

## AN ASSESSMENT OF MARINE RESERVOIR CORRECTIONS FOR RADIOCARBON DATES ON WALRUS FROM THE FOXE BASIN REGION OF ARCTIC CANADA

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**ABSTRACT.** Archaeological sites in the Canadian Arctic often contain substantial quantities of marine mammal bones and in some cases completely lack terrestrial mammal bones. A distrust of radiocarbon (<sup>14</sup>C) dates on marine mammal bones among Arctic archaeologists has caused many sites to be insufficiently dated. The goal of this study was to investigate the marine reservoir effect on Atlantic walrus in the Foxe Basin region of the Canadian Arctic through a two-pronged approach: dating of live-harvested specimens of known age collected prior to AD 1955 and dating of pairs of animal remains (walrus and caribou) from stratigraphically contemporaneous levels within archaeological features. <sup>14</sup>C dates on pre-bomb, live-harvested walrus indicate that a  $\Delta R$  value of  $160 \pm 50$  yr be used in calibrating dates on walrus from this region. These results differed significantly from a similar set of pre-bomb mollusks, which argues against applying mollusk-based corrections to marine mammals. The results of comparative dating of caribou and walrus from archaeological features provided maximum estimates of reservoir ages that were more varied than the directly measured ages. Although about half of inferred  $\Delta R$  values overlap the museum specimen results, the others indicate that the assumption of contemporaneity does not hold true.

**KEYWORDS:** archaeology, arctic, marine mammals, marine reservoir effect.

### INTRODUCTION

Archaeological sites in Arctic North America and Greenland are rich in sea mammal remains. Sites in the region encompassing Foxe Basin, northern Hudson Bay, and Hudson Strait are particularly rich in walrus (*Odobenus rosmarus*) remains, and many of them contain little or no terrestrial mammal remains. Despite excellent preservation, many sites go undated because of a pervasive distrust of dates on marine materials among archaeologists. The distrust arises from inadequate documentation of the size and regional variation of the marine reservoir effect on radiocarbon (<sup>14</sup>C) dates across the Arctic. This effect, on the order of centuries, is large with respect to site ages (less than 5000 years). For example, using the data presented by McNeely et al. (2006), Coulthard et al. (2010) calculated average  $\Delta R$  values (<sup>14</sup>C yr BP) with some minor modifications to the original geographic boundaries as presented in the McNeely et al. dataset:  $65 \pm 60$  yr (Hudson Strait),  $110 \pm 65$  yr (Hudson Bay),  $145 \pm 95$  yr (Ungava Bay),  $150 \pm 60$  yr (SE Baffin Island),  $220 \pm 20$  yr (NE Baffin Island),  $310 \pm 90$  yr (Foxy Basin),  $335 \pm 85$  yr (NW Canadian Arctic Archipelago), and  $365 \pm 115$  yr (James Bay). Following an early period of indiscriminate dating of various materials and inconsistent normalization of dates for isotopic fractionation among laboratories, McGhee and Tuck (1976) exposed inconsistencies in archaeological site age assessments that resulted when dates on terrestrial materials and uncorrected dates on marine materials were considered to be equally valid. The subsequent decision to avoid the dating of marine materials by most archaeologists, while perhaps rendering site ages less problematic, has hampered the ready dating of many sites, especially those not extensively excavated, resulting in poorly controlled chronologies in many instances.

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A marine reservoir age is essentially the residence time of CO<sub>2</sub>, and the bicarbonate formed from it, in the ocean. It is the length of time spent in transit in the ocean between the time of absorption of the gas from the atmosphere and its ventilation back to the atmosphere at sites of ocean upwelling. During this transit time, the radioactive isotope of carbon (<sup>14</sup>C) decays, imparting an “apparent age” to the carbon in the water. The global average surface ocean (0–75 m depth, which mixes by wave action) reservoir age was about 400 years in AD 1830, the value R in <sup>14</sup>C calibration programs for marine samples (Stuiver and Braziunas 1993a), but it varied through time because of variable <sup>14</sup>C production rates. Therefore, a marine reservoir age refers to the difference in the <sup>14</sup>C content of a particular area of the surface ocean and the atmosphere at a specific point in time. Temporal variations in <sup>14</sup>C content, including that caused by nuclear testing, are much larger in the atmosphere than they are in the ocean (Stuiver and Braziunas 1993b) because new <sup>14</sup>C is produced only in the troposphere and stratosphere. As a result, a simple subtraction of a single value for marine samples of any type will not accurately reflect the age of the samples. The time variance of R in the ocean is calculated in the Stuiver and Braziunas (1993a) carbon cycle model based on the tree-ring derived variance of <sup>14</sup>C levels in the atmosphere. R also varies regionally due to oceanographic processes, including downwelling, upwelling and the seasonal duration of sea ice cover (Southon et al. 2002; Ascough et al. 2005). The regional departure from R is termed ΔR. Prior to 2006 there were only a limited number of measurements of reservoir ages available from the Canadian sectors of the Northwest Atlantic and Arctic Oceans (Mangerud and Gulliksen 1975). Then McNeely et al. (2006) used a large sample of marine mollusks to define the reservoir age and ΔR in Canadian waters. They established mean values for 12 major oceanographic regions based on coastal current configurations. These results showed a decreasing cascade of ages from the Pacific coast, through the Arctic Archipelago, to the Atlantic Coast. The rounded value of ΔR for the Foxe Basin region (including northern Hudson Bay), where surface water is derived mainly from the Arctic Ocean, is 295 ± 130 yr based on 34 samples (see below).

Marine mollusk shells are ostensibly the most straightforward materials to use in assessing reservoir ages, because bivalve mollusks are non-migratory and the carbonate in their shells is deposited in isotopic equilibrium with the bicarbonate in the surrounding water. The shells therefore capture the local reservoir age. Measuring reservoir ages should be a simple task of collecting shells and <sup>14</sup>C dating them to determine “apparent ages.” Unfortunately, nuclear bomb testing has spiked the atmosphere and the oceans with large amounts of artificial <sup>14</sup>C, beginning around AD 1955 in the Northern Hemisphere, with the effect becoming strong after AD 1958 (Hua and Barbetti 2004; Hua et al. 2013). Hence, today local reservoir ages can be determined only on materials collected live prior to significant bomb testing. Therefore, only material in museum collections can be used to resolve this problem.

A by-product of the work by McNeely et al. was the finding that deposit-feeding mollusks have larger apparent ages on calcareous substrates (not on acidic substrates) than do suspension-feeding mollusks. The ΔR values derived by McNeely et al. (2006), therefore, excluded dates on deposit feeders. England et al. (2013) examined this question further by comparing coupled dates on deposit and suspension feeders, and concluded that ages on deposit feeders can be as much as 2000 years too old. They speculated that the enhanced effect in deposit feeders was due to assimilation of bicarbonate from pore water derived in part from the carbonate sediment, which is <sup>14</sup>C dead, rather than entirely from freely circulating ocean water. Foxe Basin and Hudson Strait are floored by Paleozoic carbonate rocks. These issues aside, mollusks have never been used in the Arctic to directly date archaeological sites, because they are not known to be present as food refuse.

Establishing reservoir corrections for marine mammals is potentially more problematic. Firstly, mammals may be migratory (e.g., Sergeant 1965; Richard et al. 2001; Heide-Jørgensen et al. 2003), hence averaging spatial reservoir effects. Secondly, they may dive to feed at depths greater than the 75 m surface water layer (e.g., Boeuf et al. 1988; Laidre et al. 2003), although walrus usually forage at depths of less than 80 m (Wiig et al. 1993; Born and Knutsen 1997; Jay et al. 2001). Thirdly, different species have different diets (primarily shrimp, clams, and gastropods for bearded seals; primarily fish for ringed seals; primarily benthic invertebrates for walrus; see Naughton [2012] for details) and tissues are assimilated from food, with potentially different carbon fractionation effects. Walrus may also indiscriminately include deposit-feeding mollusks in their food. Hence, we need to determine if reservoir ages established for mollusks can be applied to mammals to apply these techniques to improve archaeological chronologies. This paper is the first focused attempt to establish reservoir corrections for a specific marine mammal taxon from the North American Arctic based on live-collected, pre-bomb specimens. Most attempts to quantify local marine reservoir effects in the Arctic have utilized mollusks of known age, but several have attempted to utilize marine mammals. Olsson (1980) estimated the marine reservoir effect for Sweden using a number of different pinniped and cetacean taxa with known dates of death. Dumond and Griffin (2002) used a paired-dating approach in which they examined marine mammal bone collagen and charcoal from contemporaneous archaeological contexts to derive local  $\Delta R$  values for the eastern Bering Sea region. Furze et al. (2014) estimated  $\Delta R$  for beluga on the basis of nine  $^{14}\text{C}$  dates (originally presented by Stewart et al. [2006]) from individuals distributed across the central and eastern Canadian Arctic. We present 28 new dates on walrus from the eastern Canadian Arctic and compare them to the previous dates on mollusks. We also present 35 coupled dates on caribou (*Rangifer tarandus*) and walrus from three archaeological site complexes in the region to see if the two methods produce compatible results.

## MATERIALS AND METHODS

We obtained a list of live-harvested walrus samples in the collections of the Canadian Museum of Nature and subsequently permission to examine them. Of the samples available to us, three were from Prince Charles Island in central Foxe Basin, 21 were from islands in northern Hudson Bay and western Hudson Strait, within or directly adjacent the Foxe Basin region of McNeely et al. (2006), and three were from farther east in central Hudson Strait (Figure 1). The three walrus from Prince Charles Island correspond with the modern distribution of the Foxe Basin stock or population (Born et al. 1995; COSEWIC 2006; Stewart 2008). These walruses are widely distributed throughout this area with their movements occurring on a seasonal basis due to changing ice conditions; the Foxe Basin stock appears to be isolated (Born et al. 1995). On the basis of lead isotopic measurements consistent with divergent long-term residency patterns, there may be some internal stock distinctions within Foxe Basin (Outridge and Stewart 1999), but genetic studies indicate a single interbreeding population (de March et al. 2002). The majority of the walrus samples were collected from individuals from areas associated with the presently recognized Northern Hudson Bay-Davis Strait (NHB-DS) stock (Born et al. 1995; COSEWIC 2006; Stewart 2008). Walruses occur throughout the year in Northern Hudson Bay and western Hudson Strait, but recent genetic and satellite tracking demonstrate that some of these animals move to west Greenland (Andersen et al. 2009; Dietz et al. 2014). As with the Foxe Basin stock, there may be some internal divisions in the NHB-DS stock (Outridge et al. 2003), but the internal population dynamics remain too poorly studied to formally define sub-groups (Stewart 2008).

We extracted subsamples of about 1 g of bone or ivory from 28 specimens, which are all available specimens collected at known localities and years before AD 1955. We also recorded information on sex, age, and size from archival notes and measured selected bone elements.

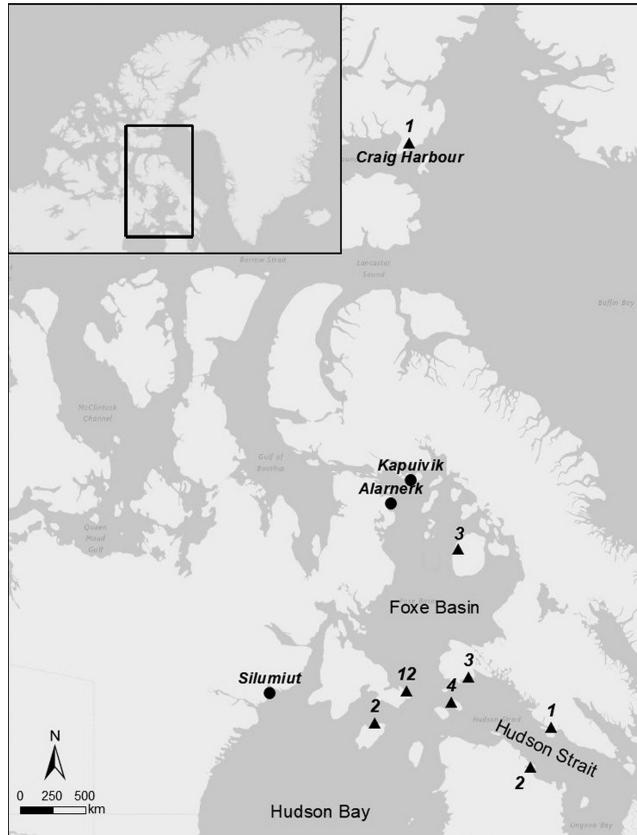


Figure 1 Map of the study area. Triangles denote areas where museum walrus specimens were collected; numbers adjacent to the triangles indicate the number of samples from each location. Archaeological sites from which paired caribou-walrus samples were obtained are indicated as circles.

All samples had been stored in plastic wraps in climate-controlled rooms and were in an excellent state of preservation. Collagen was extracted from bones (museum and archaeological) at either the Archaeology Isotope Laboratory at The University of British Columbia (UBC) or at the University of California Irvine (UCI) following Beaumont et al. (2010), ultrafiltered to obtain the >30 kDa fraction, converted to CO<sub>2</sub> and graphitized for AMS dating at The University of California at Irvine. Collagen quality was assessed from collagen yields (wt % of total bone mass) and by measuring C:N ratios (see Appendix in Supplementary Material). Carbon and nitrogen isotopic and elemental compositions were determined by CF-EA/IRMS using an IsoPrime continuous flow isotope-ratio mass spectrometer coupled to a Vario Micro elemental analyzer (Elementar, Hanau, Germany) (UBC) or a Finnigan Delta Plus isotope ratio mass spectrometer/Fisons NA1500NC elemental analyzer (UCI). Carbon and nitrogen isotopic compositions were calibrated relative to VPDB and AIR, respectively, using multiple well-characterized standard references materials. Measurement precision based on long-term reproducibility of standards was <0.1‰ for δ<sup>13</sup>C and <0.2‰ for δ<sup>15</sup>N at UCI. Standard uncertainty at UBC was determined to be ±0.20‰ for δ<sup>13</sup>C and ±0.17 for δ<sup>15</sup>N based on repeated analysis of internal and international standards, and sample replicates (Szpak et al. 2017). Elemental compositions

were calibrated using glutamic acid (USGS40) and internal methionine standards at UBC and known USGS24/IAEA N1 mixtures plus internal yeast and lysine standards at UCI. All  $^{14}\text{C}$  dates were conventionally normalized to  $\delta^{13}\text{C} = -25\text{‰}_{\text{VPDB}}$ .

During the pre-bomb collection period (AD 1885–1955),  $^{14}\text{C}$  ages were affected by changing levels of  $^{14}\text{C}$  in the atmosphere as a result of (1) short-term natural variations that were characteristic of the Holocene (cosmic ray flux modulated by varying earth and sun magnetic fields), and (2) since about AD 1900, the burning of fossil fuels, the industrial effect, which reached 100 years in 1955 (Mangerud et al. 2006). These effects are combined in the Stuiver and Braziunas (1993a) model to produce “model ages” representing the apparent  $^{14}\text{C}$  age of marine biomass for a given collection year with  $1\sigma$  errors (their Fig. 14; see also Stuiver et al. [1986:982]). The regional offset  $\Delta\text{R}$  is then the  $^{14}\text{C}$  age of a walrus minus the “model age” (McNeely et al. 2006). Here we use model ages from IntCal13 (Reimer et al. 2013), downloaded from the radiocarbon.org website. The error on  $\Delta\text{R}$  is the square root of the sum of squares of the individual  $1\sigma$  errors.

Another approach in estimating the reservoir effect is the comparison of dates on terrestrial and marine materials from contexts, either geological (e.g., Vickers et al. 2010; Ross et al. 2012) or archaeological (e.g., Southon et al. 1990; Southon et al. 1995; Yoneda et al. 2001; Dumond and Griffin 2002), wherein the materials are assumed to be contemporaneous. Following Stuiver and Braziunas (1993a), the terrestrial  $^{14}\text{C}$  ages are converted to equivalent surface water marine model ages (their Fig. 15 [updated in IntCal13]). The difference between the  $^{14}\text{C}$  age of the marine sample and the “marine model age” of the calibrated terrestrial sample is the estimated local  $\Delta\text{R}$ . Savelle and Dyke (2014) presented caribou (*Rangifer tarandus*) dates from Alarnerk, a major archaeological site complex in northwestern Foxe Basin; the site was further excavated by Howse in 2015. Here we present walrus dates from 12 Dorset features and 2 Thule Inuit features from the site complex and compare them to the caribou dates (18 pairs). We also present 12 pairs of dates on caribou and walrus from Dorset winter houses and associated middens at the Kapuvik site on Jens Munk Island. The Jens Munk sites were surveyed by Dyke in 2002 and Savelle and Dyke in 2003 (Savelle et al. 2009; Savelle and Dyke 2014). The new dates are from excavations by Desrosier and Kotar conducted in 2016. Finally we present five pairs of dates from the Thule Inuit Silumiut (KkJg-1) site in northwestern Hudson Bay (McCartney 1977). The caribou dates were calibrated using the CalPal online calculator and the error of a calibrated date is 68% of its range. The  $\Delta\text{R}$  values were calculated using Calib  $\Delta\text{R}$  application (<http://calib.org/deltar>) (Reimer and Reimer 2017). Where average  $\Delta\text{R}$  values are reported, these represent error-weighted means (Bevington 1969).

## RESULTS

### Museum Walrus Specimens

The  $^{14}\text{C}$  ages and the calculated  $\Delta\text{R}$  values for the live-collected walrus specimens are presented in Table 1, with co-ordinates for sample locations provided in the Appendix (see Supplementary Material). Two other samples, not used to calculate  $\Delta\text{R}$ , are statistical outliers. One is from Craig Harbour on SE Ellesmere Island at the head of Baffin Bay, far distant from the other samples considered here. The other is from western Hudson Strait. Its large apparent age and marginal C:N ratio led us to date a second collagen extract from bone chunks that were sonicated in acetone and methanol prior to decalcification, leading to a result accordant with the others. We suspect that a conserving agent was inadvertently included in the first collagen extract but was successfully removed by the solvent treatments applied to the second.

Table 1  $^{14}\text{C}$  ages of live-collected walrus from Foxe Basin and vicinity. Marine model ages from IntCal13.

UCIAMS #	$^{14}\text{C}$ age	Year collected	Marine model age	$\Delta\text{R}$
Central Hudson Strait				
168836	670 $\pm$ 15	1885	468 $\pm$ 23	202 $\pm$ 27
168857	630 $\pm$ 15	1886	467 $\pm$ 23	163 $\pm$ 27
168850	665 $\pm$ 15	1885	467 $\pm$ 23	198 $\pm$ 27
Average				188 $\pm$ 21
Western Hudson Strait and Northern Hudson Bay				
168831	705 $\pm$ 15	1928	452 $\pm$ 23	253 $\pm$ 27
168832	635 $\pm$ 15	1945	464 $\pm$ 23	171 $\pm$ 27
168837	615 $\pm$ 15	1953	472 $\pm$ 23	143 $\pm$ 27
168838	615 $\pm$ 15	1953	472 $\pm$ 23	143 $\pm$ 27
168839	585 $\pm$ 15	1953	472 $\pm$ 23	113 $\pm$ 27
168840	595 $\pm$ 15	1953	472 $\pm$ 23	123 $\pm$ 27
168841	570 $\pm$ 15	1953	472 $\pm$ 23	98 $\pm$ 27
168842	585 $\pm$ 15	1953	472 $\pm$ 23	113 $\pm$ 27
168843	575 $\pm$ 15	1953	472 $\pm$ 23	103 $\pm$ 27
168844	625 $\pm$ 15	1953	472 $\pm$ 23	153 $\pm$ 27
168845	585 $\pm$ 15	1953	472 $\pm$ 23	113 $\pm$ 27
168846	620 $\pm$ 15	1953	472 $\pm$ 23	148 $\pm$ 27
168847	590 $\pm$ 15	1953	472 $\pm$ 23	118 $\pm$ 27
168848	580 $\pm$ 15	1953	472 $\pm$ 23	108 $\pm$ 27
168849	590 $\pm$ 15	1954	473 $\pm$ 23	117 $\pm$ 27
168851	595 $\pm$ 15	1923	450 $\pm$ 23	145 $\pm$ 27
168852	660 $\pm$ 15	1924	450 $\pm$ 23	210 $\pm$ 27
168853	690 $\pm$ 15	1924	450 $\pm$ 23	240 $\pm$ 27
168855	670 $\pm$ 15	1924	450 $\pm$ 23	220 $\pm$ 27
168856	695 $\pm$ 15	1924	450 $\pm$ 23	245 $\pm$ 27
185717	625 $\pm$ 15	1928	452 $\pm$ 23	173 $\pm$ 27
Average				155 $\pm$ 50
Foxe Basin				
168833	685 $\pm$ 15	1949	468 $\pm$ 23	217 $\pm$ 27
168834	655 $\pm$ 15	1949	468 $\pm$ 23	187 $\pm$ 27
168835	595 $\pm$ 15	1949	468 $\pm$ 23	127 $\pm$ 27
Average				177 $\pm$ 46
Overall average				161 $\pm$ 48
Craig Harbour, Ellesmere Island				
168854	755 $\pm$ 15	1924	450 $\pm$ 23	305 $\pm$ 27
Outlier, Cape Dorset, Hudson Strait (redated 185717 above)				
168830	810 $\pm$ 15	1928	452 $\pm$ 23	358 $\pm$ 27

Stable isotope ratios, carbon and nitrogen contents and C:N ratios were tightly clustered, in expected ranges for well preserved collagen (DeNiro 1985; Ambrose 1990; van Klinken 1999) and for walrus in the Canadian Arctic (Coltrain et al. 2004; Jaouen et al. 2016). There were no indications of sample (collagen) quality issues other than the one noted above.

The  $\Delta\text{R}$  results (Table 1) suggest that all samples can be treated as a single population. The small subset of samples (3) from Central Hudson Strait averages  $190 \pm 20$  yr, and the small

subset (3) from central Foxe Basin averages  $175 \pm 45$  yr. The large subset (21) from western Hudson Strait and northern Hudson Bay averages  $155 \pm 50$  yr, necessarily close to the overall average of  $160 \pm 50$  yr. The spread of only 35 years in subset means and standard deviation of 50 years indicates that all results overlap at two standard deviations and differences are not significant (One-way ANOVA,  $F_{[2,24]} = 0.80$ ,  $p = 0.46$ ). Therefore  $160 \pm 50$  yr is our best estimate of  $\Delta R$  for walrus from this region.

The  $\Delta R$  results (Table 2) for suspension feeding mollusks from the Foxe Basin region yield a rounded error-weighted average of  $295 \pm 130$  yr ( $n = 32$ ). This distribution is significantly different from that for walrus (Mann-Whitney  $U$  test,  $U = 212$ ,  $p = 0.001$ ).

### Comparative Dating

Of the 12 Dorset Paleoeskimo features pair-dated at Alarnerk (total 16 pairs), 11 are dug-in, shallow winter house depressions or their associated middens, and the other (08DCA78F1) is Meldgaard's ocher grave (Lynnerup et al. 2003). The five pair-dated features (12 pairs) at Jens Munk Island are three dug-in Dorset winter houses and associated middens and a nearby diffuse discard area (03DCA205). All features appear to represent single cultural layers overlying raised beach gravel. The two Thule Inuit features at Alarnerk are semisubterranean sod dwellings. Those at Silumiut are similar Thule dwellings and associated middens. Such dwellings, and perhaps also the Dorset winter dwellings, may represent multigenerational occupations (see below).

In all cases, as expected, walrus  $^{14}\text{C}$  ages exceed caribou ages (Table 3). The  $\Delta R$  estimates range from  $82 \pm 34$  to  $438 \pm 23$  yr with two additional outliers at  $540 \pm 34$  and  $698 \pm 27$  yr. Excluding the outliers, the rounded mean and standard deviation are  $270 \pm 90$  yr ( $n = 33$ ).

### DISCUSSION

The significant difference between the  $\Delta R$  results for museum specimens of walrus and mollusks indicates that results from one cannot be applied to the other. It is possible that the difference will disappear with much larger sample sizes, but these will be difficult to attain given the limited availability of museum walrus collections. Because walrus feed almost exclusively on mollusks, predominantly *Mya* spp. in Foxe Basin (Fisher and Stewart 1997), and yet have a smaller  $\Delta R$ , the difference between the distributions is problematic. We suggest several possible interpretations. First, the molluscan taxa represented in Table 2, although excluding obligate deposit feeders (*Portlandia* and *Macoma*), differ from those consumed by walrus and the difference in estimated  $\Delta R$  (walrus vs. mollusk) may reflect taxonomic differences among the mollusks in the relative proportions of carbon in their shells that are derived from dissolved organic and dissolved inorganic sources, which is known to occur (Lorrain et al. 2004; Gillikin et al. 2006).

The four largest molluscan  $\Delta R$  values in Table 2 are for *Astarte* spp. and an additional five *Astarte* values exceed the molluscan mean. Thus, *Astarte*, an infaunal suspension feeder (McNeely et al. 2006), may also derive some of its shell carbonate from pore water. The single *Mya* measurement in Table 2 has a  $\Delta R$  of only  $110 \pm 40$  yr, which overlaps with the lower end of the walrus values. Secondly, the mollusk meat consumed by walrus may differ in  $^{14}\text{C}$  content from that of the mollusk shells. Walrus ingest primarily meat, because they use their tongues to suction the meat and discard shells (Kastelein et al. 1994) and the carbon that would be eventually incorporated into the collagen in the walrus bones or teeth would have been derived from dietary protein (Jim et al. 2004). Table 4 presents  $\delta^{13}\text{C}$  values for both shells and body tissue of several mollusk species collected live from Resolute Bay in the Canadian Arctic Archipelago (R. McNeely unpublished data). In all species, the tissue discriminates against the heavier

Table 2  $^{14}\text{C}$  ages of live-collected mollusks from Foxe Basin region (McNeely et al. 2006). See original for species. Results differ slightly from original because marine model ages are from IntCal3 and errors are not rounded.

Lab code	Genus	$^{14}\text{C}$ age	Year collected	Marine model age	$\Delta\text{R}$
CAMS-35484	<i>Clinocardium</i>	620 ± 40	1956	474 ± 23	146 ± 46
CAMS-47240	<i>Serripes</i>	700 ± 40	1956	474 ± 23	226 ± 46
CAMS-47242	<i>Hiatella</i>	670 ± 40	1956	474 ± 23	196 ± 46
CAMS-35487	<i>Hiatella</i>	640 ± 50	1956	474 ± 23	166 ± 55
TO-8033	<i>Musculus</i>	740 ± 50	1956	474 ± 23	266 ± 55
CAMS-33145	<i>Clinocardium</i>	780 ± 50	1956	474 ± 23	306 ± 55
CAMS-33147	<i>Hiatella</i>	790 ± 50	1956	474 ± 23	316 ± 55
CAMS-34657	<i>Balanus</i>	700 ± 50	1956	474 ± 23	226 ± 55
TO-8040	<i>Hiatella</i>	820 ± 60	1956	474 ± 23	346 ± 64
CAMS-35486	<i>Musculus</i>	710 ± 50	1956	474 ± 23	236 ± 55
TO-8039	<i>Astarte</i>	960 ± 60	1956	474 ± 23	486 ± 64
UCIAMS-6016	<i>Astarte</i>	930 ± 20	1956	474 ± 23	456 ± 30
8039/6016*	<i>Astarte</i>				461 ± 16
CAMS-34651	<i>Serripes</i>	690 ± 50	1956	474 ± 23	216 ± 55
CAMS-35483	<i>Astarte</i>	800 ± 40	1956	474 ± 23	326 ± 46
CAMS-35485	<i>Musculus</i>	730 ± 50	1956	474 ± 23	256 ± 55
GSC-6098	<i>Astarte</i>	740 ± 80	1956	474 ± 23	266 ± 83
TO-8037	<i>Astarte</i>	910 ± 50	1955	473 ± 23	437 ± 55
UCIAMS-6015	<i>Astarte</i>	880 ± 20	1955	473 ± 23	407 ± 30
8037/6015*	<i>Astarte</i>				414 ± 18
TO-8038	<i>Astarte</i>	730 ± 50	1956	474 ± 23	256 ± 55
UCIAMS-6564	<i>Hiatella</i>	735 ± 20	1956	474 ± 23	261 ± 30
CAMS-47243	<i>Hiatella</i>	720 ± 50	1955	473 ± 23	247 ± 55
CAMS-47239	<i>Musculus</i>	700 ± 40	1956	474 ± 23	226 ± 46
CAMS-47241	<i>Musculus</i>	810 ± 40	1955	473 ± 23	337 ± 46
UCIAMS-6014	<i>Astarte</i>	765 ± 20	1955	473 ± 23	292 ± 30
CAMS-33146	<i>Musculus</i>	690 ± 50	1955	473 ± 23	217 ± 55
CAMS-33144	<i>Astarte</i>	760 ± 50	1953	472 ± 23	288 ± 55
CAMS-46552	<i>Musculus</i>	560 ± 40	1953	472 ± 23	88 ± 46
CAMS-34647	<i>Clinocardium</i>	480 ± 50	1953	472 ± 23	8 ± 55
CAMS-46560	<i>Hiatella</i>	670 ± 50	1953	477 ± 23	190 ± 55
CAMS-33149	<i>Clinocardium</i>	690 ± 50	1954	472 ± 23	218 ± 55
CAMS-46549	<i>Mya</i>	590 ± 40	1954	472 ± 23	118 ± 46
CAMS-46557	<i>Clinocardium</i>	530 ± 50	1954	472 ± 23	58 ± 55
CAMS-47244	<i>Clinocardium</i>	600 ± 40	1954	472 ± 23	128 ± 46
CAMS-46547	<i>Cyclopecten</i>	510 ± 40	1954	472 ± 23	38 ± 46
Average					297 ± 130

\*Error weighted mean of two replicate samples.

isotope(s). That should, therefore, increase the walrus  $\Delta\text{R}$  with respect to that of the mollusk shell. However, these fractionation effects are corrected in conventional  $^{14}\text{C}$  age calculations. Thirdly, the walrus may have fed in areas with large terrestrial carbon input. However, the walrus collagen  $\delta^{13}\text{C}$  values are not consistent with this suggestion and there are no large estuarine areas or drainage basins near the sample sites, many of which are on islands. Fourthly,

Table 3 Comparative dating of caribou and walrus remains from Alarnerk (08DCA) and Jens Munk (03DCA) archaeological sites, northwestern Foxe Basin and from Silumiut site (KkJg<sup>-1</sup>) in northwestern Hudson Bay. Marine model ages from IntCal13. An asterisk indicates dates on tooth/tusk.

UCIAMS #	Field site	Taxon	<sup>14</sup> C age	Calibrated age ± 1σ	Marine model age	ΔR
Group 1						
53028	08DCA61F5	<i>Rangifer</i>	1900 ± 15	1854 ± 18	2244 ± 26	
175939	08DCA61F5	<i>Odobenus</i>	2470 ± 20			222 ± 26
53034	08DCA78F1	<i>Rangifer</i>	1705 ± 15	1628 ± 49	2068 ± 26	
175944	08DCA78F1	<i>Odobenus</i>	2285 ± 20			240 ± 34
185088	08DCA62F9	<i>Rangifer</i>	1990 ± 15	1854 ± 18	2244 ± 26	
185087	08DCA62F9	<i>Odobenus</i>	2495 ± 15			167 ± 27
188998	08DCA62F9	<i>Rangifer</i>	2030 ± 15	1978 ± 21	2357 ± 25	
188997	08DCA62F9	<i>Odobenus</i>	2575 ± 15			214 ± 24
189002	08DCA68F5	<i>Rangifer</i>	1085 ± 15	1006 ± 36	1469 ± 26	
189001	08DCA68F5	<i>Odobenus</i> *	1690 ± 15			232 ± 31
189004	08DCA68F5	<i>Rangifer</i>	1085 ± 15	1006 ± 36	1469 ± 26	
189003	08DCA68F5	<i>Odobenus</i> *	1695 ± 15			237 ± 31
189566	03DCA204F5	<i>Rangifer</i>	2020 ± 15	1971 ± 20	2351 ± 26	
189565	03DCA204F5	<i>Odobenus</i>	2575 ± 15			220 ± 23
189568	03DCA204F5	<i>Rangifer</i>	1875 ± 15	1839 ± 20	2232 ± 27	
189567	03DCA204F5	<i>Odobenus</i>	2460 ± 20			226 ± 32
189570	03DCA204F13	<i>Rangifer</i>	1935 ± 20	1890 ± 23	2287 ± 26	
189569	03DCA204F13	<i>Odobenus</i>	2535 ± 15			254 ± 27
189572	03DCA204F13	<i>Rangifer</i>	1985 ± 15	1940 ± 27	2328 ± 25	
189571	03DCA204F13	<i>Odobenus</i>	2580 ± 20			258 ± 30
185094	03DCA204F18	<i>Rangifer</i>	1970 ± 15	1919 ± 18	2310 ± 26	
185093	03DCA204F18	<i>Odobenus</i>	2545 ± 15			237 ± 23
189560	03DCA204F18	<i>Rangifer</i>	2015 ± 20	1967 ± 23	2348 ± 25	
189559	03DCA204F18	<i>Odobenus</i>	2590 ± 20			239 ± 29
185097	03DCA205	<i>Rangifer</i>	1870 ± 15	1833 ± 24	2229 ± 27	
185098	03DCA205	<i>Odobenus</i>	2450 ± 15			222 ± 32
189562	03DCA205	<i>Rangifer</i>	1810 ± 20	1760 ± 36	2162 ± 27	
189562	03DCA205	<i>Odobenus</i>	2345 ± 15			181 ± 33
188986	KkJg1 H13 Sq H	<i>Rangifer</i>	365 ± 15	413 ± 64	748 ± 23	
188987	KkJg1 H13 Sq H	<i>Odobenus</i>	900 ± 15			82 ± 34
188990	KkJg1 H13 ML2	<i>Rangifer</i>	390 ± 15	481 ± 16	844 ± 25	
188995	KkJg1 H13 ML2	<i>Odobenus</i>	970 ± 15			118 ± 28
188991	KkJg1 H13 ML3	<i>Rangifer</i>	425 ± 20	502 ± 9	888 ± 25	
188992	KkJg1 H13 ML3	<i>Odobenus</i>	1115 ± 15			232 ± 24
188993	KkJg1 H8 ML3	<i>Rangifer</i>	400 ± 15	490 ± 12	860 ± 25	
188994	KkJg1 H8 ML3	<i>Odobenus</i>	1005 ± 15			142 ± 26
Average Group 1						209 ± 46
Group 2						
185090	03DCA204F5	<i>Rangifer</i>	1975 ± 15	1924 ± 21	2315 ± 26	
185089	03DCA204F5	<i>Odobenus</i>	2575 ± 15			264 ± 24
189000	08DCA62F9	<i>Rangifer</i>	1845 ± 15	1780 ± 32	2175 ± 27	
188999	08DCA62F9	<i>Odobenus</i> *	2510 ± 15			323 ± 30
185092	03DCA204F13	<i>Rangifer</i>	1905 ± 15	1856 ± 19	2246 ± 26	
185091	03DCA204F13	<i>Odobenus</i>	2565 ± 15			314 ± 24

Table 3 (Continued)

UCIAMS #	Field site	Taxon	$^{14}\text{C}$ age	Calibrated age $\pm 1\sigma$	Marine model age	$\Delta\text{R}$
53030	08DCA61F7	<i>Rangifer</i>	1840 $\pm$ 20	1777 $\pm$ 34	2173 $\pm$ 27	
175941	08DCA61F7	<i>Odobenus</i>	2510 $\pm$ 20			326 $\pm$ 34
53029	08DCA61F6	<i>Rangifer</i>	1910 $\pm$ 15	1859 $\pm$ 19	2251 $\pm$ 26	
175940	08DCA61F6	<i>Odobenus</i>	2560 $\pm$ 20			306 $\pm$ 28
189564	03DCA205	<i>Rangifer</i>	1830 $\pm$ 15	1771 $\pm$ 32	2169 $\pm$ 27	
189563	03DCA205	<i>Odobenus</i>	2445 $\pm$ 15			266 $\pm$ 29
53031	08DCA67F1	<i>Rangifer</i>	1745 $\pm$ 15	1663 $\pm$ 31	2079 $\pm$ 26	
175942	08DCA67F1	<i>Odobenus</i>	2370 $\pm$ 20			280 $\pm$ 30
53038	08DCA85F1	<i>Rangifer</i>	1690 $\pm$ 15	1590 $\pm$ 22	2030 $\pm$ 27	
175948	08DCA85F1	<i>Odobenus</i>	2455 $\pm$ 20			423 $\pm$ 28
53033	08DCA70F4	<i>Rangifer</i>	905 $\pm$ 15	846 $\pm$ 47	1292 $\pm$ 24	
175943	08DCA70F4	<i>Odobenus</i>	1680 $\pm$ 20			373 $\pm$ 37
185084	08DCA72F5	<i>Rangifer</i>	940 $\pm$ 15	862 $\pm$ 43	1297 $\pm$ 25	
185083	08DCA72F5	<i>Odobenus</i>	1750 $\pm$ 15			427 $\pm$ 48
185085	08DCA66	<i>Rangifer</i>	1860 $\pm$ 15	1794 $\pm$ 36	2185 $\pm$ 27	
185086	08DCA66	<i>Odobenus</i>	2635 $\pm$ 15			433 $\pm$ 35
189005	08DCA66	<i>Rangifer</i>	1825 $\pm$ 15	1769 $\pm$ 32	2168 $\pm$ 27	
189006	08DCA66	<i>Odobenus</i> *	2595 $\pm$ 15			420 $\pm$ 30
189008	08DCA66	<i>Rangifer</i>	1925 $\pm$ 15	1872 $\pm$ 19	2267 $\pm$ 26	
189007	08DCA66	<i>Odobenus</i> *	2630 $\pm$ 15			360 $\pm$ 24
53035	08DCA82H10	<i>Rangifer</i>	260 $\pm$ 15	305 $\pm$ 5	686 $\pm$ 23	
175946	08DCA82H10	<i>Odobenus</i>	1115 $\pm$ 20			438 $\pm$ 23
188988	KkJg1H4 Sq A	<i>Rangifer</i>	310 $\pm$ 15	375 $\pm$ 48	722 $\pm$ 23	
188989	KkJg1 H4 Sq A	<i>Odobenus</i>	1090 $\pm$ 15			348 $\pm$ 32
Average Group 2						349 $\pm$ 64
Overall average Groups 1 and 2						270 $\pm$ 88
<b>Outliers</b>						
53037	08DCA82H8	<i>Rangifer</i>	120 $\pm$ 15	140 $\pm$ 96	527 $\pm$ 23	
175947	08DCA82H8	<i>Odobenus</i>	1225 $\pm$ 20			698 $\pm$ 27*
189558	03DCA204F18	<i>Rangifer</i>	1720 $\pm$ 20	1638 $\pm$ 46	2073 $\pm$ 26	
189557	03DCA204F18	<i>Odobenus</i>	2610 $\pm$ 15			540 $\pm$ 34

\*Note that this value was calculated using the method presented by Southon et al. (1995) as the Reimer and Reimer (2017) application cannot calculate  $\Delta\text{R}$  values when the  $^{14}\text{C}$  age is  $<199$  yr.

there should be differences in the  $^{14}\text{C}$  content of mollusk shells and soft tissue because the shell is inert once precipitated, while the soft tissue continuously turns over. Studies quantifying the rate of soft tissue turnover in marine invertebrates are generally lacking, but the organic carbon in the soft tissue of blue mussel was found to have a half-life of 8.9 days (Dubois et al. 2007).

Some mollusks can live for well over one hundred years (Weidman et al. 1994; Sejr et al. 2002). Consequently, depending on the position along the shell's growth axis from which the sample was taken, there may be a significant difference in the period of time reflected in the  $^{14}\text{C}$  content of the shell and the soft tissue that would have been consumed by the walrus. Finally, the sampled walrus may have fed over a region large enough to significantly sample areas with lower reservoir ages, in this case margins of the Labrador Sea. However, Foxe Basin walrus are thought to reside year-round in the region, retreating to tidally induced polynyas (open water) in the winter (Born et al. 1995). The first interpretation, therefore, seems the most reasonable.

Table 4 Comparison of stable isotope values in shells and tissues of marine mollusks from Resolute Bay.

Lab code	Taxon	Material	$\delta^{13}\text{C} \text{‰}$
GSC-4851	<i>Serripes groenlandicus</i>	Shell	+ 1.12
GSC-4988	<i>Serripes groenlandicus</i>	Tissue	- 19.40
GSC-4852	<i>Mya truncata</i>	Shell	+ 1.02
GSC-4992	<i>Mya truncata</i>	Tissue	- 20.00
GSC-4853	<i>Astarte borealis</i>	Shell	+ 1.40
GSC-4993	<i>Astarte borealis</i>	Tissue	- 20.40
GSC-4854	<i>Hiatella arctica</i>	Shell	+ 0.87
GSC-4993	<i>Hiatella arctica</i>	Tissue	- 21.20
GSC-4855	<i>Macoma calcarea</i>	Shell	+ 0.03
GSC-5845	<i>Macoma calcarea</i>	Tissue	- 16.27

The variance of the mollusk  $\Delta R$  values is much greater than that of the walrus values despite similar sample sizes; standard deviations of 130 and 50, respectively. The smaller variance for walrus may reflect their choice of a small number of mollusk taxa as food, primarily *Mya* sp. (see above). The greater variance for mollusks may reflect poorly documented species dependencies of  $\Delta R$  alluded to above. However, it may be due in part to the greater standard errors (up to 80 years vs. 15 years for walrus) in these somewhat earlier  $^{14}\text{C}$  analyses. McNeely et al. (2006) excluded several dates from their data set based on non-reproducible paired datings of individual specimens by different laboratories.

The  $\Delta R$  for the Craig Harbour walrus ( $300 \pm 15$  yr) is much larger than any of the  $\Delta R$  values from the Foxe Basin-Hudson Strait region. Craig Harbour is in the Canadian Arctic Archipelago region of McNeely et al. (2006), which has a larger  $\Delta R$  for mollusks than does the Foxe Basin Region. The larger  $\Delta R$  for Craig Harbour is thus in the right direction and it is indistinguishable from that for suspension feeding mollusks ( $300 \pm 95$  yr) in its region.

The results from comparative dating of caribou and walrus are consistent in the sense that the caribou dates are all younger but the inferred  $\Delta R$  values are discordant with the results from the museum specimens of walrus (Figure 2). Ignoring 2 outliers, the mean  $\Delta R$  value of  $270 \pm 90$  yr is significantly larger than the value for museum walrus ( $160 \pm 50$  yr) (Mann-Whitney  $U$  test;  $U = 119$ ,  $p < 0.001$ ) but statistically indistinguishable from the mollusk value of  $295 \pm 130$  yr ( $U = 408$ ,  $p = 0.12$ ). The outlier samples cannot represent time variance in  $\Delta R$  because other samples of similar ages are not outliers. We suggest instead that the outliers represent the clearest cases where the walrus remains truly are much older than the caribou. Note that the differences cannot be due to differential temporal spans of carbon integration by walrus and caribou. The average walrus lifespan is 20–30 yr (Fay 1985), while caribou average 7–12 yr, with antlers being shed annually and therefore only representing a single year of growth (Naughton 2012:570).

Aside from the outliers, we suggest that the discordance between the archaeological and museum  $\Delta R$  values is best interpreted in light of the museum results. In short, the assumption that the archaeological walrus and caribou from each feature are strictly contemporaneous is not supported in at least half the cases (Group 2 of Table 3). The assumption is based on the fact that all of the archaeological samples came from what appear to be single cultural layers and on the notion that close proximity of bones indicates closeness in age. Nevertheless, it is possible that these layers, be they house floors or midden layers, accumulated over considerable intervals, the

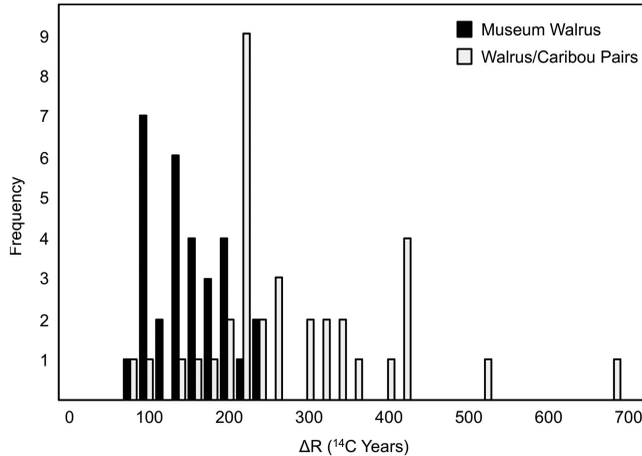


Figure 2 Histogram of  $\Delta R$  values measured from museum specimens of walrus and estimated from paired datings of walrus and caribou from archaeological features.

well-known palimpsest effect in archaeology (the studies of palimpsest formation among the Nunamiut of Alaska by Binford (1982, 1983) are especially relevant in this context). This would be especially true of semisubterranean dwellings, from which all three sites samples were collected. Because these dwellings represent considerable costs in time and energy, they would invariably be reused on a seasonal basis, often for several generations, as were similar dwellings in Alaska (Rainey 1947; Spencer 1959; see also Savelle 1987). Within such dwellings, individual occupation layers, especially in the case of shallow Paleoeskimo dwellings, are rarely identifiable, with material from multiple occupations often simply swept aside into common refuse areas (see e.g., Habu and Savelle 1994; Savelle and Habu 2004). Poor soil development further exacerbates the palimpsest problem, especially at Paleoeskimo sites in Foxe Basin, where very thin horizontal sheet middens predominate, in contrast to the more substantial mound middens in, for example, southern Hudson Bay (e.g., Taylor 1968) and Labrador (e.g., Jordan 1980). In the eight features where two or more caribou bones were dated, only one returned dates within error. The others indicate minimum occupation intervals of 48–329 yr, based on the mean calibrated ages.

Considering the miniscule percentage of available bones that have been dated, it is improbable that the oldest and youngest were included. Nevertheless, even that duration can account for the discrepancy between the two sets of results. Furthermore, some of the walrus remains may have been scavenged from older sites. In that regard, we noted that walrus skulls, mandibles and rarely tusk fragments are conspicuous on the surface of Dorset middens, which are extensive but typically are only a decimetre or so thick. However, most of the maxillae and mandibles lack teeth, suggesting that well preserved bone, especially ivory, may have been a preferred raw material for later occupants. A tooth (tusk) can be removed far more easily from a walrus skull that is decades or more old than from the skull of a freshly killed animal, where thick maxillary bone has to be removed (see e.g., LeMoine and Darwent [1998] for the various steps in the removal of tusks from fresh skulls by Paleoeskimos). Our paired datings included five on walrus teeth/tusks (Table 3). All of these yielded  $\Delta R$  estimates above the mean of Group 1 and three of them are above the overall mean. Nevertheless, the high  $\Delta R$  estimates derived from other walrus bone elements indicates that they too have long residency at sites in comparison to caribou. The middens and floor refuse may have accumulated in highly spotty patterns, leading to age differences

between nearby specimens. Specifically, therefore, we propose that only samples with apparent  $\Delta R$ s of <260 years (Figure 2; 18 out of 35 samples; Group 1, Table 3) represent approximately contemporaneous pairs. The mean  $\Delta R$  of that group is  $209 \pm 46$  yr, which is still significantly older than the museum results (Mann-Whitney  $U$  test;  $U = 119$ ,  $p = 0.004$ ).

## CONCLUSION

$^{14}\text{C}$  dates on museum specimens of walrus from the Foxe Basin-Western Hudson Strait region of the eastern Canadian Arctic indicate that a  $\Delta R$  value of  $160 \pm 50$  yr be used in calibrating dates on walrus from this region. The significant difference between these dates and a similar set of dates on museum specimens of mollusks argues against applying mollusk based corrections to sea mammals at this time. The results of comparative dating of caribou and walrus from archaeological features provide maximum estimates of reservoir ages that are more varied than the directly measured ages. Although about half of inferred  $\Delta R$  values overlap the museum specimen results, the others indicate that the assumption of contemporaneity does not hold true.

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## SUPPLEMENTARY MATERIAL

To view supplementary material for this article, please visit <https://doi.org/10.1017/RDC.2018.50>

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