented a decline in scale and spine length with increasing temperature (Martin-Wagenmann and Gutowski 1995, Gutowski 1996, Rezácová-Skaloudová et al. 2010), noting a shift toward more oval to elongate shapes (Pichrtová and Němcová 2011). Long-term cultivation under both lower temperature and light conditions can also result in a decrease in scale size (Němcová et al. 2010), and cells growing for extended periods in culture often begin to produce less silicified scales (Leadbeater 1986, Martin-Wagenmann and Gutowski 1995). Suboptimal pH conditions have also been shown to impact scale morphology (Gavrilova et al. 2005), including the degree of secondary ornamentation and width of the upturned rim (Němcová and Pichrtová 2012).

Another factor influencing scale shape and design is the position of the scale on the cell surface (Asmund 1968, Siver 1987). Scales can be divided into apical, body, and caudal scales depending on where they are found along the longitudinal axis of the cell (Kristiansen 1979). The majority of scales forming the scale coat are body scales, found over most of the length of the cell and consisting of similar features. Apical and caudal scales, found on the flagellar and posterior ends of the cell, respectively, are shaped differently from body scales in order to more effectively cover the changing shape of the cell. Often, apical scales are smaller, more rounded, and with longer spines, whereas caudal scales become more elongated with shorter spines (Kristiansen 1979, Siver 1987).

Despite small degrees of scale variability, the overall morphological structure of scales is a conservative, species-specific, and taxonomically relevant trait that has been successfully used to delineate between *Synura* taxa (Jo et al. 2016). Petersen and Hansen (1956) originally proposed to divide the genus into two sections, *Peterseniaceae* and *Spinosae*, according to the presence or absence of the median keel. Despite later taxonomic rearrangements within the genus, the presence of a keel versus a projecting spine has proven a most valuable character in understanding the phylogenetic arrangement of species (Skaloud et al. 2014, 2020, Siver et al. 2015, Jo et al. 2016). Péterfi and Momeu (1977) proposed a third section, *Lapponica*, to accommodate one species with very different scales that lacked both a keel and a spine. These authors also proposed the section *Synura* to replace section *Spinosae (=Uvellae*) and further divided section *Synura* into two series, *Synura* and *Splendidiae* (Péterfi and Momeu 1977). Additional phylogenetic relationships were suggested by Wee (1997), and Kristiansen and Preisig (2007) noted three sections within the genus, *Lapponica*, *Peterseniaceae*, and *Synura*. More recently, section *Lapponica* consisting solely of *Synura lapponica* was revealed to belong in the genus *Tessellaria* (Skaloud et al. 2013), which was later renamed as *Nootessella* (Jo et al. 2016). Multiple studies using a combination of ecological, physiological, morphological, and molecular approaches have further improved our understanding of relationships between species in *Synura* (Boo et al. 2010, Skaloud et al. 2012). Skaloud et al. (2013) confirmed section *Peterseniaceae (=Peterseniana)*

Fig. 1. The outline of siliceous *Synura* scales showing its major morphological features. Typical body scales with the spine (left) and with the median keel (right) are presented.
as clearly being monophyletic, but pointed out that the section containing species with projecting spines, namely *Uvellae* or *Synura*, was phylogenetically and morphologically diverse. As a result, Skaloud et al. (2013) proposed further classification of taxa with spines into the four sections, *Spinosa*, *Echinulaetae*, *Splendidae*, and *Uvellae*. In other recent phylogenetic works, *Synura* was divided into either two sections *Synura* and *Peterseniana* (Siver et al. 2015), or into three sections *Synura*, *Curtisipinae*, and *Peterseniana* (Jo et al. 2016). In summary, formation of a scale bearing a spine versus a keel, as originally proposed by Petersen and Hansen (1956), was an important event in the evolution of *Synura* that is a primary character used to distinguish between the currently 55 recognized taxa (Némcová et al. 2008, Jo et al. 2016, Pusztai et al. 2016, Siver and Lott 2016, Siver et al. 2018).

The goal of the current study was to uncover evolutionary patterns in scale design within *Synura* using a suite of morphological characters and fossil evidence in relation to a molecular-based phylogenetic framework. Specifically, our goal was to uncover overarching principles and trends shaping the evolution of scale morphology over geologic time. As part of our study, we have included new information on a key species, *Synura punctulosa*, a potential missing link in the evolution of the genus, and all known fossil species.

**MATERIAL AND METHODS**

*Collection, isolation, and cultivation.* *Synura punctulosa* was sampled in Chanty-Mansiysk (Хантый-Мансийский) region in south-western Siberia, Russia, during the beginning of June 2018. Specifically, the strains were collected in a flooded valley of the River Matkinskaya (Матkinsкая, 61°01'10.1'' N 68°05'02.4'' E). Standard measurements of abiotic factors of the sampling site were performed using a combined pH/conductivity meter (WTW 340i; WTW GmbH, Weilheim, Germany). At the time of collection, the temperature was 11.9°C, pH was 7.1, and the specific conductivity measured 65 µS·cm⁻¹. In the effort to establish monolcal algal cultures, individual colonies were isolated by micropipetting and transferred into a 96-well plate filled with WC liquid medium (Boenigk et al. 2006). The algae were cultivated at approximately 15°C under a constant illumination of 40 µmol·m⁻²·s⁻¹ (TLD 18W/33 fluorescent lamps, Philips, Amsterdam, the Netherlands). After 14 d, the cultures were inoculated into 50 mL Erlenmeyer flasks filled with the same medium (Pusztai et al. 2016).

*Sequencing and phylogenetic analyses.* For DNA isolation, 200 µL of cultured algae was harvested using centrifugation and frozen in PCR strips at −80°C. Then, 30 µL of InstaGene matrix (Bio-Rad Laboratories, Hercules, CA, USA) was added to the pellet. The samples were vortexed, incubated at 56°C for 30 min, and heated at 99°C for 8 min. Afterward, the supernatant was directly used as a PCR template.

We amplified six molecular markers (nu SSU rDNA, nu LSU rDNA, nu ITS rDNA, pt LSU rDNA, pt rbcL, and pt *psaK*) to infer a robust phylogenetic analysis, using the following primers. The nu SSU rDNA gene was amplified with the primers 18S_F and 18S_R (Katana et al. 2001), and the nu LSU rDNA using the combination of primers 28S_25F, 28S_2812R (Jo et al. 2011), and 28S_1435R (Pusztai et al. 2016). The pt LSU rDNA was amplified with primers 23S_Syn_AF and 23S_Syn_928R (Skaloud et al. 2020). The amplification of nu ITS rDNA was performed using newly designed primers *Chryso* _ITS_F (5'–ATC ATT TAG AGG GTG A-3') and *Chryso* _ITS_R (5'– GCT TCA CTC GGC GTT ACT-3'). The pt *rbcL* gene was amplified using primers *rbcL_R3* (Jo et al. 2011) and *rbcL_chrys_F2* (Skaloudová and Skaloud 2013). Finally, the pt *psaK* gene was amplified using the primers *psaA1390F* and *psaA1760R* (Jo et al. 2011).

All PCRs were prepared in a 10 µL volume consisting of 6.5 µL H₂O, 2 µL buffer, 0.2 µL of each forward and reverse primers, 0.1 µL MyTaq polymerase, and 1 µL DNA template. The PCR products were quantified on 0.8% agarose gel stained with ethidium bromide and purified using MagJET Magnetic Bead-based Nucleic Acid Purification (Thermo-Fisher Scientific, Waltham, MA, USA). The purified amplification products were sequenced in Macrogen Europe (Amsterdam, Netherlands).

New sequences were manually checked using SeqAssem vs. 9 (Hepperle 2004), added to the alignment published by Jo et al. (2016), and supplemented by several sequences deposited in GenBank database. Accession numbers of all analyzed sequences are listed in Supplementary information (Table S1 in the Supporting Information). Multiple alignment was built for the following analyses using MEGA5 (Tamura et al. 2011), including 64 strains of *Synurales* and two strains of *Neotessella* that were used as the outgroup. The sequences were aligned using MAFFFT version 7 under the Q-INSI strategy (Katoh et al. 2019) with the only exception of nu ITS2 rDNA, which was aligned and built according to its secondary structure. The sequences of nu ITS2 rDNA were modeled with the ITS2 Database V (Ankenbrand et al. 2015) to assure its homology and again manually checked for obvious aligning errors. DNA alignments are freely available on Mendeley Data: [http://dx.doi.org/10.17632/rh2ztg7pz.1](http://dx.doi.org/10.17632/rh2ztg7pz.1).

Suitable partition-specific substitution models were selected using the Bayesian information criterion (BIC) implemented in jModelTest 2.1.10 (Darriba et al. 2012). The following models with the lowest BIC scores were selected: (1) HKY + F for nu SSU rDNA and nu ITS 1 rDNA; (2) GTR + I+F for nu LSU rDNA, pt LSU rDNA, and nu ITS2 rDNA; (3) GTR + I for the first and third codon position of the pt *rbcL* and pt *psaK* genes, respectively, (5) JC + I for the second and (4) GTR + G for third codon position of the pt *rbcL* gene, (5) GTR + I for the first and (6) HKY + I for the second codon position of the pt *psaK* gene, and (7) SYM + G for nu 5.8S rDNA. Each part of the alignment was checked and trimmed with Gblocks software (Castresana 2000). The loci of nu SSU rDNA, nu LSU rDNA, pt LSU rDNA, nu ITS1 rDNA, nu 5.8S rDNA, nu ITS2 rDNA, pt rbcL, and pt *psaK* were concatenated, yielding a robust alignment of 9,680 bases.

The phylogenetic tree was first inferred using RaxML BlackBox with 250 replicates under the GTR Gamma + model with partitions (Stamatakis 2014) and rapid bootstrapping as implemented in the CIPRES Science Gateway (Miller et al. 2010). Second, we repeated the phylogenetic analysis with Bayesian inference (BI) using MrBayes version 3.2.1 (Ronquist et al. 2012). Three independent Markov Chain Monte Carlo (MCMC) chains were run for 3 × 100 million generations, sampling every 1,000 generation after 25% burn-in and checked for stationarity and convergence of independent chains. The resulting tree was then compared with the tree based on Bayesian framework in BEAST v1.10.4 (Suchard et al. 2018) used for final phylogenetic analyses. Lognormal relaxed clock models were applied for the partitions and a birth–death diversification process
was selected as a prior on the distribution of node heights. For temporal calibration of the phylogeny, we used time constraints based on well-preserved fossil scales found in geological deposits in Northern Canada, the Giraffe (Siver et al. 2015) and Wombat (Siver et al. 2013a) cores, as follows: (i) the lineage consisting of Synura uwell and S. splendida (Giraffe core), (ii) the lineage of S. curtispina and S. longitubularis (Giraffe core), and (iii) the stem of all Petersenianae taxa including S. macrocantha (Wombat core). The splits were adjusted on an offset of either 48 (Giraffe core) or 83 Ma (Wombat core) with the mean of 80 and a standard deviation of 6.0, which represent the minimal estimated age of fossils. The MCMC analyses were run for 5 × 100 million generations, sampling every 500,000 generation after 1.5 million generations removed as a burn-in. We checked the parameter-estimated convergence with Tracer v1.7.1. (Ram-lion generations removed as a burn-in. We checked the parameter-estimated convergence with Tracer v1.7.1. (Ram-baut et al. 2018) and then constructed the final chronogram with age estimation for all nodes. Trees were visualized using FigTree ver.1.4.2. (Rambaut et al. 2016).

Morphological investigation of silica scales. A total of 30 Synura species that are well defined using both molecular and morphological characters were used for the morphologi-cal study, including Synura americana, S. asmunidae, S.bjorkii, S.borealis, S.conopea, S. curtispina, S. echinulata, S. glabra, S. heteropora, S. hibernica, S. kristiansenii S. lanceolata, S. laticarinia, S. leptorhabda, S. longitubularis, S. macrocantha, S. macropora, S. mammillosa, S. mollissipa, S. multidentata, S. petersonii, S. punctulosa, S. soroconeae, S. sphagnicola, S. spinosa, S. splen-dida, S. sungminbooi, S. synuroidea, S. truttae, and S. uwell (Fig. 2). Images of scales from 29 of the Synura species (all except S. punctulosa) were taken largely from our personal database or from records contained in the chrysophytes.eu database (Skaloud et al. 2013) that included verified species identifications. For most species, we used images of actively growing specimens from the same cultures used for molecular analyses. Cultures and field collections of Synura punctu-losa were initially examined with an Olympus BX 51 light microscope equipped with Nomarski interference contrast, and then, the scales were characterized using images taken with a JEOl 1011 transmission electron microscope (TEM) equipped with a Veleta CCD camera and operated with Olym-pus Soft Imaging Solution Software GmbH (Mienster, Germany). Samples of S. punctulosa were prepared for observation with TEM by adding several drops from actively growing cultures onto Formvar-coated copper grids, allowed to dry, gently washed with distilled water, and finally redried.

We analyzed 44 morphological traits for all 30 species as defined in Skaloud et al. (2014), forty of which are quantita-tive (Table S2 in the Supporting Information). In an effort to more fully cover all major morphotypes, we also included scales from fossil specimens of Synura macracantha, S. nygaar-di, S. recurvata, and S. cronbergiae uncovered from the Giraffe Pipe and Wombat fossil sites in Northern Canada (Siver and Wolfe 2005, Siver et al. 2013a,b). For consistency across spe-cies, we selected only body scales for all morphometric analy-ses. All measurements were obtained using ImageJ version 1.46r (Rasband 1997) on a minimum of ten scales per spe-cies, and all traits were analyzed only if the homology derived from the similar biogenesis was reliable. We included three measurements of mean pore size depending on their position on the scale: (1) base plate pores excluding those under the keel and the hexagonal meshwork pattern (labeled as base plate pores on Fig. 1); (2) base plate pores within the hexagonal meshwork pattern (Fig. 1); (3) pores on the base of the keel (labeled as keel pores on Fig. 1). In addition to width, the length of the keel was divided into (i) that portion attached to the base plate; (ii) an estimate of the length of the portion of the keel projecting beyond the base hole; and (iii) the total length (Fig. 1). A few morphological characters, such as the unique anastomosing ribs of S. punctulosa or the ribs under the upturned rim on scales of S. uwell, were not included in the analyses.

Tracing the morphotype evolution of silica scales. Morphological features of Synura scales used to investigate ancestral charac-ter states were selected using a combination of scale structure (Fig. 1) and the Synuralean phylogeny, similar to the method used by Cerntnerow et al. (2019). A Spearman correlation matrix of all morphological characters was utilized to help select a subset of characters to use in the analysis. Highly cor-related characters were identified and removed, and we excluded morphological characters that were not evaluated for at least ten species. All analyses were performed using R v.3.6.1 (R Core Team 2019), and the Bayesian tree trimmed to 30 Synura taxa with associated morphological data.

The dataset of non-correlated morphological characters was analyzed by NMDS (non-metric multidimensional scaling) indirect analyses, using the Gower dissimilarity matrix. In addition to the 30 extant Synura species, measurements of fossil specimens of S. macracantha, S.nygaardii, S. cronbergiae, and S. recurvata were added to the dataset. We projected phylogenetic relationships among Synura species into the ordina-tion space to construct a phylomorphospace plot, using the Phytools package in R (Revell 2012). Ancestral states of standard-ized average values of morphological features were reconstructed using the densityMap, make.simmap, and con-Map functions in the Phytools package (Revell 2012), and assuming a Brownian motion model.

RESULTS

Phylogenetic analyses. A time-calibrated phyloge-netic tree (Fig. 3) dated the origin of the genus Synura in the Early Cretaceous, about 145 mya. Later in the Cretaceous, the genus radiated into the three sections recognized today as Synura, Curtispinae, and Petersenianae. The section Synura, consisting solely of the spine-bearing species, S. uwell, separated from the rest of the Synura species about 111 [100–126] mya. The sections Curtispinae and Petersenianae split slightly later at about 107 [96–120] mya. Species in section Curtispinae retained scales bearing spines, while taxa in section Petersenianae developed scales with a keel. Synura punctulosa, a species originally described by Balonov (1976), is an ancient lineage most closely aligned with S. splendida that originated approximately 79 [53–100] mya, prior to the radiation of the most extant Synura species.

The scale architectural principles. Correlations between all combinations of morphological traits (Table S3 in the Supporting Information) were used to initially examine constraints of scale forma-tion and to reveal principles of scale structure. Scale characters related to area, perimeter, and length were highly correlated, and the length of the upturned rim was also significantly related to scale perimeter and area. Spine length was related to the length, perimeter, and area of the scale and to the length of the upturned rim, while spine width was clearly linked to scale width. The total length of the keel was additionally correlated with scale perime-ter, area and length, the length of the upturned rim, and to the length of the portion of the keel
Fig. 2. The scales of Synura species evaluated in this study: (A) S. uvella, (B) S. splendida, (C) S. punctulosa, (D) S. multidentata, (E) S. leptorrhoda, (F) S. mammillata, (G) S. echinulata, (H) S. synuroidea, (I) S. sphagnicola, (J) S. sparsa, (K) S. mollispina, (L) S. longitubularis, (M) S. curtispina, (N) S. macracantha, (O) S. kristiansenii, (P) S. bjoerkii, (Q) S. asmundiae, (R) S. glabra, (S) S. hibernica, (T) S. lanceolata, (U) S. heterpora, (V) S. truttae, (W) S. sungminbooi, (X) S. soroconopea, (Y) S. conopea, (Z) S. laticarina, (AA) S. borealis, (AB) S. petersenii, (AC) S. macropora, and (AD) S. americana. The scale bars represent 1 μm.
FIG. 3. Multigene time-calibrated phylogenetic tree of the genus Synura. Newly molecularly characterized S. punctulosa is shown in bold. Values on the tree branches indicate statistical support; posterior node probability inferred with MrBayes (left), BEAST (middle), and maximum likelihood bootstrap (right). Asterisks mark the branches with the highest statistical support (1.00/1.00/100). Geologic time axis is presented in millions of years (Mya). The error bars at the tree nodes represent 95% confidence intervals longer than 2.5 Mya. Landmarks of the Synura scale morphology are shown along appropriate branches.
attached to the scale. Although keel width was variable, keel area was related to scale area, and the length of the keel tip was dependent on the scale width. Finally, the number of struts connected to the keel was related to the scale roundness.

**Trends in silica scale evolution.** We combined the NMDS ordination plot of morphological traits with phylogenetic structure to initially investigate trends of scale structure over geologic time within *Synura* (Fig. 4). The genus was clearly divided into two groups, based on the formation of scales with either a projecting spine or a keel, which supports splitting the genus into sections *Petersenianae* and *Synura + Curtispinae*. Fossil specimens of *S. macracantha*, *S. nygaardii*, *S. cronbergiae*, and *S. recurvata* further supported this primary split, but their positions within the ordination plot indicate slight shifts in morphology relative to their closest related modern counterparts, most notably traits related to scale, spine, and keel size (Fig. 4, A and C). In addition, scale morphologies in the sections *Synura* and *Curtispinae* were more variable than those in the section *Petersenianae* along the first two NMDS axes.

To analyze morphological variation in more detail, the NMDS ordination plots were constructed separately for section *Petersenianae* and for sections *Synura + Curtispinae* using all species included in the study (Fig. 5, A and B). In general, closely related species based on molecular data also had similar morphology. For example, scale morphology for closely related species *S. mammillosa*, *S. echinulata*, and *S. leptorrhagda*, and for *S. spinosa* and *S. mollispina*, overlap in the plot. It is important to remember that the projected morphospaces show scale variability for a limited number of strains and that scale morphospaces are likely larger than indicated by the analyses.

**FIG. 4.** Phylomorphospace plots representing the projection of phylogenetic relationships among selected *Synura* species onto the ordination diagram (NMDS) based on 32 morphological characters of siliceous body scales. The plots are displayed for NMDS 1 on NMDS 2 (A) and NMDS 1 on NMDS 3 (B). On both plots, species are differentiated according to their section membership. The fossil scales of *S. macracantha*, *S. nygaardii*, *S. recurvata*, and *S. cronbergiae* with unknown phylogenetic classification are added into the ordination diagram. The morphological variables strongly correlated with NMDS axes ($R^2 > 0.7$) are shown in (C) and (D), corresponding to plots (A) and (B), respectively.
In order to trace changes in scale morphology over time, non-correlated morphological traits (Table S2) were mapped directly onto the phylogenetic tree (Figs. 6, 7, S1 in the Supporting Information). Some morphological characters, including formation of the keel (section Petersenianae) and the labyrinth pattern found in Synura leptorrhabda, S. mammillosa, S. echinulata (Figs. 3, S1, A and B), clearly evolved once. Other features, such as the meshwork pattern in S. spinosa, S. mollispina, S. longitubularis, S. curtispina, and S. uvella (Figs. 3, S1C) appeared multiple times in the evolution of the genus.

Other trends in the shapes and the sizes of specific structures were noted. Scale roundness and circularity were greatest in more basal lineages, and especially declined in recent crown lineages within section Petersenianae (Fig. 6, A and D). The width of the upturned rim also tended to decline in recent lineages within section Petersenianae (Figs. 6F, S1E). Interestingly, the longest scales were produced by some of the more ancestral species inferred at the base of both the Petersenianae and Synura + Curtispinae sections, as noted in the lineages represented by S. macracantha, S. uvella, and S. splendida, respectively (Fig. 6C). In addition, the upturned rim encircled a larger percentage of the scale perimeter in the ancestral section Curtispinae lineage and gradually less of the perimeter during evolution of most taxa within section Petersenianae (Fig. 6, E and F). Larger base plate holes evolved in species with a keel (Fig. 7C), while species with spine-bearing scales tended to evolve larger base plate pores than those found on scales with a keel (Fig. 7A). Synura macracantha, at the base of section Petersenianae, has the longest keel with the highest number of attached ribs (Fig. 7D). The length of the diverticulum forming the spine or the keel was found to be quite variable (Fig. 7E), while a wider width of the diverticulum clearly evolved within species with a keel (Fig. 7F).

**DISCUSSION**

*Phylogenetic assessment of the genus Synura.* Since the 18th century, organisms with similar morphologies have often been considered to be evolutionary closely related, descendants of a common ancestor. Such is the case for Synura, as the phylogenetic relationships based on molecular analyses are mirrored in the morphological similarity of their silica scales (Fig. 3). Synura is divided into two dominant groups based on species possessing scales with either a median keel or a forward projecting spine, a character trait well supported by molecular data. Such a morphological distinction supports the division of the genus into two sections Petersenianae and Spino-sae, originally proposed by Petersen and Hansen (1956), and later revised as section Petersenianae and section Synura by Siver et al. (2015). More recently, Jo et al. (2016) proposed a third section, splitting section Synura into sections Synura and Curtispinae. Under this framework, section Synura contains S. uvella, and section Curtispinae all other species with spine-bearing scales. In our study, additional clusters of species with similar scale characters can be traced within the phylomorphospace, indicating an even finer separation of taxa bearing scales with spines as proposed by Skaloud et al. (2013). In a few cases, it has been more difficult to separate closely related taxa using scale morphology, for example, among a number of Petersenianae species (Fig. 5A) or closely related strains of S. mammillosa, S. echinulata, and S. leptorrhabda (Fig. 5B). This problem has been
aided by adding some morphological characters not previously used in delineating between species (Skaloud et al. 2014).

Combining the phylogenetic relationships of the species with their morphological traits provides further insights into the evolution of the genus Synura. Wee (1997) depicted a phylogenetic tree using scale development and morphological traits. He suggested that the most ancestral scales were those with bilateral symmetry and lacking secondary structures (such as S. sphagnicola and S. splendida). In addition, according to Wee (1997) and Lavau et al. (1997), S. uvella should be closely related to S. spinosa and S. curtispina. However, molecular studies presented.
by Siver et al. (2015), Jo et al. (2016), and confirmed in our study, indicate that *S. uvella* represents the most ancestral species in the genus. In addition, we found *S. sphagnicola* to be related to species possessing a meshwork on their scales (*S. mollispina*, *S. spinosa*, *S. curtispina*, and *S. longitubularis*). Consequently, we consider the simple appearance of a *S. sphagnicola* scale as an evolutionary simplification of a more complex scale that possessed a meshwork type of ornamentation.

Our multigene phylogenetic tree (Fig. 3) is highly congruent with the analyses of Siver et al. (2015)
and Jo et al. (2016), with a few minor modifications. The phylogenetic position of *Synura punctulosa* with its morphologically unique scales was previously unknown and thought to possibly represent a separate section of the genus (Skaloud et al. 2013). Although scales of *S. punctulosa* possess a spine, they have exceptionally small base plate perforations, less rounded scales than other spine-bearing species, and possess a unique pattern of anatomizing ribs across the scale surface. Our molecular analyses place *S. punctulosa* together with *S. splendida* at the base of section *Curtispinae* as formerly proposed by Jo et al. (2016), and separating from other members of section *Curtispinae*, as well as from each other, in the Upper Cretaceous. Scales of *Synura punctulosa* and *S. splendida* have a simple appearance of the base plate with small perforations, and a longer than average upturned rim. We hypothesize that in the course of evolution, *S. splendida* invested into enlarging its scales and the spine, while *S. punctulosa* strengthened its scales with ribs.

**Scale architectural principles.** Scales are fit into a highly organized pattern that effectively covers the changing contour of the cell. They are positioned in spiral overlapping rows that are oriented at an angle with respect to the longitudinal axis of the cell. Each spiral row contains a few apical scales, mostly body scales, and ends with caudal scales. Although the largest variations in scale structure are with the apical and caudal scales found on the ends of the cell, only slight and less pronounced changes occur among the body scales that surround the majority of the cell. Body scales are slightly concave and more or less bilaterally symmetric (Leadbeater 1986). The left side of the upturned rim is often slightly longer than the right side, perhaps to accommodate the titling of the spiral rows around the cell.

Each scale is formed endogenously within a specialized vesicle, the silica deposition vesicle (SDV). The process by which the SDV is involved in scale biogenesis was demonstrated first by Greenwood (1967), followed by a number of works each adding to our understanding of scale development (Schnepl and Deichgräber 1969, McGrory and Leadbeater 1981, Mignot and Brugerolle 1982, Brugerolle and Bricheux 1984, Leadbeater 1984, Sandgren et al. 1996). The collection of studies demonstrated that during scale biogenesis, the SDV originates near the anterior and outer surface of one plastid, and becomes attached to the periplastid endoplasmic reticulum (PER) with a series of microtubules. The SDV is then molded into the shape of a scale as it is moved down and along the outer surface of the plastid by the microtubules. Once molded into final shape, amorphous silica is deposited and polymerized forming the finished scale. During the molding process, an invagination of the SDV referred to as the cytoplasmic diverticulum is made, marking the position of the base pore and used to form either the median keel or the projecting spine. If the invagination bends forward and away from the body of the SDV, it will form a spine. In contrast, if it bends backwards and comes to rest on the SDV it will form a keel.

The size of the base plate hole is related to the size, and most likely the width, of the spine or keel that it forms (Fig. 7C). Because the keel comes to rest on and fuses to the base plate, it strengthens the scale which, in turn, decreases potential breakage. The proportions of the keel tip are highly variable among section *Peterseninanae* species (Fig. 2, N–AD). There are taxa with either very short (*Synura truttae*, *S. lanceolata*, *S. heteropora*) or very wide keel tips (*S. bjorkii*, *S. borealis*, *S. asmundiae*, *S. kristianseni*). However, keel tip length is significantly correlated with scale width, suggesting an architectural constraint in scale construction. Although the reason remains unclear, pores associated with the keel are always larger than those on the base plate.

Siliceous ribs on the scale surface can also serve to strengthen the scale. The number of ribs, or struts, that radiate from the keel onto the base plate was negatively correlated with the roundness of the scale. Thus, species with less rounded scales, such as *Synura macracantha*, possessed a high number of struts (Fig. 7D). The struts that radiate from the keel are likely produced from extensions of the portion of the diverticulum that comes to rest on the base plate. It is interesting that spine-bearing taxa may also form short ribs emanating from the base of the spine near the base hole where the diverticulum is connected to the scale. These short ribs, especially visible on *S. splendida*, were pointed out by Nicholls and Gerrath (1985), and later proposed to be homologous to the struts radiating from the keel (Wee 1997). Other ribs, such as those forming a labyrinth pattern (*e.g.*, *S. mamillillosa*, *S. echinulata*, and *S. leptorrhabda*), a hexagonal meshwork (*e.g.*, *S. mollispina*, *S. curtispina*, and *S. spinosa*), those under the upturned rim of *S. uvella*, as well as the ribs on *S. punctulosa*, are probably directly formed by the main portion of the SDV producing the base plate.

For many spine-bearing species, the anterior portion of the scale contains secondary structure, whereas the posterior part within the confines of the upturned rim may not. The upturned rim is associated with a portion of the SDV that bends up and over the base plate. Moreover, the outer surface of the rim is coated with an organic adhesive substance which aids to cement the scales in place on the cell surface (Leadbeater 1986, 1990, Beech et al. 1990). The upturned rim may serve to precisely align the scales in a similar manner to the V-rib in *Mallomonas* scales (Siver and Glew 1990). The length of the upturned rim has also been shown to vary with the degree of secondary structure found on the anterior portion of the scale (Gutowski 1996, Némcová et al. 2010). For example, Némcová et al. (2010) reported an increase in the percentage of
the scale margin covered by the upturned rim in *Synura echinulata* concurrent with a reduction in the labyrinth ribbing pattern when grown under increasing temperatures. This suggests that the upturned rim may compensate for a reduction in secondary structures, perhaps to help maintain scale strength. Because the length of the upturned rim was correlated with the area and perimeter of the scale, the length of the spine, and the length of the keel, we further conclude that it represents an evolutionary conservative feature. It is interesting to note that silica scale-bearing chrysophytes may have an advantage over other groups of eukaryotes that form cell coverings out of silica (e.g., diatoms) in that even though Si(OH)₄ is needed for scale production, it is not essential for cell growth (Sandgren et al. 1996). In culture studies where silica was depleted, cells initially formed less silicified scales, then aberrant formed scales, and eventually cells lacking scales altogether (Klaveness and Guillard 1975, Sandgren et al. 1996).

**Evolution trends of Synura scale case.** Intracellular deposition of silica is common in eukaryotes, occurring in cercozoa, alveolates, amoeboboa, rhizaria, archaeoplastida, and stramenopiles (Marron et al. 2016). Many hypotheses have been proposed to explain the potential benefits of a siliceous cell covering. First, it may relate to conserving resources since it requires more energy to build a polysaccharide cell wall than a silica scale case (Raven 1983). Second, the scales are evenly silicified and regularly perforated, which may increase light diffraction into the cell interior and simultaneously increase photosynthetic efficiency (De Tommasi et al. 2010). Third, the siliceous scales may contribute to reducing dangerous UV-B and UV-A radiation reaching the cell interior (Beardall and Raven 2004, De Stefano et al. 2007, Bismuto et al. 2008). Fourth, the silica scale case may serve to protect against grazing and parasitism, although a wide range of organisms are able to overcome such a barrier (Raven and Waite 2004, Spillane 2016, Panić et al. 2019). For example, some viruses are capable of penetrating through pores about 0.1 µm wide, and others may use an enzymatic digestion of the organic cement to disrupt the scale case (Brussaard 2004, Metreveli et al. 2014, Herringer et al. 2019). The reduction in scale pore size observed over the evolution of the genus *Synura* (Figs. 7, A and B, S1F), may have been a response to improving the protective barrier against viruses and parasites.

The reduction in pore size could also be a function of the concurrent reduction in cell size reduction observed over geologic time. The oldest lineages of both spine and keel-bearing *Synura* species (e.g., *Synura splendidia* and *S. macracantha*) have bigger scales than more recently diversified lineages (Fig. 6, A and B). Indeed, scales of the extinct species *S. cronbergiae* and fossil specimens of *S. macracantha* uncovered from the Giraffe core possessed large scales compared with modern taxa (Siver et al. 2013a). Perhaps, cells reduced the mass of heavier scales by producing bigger pores, as a method to reduce the energy needed by the cell to remain near the surface and to help prevent it from sinking out of the euphotic zone. Smaller cells with smaller scales were also thought to be related to the global warm temperatures found during the Paleocene–Eocene Epochs (Siver et al. 2015). This hypothesis was based on the general temperature-size rule proposed for protists (Atkinson et al. 2003), as well as on short-term studies of chrysophytes in culture (Pichrtová and Němcová 2011).

In addition to pore size, other traits that are associated with the scale case can be traced during the evolution of the genus. Scales have become less round, partially due to the formation of the keel for species in section *Peterseniinae* (Fig. 6, A and D). More elongated scales might fit easier around elongated cells with reduced volume to surface ratio, which are selected by the competition for nutrients (Karp-Boss and Boss 2016). Based on previous phylogenetic analyses (e.g., Siver et al 2015, Skaloud et al. 2020), and confirmed in our study, evolution of the median keel occurred only once. Interestingly, despite the fact that the keel on fossil *Synura macracantha* scales from Eocene deposits has a series of perforations along the sides, it remained well secured to the scale surface (Siver 2013). In addition, *S. macracantha* has the longest known keel with the highest number of struts among *Synura* taxa (Fig. 2N). Keel length gradually shortened over geologic time (Fig. 7E), concurrent with a decline in the number of struts associated with the keel (Fig. 7D) and scale size, possibly as a consequence of better fitting into the cell covering as cell size decreased.

Although spines are proposed to be advantageous as a defense against predators (van Tol et al. 2012, Panić and Kiørboe 2018) or to possibly reduce sinking in the water column (Laurenceau-Cornec et al. 2015, Walker 2019), such benefits seem of limited value for *Synura* species, given they are actively swimming colonial flagellates where the spines do not effectively increase the size of the colony. We propose that the benefit of a median keel for reinforcing scale strength is more beneficial for *Synura* than a protruding spine. Indeed, the keel-bearing *Synura* species are often more abundant in water bodies than the spine-bearing taxa (Kiss and Kristiansen 1994, Pichrtová et al. 2007, Kynclová et al. 2010). In addition, there has been a greater degree of species diversification within section *Peterseniinae* than section *Synura* since the late Neogene (Skaloud et al. 2020).

The impact of some morphological scale features on cell fitness is not always clear and may in fact represent a consequence of past evolutionary processes. For example, the upturned rim is longest in the more ancestral *Synura* species, including *S.
uvella, S. punctulosa, and S. splendida (in this species it even encircles the whole scale; Figs. 6E, S1D), as well as in the extinct species S. cronbergiae and S. recurvata. Interestingly, the rim also encircles the whole scale of both known species of Neotessella, a most ancestral genus of the Synurales (Siver et al. 2015). The precise hexagonal meshwork secondary structure commonly found on modern scales of S. spinosa, S. mollisperma, S. longitubularis, S. curtipina, and S. uvella (Fig. 3C), as well as on scales of fossil specimens of S. cronbergiae, S. recurvata, S. nygaardii, and S. recurvata (Siver and Wolfe 2005, Siver et al. 2013a), has remained a stable feature over time. This feature would also strengthen the scale. In fact, S. cronbergiae, which has scales mostly covered with a hexagonal meshwork, is imagined to be a possible prototype of the last common ancestor of both spine and keel-bearing Synura.

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