Population comparison of right whale body condition reveals poor state of the North Atlantic right whale

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ABSTRACT: The North Atlantic right whale Eubalaena glacialis (NARW), currently numbering <410 individuals, is on a trajectory to extinction. Although direct mortality from ship strikes and fishing gear entanglements remain the major threats to the population, reproductive failure, resulting from poor body condition and sublethal chronic entanglement stress, is believed to play a crucial role in the population decline. Using photogrammetry from unmanned aerial vehicles, we conducted the largest population assessment of right whale body condition to date, to determine if the condition of NARWs was poorer than 3 seemingly healthy (i.e. growing) populations of southern right whales E. australis (SRWs) in Argentina, Australia and New Zealand. We found that NARW juveniles, adults and lactating females all had lower body condition scores compared to the SRW populations. While some of the difference could be the result of genetic isolation and adaptations to local environmental conditions, the magnitude suggests that NARWs are in poor condition, which could be suppressing their growth, survival, age of sexual maturation and calving rates. NARW calves were found to be in good condition. Their body length, however, was strongly determined by the body condition of their mothers, suggesting that the poor condition of lactating NARW females may cause a reduction in calf growth rates. This could potentially lead to a reduction in calf survival or an increase in female calving intervals. Hence, the poor body condition of individuals within the NARW population is of major concern for its future viability.

KEY WORDS: Baleen whale · Bioenergetics · Eubalaena · Morphometrics · Photogrammetry · Unmanned aerial vehicles

1. INTRODUCTION

Right whale populations around the world were severely depleted (some reduced to less than 5% of the original population size) by commercial whaling operations from the 11th to the 20th century in the North Atlantic (Aguilar 1986, Reeves et al. 1999) and during
the 19th to 20th century in the Southern hemisphere (Dawbin 1986, Jackson et al. 2016), as well as by illegal Soviet whaling in the Southern hemisphere in the 1950s to early 1970s (Yablokov 1994, Tormosov et al. 1998). Since the cessation of commercial whaling, the southern right whale *Eubalaena australis* (SRW) has been recovering at a relatively rapid pace throughout most of its range, and currently numbers in the tens of thousands globally (IWC 2013). The population growth rate during this time has been as high as 5.55% for Australia (Bannister 2016), 5–7% for New Zealand (Carroll et al. 2013) and 6.5% for Argentina (Cooke et al. 2015), although the growth rate of the latter has slowed down substantially (Crespo et al. 2019).

In contrast, the recovery of the North Atlantic right whale *E. glacialis* (NARW) has been considerably slower, with a mean annual growth rate of 2.8% between 1990 and 2010 (Pace et al. 2017). More recent abundance estimates, between 2010 and 2015, indicate that the population has been declining at a rate of just under 1% per year (Pace et al. 2017). The rate of decline has been higher for females, which dropped at approximately 7% between 2010 and 2015, compared to about 4% for males over the same period (Pace et al. 2017). The situation for NARWs was further worsened by an unusual mortality event between 1 November 2016 and 31 December 2017, when at least 17 juvenile and adult right whales died as a result of entanglements and vessel strikes (NARWC 2018). In December 2015, prior to the 2017 mortalities, the species’ abundance was estimated at 451 individuals, of which 186 were females. The best estimate as of the end of 2017 was 411 animals (Pettis et al. 2018). While fishing gear entanglements and ship strikes are the largest direct anthropogenic threats to the NARW population (Moore et al. 2004, Knowlton et al. 2012, van der Hoop et al. 2013, Kraus et al. 2016), reduced reproductive rate resulting from nutritional stress (i.e. poor body condition) has been hypothesised as a factor further contributing to the population decline (Kraus et al. 2001, Reeves et al. 2001, Schick et al. 2013, Rolland et al. 2016). The sublethal impacts of entanglement have also been modelled to significantly impact reproductive success (van der Hoop et al. 2017).

The effect of body condition on reproduction is well documented in both terrestrial (Albon et al. 1983, Loudon et al. 1983, Skogland 1984, Atkinson & Ramsay 1995, Festa-Bianchet 1998) and marine mammals (Arnbom et al. 1997, Boltnev & York 2001, Bowen et al. 2001, Wheatley et al. 2006). In baleen whales, female body condition influences fecundity (Lockyer 2007, Williams et al. 2013), foetal growth (Christiansen et al. 2014) and calf body condition (Christiansen et al. 2016a). Like most baleen whales, right whales make annual migrations between high-latitude feeding grounds in summer and low-latitude breeding grounds in winter (Bannister et al. 1999). Females become sexually mature at around 9 yr old and give birth to a single calf at a time (Kraus et al. 2001, Cooke et al. 2003, Burnell 2008). They are ‘capital’ breeders, fasting during the winter breeding season, and thus have a finite amount of energy to invest in late pregnancy and lactation (Lockyer 1987, Stephens et al. 2009). Christiansen et al. (2018) showed in SRWs that maternal size (body length and condition) has a direct effect on the amount of energy that lactating females invest in their calves, which in turn dictates calf growth rates. When conditions are favourable, females generally have a 3 yr reproductive cycle consisting of 1 yr of gestation, 1 yr of lactation and 1 yr of resting (to recover energy stores) (Best 1994). The mean calving interval for SRWs is close to this 3 yr minimum, at 3.33 yr in Australia (Burnell 2001), 3.31 yr in New Zealand (Davidson et al. 2017) and between 2.96 and 3.24 yr in Argentina (Marón et al. 2015). In contrast, since 2015, the mean calving interval for NARW females is >7 yr (Pettis et al. 2020), suggesting that they need several years longer to recover from a reproductive event. Apart from body condition having a direct effect on female reproductive success, it can also influence juvenile growth rates (Douhard et al. 2017) and the age of sexual maturation (Sigurjónsson et al. 1990), which could negatively influence population growth.

The aim of this study was to assess the body condition of the NARW. Although the population’s body condition (based on visual assessment) has declined during the last 3 decades (Rolland et al. 2016), a comparison to healthy (growing) right whale populations is needed to assess its current status. Unfortunately, no historical data on NARW body condition exist to allow such a comparison. Instead, the best opportunity to assess the relative body condition of NARW comes from a comparison with their closest living relative, the SRW. We therefore compared the body condition of NARWs with 3 seemingly healthy (i.e. growing) populations of SRW in Australia, New Zealand and Argentina. Although we were comparing 2 different species of right whales (Rosenbaum et al. 2000, Gaines et al. 2005), which might differ in their body condition due to genetic differences, our rationale was that body condition, similar to most traits closely associated with fitness, shows low genetic variance relative to environmental variance (Mousseau & Roff 1987, Kruuk et al. 2000). We also show that NARWs and SRWs are very similar in body shape, size and life
history characteristics, which should facilitate comparison. Based on the lower population growth rate and longer calving interval of the NARW, our main hypothesis is that NARWs are in poorer body condition compared to SRWs. To help infer the potential effects of reduced body condition on different life history parameters, we split our analysis into different reproductive classes (calves, juveniles, adults and lactating females). We expected lactating females to overall have a higher body condition relative to the other reproductive classes, since they must have had sufficient energy reserves to complete gestation (Lockyer 1981, Christiansen et al. 2014). However, with NARWs being affected by numerous anthropogenic factors, we expected the body condition of lactating females to be significantly lower than for SRWs. In baleen whales, a lower maternal body condition has been shown to negatively influence calf growth rates (Christiansen et al. 2018) and body condition (Christiansen et al. 2016a). We therefore anticipated NARW calves to have a poorer body condition and/or a smaller body size (i.e. length) compared to SRW calves. Finally, with fishing gear entanglements affecting both juvenile and adult NARWs (NARWC 2018), we projected both reproductive classes to have a lower body condition compared to SRWs.

2. MATERIALS AND METHODS

2.1. Data collection

Aerial photographs of right whales were taken using non-invasive (Christiansen et al. 2016b) unmanned aerial vehicles (UAVs) in 4 locations: the North Atlantic, Australia, New Zealand and Argentina (Fig. 1). Photographs of NARW lactating females and calves were collected on their calving grounds in Florida, USA, between 12 January and 22 February 2016 and 2017, while juvenile and adult NARWs were photographed on their feeding grounds in Cape Cod Bay, USA, between 21 March and 27 April 2016 and 2017 (Fig. 1). All SRWs were measured on their breeding grounds at the Head of Bight, Australia, between 25 June and 25 September 2016, the Auckland Islands, New Zealand, between 28 July and 14 August 2016, and in Peninsula Valdés, Argentina, between 3 August and 12 November 2018.

In each location, multirotor UAVs were flown from either land (Australia and Argentina) and/or boats (North Atlantic, New Zealand and Argentina) above a surfacing whale at altitudes between 17.8 and 55.1 m (mean = 31.3 m, SD = 8.01; Argentina = 17.8–37.0 m; Australia = 27.9–46.6 m; New Zealand = 17.9–51.3 m; North Atlantic = 26.8–55.1 m), and vertical photographs were taken of the dorsal side of the whale (Fig. 2A). For the North Atlantic study site, an APH-22 hexacopter with an Olympus E-PM2 camera was used, while modified DJI Inspire 1 Pro quadcopters with Zenmuse X5 cameras were used in Australia, Argentina and New Zealand. Both UAV types were equipped with an Olympus M Zuiko 25 mm f1.8 lens to minimize picture distortion. Measurement accuracies of both the APH-22 and the Inspire 1 Pro systems have been estimated at 99.9% (Durban et al. 2015, Dawson et al. 2017, Christiansen et al. 2018). Christiansen et al. (2018) further quantified the measurement errors of the Olympus 25 mm lens when flying at different altitudes ranging from 5 to 120 m and measuring a known sized object on land.
Their results showed that within the altitude range used in this study (17.8 and 55.1 m) the mean measurement error was 0.7 cm (SD = 0.5, n = 50) with a maximum of 1.6 cm. Since the measurement errors were not influenced by the altitude of the UAV, differences in sampling altitude between locations did not bias measurements. When photographing whales, a camera gimbal ensured that the camera of the UAV was always facing down at a near-perfect 90° angle. Only photographs of adequate quality, when the whale was lying flat at the surface, were used for analyses (for details, see Christiansen et al. 2018). From the aerial photographs, individual whales were identified based on the unique callosity patterns on their heads (Payne et al. 1983). The Australian, New Zealand and North Atlantic data sets included multiple measurements from the same individuals. To avoid pseudo-replication, only a single measurement for each whale was used, which was selected randomly (using a random number generator in R) from the best photographs of that individual. We judged this to be a less biased approach compared to using the average measurements from repeated photographs of the same individuals (taken over several days), as the latter might introduce temporal variation in body measurements, and also cause heterogeneity in measurement errors between individuals (animals with single or multiple measurements) and populations.

Following the protocol of Christiansen et al. (2018), each photograph was graded (given a score of 1 [good quality], 2 [medium quality] or 3 [poor quality])
on several attributes, including degree of body roll, degree of body arch, body pitch (vertically), body length measurability and body width measurability. Only photographs with a roll, arch or pitch <3 were used in the analyses. To account for variation in body length and width measurability between photographs, we ran a sensitivity analysis where the body length and width of each individual whale were randomly varied within the confidence interval given by its length and width measurability scores (for details, see Christiansen et al. 2018). By repeating this process 1000 times, and refitting the final models in the analyses, the effect of the length and width measurement errors on the model parameters could be evaluated (see Fig. S1 in the Supplement at www.int-res.com/articles/suppl/m640p001_suppl.pdf).

### 2.2. Morphometric measurements and classification of reproductive classes

The total body length of the whales (tip of the lower jaw to the notch of the tail fluke) and their body widths (at 5% increments along the entire body axis of the animal), were measured (Fig. 2A), using a custom written graphical user interface (GUI) in MATLAB (Dawson et al. 2017). The GUI accounted for distortion of the camera lens through photogrammetric calibration (for details, see Dawson et al. 2017). All photographs were measured by a single experienced researcher, thus minimizing any potential interobserver bias. Image scale was established from the known focal length in combination with precise altitude data. For the North Atlantic data, altitude of the APH-22 was measured using an inbuilt GPS (mean error = 0.05 m; Durban et al. 2015) or a LightWare SF11/C laser range finder (mean error = 0.02 m; Dawson et al. 2017). For the SRW populations, the altitude of the Inspire 1 Pro was measured using the same type of range finder. While the accuracy of the altimeters used might have differed slightly, the width to length ratio of the whales was not affected by this, and hence this did not bias the body condition estimates.

Each whale was classified into 1 of 4 reproductive classes: calves (<4 mo of age), immature (juveniles), mature (non-lactating adults) and lactating females. Calves and lactating females were distinguished based on their close association with each other on the calving/breeding grounds. Immature and mature whales were separated based on their body length, using a threshold value of 12.0 m, which was based on the body length of the smallest lactating female measured in this study (11.72 m). A NARW mother with an older calf (>4 mo old), measured on the Cape Cod Bay feeding ground, was removed from the analyses. Similarly, adults with body volumes similar to or exceeding that of lactating females with newly born calves (calf body length ~5 m), were removed from the analyses as these likely represent late-stage pregnant females. The absolute and relative body width (body width/body length) of right whales at each measurement site was compared between locations and for each reproductive class, using linear models in R 3.5.3 (R Core Team 2019) (Figs. S2 & S3).

From the body length and width measurements, body volume was calculated using the methods of Christiansen et al. (2018). By assuming a circular cross-sectional body shape, the body volume of the whales was estimated by dividing the body of the whale into 18 frustum segments (1 between each width measurement) and calculating the volume ($V_s$) of each segment $s$ using the formula of a truncated cone (Christiansen et al. 2018):

$$V_s = \frac{1}{3} \pi h \left( r^2 + rR + R^2 \right)$$

where the height ($h$) is given by the distance between width measurements (5% of the body length), and the smaller ($r$) and larger radii ($R$) correspond to half of the smaller and larger width measurements, respectively (for details, see Christiansen et al. 2018). Total body volume ($V_{\text{Total}}$) of the whales was then estimated by summing the volumes of the different frustum segments (Christiansen et al. 2018):

$$V_{\text{Total}} = \sum_{s=1}^{S} V_s$$

Similar to Christiansen et al. (2018), the body volume of immature, mature and lactating females was calculated between 25 (the end of the head region) and 80% of their body length, which corresponds to the metabolically most active region of baleen whales (Lockyer et al. 1985, Miller et al. 2012, Christiansen et al. 2013, 2016a, 2018). Since the width to length ratio of calves is known to increase across their entire body axis during the first month of their lives (Christiansen et al. 2018), the body volume of calves was calculated from the tip of their rostrum down to 80% of their body length.

### 2.3. Body condition index

An animal’s body condition provides a measure of its energy balance, health and quality (Jakob et al.
Although body condition can be expressed through any physiological index that represents an individual’s energy reserves (Hanks 1981, Millar & Hickling 1990), it generally refers to the relative size of energy stores compared with structural components (commonly the length) of the body (Green 2001). Consequently, an individual’s body condition strongly influences its survival and reproductive success (Gaillard et al. 2000, Clutton-Brock & Sheldon 2010). Christiansen et al. (2018) showed that the body condition index (BCI) of individual right whales from the residuals of the log-log relationship (to account for non-linear relationships) between body volume and body length, divided by the expected (or predicted) body volume for the individual (to standardize BCI across body size, Christiansen et al. 2013, 2016a, 2018):

\[
\text{BCI}_i = \frac{BV_{\text{obs},i} - BV_{\text{exp},i}}{BV_{\text{exp},i}}
\]

where \(BV_{\text{obs},i}\) is the observed body volume of whale \(i\) in \(m^3\), and \(BV_{\text{exp},i}\) is the expected body volume of whale \(i\) in \(m^3\), given by the log-log relationship between body volume and body length:

\[
\log(BV_{\text{exp},i}) = \alpha + \beta \times \log(BL_i)
\]

where \(BL_i\) is the body length of whale \(i\), and \(\alpha\) and \(\beta\) represent the intercept and slope parameters, respectively, of the linear relationship between body volume and body length for all locations combined.

A positive BCI means that an individual was in relatively better condition than an average individual of the same body length, whereas a negative BCI means that the individual was in relatively poorer condition. To demonstrate that our BCI was independent from the absolute size (length) of the individual, we calculated the body condition of all measured whales using both the absolute (body length and widths, in metres) and relative body morphometrics (body length and widths, in pixels). The 2 approaches yielded nearly identical BCIs (\(F_{1,521} = 739902, p < 0.001, R^2 = 0.999\), Fig. S4), and showed that our metric accounted for potential structural differences (i.e. body length) between animals. The BCI of each individual was calculated from the length-to-volume relationship of all measured individuals. If we had modelled BCI separately for each reproductive class (i.e. fitting a separate length-to-volume model for each reproductive class) it would have resulted in a slight shift in the intercept (mean body condition) for each reproductive class, but would not have influenced the effect of location within each reproductive class (Fig. S5). Finally, although our BCI was based on a cross-sectional sample of the population (a single measurement representing a single whale), it correlated strongly with the BCI calculated from repeated measurements of the same whales (available for the Australian data set, see Fig. S6).

### 2.4. Differences in body condition between locations

To determine if NARWs were in poorer body condition compared to the 3 southern populations, we developed linear models in R 3.5.3. Right whale body condition (response variable) was modelled as a function of location (explanatory variable). Separate models were run for each reproductive class (calves, immature, mature and lactating females). During the breeding season, lactating females have finite energy reserves to support their own metabolic needs and the growth of their calf (Lockyer 2007, Christiansen et al. 2018). Consequently, lactating females decline in body condition through the breeding season as their calves grow in size (Christiansen et al. 2016a, 2018). To account for the temporal variation in body condition of lactating females (i.e. female body condition declining with increased calf length), calf body length was included as a covariate in the model. Similarly for calves, the effect of maternal body length and condition on calf body condition was investigated. Other covariates included day of the year (DOY; with the North Atlantic data converted to austral DOY by adding 183 d) and body length. However, collinearity (high correlation) between location and DOY, as well as between location and body length, resulted in only location being included in the final model for juvenile and adult right whales. To investigate the effect of location on body length, separate linear models were developed for each reproductive class.

Model validation included testing for homogeneous residuals (by plotting model residuals against the fitted model values), examining normality of residuals (from frequency histograms of residuals) and influential points and outliers (by calculating leverage scores and Cook’s distance, respectively).
2.5. Validation of cross-species comparison

To enable comparison of the body condition of the NARW and the SRW, their physiology and life history need to be very similar, so that the genetic variance does not exceed the environmental variance. To validate our cross-species comparison, we therefore compared the body shape, size and life history characteristics of NARWs and SRWs (Table 1).

To compare the structural body shape of NARWs and SRWs, we first measured the relative body width of the whales, and compared their head (0–25 % body length from the rostrum) and tail regions (80–100 % body length from the rostrum). With both these areas being mainly structural (Brodie 1975), and not part of the metabolically active body area for right whales (between 25 and 80 % of body length, Christiansen et al. 2018), any genetic difference in the external body width between species should be visible in those areas. We found no difference in the body width of the head or tail region of the whales (Table 1, Fig. S3). Further, although the site-specific body widths of NARWs and SRWs differed across the metabolically active body area of the whales, their overall body shapes were very similar (Fig. S3). Finally, Christiansen et al. (2019) showed that the relationship between body mass (or body volume) and length was very similar between NARWs (based on Fortune et al. 2012) and SRWs (from the Argentina population) (Table 1).

In regards to life history characteristics, Soviet catch data between 1951 and 1971 showed that female SRWs reach sexual maturity around 12.5 m in body length (Tormosov et al. 1998). Similarly, Sharp et al. (2019) classified NARW adults as individuals >9 yr of age, which, based on recent age-to-length curves, corresponds to a body length around 12.5 m (Table 1). Further, we found that the minimum body length of lactating SRWs in this study was 11.72 m, which was very similar to the minimum body length of lactating NARWs, which was 11.86 m (Table 1). In regards to birth size, our smallest measured NARW calf was 3.9 m in body length (which is within their predicted birth range of 4.22 ± 0.4 m; Fortune et al. 2012), which was very similar to the smallest SRW calf at 4.1 m body length (Table 1). Huang et al. (2009) also presented similar calf weaning lengths (8.78 m vs. 8.26 m) and female asymptotic lengths (17.8 m vs. 16.6 m) for NARWs and SRWs (Table 1).

3. RESULTS

The body volume of 523 right whales was successfully measured between 2016 and 2018 in the 4 study locations (Fig. 1). There was a strong linear relation-

Table 1. Comparison of body shape, size and life history characteristics between the North Atlantic right whale (NARW) and the southern right whale (SRW). BL: body length

<table>
<thead>
<tr>
<th>Structural body shape/ life history characteristic</th>
<th>NARW</th>
<th>SRW</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relative width (%BL ± SE) of head (20% BL from rostrum)</td>
<td>Calf = 20.0 ± 0.67</td>
<td>Calf = 20.4 ± 0.28</td>
<td>This study (Fig. S3)</td>
</tr>
<tr>
<td></td>
<td>Immature = 20.1 ± 0.21</td>
<td>Immature = 20.8 ± 0.18</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mature = 19.5 ± 0.15</td>
<td>Mature = 20.8 ± 0.22</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lactating = 19.8 ± 0.45</td>
<td>Lactating = 20.9 ± 0.13</td>
<td></td>
</tr>
<tr>
<td>Relative width (%BL ± SE) of tail (80% BL from rostrum)</td>
<td>Calf = 5.5 ± 0.15</td>
<td>Calf = 5.0 ± 0.16</td>
<td>This study (Fig. S3)</td>
</tr>
<tr>
<td></td>
<td>Immature = 4.2 ± 0.16</td>
<td>Immature = 4.0 ± 0.08</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mature = 4.3 ± 0.10</td>
<td>Mature = 3.8 ± 0.13</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lactating = 3.6 ± 0.19</td>
<td>Lactating = 3.9 ± 0.10</td>
<td></td>
</tr>
<tr>
<td>Female length (m) at sexual maturity</td>
<td>12.5</td>
<td>12.5</td>
<td>Tormosov et al. (1998), Sharp et al. (2019)</td>
</tr>
<tr>
<td>Minimum length (m) of lactating females</td>
<td>11.7</td>
<td>11.9</td>
<td>This study</td>
</tr>
<tr>
<td>Female asymptotic length (m)</td>
<td>17.8</td>
<td>16.6</td>
<td>Huang et al. (2009)</td>
</tr>
<tr>
<td>Minimum length (m) at birth</td>
<td>3.9</td>
<td>4.1</td>
<td>This study</td>
</tr>
<tr>
<td>Length at weaning (m)</td>
<td>8.8</td>
<td>8.3</td>
<td>Huang et al. (2009)</td>
</tr>
<tr>
<td>Weight (kg) at birth (BL = 4 m)</td>
<td>940</td>
<td>870</td>
<td>Christiansen et al. (2019), Fortune et al. (2012)</td>
</tr>
<tr>
<td>Weight (kg) at weaning (BL = 8.5 m)</td>
<td>7,830</td>
<td>7,970</td>
<td>Christiansen et al. (2019), Fortune et al. (2012)</td>
</tr>
<tr>
<td>Weight (kg) at sexual maturity (BL = 12.0 m)</td>
<td>20,680</td>
<td>21,940</td>
<td>Christiansen et al. (2019), Fortune et al. (2012)</td>
</tr>
</tbody>
</table>
ship between body volume (BV) and body length (BL) on the log-log scale ($F_{1,521} = 28953$, $p < 0.001$, $R^2 = 0.982$, Fig. 2C):

$$\log(BV_{exp.}) = -4.38 + 3.01 \times \log(BL)$$  (5)

Lactating females in the North Atlantic were in poorer body condition compared to the southern populations ($F_{3,156} = 5.11$, $p = 0.002$, $R^2 = 0.072$, Fig. 3; and see Fig. S7 and model 6 in Table S1). The body condition of lactating females from all 4 populations decreased as the calf grew in size (i.e. body length) through the breeding season ($F_{1,156} = 42.02$, $p < 0.001$, $R^2 = 0.197$), at a rate of 6.62% BCI m$^{-1}$ calf length (Fig. 4D, model 6 in Table S1). If we account for the body length of their calves (i.e. fix calf length to 6.0 m in the model), the body condition of lactating NARW females (mean = $-9.4\%$, SE = 4.8) was 20.5, 24.9 and 17.6% units lower compared to lactating SRW females in Australia (mean = 11.2%, SE = 2.1), New Zealand (mean = 15.6%, SE = 3.4) and Argentina (mean = 8.3%, SE = 6.5), respectively (Figs. 3 & 4D). The body length analyses showed that lactating NARW females (mean = 13.2 m, SE = 0.29) were on average 96, 54 and 48 cm shorter ($F_{3,157} = 5.07$, $p = 0.002$, $R^2 = 0.088$) than Australian (mean = 14.2 m, SE = 0.32), Argentinian (mean = 13.8 m, SE = 0.30) and New Zealand females (mean = 13.7 m, SE = 0.36), respectively (Fig. 4C). Consequently, the mean absolute body volume (mean = 27.9 m$^3$, SE = 2.90) of lactating NARWs was significantly ($F_{3,157} = 6.10$, $p < 0.001$, $R^2 = 0.104$) lower than for SRWs in Australia (mean = 39.9 m$^3$, SE = 3.11), New Zealand (mean = 38.1 m$^3$, SE = 3.56) and Argentina (mean = 36.1 m$^3$, SE = 2.99), at a magnitude of 11.99, 10.22 and 8.22 m$^3$, respectively.

![Diagram](image_url)

Fig. 3. (A) Predicted body condition values from the best fitting models for right whale calves (model 1 in Table S2 in the supplement), immature whales, mature whales and lactating females (model 6 in Table S1), as a function of location. (B) Predicted body condition values for right whales from Argentina, Australia, New Zealand and the North Atlantic, as a function of reproductive class. Error bars represent 95% confidence intervals. All whales were measured on their calving/breeding grounds, except for immature and mature North Atlantic right whales, which were measured on their feeding grounds. For lactating females, the full model also included calf body length as an explanatory variable, with maternal body condition declining significantly with calf body length (Fig. 4C). In the partial effect plot shown here, calf length was fixed at 6 m, which represents the mean body length of calves measured in this study. Sample sizes for all reproductive classes are given in Fig. 1.
NARW calves did not show signs of being in poorer condition compared to SRW calves, and the body length of calves (a rough proxy for time since birth) did not vary significantly between locations. Further, the day of sampling did not vary significantly between NARWs (after correcting the time of year with 183 d between the Northern and Southern hemisphere) and SRWs in Australia and New Zealand. Instead, we found that Australian calves (mean = −5.7%, SE = 2.2) were significantly thinner than New Zealand (mean = 13.5%, SE = 3.7) and Argentinian calves (mean = 3.7%, SE = 1.2).

Body length of calves was positively related to the length of their mothers (\(F_{1,158} = 12.5, p < 0.001, R^2 = 0.061\)) at a rate of 0.302 m (SE = 0.082) per m increase in maternal length (Fig. 4A; model 6 in Table S3). In addition, maternal body condition was negatively correlated (slope parameter = −2.82 m, SE = 0.474) with calf body length (\(F_{1,158} = 35.5, p < 0.001, R^2 = 0.172\)), since maternal body condition decreased as the calf grew in body length (Fig. 4B; model 6 in Table S3). The full model explained 23.3% of the variance in the data.

There was a difference in body condition of mature right whales (males and non-lactating females) between locations (\(F_{3,90} = 25.06, p < 0.001, R^2 = 0.455\), Fig. 3; Figs. S7 & S8), with North Atlantic adults (mean = −16.7%, SE = 2.0) being 27.9, 18.9 and 8.9% units lower in condition compared to individuals from Argentina (mean = 11.2%, SE = 3.3), New Zealand (mean = 2.2%, SE = 4.0) and Australia (mean = −7.8%, SE = 3.6), respectively (Fig. 3). In addition to being in poorer condition, the average body length of mature NARW (mean = 12.9 m, SE = 0.13) was lower (\(F_{3,90} = 6.07, p < 0.001, R^2 = 0.168\)) compared to Argentina (mean = 13.9 m, SE = 0.22), New Zealand (mean = 13.4 m, SE = 0.26) and Australia (mean = 13.3 m, SE = 0.24) (Fig. S9B).

We found that immature NARWs (mean = −13.1%, SE = 2.9) were in significantly poorer condition (\(F_{3,103} = 4.30, p = 0.007, R^2 = 0.111\), Fig. 3; Fig. S7) than juveniles in New Zealand (mean = −1.2%, SE = 3.4), Australia (mean = −2.4%, SE = 4.2) and Argentina.
(mean = −2.9%, SE = 3.4). On average, the BCI of immature NARWs was 11.9, 10.7 and 10.2% units lower than juveniles from New Zealand, Australia and Argentina, respectively (Fig. 3). Juvenile NARWs (mean = 11.2 m, SE = 0.19) were on average longer (F_{3,103} = 7.03, p < 0.001, R^2 = 0.170) than juveniles in Argentina (mean = 10.5 m, SE = 0.22) and New Zealand (mean = 10.5 m, SE = 0.22), and similar in size to Australian juveniles (mean = 11.3, SE = 0.27) (Fig. S9A).

The results from our sensitivity analysis showed that all body condition model parameter values were robust to measurement errors resulting from differences in picture quality (body length and width measurability) (Fig. S1).

4. DISCUSSION

Like most baleen whales, right whales rely heavily on stored energy for reproduction, particularly during lactation (Lockyer 1981, Miller et al. 2012, Christiansen et al. 2018). While the body condition of NARWs has declined during the last 3 decades (Rolland et al. 2016), this study provides the first comparison with healthy (i.e. growing) SRW populations. In agreement with our main hypothesis, we found that NARW juveniles, adults and lactating females were all in significantly poorer body condition compared to the SRW populations. Our results were robust to measurement errors resulting from variation in picture quality (body length and width measurability).

The largest difference in body condition was for lactating females, with NARW females being on average 21% units lower than the 3 SRW populations. To put this into perspective, the body condition of lactating females decreased by about 19% units during the first 3 mo of lactation, assuming a calf growth rate of 3.2 cm d⁻¹ (Christiansen et al. 2018). This early lactation period is considered the most energetically costly part of the reproductive cycle in baleen whales, since females are still relying on stored energy reserves during this time, while their calf is growing rapidly in size (Lockyer 1981, Miller et al. 2012, Christiansen et al. 2016a, 2018). In support of this, we found that the body condition of lactating females was generally better than that of juveniles and adults (Fig. 3). A compromised body condition during this critical time period means that NARW females may have considerably less energy available to invest in their calves, which is known to negatively influence calf growth rates (Christiansen et al. 2018). While we did not have data to directly investigate calf growth rates for NARWs, we were able to investigate the relationship between maternal body condition and calf length (Fig. 4D). We found that the absolute maternal cost of producing a similar sized calf (the slope parameter) was similar across populations, while the absolute maternal body condition at a given calf length (the intercept parameter) was significantly lower for NARW females. Assuming that NARW calves were growing at a slower rate compared to SRW calves, the observed difference in maternal condition could be due to a difference in the age of calves, with NARW calves being relatively older at a given body length compared to SRW calves. With NARW females having less energy reserves available to invest in their calf, this could result in them having to wean their calf at a smaller size. While weaning size is positively correlated to pup survival in pinnipeds (McMahon et al. 2000), this relationship is unknown in baleen whales. Alternatively, NARW females might compensate for their lower rate of offspring investment by extending the lactating period longer into the succeeding feeding season, when they are able to supplement their own body condition (and hence also their offspring investment) by concurrent feeding. While this strategy would likely result in a longer inter-calving interval for NARW females, since they would need more time to replenish their energy stores, it would not lead to a reduction in calf survival.

Lactating NARW females were also shorter in body length than the 3 southern populations. This was not the result of morphological differences (different asymptotic body lengths) between the 2 species, since whaling and stranding records show no species difference in body length (Tormosov et al. 1998, Moore et al. 2004, Huang et al. 2009, Fortune et al. 2012). With the absolute body volume of right whales being largely determined by their body length (Christiansen et al. 2018), lactating NARW females likely have less energy available to invest in their calves, which again will negatively affect calf growth rates. If we use Christiansen et al.’s (2018) relationship between maternal investment (rate of body volume loss) and maternal body length and condition for SRW in Australia, the magnitude of difference for lactating NARW (20.5% unit lower body condition and 96 cm shorter body length) equates to a loss in maternal rate of investment of 50% (rate of decline in maternal body volume: North Atlantic = 0.063 m³ d⁻¹; Australia: 0.126 m³ d⁻¹). Determining the lower threshold in body condition at which lactating females will no longer be able to energetically support their calves should be the aim of future research, as well as identifying the threshold below which fertility
whales and maternal body condition in humpback whales reported a positive relationship between calf body condition and maternal body condition in humpback whales *Megaptera novaeangliae*. While this lack of effect could be due to a difference in the timing of sampling between locations, this is unlikely since the body length of calves (a rough proxy for time since birth) did not vary significantly between locations. It hence seems that a reduction in maternal body condition in right whales does not lead to a reduction in calf body condition, although it could still be suppressing calf growth in length.

We found that Australian calves had a significantly lower body condition compared to the other SRW populations. This was unexpected, since lactating females in Australia had similar body condition as females in New Zealand and Argentina (Fig. 3), and so we anticipated their calves to have similar condition. However, the lower body condition of Australian calves does not seem to be correlated with lower calf survival, since the population is growing at a similar rate (5.55%, Bannister 2016) to the New Zealand population (5–7%, Carroll et al. 2013), and also has similar inter-calving intervals (Australia: 3.33 yr, Burnell 2001; New Zealand: 3.31 yr, Davidson et al. 2017). Further, only 4.2% of all measured calving intervals in Australia were 2 yr (Charlton 2017), an indication that females lost their calf early in lactation (Marón et al. 2015), compared to 8.9% in New Zealand (Davidson et al. 2017). It thus seems that the body condition of calves, within the range of values observed in this study, is not linked to their survival. Logically, calves should starve to death if their body condition falls below a critical limit where they can no longer afford to maintain homeostasis. However, assuming that their mothers can support them with sufficient energy, in the form of milk, to support their basic metabolic needs, baleen whale calves do not necessarily need to build up large fat reserves to survive. Instead, the lower body condition (higher surface area to volume ratio) of Australian calves might be an adaptation to the relatively warmer waters (lower heat loss) experienced on their breeding grounds (15°C, sea surface temperature on 1 August, www.meteoblue.com) compared to New Zealand (6.1–7.7°C, Rayment et al. 2015) and Argentina (11°C). In contrast, juveniles, adults and lactating females all rely on their own energy reserves during the breeding season, and so their survival and reproductive success is likely to be more closely linked to their body condition, whereas heat loss is likely to be less important due to their overall larger body size (lower surface area to volume ratio) and thicker blubber layer (more insulation).

Similar to lactating females, mature NARWs were in poorer body condition and smaller in size (i.e. body length) than the 3 SRW populations. Miller et al. (2011, 2012) found a similar difference in blubber thickness and body width of NARWs during the summer feeding season and SRWs in South Africa during the winter breeding season. The observed difference could be due to variations in the timing of sampling, with NARW adults being sampled early in their summer feeding season (when their energy reserves are still low from the previous breeding season) while SRW adults were measured during their winter breeding season (when they still have much of their energy reserves remaining). However, the magnitude of the difference in body condition between NARW and SRW adults in our study (Argentina = 27.9% units, Australia = 8.9% units, New Zealand = 18.9% units) was similar or exceeded the observed variation in body condition within locations (95% confidence range: Argentina = 15.2%, Australia = 9.3%, New Zealand = 13.6%, North Atlantic = 5.6%), suggesting that variation in the time of sampling alone cannot explain the observed difference in body condition of adults between locations (Fig. 3). Further, had the NARW adults been measured towards the end of the feeding season (when they are at their peak body condition), the fact that they are still in poorer BCI compared to SRW adults is even more alarming: Repeated sampling of NARW and SRW adults on their feeding grounds, to determine the rate of fattening, is needed to accurately quantify the magnitude of the difference in BCI between adults from the 2 species.

From the measured mature NARWs of known sex, 36.1% (13 of 36) were females, which, based on their reproductive cycle, should have been either in a pregnant or resting state (non-pregnant, non-lactating). A reduction of body reserves in pregnant females can result in less energy available for the foetus, which in minke whales *Balaenoptera acutorostrata* has been shown to result in smaller (i.e. shorter) foetuses (Christiansen et al. 2014). However, given that no NARW calves were born in the 2017–2018 breeding season (NARWC 2018), we can conclude that the measured adult females in this study were either...
resting, had a failed pregnancy or lost their calf shortly after giving birth (before they could be sighted). In resting females, lower body condition suggests that females are taking longer to recover from reproduction and nursing than right whales in other populations. This could help explain the substantially longer calving interval of NARW females (>7 yr) versus the SRW populations (~3.3 yr) (Burnell 2001, Cooke et al. 2003, Davidson et al. 2017, Pettis et al. 2020). By combining aerial photogrammetry (to determine BCI) and breath sampling (to determine reproductive status), future research should aim to assess whether a reduction in the BCI of NARW females is negatively affecting their fertility (their ability to become pregnant), pregnancy (their ability to complete gestation), offspring survival (their ability to energetically support their calf) and/or the time of recovery (their ability to deposit energy) from calving.

As for the adult whales, the lower body condition of immature NARWs could be partly due to variations in the timing of sampling between the North Atlantic and the southern populations. Younger NARW juveniles (1−4 yr) have also been found to have lower body condition (i.e. blubber thickness) compared to older juveniles (5−8 yr) (Miller et al. 2011). Potential age differences between locations are unlikely to explain the lower body condition of juvenile NARWs in this study, which on average were longer than juveniles in Argentina and New Zealand, and similar in size to Australian juveniles. Although the implication for vital rates is hard to determine, poorer body condition in juvenile NARWs could reduce the energy available for growth. This, in turn, could delay sexual maturation, which in baleen whales is strongly influenced by body size (Sigurjónsson et al. 1990). All else being equal, delayed sexual maturation would act to slow the population growth rate. A comparison of length-at-age growth curves between locations would help determine if NARWs are growing at a slower rate compared to the southern populations. This highlights the value of long-term monitoring projects with well-studied photo-identified individuals, for which age can be accurately determined.

With the NARWs being genetically isolated from the SRWs (Rosenbaum et al. 2000, Gaines et al. 2005), it is possible that some of the observed difference in BCI between the 2 species derives from genetic divergence and local adaptations to different environmental conditions (e.g. different water temperature and prey availability). Although traits closely associated with fitness, such as body condition, generally show low heritabilities (Mousseau & Roff 1987, Kruuk et al. 2000), the genetic component of variance in body condition can still be significant (Réale et al. 1999, Merilä et al. 2001). Further, differences in salinity and prey depth between locations (feeding grounds) could influence the optimal body shape (fat to muscle ratio) to achieve neutral buoyancy during foraging (Narazaki et al. 2018). However, our findings show that the structural body shape of NARWs and SRWs is very similar, while published records demonstrate similar body sizes and life history characteristics of the 2 species (Tormosov et al. 1998, Huang et al. 2009, Sharp et al. 2019). Further, our data show that the body condition (and hence volume) needed to produce a similar-sized offspring was the same for lactating females across populations (Fig. 4D), suggesting that the body energy content was similar across species and locations. Finally, differences in water temperature between locations (both on the breeding and feeding grounds) could influence the optimal body shape (and hence BCI) for minimizing heat loss. While this could explain the observed difference in body condition between right whale calves, heat loss is unlikely to lead to a population difference in body condition of juveniles and adult whales, due to their significantly larger body size (lower surface area to volume ratio and thicker blubber layer), and ability to tolerate a wide variation in temperatures across their spatial range (between subtropical and subpolar zones).

The observed differences in body condition between the NARW and the SRW populations are most likely to result from differences in the exposure to anthropogenic factors. While the 3 SRW populations examined reside in relatively remote and unimpacted environments, the home range of the NARW overlaps with heavily developed coastal areas, the greatest lobster and crab trap and line densities and some of the world’s busiest shipping lanes (Moore 2019). Despite management actions, ship strikes remain responsible for ongoing right whale mortalities in the North Atlantic (Moore et al. 2004, Knowlton et al. 2012, van der Hoop et al. 2013, Sharp et al. 2019); however, morbidity and mortality due to entanglement has become the predominant source of diagnosed trauma to NARWs since 2010 (NOAA 2018). This increasing entanglement in fishing gear is a major threat to NARWs; more than 83% of individuals carry scars from at least 1 entanglement, and 15.5% of the population is entangled every year (Knowlton et al. 2012). The additional drag, buoyancy and impeded foraging ability caused by various fishing gear leads to significant increases in the energy expenditure of right whales (Cassoff et al. 2011, van der Hoop et al. 2016, 2017). The cumula-
tive and prolonged interactions with fishing gear can lead to substantial reductions in body condition (Rol
land et al. 2012, Schick et al. 2013, Pettis et al. 2017, van der Hoop et al. 2017), which could result in
reproductive failure and even death (Moore et al. 2004, Robbins et al. 2015, Rolland et al. 2016). Anthro-
pogenic noise (e.g. from shipping) increases stress in NARWs, which carries energetic costs (Rolland et al.
2012).

Finally, climate-associated changes in right whale prey (the copepod *Calanus finmarchicus*) availability
and distribution in the North Atlantic are believed to reduce the rate of energy intake, body condition
and consequent calving rates (Miller et al. 2011, Meyer-Gutbrod et al. 2015, Meyer-Gutbrod & Greene 2018).
The smaller absolute body size (i.e. body length) of NARW adults and lactating females further suggests
that the high mortality rate of females in the North Atlantic might have skewed their age range to a sig-
nificantly lower average age than in SRWs (Pace et al. 2017). This would deprive NARW females of the
advantage of larger size as shown by Christiansen et al. (2018). The 2017–2018 NARW breeding season
highlighted the severity of the situation, when not a single calf was born into the population (NARWC
2018). Unless their situation improves soon, the ongoing decline of NARWs will result in them becom-
ing another of the growing list of cetaceans (including vaquita *Phocoena sinus*, Pennisi 2017;
Maui dolphin *Cephalorhynchus hectori maui*, Pala 2017; Gulf of Mexico Bryde’s whale *Balaenoptera
deni*, Corkeron & Kraus 2018) at serious risk of extinction.

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