First partial skeleton of *Delphinornis larseni* Wiman, 1905, a slender-footed penguin from the Eocene of Antarctic Peninsula

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ABSTRACT

The oldest fossil record of Antarctic penguins comes from Seymour Island (Antarctic Peninsula) and dates to the Paleocene and Eocene. The Paleocene bones are extremely rare, whereas specimens from the latter epoch are numerous. Despite the recent discoveries of incomplete skeletons assignable to the giant penguins from the Eocene of Antarctic Peninsula, the reliable systematics of their smaller contemporaneous relatives, known from isolated bones, have remained dependent on the tarsometatarsus. Here, new data on the skeleton of *Delphinornis larseni*, the most abundant among non-giant Eocene penguins, are reported. The specimen, collected from the Submeseta Formation on Seymour Island, comprises the incomplete pelvis and numerous bones from the hind-limb skeleton, including a well-preserved (diagnostic) tarsometatarsus. The acetabular foramen is, like in larger fossil penguins, clearly smaller than the elongated ilioischiald foramen. The area of the latter opening, not occupied by the connective-tissue sheet, supposedly accounted for one-third of the foramen. We propose that the ischiadic artery was, unlike in present-day penguins, the main blood vessel supplying most of the hind limb. The proximal fovea of the femoral head is uniquely preserved, revealing an osteological aspect of the bone-ligament interface. We surmise that the individual was similar, in terms of body size, to extant *Pygoscelis papua*, but was characterized by more elongate feet. In our opinion, it was probably a young bird, up to several years old.
INTRODUCTION

Present-day penguins (Aves, Sphenisciformes) are the most distinct among orders of birds. All of them are flightless, wing-propelled marine divers that can be found across the Southern Hemisphere (e.g., Williams, 1995). Such a characteristic held true also for their Paleogene (66-23 Ma) antecedents, which is unambiguously documented by their fossil record (Slack et al., 2006; Jadwiszczak, 2009; Ksepka and Ando, 2011; Mayr et al., 2018). These highly derived seabirds presumably originated in the Cretaceous (145-66 Ma) (e.g., Slack et al., 2006) or the Paleocene (66-56 Ma) (e.g., Jarvis et al., 2014). The fossil record of Eocene (56-34 Ma) penguins is by far more abundant. During this epoch, Sphenisciformes became widespread, expanding their ranges to lower latitudes, ultimately reaching the equatorial region of western South America (Clarke et al., 2017a) (Figure 1.2, 1.4). Apart from penguin bones, this locality has produced remains of teleost fishes, a single *Carcharocles* shark tooth, a rostrum of a “pseudotooth” bird (Pelagornithidae), and whale bones (Wiman, 1905; Bargo and Reguero, 1998; Kriwet et al., 2016). The plateau exposes the middle part of the Submeseta Formation (Submeseta II = level 38) according to Montes et al. (2013), or Telm 7 of the La Meseta Formation according to Sadler (1988) (Figure 1.2-4). Most authors regard this part of the depositional succession on Seymour Island to be late Eocene (Pria-
bonian) in age (e.g., Marenssi, 2006; Douglas et al., 2014; Buono et al., 2016).

The fossil material is housed in the paleozoological collections of the Swedish Museum of Natural History, Stockholm. It was partially (manually) purged of matrix remnants in 2017. The comparative material studied directly comes from the above-mentioned institution (fossils), the University of Białystok, Poland (fossils and a small number of recent bones), the Natural History Museum at Tring, UK (a vast collection of recent bones), and the Natural History Museum, London, UK (fossils). TM and PJ have been curators of Swedish and Polish collections (respectively); bones from Lon-

**FIGURE 1.** Location of Seymour Island (Antarctica) and stratigraphy of its NE part. 1. Map of Antarctica showing the position of Seymour Island. 2. Map of Seymour Island showing the location of the Eocene La Meseta and Submeseta formations, and position of the locality DPV 13/84 with sampling site. 3. Simplified stratigraphic column of the Submeseta Formation with an arrow pointing at the sampling horizon, based on data from Montes et al. (2013). 4. View of the sampling site location from under the plateau of the Weddell Formation; photograph was taken in 2011 by TM. Abbreviations: cg, conglomerates and sandstones; t, *Turritella* (a gastropod).
London and Tring were studied by PJ in 2011 and 2015, as part of the SYNTHESYS grants GB-TAF-987 and GB-TAF-4610 (respectively).

Photographs of penguin bones were taken using a digital single-lens reflex camera (20.9 megapixel image sensor, devoid of a low-pass filter) paired with 40 mm macro lens. Raw X-ray images of the new fossil material assignable to *D. larseni* were captured at 58 kV (kilovolts), 3.2 mAs (milliamperie-seconds), distance from the (anode) focus of 100 cm, without anti-scattering grid, and digitally pre-processed following the protocol for canine limb joints (CRYsTestView software; Alara Team, 2005-2013). This “mammalian” protocol turned out to be the most suitable in the case of studied penguin remains. The heat map of the acetabular socket was created from an X-ray by means of Fiji (a redistribution of ImageJ program; Schindelin et al., 2012), using the 3D Surface Plot module (Thermal LUT mode). The heat map for a portion of the left tarsometatarsus was generated from an X-ray, using Sante DICOM Viewer Free program in NIH palette mode (Santesoft Team, 2018).

Shape analyses of outlines of major hip bone foramina were conducted using the Momocs package (version 1.2.9; Bonhomme et al., 2014) for R environment (version 3.4.3; R Core Team, 2017). This is a relatively easy-to-use, well-designed toolkit, with implemented methods for analysing the landmark-poor configurations. Shapes of acetabular and ilioischiadic foramina belonging to representatives of eight present-day penguins (specimens from Bialystok and Tring) (Appendix 1), supplemented by those from the studied Eocene specimen, were extracted (black silhouettes, white background) from photographs taken by a single person (PJ; to control the viewing angle) and saved as jpg files (an acetabular-ilioischiadic pair per file) (Appendix 2.1). To ensure each configuration is treated by a program as a single shape, a thin black line connecting extreme points of foramina was drawn prior to file creation. In the case of NRM-PZ A.994, because of imperfect specimen preservation, such a pair was based on combined images from left and right hip bones.

Configurations for Paleocene and Oligocene penguins, used solely in a cladogram, were assessed based on published images (Slack et al., 2006, figure 1.A; Ksepka et al., 2012, figure 6.E-F).

The set of files was imported into the R/Momocs environment, converted into an Out-class object (a list of closed outlines) by means of Out function and inspected for errors using panel function (Appendix 2.2-3). To visualize the location of the Eocene specimen relative to the sampled extant-penguin morphospace, the contents of above object was aligned, centered, and scaled using package-supplied helper procedures, and plotted by means of stack function (Appendix 2.3).

Outline data were also used to compute elliptic Fourier transform (EFT) by means of efourier function, using 16 harmonics and other parameters set to their defaults. The number of harmonics (their sum approximates the outline) was based on a harmonic power analysis and visual inspection of reconstructed forms. The procedure resulted in normalized elliptic Fourier coefficients (outline-form descriptors; four per harmonic; Appendix 3) in a form of an OutCoe-class object. This normalized dataset was utilized, via PCA function, in Principal Component Analysis (PCA). It was also used, after reverse transformation from coefficients to shapes using mshapes to calculate and plot deformation grids between two (polarized) configurations. The latter step was performed by means of tps_grid function, and the choice of configurations was based on visual inspection of outlines and results of PCA (Appendix 4). For more details, see Appendix 2.3 as well as consult Bonhomme et al. (2014) and Momocs documentation.

The bubble chart for tarsometatarsal measurements was created using functionality of the ggplot2 package for R (Wickham, 2009). Measurements of comparative material (Appendix 1) were obtained both directly from specimens belonging to European collections (Bialystok, Stockholm and Tring) and from literature (Myrcha et al., 2002, table 1).

The anatomical nomenclature, in a form of English equivalents to Latin terms, follows Baumel and Witmer (1993) and Baumel and Raikow (1993). In some cases, in order to reflect peculiarities of penguin osteology, the terminology had to be adjusted. Body-size categories follow Williams (1995), and Jadwiszczak and Chapman (2011). Linear measurements were taken by means of a digital caliper and rounded to the nearest 0.1 mm.

**Institutional Abbreviations**

CM, Canterbury Museum, New Zealand; IB/P/B, University of Bialystok, Bialystok, Poland; MUSM, Natural History Museum of the National University of San Marcos, Lima, Peru; NHMUK, Natural History Museum, London, UK; NHMUK/T, Natural History Museum at Tring, UK; NRM-PZ, Museum of Natural History, Stockholm, Sweden; OU, Geology Museum, University of Otago, New Zealand.
SYSTEMATIC PALEONTOLOGY

Class AVEST Linnaeus, 1758
Order SPHENISCIFORMES Sharpe, 1891 (also sensu Clarke et al., 2003)
Family SPHENISCIIDAE Bonaparte, 1831 (but not sensu Clarke et al., 2003)
Genus DELPHINORNIS Wiman, 1905
  Delphinornis larseni Wiman, 1905
  Delphinornis gracilis Wiman, 1905
Figures 2.1-7, 2.15, 2.17-19, 3, 4.2-6, 4.8-17, 5.3-4, 5.6, 5.8-9, 5.13, 5.16, 6

Material. Incomplete pelvis and leg skeleton including femur, tibiotarsus, fibula, tarsometatarsus and phalanges (NRM-PZ A.994); both body sides represented, partly in matrix.

Taxonomic Remarks

Suprageneric position of NRM-PZ A.994. The hierarchical system of classification places all penguin species, both extinct and extant, in a single family, Spheniscidae, within a monotypic order - Sphenisciformes (Simpson, 1946, 1971; Myrcha et al., 2002; Jadwiszczak, 2006, 2009). Therefore, within this framework, D. larseni is a member of Spheniscidae. Clarke et al. (2003) introduced a new context to the penguin systematics, treating these terms as the names of clades. The term Spheniscidae was suggested for the crown group, and this approach can be found in many later papers (e.g., Clarke et al., 2007; Ksepka and Ando, 2011; Ksepka et al., 2012). All extant genera of Sphenisciformes as well as other crown penguins are Miocene in age or younger (e.g., Ksepka and Ando, 2011; Gavryushkina et al., 2017), whereas all fossils assignable to pre-Miocene genera (including Eocene Delphinornis) and a number of younger findings represent the stem group (e.g., Ksepka and Clarke, 2010). In our work, we do not consider the cladistic/phylogenetic view as opposed to the traditional rank-based classification, but rather as another (informative) layer.

Generic and specific classification of NRM-PZ A.994. The complete tarsometatarsus discussed here (Figure 2.1-7, 2.18) is a small specimen, within the size range of non-giant penguins (Figure 3). Its hypotarsus, unlike in similarly-sized Eocene Marambiornis Myrcha et al., 2002 and Mesetaornis Myrcha et al., 2002 from Antarctica, is devoid of a groove separating the intermediate hypotarsal crest, and the medial crest slopes towards the medial margin of the bone (Figure 2.5-6). Considering the position of the medial vascular foramen (plantar view) and medial hypotarsal crest relative to the tarsometatarsal longitudinal axis, the former is abaxial (unlike in said genera) (Figure 2.5). This set of traits is in line with the revised diagnosis for the genus Delphinornis enunciated by Myrcha et al. (2002).

The following combination of character states found in the bone in question is, according to Myrcha et al. (2002), diagnostic of Delphinornis larseni. The specimen is more elongated (elongation index of 2.76; a measure defined in Myrcha et al., 2002) than its counterparts in Delphinornis arctowskii (respective values do not exceed 2.5). It is also longer, in terms of absolute dimensions, than tarsometatarsi assignable to D. arctowskii and Delphinornis gracilis (Figure 3). The medial hypotarsal crest slopes quite gently (unlike in both congeners species), gradually decreasing its steepness towards the medial margin of the bone. This is accentuated by the abrupt, albeit short, vertical drop to the edge of the second metatarsal (Figure 2.6). All three trochleae are relatively stout (Figure 2.2., 2.7), and the intercotylar eminence is quite wide and rounded (moderately pronounced) in dorsal view (Figure 2.1-2, 2.6) (unlike in D. gracilis). The preservation of both tarsometatarsi described above precludes decided assessments regarding the distal vascular foramen, which is well developed in the type specimen (e.g., Wiman, 1905, figure II.2) and some other specimens assignable to this species (e.g., Figure 2.9; see also revised diagnosis in Myrcha et al., 2002). We can state, however, that it is present in NRM-PZ A.994.

Comparative Description

Synsacrum. The synsacral part of the pelvis is represented by two small fragments of the side wall of the canal of ankylosed vertebrae with conspicuous intervertebral foramina (Figure 4.3-6). Two best preserved foramina are indistinguishable, no separate openings for the dorsal and ventral roots of the spinal nerves are present. Such a condition can be observed in caudal, sacral and lumbar vertebral series of other synsacra assignable to Eocene Antarctic penguins (Figure 4.7; Jadwiszczak, 2006, figure 18.E, 2014, figure 1.C) and Oligocene Kairuku waitaki Ksepka et al., 2012 from New Zealand (Ksepka et al., 2012, figure 6.B). In lumbar part and lumbosacral transition, both types of openings can exist, and left and right side of the synsacrum may differ in this respect, even at the same level along the bone longitudinal axis (e.g., in IB/P/B-0149; Figure 4.7).

One of those foramina, belonging in a fragment connected through a piece of matrix with a cranial portion of the wing (ala) of the left ischium, is clearly larger and more rounded (Figure 4.5-6).
FIGURE 2 caption next page.
The other opening represents an element that diagenetically altered sediment associated with the body (corpus) of the left ischium and cranialmost pubis (accompanied by unrecognizable elements). It is elongated dorsosventrally and located between two protrusions of a supposedly inner surface of the wall (Figure 4.3-4).

**Hip bones.** The aforementioned two non-adjoining hip bone fragments contribute to partially preserved rims of three major openings: the acetabular, ilioischial, and obturator foramina (Figure 5.7). The first structure is best preserved, showing approximately two-thirds of a roughly circular margin bounded by all three coxal/hip bones (Figure 4.2-3). Its diameter, estimated along the main bone axis in medial view, is ca. 13.6 mm, i.e., 118% of the value obtained by us for an unnumbered specimen (IB/P/B) of extant *Pygoscelis papua* (Forster, 1781) (Figure 5.12), 64% of the value reported for the largest living penguin, *Aptenodytes forsteri* Gray,1844, and the Eocene individual from Tierra del Fuego, Argentina (Clarke et al., 2003, table 1), but merely 48% of that measured (by us) for the unassigned specimen (IB/P/B-0488; Figure 5.10-11) from the Eocene of Antarctica. The missing section corresponds to the preacetabular ilium, although a fragmentary iliac crest of the acetabular socket is retained. The acetabular foramen is “more or less regularly rounded in outline” in all birds (Kuznetsov and Sennikov, 2000, p. 441), including modern penguins (Zusi, 1975, figure 4.1; Stephan, 1979, figure 42). This is also evidenced for early Sphenisciformes from the Paleocene (Slack et al., 2006, figure 1A), Eocene (Clarke et al., 2003, figure 4B; Jadwiszczak, 2006, figure 4.G; Acosta Hospitaleche and Olivero, 2016, figure 4.A), whereas relevant smaller fossil specimens (e.g., IB/P/B-0211; Figure 5.5) are too incomplete to compare. The considered surface is also (relatively) wide in similarly-sized modern pygoscelid penguins (unnumbered specimens from IB/P/B; Figure 5.12). The antitrochanter (Figure 5.6) is well preserved, located expectedly caudodorsal to the lateral opening of the acetabulum, tongue-shaped (common for penguins), observed even in Paleocene *Waimanu manneringi* Jones et al., 2006 (in Slack et al., 2006; see Slack et al., 2006, figure 1A) and moderately inflected laterally. The inflection varies in modern

**FIGURE 2 (continued from previous page).** Partial foot skeleton of Eocene *Delphinornis larseni*, NRM-PZ A.994. 1. Right tarsometatarsus in dorsal view. 2. Left tarsometatarsus in dorsal view. 3. Same in medial view. 4. Same in lateral view. 5. Same in plantar view. 6. Same in proximal view. 7. Same in distal view. 8. Left tarsometatarsus of *D. larseni*, IB/P/B-0547 (Eocene, Seymour Island, Antarcctica), in lateroplantar view. 9. Left tarsometatarsus of *D. larseni*, IB/P/B-0062 (Eocene, Seymour Island, Antarcctica), in dorsal view. 10. Same in medial view. 11. Same in lateral view. 12. Same in plantar view. 13. Same in proximal view. 14. Same in distal view. 15. Reconstruction of right-foot skeleton of *D. larseni* (NRM-PZ A.994) in dorsal view. 16. Outline of left-foot skeleton of extant *Pygoscelis papua* (unnumbered specimen from IB/P/B) in same view. 17. Reconstruction of left-foot skeleton of *D. larseni* (NRM-PZ A.994) in same view. 18. Schematic view of left-foot skeleton of *D. larseni* (NRM-PZ A.994) in dorsal view, as preserved in matrix, superimposed over X-ray image of same specimen. 19. X-ray-based heat map of portion of above fossil indicating location of tarsometatarsal dvf. 20. Left tarsometatarsus of *Pygoscelis papua* (unnumbered specimen from IB/P/B) in dorsal view. 21. Same in plantar view. Abbreviations: d2-4, digit 2-4; df, dermal flange; dvf, distal vascular foramen; ft, flexor tubercle; hs, hypotarsal sulcus; ice, intercotylar eminence; icf, infracotylar fossa; ire, impressions of extensor retinaculum; ihc, intermediate hypotarsal crests; in, intertrochlear notch; itc, insertion of tibial cranial muscle; lc, lateral cotyle; lhc, lateral hypotarsal crest; lis, lateral intermetatarsal sulcus; lm, lateral margin; lt, lateral (hypotarsal) tuberosity; ltm, left tarsometatarsus; lvf, lateral vascular foramen; m2-4, metatarsals 2-4; mc, medial cotyle; mhc, medial hypotarsal crest; mis, medial intermetatarsal sulcus; mm, medial margin; mvf, medial vascular foramen; ns, neovascular sulcus; ota, origin surface for toe abductor; p2-4, phalanx 2-4; pu, ungual (terminal) phalanx; rtm, right tarsometatarsus; sd, supratrochlear depression; ta, tarsus; t2-4, trochlea 2-4. Asterisks denote comparative material. Scale bar equals 1 cm.
penguins (e.g., Stephan, 1979, figure 41), even within the genus (pers. obs., based on unnumbered modern-pygoscelid specimens from IB/P/B). Considering the antitrochanteric inflection in the above genus of medium-sized Sphenisciformes, NRM-PZ A.994 resembles *P. papua* most.

The ilioischiadic foramen is much larger than the acetabular opening, oval (which is clear, despite the incomplete iliac margin), with its major axis clearly longer than the minor one (30.4 mm vs. 14.1 mm). In this regard, the new specimen resembles other early Sphenisciformes, especially Paleocene *Waimanu manneringi* (Slack et al., 2006, figure 1.A) and unassigned Eocene IB/P/B-0488 (Figure 5.10-11) as well as even some extant volant aquatic birds, e.g., loons and albatrosses (Wilcox, 1952, figure 11; Klemm, 1969, figure 38; see also Ibáñez and Tambussi, 2012, figure 1). The opposite part of the spectrum includes, among others, Miocene *Madrynornis mirandus* Acosta Hospitaleche et al., 2007 (early crown penguin; Acosta Hospitaleche et al., 2007, figure 5.B) and modern penguins (e.g., Figure 5.7, 5.12; Zusi, 1975, figure 4.1; we discuss it later in our work). In Plotopteridae, extinct penguin-like seabirds from the Northern Hemisphere, the ilioischadiac foramen, albeit large, is clearly tapered cranially (unlike in penguins; Mayr and Goedert, 2018, figure 3.B-C, 3.F). In the majority of present-day birds, said opening is roughly round or short oval (Baumel and Witmer,
FIGURE 4 caption next page.
The major axis of the oval (which is common in birds; Baumel and Witmer, 1993) obturator foramen is ca. 10.8 mm long and devoid of a large portion of its ventral pubic margin. The ischiadic attachment site for the ischiopubic ligament is spindle-shaped (like in Eocene IB/P/B-0488, subcircular in pygoscelids; Figure 5.9, 5.11-12).

The body of the ischium is characteristically angular in medial view (Figure 5.4). The wider, medial surface of the cranial ischium and its virtually ventral counterpart meet at right angles and the resulting edge is parallel to the main axis of the hip bone. Such a sharply defined angularity appears to be not traceable in present-day penguins (Figure 5.12), but it is conspicuous in Eocene IB/P/B-0488 and IB/P/B-0627; Figure 5.10). The caudal extension of the ilioischial pillar, a bony reinforcing structure, is clearly visible, due to surface damage to adjacent fossilized tissue, along the postnatal ankylosis between two main coxal constituent parts (Figure 5.4). Both, the preserved postacetabular ilium and available fragments of the ischiadic wing appear to be rather delicately built, especially their off-axial surfaces. Their joint surface is smoothly convex medially. Retained fragments suggest relative elongation of the postacetabular pelvis. The length of the preserved ilioischial part of the postacetabular region, which most likely underestimates the actual dimension, exceeds five and a half acetabular diameters (medial view; Figure 5.2, 5.4). Taking into account the above note, this indicates more pronounced elongation than in complete extant specimens used for Figure 6 (see also Appendix 1), most of them do not exceed the value of five diameters. Relatively elongated postacetabular hip bone appears to be present in Oligocene Kairuku grebetti (Ksepka et al., 2012, figure 6.D-F), and more so in some volant aquatic birds (Ibáñez and Tambussi, 2012, figure 1). In the case of fossils, however, their incompleteness precludes certainty in this respect.

The medial surface of the preacetabular ilium has a conspicuous (though shallow) facet for articulation with the transverse process of a synsacral vertebra (Figure 5.4). Such a scar has been previously documented by Clarke et al. (2003, figure 4.B; see also Acosta-Hospitalche and Olivero, 2016, figure 4.A) for another Eocene hip bone (not articulated with synsacrum). This is a part of an incomplete skeleton from Tierra del Fuego (Argentina), recently assigned to Palaeoepyptes gunnari (see Acosta-Hospitalche and Olivero, 2016). The facet is also present in a relatively small-sized Eocene specimen IB/P/B-0211 (Figure 5.5) and large IB/P/B-0488 (Figure 5.10), and is common in present-day penguins (pers. obs.). Another preserved surface for contact with the synsacrum is represented by a short fragment of the iliac part of the iliosynsacral suture located between the acetabulum and ilioischial foramen (Figure 5.4, 5.8). The right hip bone is partially covered by matrix together with three left leg bones: the femur, tibiotarsus, and fibula (Figure 5.2-4, 5.6).

Femora. The incomplete left femur comprises the poorly preserved head and proximal shaft, well-preserved diaphyseal midsection and damaged distal end with condyles. It is ca. 91.0 mm long, a distance measured between the ligament fovea of the head and medial condyle. The overall length of the bone is comparable to that of the largest individuals of Pygoscelis papua and some representatives of Aptenodytes patagonicus (pers. obs.; Stephan, 1979, table 1). It is clearly larger than IB/
FIGURE 5. Incomplete pelvic skeleton and fragmentary leg bones of Eocene Delphinornis larseni, NRM-PZ A.994, major fragment (bones encased in a single lump of matrix). 1. Location of these skeletal elements within penguin body, in ventral view, is presented in red. Other bones included in NRM-PZ A.994, and shown in Figures 2.1-7, 2.15, 2.17-19, 4.2-6, 4.8-17, are in gray. 2. X-ray-based schematic view of right hip bone and associated left leg bones, as preserved in matrix. 3. X-ray image of same. 4. Medial view of same. 5. Fragmentary right preacetabular ilium IB/P/B-0211 (Eocene, Seymour Island, Antarctica) in medial view. 6. Lateral view of right hip bone and associated left leg bones (NRM-PZ A.994). 7. Right (reversed left) hip bone of extant Spheniscus demersus (Linnaeus, 1758), NHMUK/T S/1998.23.2, in medial (left) and lateral view. 8. Dorsomedia l view of two major hip-bone foramina of NRM-PZ A.994. 9. Ventromedial view of obturator foramen of same specimen. 10. Medial view of right hip bone, IB/P/B-0488 (Eocene, Seymour Island, Antarctica). 11. Lateral view of same. 12. Ventromedial view of three major hip-bone foramina of extant Pygoscelis papua (unnumbered specimen from IB/P/B). 13. Dorsomedial view of nutrition foramen of left tibiotarsus (NRM-PZ A.994). 14. Same view of tibiotarsus IB/P/B-1033 (Eocene, Seymour Island, Antarctica). 15. Caudal view of reversed right femur IB/P/B-0130 (Eocene, Seymour Island, Antarctica). 16. Same view of middle shaft of NRM-PZ A.994. Abbreviations: aa, articular area; af, acetabular foramen; ar, adductor ridge; at, antitrochanter; cil, caudal intermuscular line; dfe, distal end of femur; fc, fibular crest; feh, femoral head; fes, femoral shaft; iif, iliobischiadic foramen; il, ilium; im, insertion of iliofibular muscle; is, ischium; ipl, ischiopubic ligament; iss, ilioschischiadic suture; lc, lateral condyle; lfb, left fibula; lfe, left femur; lt, left tibiotarsus; lp, ligamental pit; mx, matrix; nf, nutrition foramen; of, obturator foramen; pa, pronounced angularity within ischium; pb, pubis; pdp, correlate of proximal dorsal process; ptt, proximal end of tibiotarsus; rf, renal fossa; rhb, right hip bone; s, synsacrum; syn, iliosynsacral synostosis; tb, tubercle; tts, tibiotarsal shaft. Asterisks denote comparative material. Scale bar equals 1 cm.
P/B-0130 (Figure 5.15; Jadwiszczak, 2006, figure 14.A-B; hypothesized to represent Delphinornis gracilis) and slightly shorter than IB/P/B-0215 (Jadwiszczak, 2006, figure 16.A-B; hypothesized to represent Mesetaornis polaris Myrcha et al., 2002). Unlike the left head, its right counterpart (Figure 4.8-9), found in close vicinity, is in a good condition being the only proximal remnant of the respective bone. It is 11.4 mm wide and has a large proximal pit (the capital ligamental fossa/fovea), a round ligament insertion site. The pit consists of two distinct parts, a deeper, ovoid central area with a partially rough, darker floor and a peripheral zone with a smooth surface (Figure 4.15). The ligamental fossa is located (like in all but one Sphenisciformes and closely related Procellariiformes) medioproximally to proximally. In this respect, the only exception (medial placement) appears to be Paleocene Crossvallia unienwillia Tambussi et al., 2005, the poorly preserved Antarctic stem penguin (Jadwiszczak et al., 2013, figure 6.2-3, 6.5). The right femoral shaft is characterized by a well-defined curvature (similarly to P. papua, but unlike most other members of the Sphenisciformes; e.g., Figure 5.15; Ksepka and Clarke, 2010) and caudal intermuscular line that ends distally with a conspicuous tubercle (Figure 5.6, 5.16; not as prominent as in Inkayacu paracasensis; Clarke et al., 2010, figure 1.K). Clearly, there is an actual discontinuity between the above line and adductor ridge (Figure 5.16). Such a feature can be also observed in some other fossil penguin femora from the Eocene of Antarctica, both small- (e.g., IB/P/B-0108; unfigured), supposedly in Paleocene Sequiwaimanu rosiaeae Mayr et al., 2018 from New Zealand (Mayr et al., 2018, figure 11.A) as well as in Paraptenodytes antarcticus (Moreno and Mercera, 1891) - a Miocene stem penguin from Patagonia (Bertelli et al., 2006, figure 19.D). Moreover, this discontinuity is quite conspicuous in present-day Aptenodytes forsteri and A. patagonicus (pers. obs., based on NHMUK/T S/1998.55.2 and NHMUK/T S/ 1846.4.15.31, to name two). The diaphyseal midsection is essentially circular with its minimum mediolateral width of 8.8 mm (slightly wider than IB/P/B-0130 and narrower than IB/P/B-0215; Jadwiszczak, 2006, figure 16.A-B). Taking into account the minimal shaft diameter and head breadth, their values for NRM-PZ A.994 do not exceed those obtained from femora attributable to P. papua (pers. obs.; Stephan, 1979, table 1). Internally, it is characterized by a pronounced local narrowing of the medullary cavity, apparent on the X-ray image (Figure 5.3). There are also two small matching shaft-wall fragments most likely of the right femur (Figure 4.10-11), assignable to the same individual, showing such a stricture (Figure 4.11). Proximally to the narrowing, the former shaft exhibits a small nutrient foramen, its opening slightly medial to the caudal intermuscular line. Another, even smaller foramen opens distally to the aforementioned line (Figure 5.6, 5.16). Both foramina are 21.5 mm apart. The proximal foramen, likely representing the entry point of the main nutrient artery (see Midtgård, 1982), is quite easily traceable (unlike the distal one) in modern and some Eocene penguin femora (e.g., Figure 5.15; see also Acosta Hospitaleche and Olivero, 2016, figure 4.D; Mayr et al., 2018, figure 11.A).

**Tibiotarsi.** The left tibiotarsus is poorly preserved (Figure 5.6), and its overall length cannot be reliably estimated. Its proximal part is represented by lateral and medial articular surfaces and a small remnant of the adjacent diaphysis. Distally, a much longer (48.3 mm) non-joining fragment of tibiotarsal diaphysis is present. It is relatively complete solely in its distal part, whereas the bulk of the preserved shaft is limited to the caudal surface with a partial fibular crest. The crest is accompanied by a short furrow leading to a small nutrient foramen (Figure 5.6, 5.13) - a common feature in Eocene penguins of different sizes (e.g., Figure 5.14), also present in present-day Sphenisciformes (distinctness varies; pers. obs.), Paleocene Waimanu manneringi (Mayr et al., 2018, figure 11.H), and Oligocene Kairuku grebneffi (Ksepka et al., 2012, figure 7.D). The estimated diaphyseal mediolateral width just distal to the foramen is ca. 9.3 mm. Two isolated fragments of a more distal shaft section are most likely also, based on their cross-sectional shape, assignable to this bone (Figure 4.13, 4.14). The distalmost tibiotarsus is represented by two fragments from the right-limb skeleton: a fragmentary shaft showing a proximal part of the extensor sulcus (Figure 4.16) and, clearly non-joining, distal end with the supratendinal bridge, extensor canal, and condyles (Figure 4.17). The canal appears to be rather tight proximally (in comparison to some other similarly-sized Eocene bones, e.g., IB/P/B-1033; Figure 4.18), but the bridge appears to be fractured (Figure 4.17), which could affect the underlying tendinal passage. The bridge is moderately wide, narrower than in Miocene Paraptenodytes antarcticus (Bertelli et al., 2006, figure 20.C) and Madrynornis mirandus (Acosta Hospitaleche et al., 2007, figure 5.A) as well as extant Pygoscelis papua (Figure 4.19).
FIGURE 6. Shape analyses of outlines of major hip bone foramina. 1. Comparison of foramen configurations and shapes of eight extant and three stem-group penguins, including the Eocene specimen NRM-PZ A.994 (*Delphinornis larseni*); foramen pairs are not to scale, branching pattern for present-day species used in the cladogram follows Subramanian et al. (2013, figure 1). 2. Position of NRM-PZ A.994 relative to the extant-penguin morphospace, based on aligned, centered and scaled outlines. 3. Deformation grid between configurations representing NRM-PZ A.994 (*D. larseni*) and NHMUK/T S/2002.2.1 (extant *Eudyptula minor*). For other specimen IDs, see Appendix 1; for PCA plots, see Appendix 4.
opening of the passage is located centrally in relation to the condyles (Figure 4.17), unlike in Paleocene penguins (Sequoiawan rosaeae and W. manneringi; Mayr et al., 2018, figure 11.J, 11.M). The medial condyle is ca. 18.1 mm wide caudocranially, whereas the mediolateral width of the whole fragment, measured between both condyles in distal view, amounts to ca. 17.9 mm. Considering the latter measurement together with the diaphyseal mediolateral width (see above), NRM-PZ A.994 most closely resembles IB/P/B-0266 (hypothesized to represent Delphinornis arctowski; Jadwiszczak, 2006, table 20). The analyzed material also includes a single long-bone fragment, presumably representing (judging from its angular shape) a proximal tibiotarsal shaft (Figure 4.12).

**Fibula.** The left fibula, the only one available, is quite well preserved (Figure 5.6). It is 78.6 mm long and 8.8 mm wide proximally. The shaft gradually decreases in its width down to the level of the fibular crest of the collaterally located tibiotarsus, where it expands laterally. This expansion, constituting the insertion point for the iliofibular muscle (Baumel and Witmer, 1993, figure 4.17), is sharper defined than in modern penguins (pers. obs.; Stephan, 1979, figure 47.1), and apparently also in Palaeaeudyptes gunnari (Acosta Hospitaleche and Reguero, 2010, figure 4.G) and Kairuku grebneffi (Ksepka et al., 2012, figure 7.B). A distalmost one-third of this bone is very thin.

**Tarsometatarsi.** The complete left tarsometatarsus and six associated pedal phalanges have been originally located within the same lump of matrix (Figure 2.18). Bones have been in close contact, though largely displaced. The tarsometatarsus (Figure 2.2-7) is relatively elongated like many early penguins (e.g., Myrcha et al., 2002; maximum length of 45.4 mm, 44.1 mm along main axis – to the groove of the third trochlea, midwidth of 16.0 mm) and flattened dorsoplantarly (the latter is typical of all penguins; e.g., Figure 2.20-21; Stephan, 1979; Myrcha et al., 2002). The third metatarsal is 6.4 mm thick in its midsection. This bone is smaller than most well-preserved specimens attributable to D. larseni, including the holotype (for more contrasts, see Figure 3). The comparative and functional aspects of tarsometatarsal elongation are considered in Discussion and Conclusions. The location of fusion between three ankylosed metatarsals and the distal tarsal bone is well marked internally, the condition revealed by the X-ray imaging (Figure 2.18). Externally, the markings are constricted to a low dorsal crest – a partial distal frame of the infracotylar fossa, and proximal swelling of the medial margin of the bone (Figure 2.2). The wide oval medial cotyle is strongly concave, whereas its lateral counterpart resembles a clearly narrower flat shelf. The intercotylar eminence (Figure 2.2) is distinctly less prominent than in Paleocene Waimanu manneringi (Slack et al., 2006, figure 1.C) and, in medial and lateral views (Figure 2.3-4), virtually level with the intercotylar area. The proximal part of the bone is 18.9 mm wide mediolaterally. The medial hypotarsal crest and lateral hypotarsal tuberosity (see below) are separated by the well-developed sulcus (Figure 2.5-6; much shallower in present-day penguins, but clearly deeper in Eocene Marambiornis exilis Myrcha et al., 2002 and Procellariiformes; Jadwiszczak, 2015, figure 6). The medial hypotarsal crest, in terms of its mediolateral width and cross-sectional shape, subtly resembles its equivalents in some more crownward penguins, e.g., Eocene Anthropornis sp. (not all specimens from this genus; Acosta Hospitaleche and Jadwiszczak, 2011, figure 3.B-C) and Miocene Paraptenodytes antarcticus (Bertelli et al., 2006, figure 21.C-D). The lateral hypotarsus is dominated by the lateral of intermediate crests. Its medial counterpart is barely visible along the proximal section of the former ridge; the proper lateral hypotarsal crest is vestigial as well. Nevertheless, the hypotarsus described above, together with its counterparts in congenerics, appears not to be as simple as its counterparts in modern penguins (e.g., Jadwiszczak, 2015, figure 6.A, 6.D, 6.G, 6.J) and Oligocene Kairuku (Ksepka et al., 2012, figure 7.L, 7.P). The other extreme of the spectrum (i.e., notably distinct crests – like in most birds, including Procellariiformes; Baumel and Witmer, 1993; Jadwiszczak, 2015, figure 6) is represented by Paleocene (Mayr, 2017a, figure 1.C-D, 1.F, 1.H) and some Eocene Sphenisciformes (especially Marambiornis exilis; Myrcha et al., 2002, figure 15.B; Jadwiszczak, 2015, figure 6.I).

The fourth metatarsal is straight (Figure 2.2, 2.4-5; unlike in Oligocene Archaeospheniscus lopodelli Marples, 1952, and Duntroonornis parvus Marples, 1952, from New Zealand; Marples, 1952, figure 8.3-4, 8.5; post-mortem deformation cannot be excluded), whereas the second one is distally slightly curved abaxially in dorsal view, just before the respective trochlea (Figure 2.2-3; not as much as in, e.g., Palaeaeudyptes; Myrcha et al., 2002, figures 7, 8). The dorsal surface of the third metatarsal is planarly lowered in its proximal part. X-ray imaging revealed the pronounced localized narrowing of the medullary cavity in distal sections of
all three metatarsals (Figure 2.18). The insertion site of the tibial cranial muscle is fusiform in shape (Figure 2.2; wider, approximately oval in present-day and larger fossil penguins; Jadwiszczak, 2006, figure 5.A, 8.A, 2015, figure 4.A; Ksepka et al., 2012, figure 7.M, 7.Q; Mayr et al., 2017a, figure 1.A). Both proximal vascular foramina are small, like in Paleocene penguins (Slack et al., 2006, figure 1.A; Mayr et al., 2017a, figure 1.A) and Eocene *Marambiornis exilis* (Myrcha et al., 2002, figure 15.A), the lateral one is slightly more proximal (Figure 2.2; like in *M. exilis*; Myrcha et al., 2002, figure 15.A). Moreover, judging from the X-ray image (Figure 2.18), it appears to be a little better developed on the whole than the quite slit-like medial foramen, which is not that obvious during external examination. The medial foramen is medially framed by two convex impressions of the extensor retinaculum. Both plantar proximal openings are more or less less axial relative to the hypotarsal sulcus and crests (Figure 2.5; like in most penguins, Paleogene exceptions are represented by *Marambiornis* and *Mesetaornis*; Myrcha et al., 2002, figures 13.B, 14.B, 15.B). The distal vascular foramen is present (a plesiomorphism; e.g., Mayr et al., 2017a), evidenced by both the X-ray (Figure 2.19) and plantar-aspect morphology of the lateral intertrochlear notch (Figure 2.5), but supposedly not fully enclosed. This plesiomorphic trait (e.g., Mayr et al., 2017a) is present in Paleocene (Slack et al., 2006, figure 1.A; Mayr et al., 2017a, figure 1.C) and relatively small-sized Eocene Antarctic penguins (its distinctness varies; see Myrcha et al., 2002). The medial intermetatarsal sulcus is slight, whereas its lateral counterpart is well formed (Figure 2.2; intensity of this asymmetry varies in stem penguins; e.g., Myrcha et al., 2002). These structures are barely traceable plantarly (Figure 2.5). The plantar supratrochlear depression is restricted to the distal third metatarsal (wider mediolaterally in *M. polaris*, shifted medially in *M. exilis*; pers. obs.; Myrcha et al., 2002, figure 13.B, 15.B). The third trochlea is the largest one (Figure 2.7), its dorsoplantar width of 10.4 mm and maximum mediolateral width of 8.1 mm. The distal mediolateral width of the tarsometatarsus, measured between the midpoints of trochlear margins in distal view, amounts to 22.8 mm.

The right tarsometatarsus is represented by two isolated and non-adjoint fragments of proximal and distal parts with some portions of metatarsals attached (Figure 2.1). They are less well preserved than their left-sided counterparts. Most importantly, the latter specimen directly indicates the presence of a dorsal opening of the distal vascular foramen.

**Pedal phalanges.** Digits of the left foot are represented by eight recognized phalanges, which are largely well preserved (Figure 2.17). The second digit retained a proximal and intermediate of its three phalanges, the third digit is devoid solely of the ungual phalanx and its penultimate (third) phalanx is very fragmentary, and within the five-united fourth digit, two unambiguous proximalmost phalanges remained, most likely accompanied by the distalmost (ungual) bone. The first phalanx of the third digit, the largest unit within a penguin forefoot, is 29 mm long plantarly, measured along its main axis. The slender-built ungual phalanx of the fourth digit, devoid of its very apex, is 15.8 mm long plantarly. Both of its curved neurovascular sulci are present, the medial groove is much better developed and more dorsal. The flexor tubercle is prominent. Of all the aforementioned phalanges, only the second and third bones from the third digit were not encased by matrix together with the tarsometatarsus (Figure 2.18).

The right-foot toes are known from the first three incomplete phalanges of the fourth digit, the first two even more fragmentary phalanges of the third digit, supplemented by the well-preserved ungual unit, and a small fragment of the second phalanx from the second digit (Figure 2.15). The ungual bone has a slightly abraded proximal end and is devoid of a small portion of its distal tip. Nevertheless, it is stouter in appearance than the only known ultimate phalanx from the left foot, and its lateral neurovascular sulcus is also deeper and wider.

The associated phalanges that are assignable to a named genus or species are not common in the fossil record of early Sphenisciformes. The proximal phalanx of the third digit (i.e., the largest one) from NRM-PZ A.994 (Figure 2.17) is relatively more elongated (i.e., possesses larger midlength/midwidth ratio) than its counterpart in giant Eocene penguins from Antarctica: *Anthropornis* sp. (Jadwiszczak, 2012, figure 3.A) and *Palaeepyptes klekowskii* (Acosta Hospitaleche and Reguero, 2014, figure 7.L). In this respect, *Delphinornis larseni* resembles CM 2016.158.1, an unnamed giant penguin from the Paleocene Waipara Greensand in New Zealand (Mayr et al., 2017a, figure 1.A). The proximal phalanx from the second digit has similar proportions to that in *P. klekowskii* (Acosta Hospitaleche and Reguero, 2014, figure 7.M). The second phalanx of the third digit from NRM-PZ A.994 seems to be relatively more elongated than its
counterpart in another large-sized stem penguin, *Inkyacu paracasensis* from the Eocene of Peru (Clarke et al., 2010, figure S.3), but this difference is much smaller when compared with the Waipara Greensand penguin. Two proximalmost phalanges of the fourth digit from CM 2016.158.1 appear to be relatively more elongated than those in NRM-PZ A.994. The issue of phalangeal proportions is also considered in Discussion and Conclusions.

**DISCUSSION AND CONCLUSIONS**

The tarsometatarsus, due to the very nature of the fossil record of Antarctic penguins, has constituted the only element unambiguously assignable to *Delphinornis* (Simpson, 1971; Myrcha et al., 2002; Jadwiszczak, 2006). The proportions of the complete left bone, quite typical of the species (e.g., Figure 2.2, 2.8-9), clearly differ from their counterparts in extant penguins (e.g., Figure 2.20-21). Although discussed specimens are longer than some foot bones belonging to *Aptenodytes Miller, 1778*, the largest present-day Sphenisciformes, they are at the same time much thinner mediolaterally. Considering the latter dimension, NRM-PZ A.994 resembles some medium-sized extant penguins (Figure 3).

Relatively elongated tarsometatarsi are characteristic of all known Paleocene and many Eocene penguin species (Myrcha et al., 2002; Slack et al., 2006) as well as, much more pronouncedly, Procellariiformes (pers. obs.; Klemm 1969, figure 45) – the closest living relatives of extant penguins (e.g., Simpson, 1976; Hackett et al., 2008). Hence this is an obvious plesiomorphy. Recently, a conspicuous morphological disparity among Paleocene Sphenisciformes, also in terms of foot-bone proportions, has been reported by Mayr et al. (2017a). However, even the relatively shortened tarsometatarsus of an unnamed penguin from the mid-Paleocene of New Zealand (see Mayr et al., 2017a, figure 1) differs in this respect from its counterparts in crown-group Sphenisciformes or Spheniscidae sensu Clarke et al. (2003).

Tarsometatarsal elongation has its functional consequences, for instance, for the force output during ankle (intertarsal joint) flexion. Considering any group of geometrically similar birds, the longer the tarsometatarsus (outforce lever arm) relative to the distance between the flexor attachment and intertarsal joint, representing the inforce moment arm, the lower the magnitude of the force output (Zeffier and Norberg, 2003). The ratio of the latter to the former length, calculated for NRM-PZ A.994, amounts to 0.363. This value (hence also the relative force output) is slightly larger than the ratio in another well-preserved bone assignable to *Delphinornis larseni* (Figure 2.9-14; 0.351) as well as type specimens of *D. arctowski*, *D. gracilis*, and *Marambiornis exilis* (0.341, 0.351, and 0.334, respectively), all of them from the Eocene. However, it is slightly smaller than that of another Eocene penguin – *Mesetaornis polaris* (0.371) and more so relative to extant pygoscelid penguins (>0.4).

Present-day penguins, like most birds, are essentially digitigrade; i.e., they walk on their toes, although use whole feet at rest (pers. obs.; see also Simpson, 1946). It is most parsimonious to assume that they share this trait with their Paleogene relatives. This view has recently gained support by the observation that the alleged lumbosacral sense organ for the control of walking (Necker, 2006) does not differ substantially between the Eocene and extant Sphenisciformes (Jadwiszczak, 2014).

Distal width of the largest foot bone, the tarsometatarsus, constitutes a proxy for the span of extreme proximal-phalanx bases. In this regard, the NRM-PZ A.994 is within the range of values reported by Stephan (1979, table 1) for the extant Gentoo penguins, *Pygoscelis papua*, the largest among medium-sized modern Sphenisciformes (see Williams 1995). Interestingly, two proximalmost phalanges from each of three main digits (well preserved in the fossil skeleton) are clearly longer (and more elongated) in the Eocene specimen (Figure 2.15, 2.17). Additionally, the elongated tarsometatarsus can provide larger surface areas for the origins of toe abductors, muscles that arise fleshy from the second and fourth metatarsals in present-day penguins (Schreiweis, 1982; Jadwiszczak, 2015, figures 4, 5). One of such enlarged areas (related to the abductor of the fourth digit) is especially well delineated in IB/P/B-0547, another specimen assignable to *D. larseni* (Figure 2.8).

Considering condition of the abductor of the fourth digit, there is some variation among Procellariiformes. In this sister clade/taxon (characterized by much longer fore- and hindfoot bones), the muscle can be fleshy proximally and distally (i.e., with two origins, tendinous within the middle third; albatrosses) or fleshy along its length (shearwaters), or even originate distally (storm petrels; Klemm, 1969). The abduction produced in the first scenario was specified by Klemm (1969) as strong.

The above findings indicate the longer forefoot and theoretically wider toe spacing attainable by *Delphinornis larseni*. Such a characteristic
potentially facilitates, by spreading body weight over a wider area, walking by reducing the risk of losing balance. The said condition, together with the elongate tarsometatarsus (e.g., Figure 2.2.2.9), obviously leads to larger hydrodynamic drag. On the other hand, penguins are wing-propelled divers, using their feet as a rudder. To enlarge its area, extant (i.e., short-footed) Sphenisciformes possess additional dermal flanges on foot sides (Figure 2.16; Cornithwaite, 2013), an adaptation possibly either not present or not that essential for at least some early penguins. Moreover, like the vast majority of marine birds, penguins and their closest living relatives (Procellariiformes) have webbed feet (e.g., Schreiber and Burger, 2002). The members of the latter group, however, do not possess additional dermal flanges (pers. obs.). Those of them that dive, use their wings as oars, just like penguins (e.g., Del Hoyo et al., 1992).

The proximal fovea of the right femoral head (Figure 4.8-9, 4.15) has a heterogeneously formed floor with the ovoid central area, constituting the actual attachment site of the round ligament of the femur. The smoother and shallower peripheral area apparently (partly) served as a receptacle zone (e.g., Perumal et al., 2017). The fovea reveals an osteological aspect of the bone-ligament interface – a distribution of units reflecting higher contact intensity (separate bundles) (Figure 4.15). They are distinguishable by differences in their tinge, depth, and presence of foramina, at least some of them being vascular. No other femoral fovea in available fossil-penguin specimens represents such a degree of detail.

The pelvis of early penguins, including NRM-PZ A.994, exhibits a well-defined fixed asymmetry in the size of two main hip bone openings, the acetabular foramen being always smaller than the ilioischiadic foramen (Figure 6; see also Jadwiszczak 2006, figure 19). Extant penguins are clearly more diversified in this respect (Figure 6.1-2), the former condition present within the *Eudyptula-Spheniscus* group (Figure 6.1; Zusi, 1975). The configuration exhibited by *Delphinornis larseni* is most closely approached by much smaller *Eudyptula minor* (Forster, 1781). However, even in this case, differences are obvious (Figure 6.3). The large and elongated ilioischiadic foramen of Paleogene Sphenisciformes (Figure 6.1) supposedly represents an ancestral trait state, because it is also larger than the acetabulum and often clearly elongated in Procellariiformes (e.g., Klemm, 1969; Mayr, 2009). The oldest penguins exhibiting inverse proportions of opening sizes are known from the Miocene (e.g., Acosta Hospitaleche et al., 2007, figure 5.B).

In birds, the cranial part of the ilioischiadic foramen transmits the ischiadic nerves and vessels, the remaining part of the opening is occupied by the connective tissue sheet (a membrane), with muscles attached to its both surfaces (Baumel and Raikow, 1993). Hutchinson (2001) claimed that the proximal dorsal process of the avialan ischium is correlated with the formation of the above membrane. If so, then the characteristic point-of-change of the ischiadic-rim width, observable in at least some penguin fossils (NRM-PZ A.994, but also in large IB/P/B-0488; Figure 5.8, 5.11) and modern bones (Figure 5.12), that in our estimation corresponds to that process, marks the proximalmost attachment of the membrane (see also Hutchinson, 2001). Assuming correctness of our reasoning, the unobstructed area of the ilioischiadic foramen accounted for about a third of the entire opening in Eocene sphenisciforms. Such a fraction is typical of most birds (see Baumel and Raikow, 1993), but not necessarily of present-day penguins. In the latter group, the above-mentioned point is located closer to the caudal frame (Figure 5.12; Appendix 5), hence the membrane covers a smaller part of the opening (Appendix 5).

According to Midtgård (1982), in extant penguins (represented in his study by *Spheniscus humboldti* Meyen, 1834), unlike in a majority of birds (including Procellariiformes), the ischiadic artery is not the main blood vessel supplying most of the hind limb. If there was a link between reducing the size of the ilioischiadic foramen during penguin evolution (see Figure 6) and gaining the importance by the femoral artery, a vessel not associated with the above foramen, the early Sphenisciformes might have been more like procellariiforms in this regard.

The above shrinkage was possibly related to the ossification of the ilioischiadic membrane resulting from the assumed shortening/reinforcement of the postacetabular pelvis (see comparative description of hip bones) and associated (hypothetical) cranial shift or expansion of origins of relevant muscles, e.g., the ischiofemoral muscle. The origin of this flexor is caudal to the ilioischiadic foramen and its origin in modern penguins is “extensive” (Schreiweis, 1982, figure 41). Both in turn could have been connected to the evolution of the streamlined form (useful when diving) with legs shifted backwards, enforcing a (fully) upright posture on land (including use of shorter tarsometatarsi; in water, feet essentially serving as a rudder).
Another factor, that ought to be taken into account, is a possible selection for the better protection (by shrinking "holes") of the caudal portion of kidneys, especially during deep dives.

The spin-off of that evolutionary process could have been a higher risk of choke-point formation (see Wideman, 2016, figure 6), resulting in an increased incidence of complications such as aneurysm and ischaemia. Assuming the above scenario, increasing the importance of the femoral artery would be advantageous. Although the persistence of an ancestral character state in stem penguins appears to be plausible, the issue flagged up here requires further studies.

Penguins, typically of seabirds, are long-lived and banding records indicate that some individuals can live much longer than 20 years, and even breed after crossing their twenties (e.g., Dann et al., 2005). Results of our analysis of the specimen NRM-PZ A.994 indicate that the individual discussed here was most likely a relatively young bird. We surmise that the above penguin was up to several years old; decidedly not a chick, because the fusion of tarsometatarsal components is outwardly complete (Figure 2.2-5). Against the really advanced age of the individual testify the following foot-bone conditions: the presence of very slight traces of plantar intermetatarsal sulci (especially the lateral one; Figure 2.5), transition between the second metatarsal and tarsus marked as an abrupt bone swelling (Figure 2.2-3), the whole synostosis zone between proximal metatarsals and the tarsal component conspicuous internally (highly radiopaque; no "cross-border" remodeling marked; Figure 2.18). Moreover, the distal vascular foramen was possibly not as well developed as that in the hypodigm expanding throughout the inclusion of further tarsometatarsi. Whereas giant early Sphenisciformes, in a sense, ruled the minds of researchers studying the evolutionary history of this fascinating order of birds, fueled by the growing number of discovered skeletons, the fossil record of smaller-sized stem-group penguins has been gaining rather quantitatively than qualitatively. The specimen reported here is a game-changer in paleontology of stem penguins, offering insight into valuable details of the pelvic girdle and hind-limb bones of the latter group and hopefully reducing the existing bias towards the larger birds.

Since its first formal description (Wiman, 1905), *D. larseni* has remained quite an enigmatic bird with its hypodigm expanding throughout the inclusion of further tarsometatarsi. Whereas giant early Sphenisciformes, in a sense, ruled the minds of researchers studying the evolutionary history of this fascinating order of birds, fueled by the growing number of discovered skeletons, the fossil record of smaller-sized stem-group penguins has been gaining rather quantitatively than qualitatively. The specimen reported here is a game-changer in paleontology of stem penguins, offering insight into valuable details of the pelvic girdle and hind-limb bones of the latter group and hopefully reducing the existing bias towards the larger birds.

We currently know that modest-sized Eocene penguins assignable to *D. larseni* were osteologically "slender-footed" in respect to both (tarsometatarsal cotyle) and texture in the fossil bone preclude the complete fusion, rather being suggestive of a less rigid connection. In contrast to NRM-PZ A.994, another fragmentary ilium from the Eocene of Seymour Island (IB/P/B-0211; Figure 5.5), an isolated element belonging to a slightly larger individual from unknown species, possesses a smaller cotyle with much smoother surface. In our estimation, this difference reflects the actual condition – a less definite element connection in the latter bird. Moreover, such a cotyle is also present in some other giant penguins (see comparative description of hip bones), whereas more definite connections can be observed in the giant penguin genus *Kairuku* (Ksepka et al., 2012, 6.C, 6.F). Interestingly, the preserved iliac part of the iliosynsacral suture in NRM-PZ A.994 (Figure 5.6, 5.8) is very rough, indicating a fracture within rigidly united elements, therefore precluding a very young age of the bird.

According to Jadwiszczak and Chapman (2011, figure 1), rough estimates obtained by Jadwiszczak (2001) via the modelling of the tarsometatarsal dorsoplantar width and overall body length relationship, suggest that small individuals assignable to *D. larseni*, and NRM-PZ A.994 is not that large, could have been slightly shorter than some extant Gentoo penguins, *Pygoscelis papua*. Considering longer weight-bearing bones, femoral shaft and head dimensions are often utilized in body-mass assessments (e.g., Brassey, 2016). Taking cognizance of femoral dimensions, the studied skeleton belonged to an individual both shorter and lighter than whichever representative of so-called giant penguins (e.g., Jadwiszczak and Chapman, 2011) from the Eocene of Seymour Island.
tarsal and digital) distalmost components of the hind limb. Their postacetabular hip bones (precisely - their main, ilioischiatric parts, which are much better preserved than preacetabular ones) were apparently relatively elongated, possessing the extensive ilioischiatric foramen, largely covered with membrane. These major findings, unique for non-giant stem penguins, supplemented by a number of other observations, contribute to the known set of character configurations. Hence our report can have obvious uses in prospective refinements of published cladograms. The functional angle appears to be indispensable as well. It ranges across well-grounded considerations (the force output during intertarsal joint flexion), those slightly debatable (the role of elongated toes in reducing the risk of losing balance on land), and more speculative (though logically justified – increasing the importance of the femoral artery). Although we consider our contribution as an important step forward in penguin paleontology, the catchphrase “more research is needed” remains appropriate.

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APPENDIX 1.

Comparative material utilized in Figures 3 and 6. Other specimens used for comparisons are referenced in-text, either via their catalog numbers (if they were studied directly) or by citations of source works with relevant figure/table numbers.

1. Tarsometatarsi used in Figure 3. Data for fossil specimens from IB/P/B and MLP after Myrcha et al. (2002, table 1), CM - after Slack et al. (2006) and Mayr et al. (2017a), MUSM – after Clarke et al. (2010), OU - after Marples (1952) and Ksepka et al. (2012); Trochlear widths for *Inkayacu*, *Kairuku*, *Muriwaimanu*, *Waimanu*, and CM 2016.158.1 as well as midwidths of *Muriwaimanu*, *Waimanu* and CM 2016.158.1 were assessed based on published photographs (with scales). Other measurements were taken directly by PJ.

a. Fossil bones

| Species                  | Catalog Numbers                                                                 |
|--------------------------|----------------------------------------------------------------------------------|
| *Anthropornis grandis*   | IB/P/B-0483, MLP 95-I-10-142, MLP 94-III-15-178                                 |
| *Anthropornis nordenskjoeldi* | NRM-PZ A.45                                                                    |
| *Archaeospheniscus lopdelli* | NHMUK A4080 (cast)                                                               |
| *Archaeospheniscus wimani* | MLP 90-I-20-24, NHMUK A3331                                                     |
| *Delphinornis arctowski*  | IB/P/B-0484, MLP 93.X.1.92                                                       |
| *Delphinornis gracilis*   | IB/P/B-0279a                                                                     |
| *Delphinornis larseni*    | NRM-PZ A.21, NRM-PZ A.587, NRM-PZ A.861, NRM-PZ A.895, IB/P/B-0062, IB/P/B-0547, IB/P/B-0548, MLP 84.II.1.79, MLP 91.II.4.174, MLP 83.V.20.5 |
| *Inkayacu paracasensis*   | MUSM 1444                                                                        |
| *Kairuku grebneffi*      | OU 22094                                                                         |
| *Kairuku waitaki*        | OU 12652                                                                         |
| *Korora oliveri*         | OU C.50.63                                                                       |
| *Marambiornis exilis*    | IB/P/B-0490, MLP 93.X.1.111                                                      |
| *Mesetaornis polaris*    | IB/P/B-0278                                                                      |
| *Muriwaimanu tuatahi*    | CM zfa 34                                                                        |
| *Palaeaeudyptes antarcticus* | NHMUK A1048                                                                    |
| *Palaeaeudyptes gunnari* | IB/P/B-0072, IB/P/B-0277, IB/P/B-0487, MLP 91-II-4-222, MLP 87-II-1-45, MLP 82-IV-23-6, MLP 84-II-1-124' |
| *Palaeaeudyptes klekowski* | IB/P/B-0485, IB/P/B-0545, MLP 93-X-1-142, MLP 84-II-1-78, MLP 93-X-1-63, MLP 84-II-1-76, MLP 93-X-1-106, MLP 94-III-15-20, MLP 83-V-30-15, MLP 83-V-30-16, MLP 84-II-1-124 |
| *Palaeaeudyptes marplesi* | NHMUK A6119 (cast)                                                               |
| *Waimanu manneringi*     | CM zfa 35                                                                        |
Undetermined Paleocene penguin from New Zealand
CM 2016.158.1

b. Modern bones

Aptenodytes forsteri
NHMUK/T 1197A, NHMUK/T 1197D, NHMUK/T 1197B, NHMUK/T 1972.1.25, NHMUK/T 1905.12.30.419, NHMUK/T 1998.55.2

Aptenodytes patagonicus
NHMUK/T 1846.4.15.33, NHMUK/T 701A, NHMUK/T 701B, NHMUK/T 1952.1.28, NHMUK/T 1972.1.24, NHMUK/T 2000.12.1

Eudyptes chrysocome
NHMUK/T 1952.1.39, NHMUK/T 1898.7.1.15, NHMUK/T 1964.14.2, NHMUK/T 1898.7.1.14

Eudyptula minor
NHMUK/T 1966.51.1, NHMUK/T 2002.2.1, NHMUK/T 1896.2.16.38

Pygoscelis adeliae
NHMUK/T 1849.10.2.2, NHMUK/T 1850.9.7.1, IB/P/B – three unnumbered specimens.

Pygoscelis papua
NHMUK/T 2006.1.16, IB/P/B - an unnumbered specimen,

Spheniscus demersus
NHMUK/T 1898.7.1.8, NHMUK/T 1898.7.1.9, NHMUK/T 1998.23.2, NHMUK/T 1998.48.24

2. Hip bones specimens used in Figure 6. Outlines of foramina for Kairuku grebneffi and Waimanu manneringi were assessed based on Ksepka et al. (2012, figure 6) and Slack et al. (2006, figure 1) respectively.

a. Fossil

Kairuku grebneffi
OU 22094

Waimanu manneringi
CM zfa 35

b. Modern

Aptenodytes forsteri
NHMUK/T 1972.1.25

Aptenodytes patagonicus
NHMUK/T 1846.4.15.32

Eudyptes chrysocome
NHMUK/T 1956.14.1

Eudyptula minor
NHMUK/T 2002.2.1

Pygoscelis adeliae
IB/P/B - an unnumbered specimen

Pygoscelis papua
IB/P/B - an unnumbered specimen

Spheniscus demersus
NHMUK/T 1905.5.23.1

Spheniscus humboldti
NHMUK/T 1952.1.42
APPENDIX 2.

Details of the shape-analysis workflow.

1. **Input files preparation (one per specimen)**
   - Raw photo
   - Main foramina masked (mask indicated by its marquee)
   - Masked areas filled with black color
   - Black areas extracted, pasted onto white background, connected with thin black line and saved to file (jpg, quality set to 100%, 300 DPI)

2. **Import to R/Momocs environment**
   - Function name
   - `import_jpg()`
   - Library (Momocs)
   - List of extracted outline coordinates from multiple files
   - Form of extracted outline
   - (1544 pairs for this outline)

3. **Analyses in R/Momocs environment**
   - **PCA() plot()**
   - **Elliptic Fourier Transform**
     - Closed outlines decomposed into a series of elliptic Fourier harmonics, each described by four coefficients
     - Out/Coe class object
     - `efourier()`
   - **Calibrate harmonic power efourier()**
     - (to choose optimal number of harmonics)
   - **Harmonic power assessment**
     - (for mean shape)
     - **Morphospace visualization**
     - (aligned, centered, scaled and stacked outlines)
     - **Inspection**
     - (silhouettes also used in cladogram)
   - **Reconstructed outlines**
     - (based on 16 harmonics, hence devoid of high-frequency noise)
   - **Outline coordinates extracted**
     - 120 x-y pairs for each outline (default value)
   - **Deformation-grid plotting**
     - (showed for D. larseni and E. minor - the most similar among PCA-analyzed configurations)
APPENDIX 3.

Harmonic coefficients (normalized) calculated for outlines of major hip bone foramina (nine outlines × 16 harmonics, four coefficients for each harmonic).

Jadwiszczak and Mörs, First partial skeleton of *Delphinornis larseni* Wiman, 1905, a slender-footed penguin from the Eocene of Antarctic Peninsula.

APPENDIX 3. Harmonic coefficients (normalized) calculated for outlines of major hip bone foramina (nine outlines × 16 harmonics, four coefficients for each harmonic).

|                | A1   | A2   | A3   | A4   | A5    | A6    | A7    |
|----------------|------|------|------|------|-------|-------|-------|
| *A. forsteri*  | 1.00000 | 0.035786 | 0.133590 | -0.002903 | 0.017254 | -0.018801 | -0.010171 |
| *A. patagonicus* | 1.00000 | 0.067544 | 0.158027 | -0.019771 | 0.024652 | 0.003973 | -0.02821 |
| *D. larseni*   | 1.00000 | -0.076995 | 0.174987 | 0.012179 | -0.032708 | -0.012419 | 0.009728 |
| *E. chrysocome* | 1.00000 | 0.038634 | 0.138061 | -0.003595 | 0.028278 | -0.023276 | -0.009829 |
| *E. minor*     | 1.00000 | -0.008819 | 0.155846 | 0.004365 | -0.004620 | 0.025240 | 0.002692 |
| *P. adeliae*   | 1.00000 | -0.023009 | 0.160485 | -0.001064 | 0.019697 | -0.010874 | -0.004514 |
| *P. papua*     | 1.00000 | -0.017745 | 0.165500 | -0.017904 | 0.006324 | -0.009807 | 0.012399 |
| *S. demersa*   | 1.00000 | -0.008206 | 0.150995 | -0.003024 | 0.023549 | 0.019210 | -0.012416 |
| *S. humboldti* | 1.00000 | -0.016275 | 0.151114 | -0.003572 | 0.025483 | 0.014327 | -0.010148 |

| B1  | B2  | B3  | B4  | B5  | B6  | B7  |
|-----|-----|-----|-----|-----|-----|-----|
| 0.00000 | -0.066369 | 0.077434 | -0.046354 | 0.017405 | 0.010943 | -0.002811 |
| 0.00000 | -0.138297 | 0.099830 | -0.060066 | -0.007940 | 0.021092 | 0.033620 |
| 0.00000 | -0.294488 | 0.028276 | -0.000809 | 0.005157 | -0.068304 | -0.014133 |
| 0.00000 | -0.053677 | 0.073756 | -0.049932 | 0.012276 | 0.021648 | 0.007629 |
| 0.00000 | -0.166471 | 0.027444 | 0.009916 | -0.020699 | -0.026779 | 0.022776 |
| 0.00000 | -0.024607 | 0.032631 | -0.007803 | -0.002715 | 0.001851 | 0.016961 |
| 0.00000 | 0.056545 | 0.007133 | 0.003373 | 0.002915 | 0.005710 | 0.004768 |
| 0.00000 | -0.130001 | 0.045129 | -0.018012 | -0.015749 | -0.011905 | 0.035785 |
| 0.00000 | 0.058072 | -0.039889 | -0.003574 | 0.012208 | 0.018226 | -0.026498 |

| C1  | C2  | C3  | C4  | C5  | C6  | C7  |
|-----|-----|-----|-----|-----|-----|-----|
| 0.00000 | -0.046828 | -0.016487 | 0.008143 | -0.007220 | 0.020121 | 0.000854 |
| 0.00000 | -0.069232 | -0.059733 | 0.007309 | -0.002458 | 0.006201 | 0.008297 |
| 0.00000 | -0.041711 | -0.144774 | -0.097485 | 0.001300 | -0.027899 | 0.005981 |
| 0.00000 | -0.062813 | -0.031994 | 0.010307 | -0.001180 | 0.014191 | 0.004156 |
| 0.00000 | -0.068977 | -0.059783 | -0.025213 | 0.006385 | -0.013783 | 0.017435 |
| 0.00000 | -0.021207 | -0.050750 | 0.017365 | 0.009700 | 0.001596 | 0.006169 |
| 0.00000 | 0.025050 | -0.053472 | 0.021913 | 0.012772 | 0.006802 | -0.001025 |
| 0.00000 | -0.066140 | -0.081260 | -0.007872 | 0.008306 | -0.005627 | 0.017985 |
| 0.00000 | 0.002107 | -0.051842 | -0.007601 | 0.006532 | -0.004929 | 0.011913 |

| D1  | D2  | D3  | D4  | D5  | D6  | D7  |
|-----|-----|-----|-----|-----|-----|-----|
| -0.347374 | -0.170325 | -0.255344 | 0.095882 | 0.009473 | -0.010230 | -0.004098 |
| -0.312126 | -0.121268 | -0.270959 | 0.045558 | 0.025827 | -0.017454 | 0.004731 |
| -0.304816 | 0.163042 | -0.171745 | -0.062039 | -0.007063 | 0.016434 | -0.003140 |
| -0.353861 | -0.133744 | -0.264986 | 0.082370 | 0.023491 | -0.018382 | 0.002992 |
| -0.309237 | 0.080586 | -0.257265 | -0.054735 | 0.012556 | 0.023382 | -0.008356 |
| -0.313182 | -0.076122 | -0.231917 | 0.031058 | 0.040847 | -0.008925 | -0.030261 |
| -0.360038 | -0.126847 | -0.207726 | 0.053627 | 0.038066 | -0.023022 | -0.020622 |
| -0.296603 | 0.008250 | -0.262423 | -0.022735 | 0.032945 | 0.007891 | -0.007402 |
| 0.277739 | -0.028478 | 0.250630 | 0.040632 | -0.022270 | -0.014213 | 0.012320 |
### APPENDIX 3 (CONTINUED)

|    | A8   | A9    | A10  | A11  | A12   | A13    | A14   | A15   | A16   |
|----|------|-------|------|------|-------|--------|-------|-------|-------|
| B8 | -0.018763 | -0.015603 | -0.003158 | -0.004909 | 0.002695 | 0.001318 | 0.001283 | 0.003309 | -0.001630 |
|    | -0.032812 | 0.005399 | 0.013294 | -0.002141 | -0.000020 | 0.001276 | 0.001373 | -0.001894 | 0.002875 |
|    | 0.010811 | 0.014850 | 0.003575 | -0.001014 | -0.004365 | -0.001045 | 0.004444 | 0.000404 | -0.004292 |
|    | -0.019109 | 0.014112 | -0.000065 | -0.000862 | -0.002091 | -0.001518 | -0.000536 | 0.000557 | -0.001122 |
|    | 0.020568 | -0.003704 | -0.008676 | -0.001494 | 0.008097 | -0.00214 | -0.001406 | -0.002045 | 0.002143 |
|    | -0.013088 | -0.001741 | 0.003595 | 0.003090 | -0.006064 | 0.001524 | 0.003434 | -0.000157 | -0.003918 |
|    | -0.008252 | -0.001638 | 0.003728 | -0.001322 | -0.006307 | 0.002722 | 0.004228 | -0.004580 | -0.001935 |
|    | -0.001990 | -0.011731 | -0.005512 | 0.008245 | 0.001866 | -0.003963 | -0.000144 | 0.000332 | -0.000858 |
|    | -0.005246 | 0.007524 | 0.006890 | -0.006692 | -0.000753 | -0.000193 | -0.001932 | -0.000648 | 0.002764 |
| C8 | -0.004366 | 0.008512 | -0.007555 | -0.006564 | -0.005315 | 0.009467 | -0.001163 | -0.000402 | 0.000001 |
|    | -0.024647 | 0.013394 | 0.003694 | 0.004299 | -0.009530 | 0.004056 | 0.002838 | -0.001564 | 0.000548 |
|    | 0.012229 | 0.008024 | -0.004609 | -0.001396 | 0.002895 | -0.003295 | -0.001562 | -0.000113 | -0.000362 |
|    | -0.012077 | 0.009136 | 0.003254 | -0.002831 | -0.004192 | 0.010117 | -0.000183 | -0.000489 | -0.001362 |
|    | 0.000104 | -0.002382 | -0.005083 | -0.001823 | 0.008194 | 0.001727 | -0.001223 | -0.002544 | 0.002664 |
|    | -0.003653 | -0.000998 | -0.002546 | 0.002437 | -0.002591 | 0.002792 | 0.003591 | 0.000632 | -0.004039 |
|    | 0.010684 | 0.000741 | -0.009855 | 0.002205 | -0.000010 | -0.000711 | 0.000757 | 0.000345 | -0.000243 |
|    | -0.005916 | -0.005490 | -0.002383 | 0.006657 | 0.005134 | -0.004296 | -0.003312 | 0.004913 | 0.002217 |
|    | 0.004353 | -0.001923 | -0.007939 | 0.003999 | 0.004739 | -0.003451 | -0.006443 | 0.002955 | 0.004493 |
| D8 | -0.011272 | 0.002846 | -0.001170 | -0.000407 | 0.000812 | 0.008404 | -0.002065 | -0.001662 | 0.000248 |
|    | -0.011198 | 0.003294 | -0.000476 | -0.003636 | 0.000417 | 0.002301 | -0.002516 | 0.003440 | -0.000871 |
|    | -0.011933 | -0.001645 | -0.003214 | 0.009410 | 0.005423 | -0.000944 | -0.004952 | -0.002464 | 0.001424 |
|    | -0.011835 | 0.001669 | 0.007104 | -0.003379 | 0.001233 | -0.000818 | -0.002826 | -0.001700 | -0.001279 |
|    | -0.006632 | 0.004342 | -0.006716 | 0.002375 | -0.003592 | -0.003216 | 0.004734 | 0.001342 | -0.001863 |
|    | 0.004306 | -0.002327 | 0.001054 | 0.003229 | 0.001235 | 0.000249 | 0.000688 | 0.000810 | -0.001537 |
|    | 0.014305 | -0.004900 | -0.004491 | 0.001323 | 0.007238 | -0.001710 | -0.002310 | -0.000106 | 0.001007 |
|    | -0.000790 | -0.006040 | -0.005224 | 0.004162 | -0.000308 | 0.000077 | -0.000052 | -0.001035 | -0.001808 |
|    | 0.000358 | 0.006046 | 0.008487 | -0.003312 | -0.004243 | -0.000536 | 0.000058 | -0.002739 | 0.000267 |
APPENDIX 4.

Scatterplots of scores from the Principal Component Analysis (PCA) on harmonic coefficients calculated for outlines of major hip bone foramina (source data in Appendix 3).
APPENDIX 5.

Left hip bone of an extant penguin *Eudyptula minor* (NHMUK/T 1881.1.17.105) showing the ilio-ischiadic membrane (lateral view). Abbreviations: at, antitrochanter; iff, ilioischiadic foramen; iim, ilioischadic membrane; il, ilium; is, ischium; pb, pubis; pdp, correlate of proximal dorsal process; pfe, proximal femur (broken). Scale bar equals 1 cm.