Pinniped (Mammalia: Carnivora) fossils from Black Rock, a new late Neogene vertebrate locality in Victoria, Australia

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ABSTRACT

The fossil record of true seals (Family Phocidae) is notoriously poorly preserved, most notably in the Southern Hemisphere. This fossil record bias has made it difficult to assess whether populations of phocid species in the Southern Hemisphere remained resident in particular regions throughout deep time, with only the eastern South Pacific preserving a near-continuous fossil record. Here we report two new seal fossil remains from the late Miocene-early Pliocene of southeastern Australia, discovered from a new site, Black Rock. This is the fourth phocid fossil site in Australia, and the seventh in Australasia. The two fossils are referable to the clades Pinnipedia and Monachinae. Combined with other recent fossil phocid finds in Australasia, this suggests that true seals were continuously present in the region at least throughout the Pliocene. This lends further support for the importance of Australasia in the global evolution of true seals.

INTRODUCTION

Since the early Miocene, true seals (family Phocidae) have dispersed around the world, while the remaining groups of pinnipeds spent most of their evolutionary history in the North Pacific (Berta et al., 2018). True seal fossils have been found on the margins of the North Pacific, North Atlantic, Paratethys, South Pacific, and South Atlantic
Rule & Fitzgerald: Black Rock pinniped fossils

oceans. As a result, investigations into the biogeographic history of phocids through the expansion of this global fossil record is one of the key areas of investigation in contemporary pinniped palaeontology.

Many fossil phocids have been found in the Northern Hemisphere (Valenzuela-Toro and Pyenson, 2019), and studies of new fossils often focus on expanding the geographic range of extinct species across oceans (Koretsky and Ray, 2008; Rahmat et al., 2018). However, cosmopolitan records are usually based off fragmentary fossils (Valenzuela-Toro and Pyenson, 2019) and often are not taxonomically stable (Dewaele et al., 2018; Churchill and Uhen, 2019; Rule et al., 2020a). With such a focus on expanding the geographic range of extinct species, there has been relatively little attention on expanding the chronological range of extinct true seals; in other words, whether populations of a fossil taxon in a particular area were resident in the region throughout deep time. The best-known example of long-term occupancy is that of true seal species from Miocene sites in Peru and Chile (Muizon, 1981; Walsh and Naish, 2002; Valenzuela-Toro et al., 2016; Bosio et al. 2021), which suggest an established presence of populations of the monospecific genera *Piscophoca*, *Acrophoca*, and *Australophoca* in the eastern South Pacific. These sites are located in the Southern Hemisphere, a region that has historically been neglected in pinniped palaeontology (Valenzuela-Toro and Pyenson, 2019).

Recent research has expanded upon the fossil record of true seals in another region in the Southern Hemisphere; Australasia (Fitzgerald, 2005; Boessenecker and Churchill, 2016; Rule et al., 2020b; Rule et al., 2020c; Rule et al., 2020d; Rule et al., 2021a). It is becoming clear that Australasia was a critical region in the evolution of the monachines (the southern true seals), with many modern groups, such as monk seals and elephant seals, originating there (Rule et al., 2020c). However, the distribution of true seals in the region throughout deep time has received little attention, and it is currently unclear if true seal populations in Australasia were resident breeding populations or represent migrants and one-off dispersals to the region. Here we describe two phocid fossils from a new site, Black Rock, in south-eastern Australia, and discuss the possibility of long-term resident true seal populations in Australasia during the past.

### MATERIALS AND METHODS

The two fossils (NMV P254178 and NMV P254995) are housed in the collections of Museums Victoria (NMV), Melbourne, Victoria, Australia. Measurements were taken to the nearest 0.01 mm using digital callipers and are reported in Table 1. To assist with illustrating preserved morphology, NMV P254995 was scanned using an Artec Space Spider surface scanner and meshed using Artec Studio 15 (Artec Group, Luxembourg). Comparative specimens (Table 2) were studied from the mammalogy collection of Museums Victoria, as well as the Natural History Museum London (NHMUK, United Kingdom), the Smithsonian Institution (USNM, Washington D.C., United States of America), and the palaeontology collection of Muséum national d'Histoire naturelle (MNHN, Paris, France).

### LOCALITY AND AGE

The pinniped fossils were collected ex situ on the seabed, below low-tide mark, off Black Rock Beach, Black Rock, Port Phillip Basin, Victoria, Australia; near 37°58'26" S, 145°00'48" E (Figure 1; precise locality coordinates available from Museums Victoria). Black Rock is situated on the eastern shoreline of Port Phillip Bay, about 18 km southeast of the central business district of Melbourne, and approximately 3 km northwest of the Beaumaris Bay fossil locality (Figure 1). Along the Black Rock coastline, the shallow marine upper Miocene–lower Pliocene Black Rock Sandstone (=

| Measurement | Metric (mm) |
|-------------|-------------|
| NMV P254995 |             |
| Total length of mandible | 103.97 |
| Thickness of mandibular body at m1 | 12.74 |
| Length of preserved toothrow | 29.95 |
| Depth under m1 | 22.87 |
| Depth of mandibular body | 23.32 |
| Height of preserved coronoid fossa | 36.84 |
| NMV P254178 |             |
| Total width | 17.14 |
| Total length | 72.01 |
| Width of mid-shaft | 11.56 |
| Width of base dorsal-ventral | 13.17 |

**TABLE 1.** Measurements from the two fossil specimens from the Sandringham Sandstone, Black Rock, Victoria, Australia.
**TABLE 2.** List of comparative specimens referred to in study.

| Taxon                      | Specimen Number | Institution | Sex |
|----------------------------|-----------------|-------------|-----|
| *Acrophoca longirostris*   | MNHN-F-SAS-563  | MNHN        | ?   |
| *Cystophora cristata*      | NHMUK 1907.9.4.4| NHML        | ?   |
| *Erignathus barbatus*      | NHMUK 1908.11.26.1| NHML | ?   |
| *Hadrokirus martini*       | MNHN-F-SAS-16276| MNHN        | ?   |
| *Halichoerus grypus*       | NHMUK 1961-5-18-11| NHML | Male|
| *Hydrurga leptonyx*        | NMV C7375       | NMV         | Female|
| *Leptonychotes weddellii*  | NMV C25094      | NMV         | Male|
| *Lobodon carcinophaga*     | NMV C7385       | NMV         | Male|
| *Mirounga leonina*         | NMV C7407       | NMV         | Male|
| *Monachus monachus*        | USNM 219059     | USNM        | ?   |
| *Neomonachus schauinslandi*| USNM 243839     | USNM        | Female|
| *Neomonachus tropicalis*   | USNM 100354     | USNM        | Male|
| *Ommatophoca rossii*       | NMV C7354       | NMV         | ?   |
| *Pagophilus groenlandica*  | NHMUK 1939.12.10.2| NHML | ?   |
| *Phoca vitulina*           | NHMUK 1919.7.7.3260| NHML | ?   |
| *Piscophoca pacifica*      | MNHN-F-SAS-564  | MNHN        | ?   |
| *Pusa hispida*             | NHMUK 1938.11.26.6| NHML | Male|

**FIGURE 1.** The locality of the fossil site at Black Rock. Panels show A) Australasia, B) the Australian state, Victoria, C) Port Phillip Bay, D) Bayside, including Beaumaris and Black Rock.
Sandringham Sandstone: VandenBerg, 2016) is exposed as a shore platform at: the northern end of Black Rock Beach, southeast of Black Rock Point (37°58′16.99″S, 145°00′39.87″E); and further south at Quiet Corner (37°58′54.85″S, 145°01′04.52″E) (Gill, 1957; Ter and Buckeridge, 2012; VandenBerg, 2016). At the latter localities, the lithology in intertidal outcrop consists of sparsely fossiliferous, non-calcareous, ferruginous sandstone containing invertebrate burrows infilled by goethite (Gill, 1957; Abele et al., 1988; Ter and Buckeridge, 2012; Vandenberg, 2016). This lithology represents the top ~8.5 m of the Black Rock Sandstone (Abele et al., 1988). Stratigraphically underlying that top half of the Black Rock Sandstone, the lower ~6.5 m of the formation comprises fossiliferous calcareous sandstone, with a <0.3 m thick phosphatic nodule bed marking the base of the unit (Abele et al., 1988). This lower half of the Black Rock Sandstone is best exposed on- and offshore along Beaumaris Bay (see stratigraphic column in figure 1 of Rule et al., 2020b). The middle–upper Miocene Fyansford Formation (grey calcareous clay) unconformably underlies the Black Rock Sandstone in the Port Phillip Basin (Mallett and Holdgate, 1985).

Sedimentary matrix adhering to the fossils described here comprises Fe-stained orange-light brown fine calcareous sandstone, containing rounded quartz grains and microfossils (including mollusc shell fragments). This associated matrix is identical to lithology within the lower 6.5 m of the Black Rock Sandstone but can be differentiated from the sediments of the basal nodule bed, which are distinctly yellow, phosphatic, and rich in quartz pebbles (Dickinson and Wallace, 2009; Rule et al., 2020b). It is, therefore, deduced that: (1) the pinniped fossils originated from a horizon within the lower 6.5 m of the Black Rock Sandstone, but stratigraphically above the basal phosphatic nodule bed; and (2) that the stratigraphic section at Black Rock Beach includes at least part of the lower 6.5 m of the Black Rock Sandstone, exposed as offshore submarine outcrop.

The Black Rock Sandstone was deposited between the late Miocene (Messinian) and early Pliocene (Zanclean) (Fitzgerald and Kool, 2015; Rule et al., 2020b). Measured $^{87}$Sr/$^{86}$Sr isotope ratios of francolite in phosphates from the basal nodule bed gave estimated ages of 6.24–5.38 Ma, suggesting a maximum age for the Black Rock Sandstone of latest Miocene (Dickinson and Wallace, 2009). $^{87}$Sr/$^{86}$Sr ratios from molluscan/bra-chiopod carbonate at horizons approximately 1.0 m and 2.6 m above the basal nodule bed, produced ages of 5.98–5.59 Ma (late Miocene) and 4.91 Ma (early Pliocene), respectively (Dickinson and Wallace, 2009). These data indicate that the Miocene-Pliocene boundary lies within the lowest 2.6 m of the Black Rock Sandstone. However, due to the scarcity, if not absence, of age-indicative fossils in the upper ~12 m of the Black Rock Sandstone, its minimum age is unresolved. Within the Port Phillip Basin, basalts of the Newer Volcanics Province overlie the Neogene marine succession (including the Black Rock Sandstone) (Abele et al., 1988; Wallace et al., 2005). The minimum radiometric age for the commencement of this volcanism is 4.62 Ma (Hare et al., 2005). VandenBerg (2016) inferred that the onset of this volcanism coincided with the end of Neogene marine deposition in the Port Phillip Basin (including the Black Rock Sandstone) but cautioned that the stratigraphically lowest basalt flows have not been dated. Nevertheless, we concur with VandenBerg (2016) in using these data to conservatively estimate deposition of the Black Rock Sandstone between 6.24 and 4.62 Ma; with the Black Rock section of the Black Rock Sandstone (and pinniped fossils described herein) deposited between ~5.98 and 4.62 Ma, late Miocene to early Pliocene.

Until now, the vertebrate palaeontology of the Black Rock Sandstone has been almost exclusively represented by the rich Beaumaris Local Fauna of Beaumaris Bay, comprised of: chondrichthyan and actinopterygian fish (Kemp, 1991; Fitzgerald, 2004); marine and freshwater turtles (Fitzgerald and Kool, 2015; Rule et al., 2021b); penguins (Park and Fitzgerald, 2012a), pelagornithid birds (Fitzgerald et al., 2012), an albatross (Wilkinson, 1969), a dromornithid (Park and Fitzgerald, 2012b); marsupials (Piper et al., 2006); a sirenian (Fitzgerald et al., 2013); phocid seals (Rule et al., 2020b, 2021a); and cetaceans (Fordyce, 1982; Fitzgerald, 2004, 2012; Marx et al., 2018). The only prior documentation of a vertebrate fossil from Black Rock is a zygomaturine diprotodontid left maxilla (erroneously reported as a dentary from Beaumaris) figured by Vickers-Rich (1993:177) and Warne et al. (2003:638). In addition to the pinniped material described here, recently collected vertebrate fossils from the Black Rock locality include isolated specimens of Diprotodontidae, Cetacea, Spheniscidae, Pelagornithidae, and chondrichthyans, suggesting general equivalence between the Black Rock assemblage and the Beaumaris Local Fauna.
SYSTEMATIC PALAEONTOLOGY

Order CARNIVORA Bowdich, 1821
Suborder PINNIPEDIA Illiger, 1811

Referred specimen. NMV P254178, an isolated phalanx collected by B.S.J. Francischelli in 2019 (Figure 2).

Diagnosis. An intermediate or proximal phalanx from the manus (from either digits II, III, or IV) of a pinniped, due to the straight medial and lateral surfaces, a distal end narrower than a proximal end, and a slightly concave ventral surface.

Description. The proximal end is rounded, and is missing the articular surface (Figure 2G-J). The distal surface is trochleated (Figure 2G-H), and larger medio-laterally than dorso-ventrally. The shaft is sub-circular in cross section. Due to the lack of diagnostic features beyond Pinnipedia, we are unable to make meaningful comparisons to other species.

Family PHOCIDAE Gray, 1821
Subfamily MONACHINAE Gray, 1869

Referred specimen. NMV P254995, a fragmentary right mandible collected by B.S.J. Francischelli in 2019 (Figure 2).

Diagnosis. A fragmentary right mandible based on the position of the masseteric fossa. Referred to Pinnipedia based on a transversely flattened mandibular body, and a coronoid crest that does not ascend vertically from the toothrow. Referred to Phocidae based on a double rooted m1 with clearly separated anterior and posterior alveoli. Referred to Monachinae based on a shallow coronoid crest that ascends below a 45° angle.

Description. NMV P254995 is missing the proximal portion of the mandible, including the articular surface and a portion of the coronoid crest, and the distal portion from the p4 onwards. The body of the mandible is thick (12 mm) compared to the low height (23 mm), making the medial-lateral thickness roughly half of the dorsal-ventral length. The medial surface of the body is flat compared to the lateral surface, which is slightly dorso-ventrally curved (Figure 2C). While the preserved body of the mandible is antero-posteriorly straight, the coronoid crest curves medially. The masseteric fossa
is shallow (Figure 2E), and barely visible on the lateral surface of the coronoid. The coronoid crest ascends shallowly from the toothrow (<45°), with a short space separating the crest from the posterior margin of the toothrow. Both alveoli of the m1 and the preserved posterior alveolus of the p4 are round, with the p4 alveolus slightly larger than the m1 (Figure 2F). A short diastema separates the m1 and p4. The posterior portion of the anterior alveolus of the p4 is preserved (Figure 2F).

**Comparison.** The shallow coronoid crest of NMV P254995 is similar to that condition seen in Neomonachus, Mirounga, Acrophoca, and Lobodontini, and is therefore mostly consistent with Monachinae (Phocinae have a steep coronoid crest). The only exceptions are Monachus, Homiphoca, Piscophoca, and Hadrokirus. The toothrow is separated from the coronoid crest by a short space (less than twice the length of the m1), similar to the monachines Lobodon, Hydrurga, Hadrokirus, and tribe Monachini; and similar to phocines such as Erignathus, Halichoerus, Phoca, and Pusa.

The body of the mandible NMV P254995 below the toothrow is thick, similar to Monachini and Hadrokirus. All other phocids, including other monachines, have either thin mandibles below the toothrow (width less than half the height), or mandibles that grade from thin posteriorly to thick anteriorly. While most phocids have a mandible that is fairly straight antero-posteriorly, NMV P254995 has a slightly curved mandible similar to Ommatophoca, Neomonachus, and Erignathus. Overall, the preserved morphology of NMV P254995 has the most similarities with the genus Neomonachus.

Two other fragmentary fossil pinniped mandibles are known from Australia: NMV P251642 (Beaumaris: Rule et al., 2020b) and NMV P218465 (Portland: Fitzgerald, 2005). The shallow coronoid crest and round profile of the posterior m1 alveolus of NMV P254995 is shared with NMV P251642 and NMV P218465. Further, several additional characters are shared between NMV P254995 and NMV P218465: a posterior p4 alveolus larger than both m1 alveoli (also in common with other phocids), and an m1 alveoli at a slightly imbricated angle compared to the p4 alveoli. There are no discernible differences between NMV P254995, NMV P251642, and NMV P218465. The similarities between NMV P254995 and NMV P218465 are enough to confidently revise the identification of NMV P218465 to Phocidae cf. Monachinae, confirming the tentative diagnosis of Phocidae of Fitzgerald (2005).

**DISCUSSION**

True seals are currently thought to have originated in either the North Atlantic or Paratethys, where they are known from multiple localities covering most of the Neogene (Berta et al., 2018). However, in the North Pacific and the Global South, true seals are usually only known from very few sites. The best record in the Southern Hemisphere is the Neogene of the eastern South Pacific, with multiple fossil sites from Peru and Chile, preserving multiple species from several stratigraphic units (Muizon, 1981; Walsh and Naish, 2002; Amson and Muizon, 2014; Valenzuela-Toro et al., 2016; Benites-Palomino et al., 2022). This is potentially sufficient evidence that phocids were resident to the eastern South Pacific during the late Miocene to early Pliocene. Outside of this region, the record of phocids is sparse, (Valenzuela-Toro and Pyenson, 2019). For example, Argentina has two species from two formations (Muizon and Bond, 1982; Cozzuol, 2001) dated to the middle-late Miocene, and South Africa has many fossils of one species represented from a few early Pliocene sites (Hendey and Repenning, 1972; Muizon and Hendey, 1980; Govender, 2015). As a result, not many biogeographic regions (other than the eastern South Pacific) have phocid fossils present across a wide span of time and space.

Despite the apparent constraints on the global fossil record of true seals (Valenzuela-Toro and Pyenson, 2019), Australasia is emerging as a significant region in true seal evolutionary history due to an increase of reporting of fossil finds (Boessenecker and Churchill, 2016; Rule et al., 2020b; Rule et al., 2020c; Rule et al., 2020d; Rule et al., 2021a). These new finds span from the late Miocene to early Pleistocene, from both southeastern Australia and the north island of New Zealand. Together with the new fossils reported here from Black Rock, these records suggest that phocids may have had resident breeding populations in Australasia throughout the Pliocene.

Until recently, it was unknown which ancient true seals were present in Australasia during the Neogene, with the most specific identification being Monachinae (Fordyce and Flannery, 1983). Since then, recent studies have revealed that monk seals (tribe Monachini) existed in the Neogene of Australasia (Figure 3). Specifically, monk seal temporals are known from Beaumaris and Hamilton in Australia (Rule et al., 2021a), and skulls are known from Taranaki in New Zealand (Rule et al., 2020c). While none of the Black Rock specimens are referable to Monachini, it is notable that NMV P254995
is most similar to the genus *Neomonachus*. The presence of monk seals in Australasia, from at least three sites spanning a time period of over 3 million years, is unexpected and unprecedented, as monk seals were traditionally considered a northern clade. This implies that Austral latitudes presented ideal environmental conditions for monk seal fauna to inhabit during at least the Pliocene.

These records seem to indicate that monk seals were widespread around Australasia during the Pliocene, only to go extinct during the marine megafauna extinction (3.8–2.4 Ma; Boessenecker, 2013; Pimiento et al., 2017). This extinction coincided with the late Pliocene pinniped turnover events (Avery and Klein, 2011; Valenzuela-Toro et al., 2013; Rule et al., 2020d; Benites-Palomino et al., 2022), which saw the disappearance of most archaic phocids in the Southern Hemisphere. After this extinction, fossils representing lobodontins (Antarctic true seals) and miroungins (elephant seals) are known from the early Pleistocene of New Zealand (King, 1973; Boessenecker and Churchill, 2016). It is remarkable that the three tribes of Monachinae (monk, elephant, and Antarctic true seals) were present so close in time to each other in Australasia (Figure 3). *Eomonachus* (the New Zealand monk seal) is separated from the Napier Ross seal and Waipunga elephant seal by only 500,000–600,000 years (Figure 3; King, 1973; Boessenecker and Churchill, 2016). While this represents a very narrow separation in geological time, the two occurrences are separated from each other by the marine megafauna extinction (Pimiento et al., 2017). Furthermore, monk, ele-
phant, and Antarctic seals have distinct ecologies and life histories, which influence their geographic distribution today. As such, there is no reason to postulate that the three tribes once co-occurred in the same fauna. In addition, elephant and Antarctic seals co-occur in the Southern Ocean today, so the early Pleistocene records of Lobodontini and Miroungini in New Zealand may simply represent the earliest onset of the Southern Ocean fauna (Figure 3). There is also the possibility that this may indicate another cryptic pinniped turnover event in New Zealand during the early Pleistocene (Figure 3), although additional fossils of these phocids should be discovered to ensure they were resident in the region (either for breeding or hauling-out to rest).

By the Late Pleistocene (Figure 3), true seals were no longer the dominant pinnipeds in Australasia. By then, fur seals and sea lions had dispersed south from the North Pacific Ocean (Yonezawa et al., 2009; Churchill et al., 2014) and become resident in Australia (McCoy, 1877) and New Zealand (Churchill and Boessenecker, 2016). The only true seals present in the region were Southern elephant seals (*Mirounga leonina*) in Bass Strait (Bryden et al., 1999), which remained until they became locally extinct from overhunting. This was reflective of the post-pinniped turnover fauna present elsewhere in the Southern Hemisphere, including South Africa (Avery and Klein, 2011), South America (Valenzuela-Toro et al., 2015), and sub-Antarctic islands (Berta et al., 2018). This modern pinniped fauna relied on cooler, more productive waters near the southern continents. Without a doubt, a transition from a more archaic pinniped fauna to the modern pinniped fauna occurred sometime between the Late Pliocene and Late Pleistocene.

True seals are now known from four fossil sites in Australia and three fossil sites in New Zealand. These sites encompass the entirety of the Pliocene and the beginning of the Pleistocene, indicating that exploration of other sites representing this time period in Australasia will yield more phocid fossils. We recommend further exploration of late Miocene and early-middle Pleistocene aged marine fossil sites in Australasia, as further finds of fossil phocids would expand the chronological range of the group in this region. By extension, Pleistocene marine fossil sites in Australasia could further expand the fossil record of otariids in the region. Therefore, any pre-Holocene pinniped fossils will be critical for refining the timing of the pinniped turnover in Australasia.

**CONCLUSIONS**

The fossil record of phocids from Australasia has greatly expanded in recent years. As a result, it seems likely that true seals were resident in the region throughout the Pliocene period. This confirms Australasia as an important region for true seal evolution in the past, with the potential to inform about the diversification and ultimate extinction of ancient true seals in the Southern Hemisphere.

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