# Static inflation and deflation pressure-volume curves in excised lungs of marine mammals

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Running head: Lung mechanics in marine mammals

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**Abstract:** Excised lungs from 8 marine mammal species (harp [*Pagophilus groenlandicus*], harbor [*Phoca vitulina*], and gray seal [*Halichoerus grypus*], Atlantic white-sided [*Lagenorhynchus acutus*], common [*Delphinus delphis*] and Risso's dolphin [*Grampus griseus*], long finned pilot whale [*Globicephala melas*], and harbor porpoise [*Phocoena phocoena*]) were used to determine minimum air volume of the relaxed lung (MAV, n = 15) and the elastic properties (pressure-volume curves, n = 24) of the respiratory system, and total lung capacity (TLC). Our data indicate that mass-specific TLC (sTLC,  $1 \cdot kg^{-1}$ ) does not differ between species or groups (odontocete vs. phocid) and agree with that estimated (TLC<sub>est</sub>) from body mass ( $M_b$ ) by: TLC<sub>est</sub> = 0.135  $\cdot M_b^{0.92}$ . Measured MAV was on average 7% of TLC, with a range from 0% to 16%. The pressure-volume curves were similar among species on inflation but diverged during deflation in phocids as compared with odontocetes. These differences provide a structural basis for observed species differences in depth at which lungs collapse and gas exchange ceases.

**Keywords** Lung mechanics, total lung capacity, minimum air volume, excised lung, diving physiology

#### Introduction

In 1940, Per Scholander (Scholander, 1940) published an article that described the unusual properties of the marine mammal respiratory system. Scholander suggested that the highly compliant lung and rib cage would easily compress, the air being shunted into the rigid upper airway. This hypothesis has remained a central tenet in marine mammal diving physiology, with the assumption that this trait will reduce uptake of nitrogen (N<sub>2</sub>) and the risk of decompression sickness (DCS). Still, only a few studies have attempted to determine how pressure affects gas exchange in forced or freely diving marine mammals. These studies have assessed alveolar compression either directly by quantifying the depth-related pulmonary shunt (Kooyman and Sinnett, 1982), or indirectly by measuring N<sub>2</sub> tension in arterial and venous blood during a dive (Falke et al., 1985; Kooyman et al., 1972) or measuring N<sub>2</sub> removal from the muscle following a series of repeated dives (Ridgway and Howard, 1979). As the estimated depth at which alveolar collapse occurs differs considerably between studies, Bostrom et al., (Bostrom et al., 2008) developed a mathematical model in an attempt to understand lung compression in marine mammals. The objective was to create a theoretical framework that could be used to predict the air volumes of the various compartments of the respiratory system to the limit of collapse. The results from the model suggested that both the initial diving lung volume and the relative size and structural properties (compliance) of the upper and lower airways are important in determining the depth at which the alveoli collapse and gas exchange ceases (Bostrom et al., 2008). The model output was compared to available data for collapse depth in phocids (Falke et al., 1985; Kooyman et al., 1970; Kooyman et al., 1972; Kooyman and Sinnett, 1982) and odontocetes (Ridgway and Howard, 1979), and it was concluded that behavioral (initial diving lung volume) and structural (lung and dead space compliance) variations between species could account for the observed differences (Bostrom et al., 2008; Fahlman et al., 2009).

Static pressure-volume loops are commonly used to measure the physical properties (compliance) of the respiratory system. Published data exist on excised lungs for several terrestrial species, but only a few measurements have been made for marine mammals (Denison and Kooyman, 1973; Denison et al., 1971). It is therefore difficult to assess the accuracy of the model output without species-specific respiratory compliance estimates. In addition, common indices of respiratory capacity, e.g. minimum air volume of the relaxed lung (MAV), total lung capacity (TLC), exist only for a limited number of marine mammals of different age classes

(Kooyman, 1973). Therefore, this study was undertaken to increase the current knowledge of the mechanical properties of the respiratory system of marine mammals and to estimate respiratory capacity (namely TLC and MAV) for a larger number of marine mammals. The primary objectives of the current study were to 1) compare the mechanical properties of a range of marine mammal species and 2) to estimate TLC from the P-V curves.

#### Material and methods

Animals

Thirty-one drowned (with no sea water observed in the respiratory airways or lungs) fishery bycaught (animals that either asphyxiated in gill nets, or bottom otter trawls) or stranded seals (Harp seal, *Pagophilus groenlandicus*; Harbor seal, *Phoca vitulina*; Gray seal, *Halichoerus grypus*) or odontocetes (Atlantic white-sided dolphin, *Lagenorhynchus acutus*; common dolphin, *Delphinus delphis*; long finned pilot whale, *Globicephala melas*; harbor porpoise, *Phocoena phocoena*; Risso's dolphin, *Grampus griseus*), were obtained respectively from the NMFS Northeast Fisheries Observer Program (NEFOP) or the Marine Mammal Rescue and Research Division of the International Fund for Animal Welfare (IFAW, Table 1). Specimens accidentally caught in the regional fisheries (bycaught) were transported by road to the Marine Research Facility (MRF) at Woods Hole Oceanographic Institution (WHOI), Woods Hole, MA, USA. Stranded animals that died or were euthanized on the beach were directly transported to WHOI. Animals were kept cold during transport and then stored at 4°C until the lungs were excised and studied less than 12 hours after arrival. No animal was euthanized for the purpose of this study. At the laboratory, the sex was determined and each animal weighed (± 0.2 kg) and routine morphometric measurements were completed (Table 1).

Each specimen was examined for obvious signs of scavenging, orifice discharges and bloating. Of the 31 specimens, static pressure-volume (P-V) loops and MAV were measured successfully in 24 and 15 lung samples, respectively (Table 1). In 9 lung samples, both MAV and P-V data were collected (Table 1). Not all tests could be performed on all specimens due to a variety of factors, e.g. time-constraints, leaks during P-V experiments.

Static pressure-volume measurements

The entire respiratory tract was excised (lungs, bronchi and trachea) with the heart attached. The excised tissues were placed on a tray and the trachea intubated immediately below the pharynx using a human (Hudson RCI, Teleflex Medical Inc., Research Triangle Dr., NC, USA) or veterinary (Equine Endotracheal Tube, Jorgensen Laboratories, Loveland, CO, USA) endotracheal tube. The endotracheal tube was attached to a manifold of 3-way valves with a 3 l syringe (Series 5530, Hans-Rudolph Inc, Kansas City, MO, USA) used to inflate the lungs in 100 ml increments.

The airway opening pressure ( $P_{air}$ , cm $H_2O$ , 1 cm $H_2O$  = 98.07 Pa) was measured using a differential pressure transducer (MPX type 339/2, Harvard apparatus, Holliston, MA, USA) connected to an amplifier (Tam-A, Harvard Apparatus, Holliston, MA, USA); these data were collected on a laptop computer using an A/D card (USB 1208LS, Measurement Computing, Norton, MA, USA) sampling at 2 Hz. The ambient pressure ( $P_{amb}$ ) was used as a reference and set at 0 cm $H_2O$ .

The lungs were initially pre-conditioned by at least 3 inflations to a transpulmonary pressure ( $P_{tp}$ ) of ~ 30 cm $H_2O$  ( $P_{tp} = P_{air} - P_{amb}$ ), which in mammals is often used to define the volume at TLC (Soutiere and Mitzner, 2004). The total volume used to inflate the lung to a  $P_{tp}$  of 30 cm $H_2O$  was divided in 4-5 equal increments. The pressure-volume (P-V) relationship was determined by adding or removing air in the increments determined during the pre-conditioning step. A minimum of 3 leak-free inflation/deflation curves was recorded. After injection or removal of an aliquot of air, the volume was held constant until the pressure had reached a plateau, usually for between 15-20 sec, before the next step-wise change in lung volume (Kooyman and Sinnett, 1979).

## Minimum air volume (MAV) estimates

In a subset of samples (n =15), MAV of the relaxed ( $P_{tp} = 0 \text{ cmH}_2O$ ) excised lungs was estimated by weighing the lung and the volume of fresh water it displaced and assuming tissue specific gravity of 1 kg ·  $I^{-1}$  (Kooyman and Sinnett, 1979). Three estimates were obtained for each lung; if the inter-test variation was > 10%, tests were repeated until the variation became less than 10%.

#### Data processing and statistical analysis

Room temperature and ambient pressure were used to convert all volumes to standard temperature pressure dry (STPD). We assumed intrapulmonary air to be saturated.

We modeled the relationship between pressure and volume for the excised lung using the sigmoidal equation published by Bostrom et al. (Bostrom et al., 2008):

$$V_{l} = a \cdot (1 + e^{(c(b - P_{air}))})^{-1}$$
(1)

where a, b, c, are fitting parameters and  $V_l$  the inflated air. The parameter a corresponds to the change in volume between the lower and upper asymptotes of the sigmoid. When inflating a relaxed excised lung to TLC, a is an estimate of the difference between TLC and MAV. The parameter b corresponds to the pressure at the inflection point of the sigmoid. The parameter c is an index of the compliance (steepness) of the curve where most of the volume change occurs (i.e. near b). Parameters were fitted using the software RGui (R: A Language and Environment for Statistical Computing, R Foundation for Statistical Computing, version 2.5.1, 2007).

We used two approaches to estimate TLC. In the first, we used the MAV and a from the P-V data. For lung samples where data existed to estimate both a and MAV (n = 9, Table 1), TLC was estimated as: TLC = a + MAV. For these nine samples, the MAV was then expressed as a relative proportion of TLC as: pMAV = MAV · TLC <sup>-1</sup>. For those samples where only MAV was determined (n = 7, Table 1), pMAV was computed as: pMAV<sub>est</sub> = MAV · TLC<sub>est</sub> <sup>-1</sup>, where TLC<sub>est</sub> was computed by the equation published by Kooyman (TLC<sub>est</sub> =  $0.135 \cdot M_b^{0.92}$ , Kooyman, 1973). The average pMAV was used to estimate TLC as: TLC =  $a \cdot (1-pMAV)^{-1}$ . Thus, we used the average pMAV for data where both a and MAV had been computed.

In the second and more common approach, TLC was estimated as the lung volume at a  $P_{tp} = 30 \text{ cmH}_2\text{O}$  (TLC<sub>30</sub> (Kooyman and Sinnett, 1979). As the inflated volume at  $P_{tp} = 30 \text{ cmH}_2\text{O}$  is an estimate of the vital capacity (VC), TLC<sub>30</sub> was estimated as TLC<sub>30</sub> = VC (1-pMAV)<sup>-1</sup>. To compare the two estimates, TLC was plotted against TLC<sub>30</sub>. Finally, the computed TLC was plotted against animal body mass ( $M_b$ ) and compared to the estimated TLC<sub>est</sub>.

The P-V relationship during inflation was analyzed by fitting the parameters a, b, and c, for each lung sample. TLC was estimated as described above and plotted against  $M_b$ . The scatter plot of TLC against  $M_b$  was compared to TLC<sub>est</sub>. The appearance of the deflation curve is

dependent on the starting lung volume (Glaister et al., 1973). One option was to standardize the inflation volume for each sample; another option was to standardize the curves by expressing  $V_l$  as a percentage of the starting volume, i.e. the maximum inflation volume. As we had data to different maximum inflation volumes for each sample, we opted for the latter to increase the statistical power. Thus, when analyzing the elastic properties of the deflation curves,  $V_l$  was expressed as a fraction of the total volume inflated. To model the biological variation within a species, and not just that of our samples, we used a nonlinear random effects model that treated animal as a random factor. To compare the curves within and between species, the 95% confidence limit (CL) was computed for each fitted model.

In this study P-values  $\leq$  0.05 were considered as significant, whereas P  $\leq$  0.1 was considered a trend. Data are presented as mean  $\pm$  standard error of the mean (SEM), unless otherwise stated.

#### **Results**

Summary morphometrics for the individuals used in the study are included in Table 1. Most odontocetes were stranded individuals (samples with an IFAW case number) while seals were generally bycaught specimens (D0, H0, and A case numbers).

Figure 1 shows an example of the elastic properties of an excised lung from a harbor porpoise (IFAW10-049). The experiment was repeated 4 times, and the error bars show the reproducibility between repeated inflation/deflation cycles. Two lung samples were difficult to inflate and were very stiff (D08685 and IFAW10-062, Table 1); therefore they were not included in the statistical analyses even though their data are reported in Table 1. The pressure-volume data were used to fit the parameters in Eq. 1. For lung samples where both a water displacement measurement and inflation experiment had been done, the average pMAV was 7% (Table 1). The pMAV<sub>est</sub>, estimated from the measured MAV and TLC<sub>est</sub>, was 5% and not different from pMAV (P > 0.1, t-test). TLC for animals where P-V data had been collected, was estimated by adding the average pMAV to a, i.e. TLC =  $a \cdot (1-0.07)^{-1}$ . TLC<sub>30</sub> was estimated as VC ·  $(1-0.07)^{-1}$ . The results for the two methods of estimating TLC were plotted against each other (Fig. 2A). In addition, the relative difference between TLC and TLC<sub>30</sub>, computed as (TLC-TLC<sub>30</sub>) · TLC<sup>-1</sup> · 100, were plotted against  $M_b$ . TLC was on average 12% higher than TLC<sub>30</sub>, with a median of 8%. TLC was plotted against  $M_b$  (Fig. 3). The estimated regression parameters from our TLC

(slope: 0.81, intercept: 0.225) were not significantly different from those of the TLC<sub>est</sub> equation (slope: 0.92, intercept: 0.135, Fig. 3). There were no apparent species-related trends and the residuals were randomly distributed with  $M_b$ .

To compare the elastic properties during inflation between species, the 95% confidence limit was estimated for b and c. Data for each species were fitted to Eq. 1. For the inflation data, we used the estimated TLC (TLC =  $a \cdot (1-0.07)^{-1}$ ) expressed as a percentage of TLC<sub>est</sub> for  $V_l$  in Eq. 1. For the deflation data,  $V_l$  was adjusted as described above. The SEM for each parameter was used to determine the 95% confidence limit (CL) and differences between species were assessed by determining which samples had overlapping CL. For the inflation curves, there were no differences in a, b, or c between age groups, within or between species, nor were there any differences between odontocetes (toothed whales) as compared with phocids (seals, Table 2). For the deflation curves, both b (Odontocete:  $12.2 \pm 0.8$ ; Phocid:  $8.2 \pm 0.5$ ) and c (Odontocete:  $0.17 \pm 0.01$ ; Phocid:  $0.25 \pm 0.01$ ) differed significantly between odontocetes and phocids. Again, none of the deflation parameters differed between age groups. The data for odontocetes and phocids were pooled to determine the best fit parameters for each group and the results are plotted in Fig. 4.

#### **Discussion**

We examined the mechanical properties of lungs excised from eight different marine mammal species. Our main objectives were to estimate: 1) the elastic properties and 2) the total lung capacity and minimum air volume in excised lungs from a range of marine mammals. The results from the P-V data were compared between species and groups. The parameter estimates suggest that the structural properties cause different behaviour during deflation in seals and toothed whales. We estimated TLC, and this was plotted against  $M_b$  to assess the mass-specific TLC (sTLC,  $1 \cdot \text{kg}^{-1}$ ). The mass-exponent to predict sTLC does not differ between species or groups (odontocete vs. phocid), and is similar to TLC<sub>est</sub> (Kooyman, 1973).

P-V curves and the structural properties of marine mammal lungs

Two theoretical studies have suggested that the depth of alveolar collapse could be estimated using the physical properties (P-V relationship) of the different parts of the respiratory system (Bostrom et al., 2008; Fitz-Clarke, 2007). These models assume that as the pressure

increases and the gas becomes more compressed, the conducting airways remain open. This allows the air in the alveoli to move into the central airway during compression. Bostrom et al., (Bostrom et al., 2008) showed that their model agreed well with available data for alveolar (Falke et al., 1985; Kooyman et al., 1972; Kooyman and Sinnett, 1982; Ridgway and Howard, 1979) and tracheal volume (Kooyman et al., 1970) at different depths (Bostrom et al., 2008). However, it was also shown that varying the elastic properties of the respiratory system would significantly alter the alveolar collapse depth. As only limited information exists on the structural properties of the respiratory system in marine mammals, our aim was to compare the elastic properties in different species. Our ultimate aim is to predict the depth of alveolar collapse and improve our understanding of how gas exchange is altered during breath-hold diving. However, the data presented in this study are not suitable to make such predictions, because we could not determine the small volume changes at negative trans-pulmonary pressures crucial to alveolar collapse. Our data do provide evidence that the elastic properties during deflation of the excised lungs differ between phocids and odontocetes. Assuming that pulmonary shunt and/or ventilation perfusion mismatch increase with decreasing alveolar volume, varying elastic properties may affect the functional gas exchange at shallow depths. Future studies need to refine the P-V curves at negative P<sub>tp</sub>'s and improve the estimates for MAV and the volume of the conducting airways. In addition, measurements are needed on live animals to test the hypothesis that the chest compliance does not affect collapse depth.

The data of the current study allow a comparison of the mechanical characteristics of the excised lungs and conducting airways in a range of marine mammals. While it is generally believed that all marine mammals have a compliant rib cage that recoils inward to very low lung volumes and that lungs collapse occur at shallow depths, only a few studies have tested this hypothesis and only in a limited number of species. Our data indicate that the inflation curves in excised marine mammal lungs are similar among age groups and species and between phocids and odontocetes. The inflation curves were similar to the previously published values for the California sea lion (Denison et al., 1971) and harbor porpoise (Kooyman and Sinnett, 1979). However, in the deflation curves there was considerably more hysteresis in the P-V loop in the phocid compared to the odontocete, causing lower recoil pressures in the former. The deflation curve for the odontocete is similar to that reported for the California sea lion (Denison et al., 1971) and harbor porpoise (Kooyman and Sinnett, 1979), whereas that of the phocid is similar to

that of the dog (Denison et al., 1971). When the pressure is reduced from  $30 \text{ cmH}_2\text{O}$  to  $10 \text{ cmH}_2\text{O}$  the volume in the odontocete decreases by 60 %, but only by 24 % in the phocid. At pressures below  $10 \text{ cmH}_2\text{O}$  the deflation curve in the odontocete has greater elastance. Extending the data below MAV, the sigmoidal equation approaches  $0 \text{ volume at } -20 \text{ cmH}_2\text{O}$ . In the California sea lion, 6% of TLC remained at  $P_{tp} = -30 \text{ cmH}_2\text{O}$  so our extrapolation needs to be viewed carefully as we have no data at negative  $P_{tp}$ 's. The greater pulmonary elastance in odontocetes suggests that proportion of intrapulmonary gas remaining in the alveoli during compression may be less in odontocetes than in phocids. This agrees with data from our recent study where the lung volumes at different dive depths (pressures) were measured in post-mortem specimens and the results indicated that collapse depth appear to be greater for cetaceans as compared with phocids (Moore et al., In press).

The differences in deflation curves may be due to structural or biochemical (surfactant) differences between phocids and odontocetes. In 1971, Denison et al. (Denison et al., 1971) showed that the reduction in alveolar volume during compression was greater in California sea lions than in dogs. It was suggested that these differences could be related to the greater abundance of elastic fibers and muscle in the supportive tissue of the lung and conducting airways in the sea lion (Denison et al., 1971; Kooyman and Sinnett, 1982); these anatomical features may contribute to the elastic differences observed in our study. In addition, preliminary results indicate that collateral ventilation, i.e., gas moving laterally through the lung parenchyma, occur in odontocetes (dolphin and pilot whale) but not in phocids (Fahlman unpublished data), further suggesting anatomical differences. Alternatively, differences between marine and terrestrial mammals could be related to differences in alveolar surface tension. Early reports suggested that surfactant properties of lung lavage fluid from marine mammals do not differ from those of terrestrial mammals (Denison et al., 1971; Kooyman and Sinnett, 1982). More recent work has shown that lung surfactants of pinnipeds (elephant seal, northern fur seal, California sea lion and ringed seal) have anti-adhesive properties that aid alveolar recruitment following collapse (Foot et al., 2006), but no study has reported surfactant properties of odontocete lungs. Further work is required to determine if the differences in hysteresis and deflation are related to differences in anatomy, biochemistry or both. In either case, the elastic properties allow us to improve predictions of the air volumes in the upper and lower airways in excised lungs at the point of collapse. While our results illustrate the mechanical differences in

excised lungs among species, studies on live animals are required to understand how variation in blood flow and chest compliance alters lung mechanics. For example, Leith (1976) showed that the chest wall of a ribbon seal is highly compliant and recoils inward. Our own observations agree with this, but the chest wall in the odontocete appears to be much stiffer (Fahlman et al., unpublished observation). The greater recoil of the odontocete chest wall may help compensate for the stiffer lung and allow for the passive exhalation in odontocetes (Olsen et al., 1969). Thus, future studies are needed to confirm these observations of chest compliance.

Our experimental design was similar to those used in previous studies that described the elastic properties of lungs excised from sacrificed (< 19 hrs) harbor porpoises (Kooyman and Sinnett, 1979) and California sea lions (Denison et al., 1971). The lung samples used in this study were obtained from animals that had either asphyxiated in gill nets, or bottom otter trawls (bycaught), or died during a stranding event (stranded). Therefore, the interval between death and study was often longer (> 24 hrs) than in previous studies in the harbor porpoise (Kooyman and Sinnett, 1979) and California sea lion (Denison et al., 1971). This delay may have changed tissue properties. In addition, bycaught animals died by drowning or asphyxiation. Asphyxiation was confirmed by absence of water in the conducting airways. Despite these differences, the variability in the P-V relationship between lung samples within a species was reasonably low, except in two cases (D08685La and IFAW10-062Dd), and our results for the harbor porpoise are similar to those previously obtained from fresh tissues (Kooyman and Sinnett, 1979).

As for the possibility that over-inflation may have damaged the lungs, no subpleural blebs were observed in this study after inflations to  $P_{tp} < 30 \text{ cmH}_2\text{O}$ . Typically, these blebs develop under the visceral mesothelium after inflations to a  $P_{tp} > 50 \text{ cmH}_2\text{O}$ . In addition, our results showed little variability within an individual (Fig. 1), suggesting that the lungs had not been damaged by repeated inflation/deflation cycles. In addition, no such damage was reported in lung histology of previous studies done in fresh tissue (Denison et al., 1971; Kooyman and Sinnett, 1979) and we consider it unlikely that our results are affected by damage of lung parenchyma.

#### Estimating TLC

We used two different methods to estimate TLC. In terrestrial mammals, TLC is usually defined as the air volume in the lung at a P<sub>tp</sub> ranging between 25 to 40 cmH<sub>2</sub>O (Venegas et al.,

1998). A  $P_{tp}$  of 30 cm $H_2O$  has been used to estimate TLC in excised marine mammal lungs (Kooyman and Sinnett, 1979) and this was value used in the current study. However, some species do not appear to reach maximal lung volume at  $P_{tp}$ 's as high as 80 cm $H_2O$  (Soutiere and Mitzner, 2004), and the estimated TLC will therefore vary depending on the pressure used to define TLC. For our data, most samples did not exhibit a distinct asymptote at a  $P_{tp} = 30$  cm $H_2O$ , e.g. Fig 1, and the fitted parameter a was therefore larger than the air volume at 30 cm $H_2O$ . It is possible that a standard  $P_{tp}$  may not be valid for all species or age classes. Alternatively, a sigmoidal equation can be used to define the upper asymptote (Glaister et al., 1973; Schroter, 1980; Venegas et al., 1998), resulting in a set of physiologically meaningful parameters that can be compared between samples from P-V curves produced under different experimental and pathological conditions (Venegas et al., 1998). Using this approach, some estimates of TLC differed from TLC<sub>est</sub> by as much as 50%, as small differences in the shape of the P-V curves result in large differences in a. Because histological examination was not possible in all lung samples, we cannot exclude that some specimens had undergone pathological changes.

While our limited data did not reveal age differences in TLC (normalized by body weight), the lung capacity of juveniles may be different from adults. Our estimates of TLC are in general agreement with those from the equation published by Kooyman (Kooyman, 1973) in a range of species and age classes. In the present study, the average pMAV and pMAV<sub>est</sub> values were, respectively 7% and 5% (Table 1) and measured MAV ranged between 0 and 16% of TLC. The ratio of dead space to alveolar volume is a dimensionless constant for terrestrial mammals (Leith, 1983). Our data suggest that this ratio is also constant in marine mammals, even in different age classes. In addition, the pMAV reported in this study is similar to the estimate from other species and age classes, e.g. the harbor porpoise (Kooyman and Sinnett, 1979), the awake Pilot whale (Olsen et al., 1969), the sei and fin whale (see pers. comm. in Kooyman and Sinnett, 1979), but lower than the average in the California sea lion (Denison et al., 1971).

## Summary

Our study confirms previous observations studies in the harbor porpoise (e.g. Kooyman and Sinnett, 1979) in which the residual air volume at  $P_{tp} = 0 \text{ cmH}_2\text{O}$  is smaller than in terrestrial mammals. The inflation curves of excised lungs for all marine mammals studied are similar to those measured in the dog. By contrast, whereas the deflation curve of the phocid resembles the

dog's, that of the odontocetes is similar to the California sea lion's and shows greater recoil at mid-lung volumes. These elastic differences during deflation may be due to anatomical and/or biochemical differences, and future studies are needed to confirm these findings in live animals, so that the relationship between ambient pressure, alveolar size, and the extent of gas exchange at depth can be assessed.

### **List of Symbols and Abbreviations**

 $M_{\rm b}$  = body mass

P<sub>tp</sub> =transpulmonary pressure

 $P_{amb} = ambient pressure$ 

P<sub>air</sub> =airway pressure

TLC = total lung capacity

MAV =minimum air volume

pMAV = proportional minimum air volume

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#### References

**Bostrom**, B. L., Fahlman, A. and Jones, D. R. (2008). Tracheal compression delays alveolar collapse during deep diving in marine mammals. *Respir Physiol Neurobiol* 161, 298-305.

**Denison, D. M., Warrell, D. A. and West, J. B.** (1971). Airway structure and alveolar emptying in the lungs of sea lions and dogs. *Respir Physiol* 13, 253-260.

**Denison, D. M. and Kooyman, G. L.** (1973). The structure and function of the small airways in pinniped and sea otter lungs. *Respir Physiol* 17, 1-10.

Fahlman, A., Hooker, S. K., Olszowka, A., Bostrom, B. L. and Jones, D. R. (2009).

Estimating the effect of lung collapse and pulmonary shunt on gas exchange during breath-hold diving: The Scholander and Kooyman legacy. *Respir Physiol Neurobiol* 165, 28-39.

Falke, K. J., Hill, R. D., Qvist, J., Schneider, R. C., Guppy, M., Liggins, G. C., Hochachka, P. W., Elliott, R. E. and Zapol, W. M. (1985). Seal lungs collapse during free diving: evidence from arterial nitrogen tensions. *Science* 229, 556-558.

**Fitz-Clarke, J. R.** (2007). Mechanics of airway and alveolar collapse in human breath-hold diving. *Respir Physiol Neurobiol* 159, 202-210.

**Foot, N. J., Orgeig, S. and Daniels, C. B.** (2006). The evolution of a physiological system: the pulmonary surfactant system in diving mammals. *Respir Physiol Neurobiol* 154, 118-138.

**Glaister, D. H., Schroter, R. C., Sudlow, M. F. and Milic-Emili, J.** (1973). Bulk elastic properties of excised lungs and the effect of a transpulmonary pressure gradient. *Respir Physiol* 17, 347-364.

**Kooyman, G. L.** (1973). Respiratory Adaptations in Marine Mammals. *American Zoologist* 13, 457-468.

**Kooyman, G. L., Hammond, D. D. and Schroeder, J. P.** (1970). Bronchograms and tracheograms of seals under pressure. *Science* 169, 82-84.

**Kooyman, G. L., Schroeder, J. P., Denison, D. M., Hammond, D. D., Wright, J. J. and Bergman, W. P.** (1972). Blood nitrogen tensions of seals during simulated deep dives. *Am J Physiol* 223, 1016-1020.

**Kooyman, G. L. and Sinnett, E. E.** (1979). Mechanical properties of the harbor porpoise lung, *Phocoena phocoena. Respir Physiol* 36, 287-300.

**Kooyman, G. L. and Sinnett, E. E.** (1982). Pulmonary shunts in harbor seals and sea lions during simulated dives to depth. *Physiol Zool* 55, 105-111.

Leith, D. E. (1976). Comparative mammalian respiratory mechanics. *Physiol* 19, 485-510.

**Leith, D. E.** (1983). Mammalian tracheal dimensions: scaling and physiology. *J Appl Physiol* 55, 196.

Moore, M.J., Hammar, T., Arruda, J., Cramer, S., Dennison, S., Montie, E., and Fahlman,

**A.** (2011). Hyperbaric computer tomographic measurements of lung compression in seals and dolphins. *J Exp Biol* 214, 2390-2397.

**Olsen, C. R., Hale, F. C. and Elsner, R.** (1969). Mechanics of ventilation in the pilot whale. *Respir Physiol* 7, 137-149.

**Ridgway, S. H. and Howard, R.** (1979). Dolphin lung collapse and intramuscular circulation during free diving: evidence from nitrogen washout. *Science* 206, 1182-1183.

**Scholander**, **P. F.** (1940). Experimental investigations on the respiratory function in diving mammals and birds. *Hvalrådets skrifter* 22, 132.

**Schroter, R. C.** (1980). Quantitative comparisons of mammalian lung pressure volume curves. *Respir Physiol* 42, 101-107.

**Soutiere, S. E. and Mitzner, W.** (2004). On defining total lung capacity in the mouse. *J Appl Physiol* 96, 1658-1664.

**Venegas, J. G., Harris, R. S. and Simon, B. A.** (1998). A comprehensive equation for the pulmonary pressure-volume curve. *J Appl Physiol* 84, 389-395.

**Table 1.** Animal identification number (D0=bycaught, IFAW=stranded), group, species, sex (male/female), body mass ( $M_b$ ), animal length, age class, lung weight, minimum air volume of the relaxed lung (MAV), MAV expressed as a volume or as a percentage of the model fitted TLC in Fig. 3 (pMAV). Species abbreviation (Pg = Pagophilus groenlandicus, Pv = Phoca vitulina, Hg= Halichoerus grypus, La = Lagenorhynchus acutus, Dd = Delphinus delphis, Gm = Globicephala melas, Pp = Phocoena phocoena, Gg = Grampus griseus). Group abbreviation: Ph=phocid, Od=odontocete. D0 animals were either caught in gill nets or otter trawls (indicated by \*). Animals were grouped into age class by their reproductive age, defined as: Young of the year (YOY), juvenile (J), immature (I), and adult (A). †Specimens for which P-V curves were not collected, in which TLC<sub>est</sub> (Kooyman, 1973) was used to estimate pMAV (pMAV<sub>est</sub>).

Animal ID	Group	Species	Sex	Mb	Length	Age	Lung weight	MAV	pMAV
	•	1		(kg)	(cm)	class	(g)	(ml)	% TLC
D05952	Ph	Pg	F	27.4	99	YOY	935	20	1
IFAW09-034	Ph	Pg	F	25.5	106	YOY		273	11
IFAW09-041	Ph	Pg	M	22.7	90	YOY	591	395	15
IFAW09-047	Ph	Pg	F	24.2	91	YOY			
H0-0011	Ph	Pg	F	28.4	105	YOY	944	484	11
D01848*†	Ph	Pg	F	32.7	120	J	984	238	7
IFAW09-029†	Ph	Pg	F	15	93	YOY	419	45	3
IFAW09-076†	Ph	Pg	F		102	YOY	1023	23	<1
D06306†	Ph	Pg	M	35.5	118	J	1044	34	1
D05845	Ph	Pv	N/A	26	97	YOY			
D06318	Ph	Pv	M	25.2	94	YOY	894	442	16
D08811	Ph	Pv	F	25.6	104	YOY	704		
D08812	Ph	Pv	M	29.8	103	YOY	895		
D06322*	Ph	Hg	M	223	215	I	5278	10	0
D08883	Ph	Hg	F	21.4	89	YOY	608		
D07662*	Ph	Hg	M	208	210	I	4264	50	0
D05294*	Od	La	M	250	260	A	5850		
D08685*†	Od	La	F	69.4	171	I			
D08982*	Od	La	F	110.5	203	A	2809		
IFAW10-085	Od	La	M	154.2	217	A	3983		
IFAW09-064	Od	La	M	129.2	214	I	2981		
IFAW09-046	Od	Dd	M	57.6	174	I			

IFAW09-049	Od	Dd	N/A	59.4	177	I			
IFAW09-062†	Od	Dd	M	96.7	215	A	2859	1420	16
IFAW09-209	Od	Dd	M	106.2	223	A			
IFAW10-065	Od	Dd	M	75	190	I	1200		
DO0194*†	Od	Dd	M	102.4	210.4	A	2710	302	3
IFAW09-203	Od	Gm	M		313	J	9930		
IFAW09-214	Od	Gm	M	305	279	J	7857		
IFAW10-049	Od	Pp	M	22.1	105	YOY	682	88	3
D08417	Od	Gg	M	74.4	190	YOY	2022	332	4

**Table 2.** Parameter estimates ( $\pm$  SEM) for A) inflation and B) deflation pressure-volume data fitted to a sigmoidal equation. The sigmoidal equation results in 3 parameters, a, b and c as detailed in the Material and Methods. For fitting the inflation data, we used the estimated TLC expressed as a percentage of TLC<sub>est</sub> rather than the inflated volume for  $V_l$  in Eq. 1. Species abbreviations are:  $Pg = Pagophilus \ groenlandicus$ ,  $Pv = Phoca \ vitulina$ , Hg = Halichoerus grypus,  $La = Lagenorhynchus \ acutus$ ,  $Dd = Delphinus \ delphis$ ,  $Gm = Globicephala \ melas$ ,  $Pp = Phocoena \ phocoena$ ,  $Gg = Grampus \ griseus$ .

Table 2A.

Species	a (% TLC)	b (cmH <sub>2</sub> O)	c (cmH <sub>2</sub> O)	Number of animals	Number of data points
Pg	$101.9 \pm 1.8$	$24.9 \pm 1.0$	$0.20 \pm 0.02$	5	111
Pv	$89.9 \pm 5.1$	$22.0 \pm 0.8$	$0.22 \pm 0.02$	4	102
Hg	$98.7 \pm 10.3$	$20.1 \pm 0.3$	$0.20 \pm 0.03$	3	126
La	$96.5 \pm 1.8$	$21.1 \pm 2.6$	$0.25 \pm 0.03$	4	53
Dd	$102.9 \pm 6.6$	$22.4 \pm 2.1$	$0.20 \pm 0.02$	4	130
Gm	$103.7 \pm 7.2$	$24.0 \pm 0.6$	$0.19 \pm 0.01$	2	73
Pp	$110.4 \pm 2.4$	$21.3 \pm 0.2$	$0.22 \pm 0.01$	1	39
Gg	$108.4 \pm 2.9$	$21.0 \pm 0.2$	$0.26 \pm 0.01$	1	14
Phocid	$97.3 \pm 4.5$	$21.7 \pm 1.2$	$0.24 \pm 0.02$	12	339
Odontocete	$104.6 \pm 12.4$	$22.0 \pm 0.2$	$0.19 \pm 0.01$	12	309

Table 2B.

Species	a (% TLC)	b (cmH <sub>2</sub> O)	c (cmH <sub>2</sub> O)	Number of animals	Number of data points
Pg	$92.5 \pm 1.4$	$7.1 \pm 0.9$	$0.25 \pm 0.02$	5	85
Pv	94.1 ± 1.6	$7.8 \pm 0.3$	$0.23 \pm 0.02$	4	99
Hg	$94.8 \pm 1.5$	$7.3 \pm 1.3$	$0.26 \pm 0.03$	3	116
Dd	$100.2 \pm 1.3$	$11.7 \pm 1.3$	$0.17 \pm 0.01$	4	109
La	$101.4 \pm 4.5$	$10.0 \pm 0.7$	$0.18 \pm 0.02$	4	62
Gm	$105.0 \pm 2.5$	$16.4 \pm 0.3$	$0.14 \pm 0.01$	2	65
Pp	$98.7 \pm 1.8$	$12.3 \pm 0.4$	$0.18 \pm 0.01$	1	34
Gg	$111.1 \pm 6.3$	$15.4 \pm 1.0$	$0.15 \pm 0.01$	1	14
Phocid	$98.6 \pm 0.9$	$8.2 \pm 0.5$	$0.25 \pm 0.01$	12	300
Odontocete	$102.8 \pm 1.6$	$12.2 \pm 0.8$	$0.17 \pm 0.01$	12	284

## Figure legends

Figure 1. Mean pressure-volume data ( $\pm$  SEM, n = 4) for inflation and deflation of an excised lung of a harbor porpoise (IFAW10-049). Volume includes the estimated MAV and was normalized as a proportion of the estimated total lung capacity (TLC<sub>est</sub>) (Kooyman and Sinnett, 1979) and pressure is pressure at the airway opening ( $P_{air}$ ), which in excised lungs is the transpulmonary pressure.

Figure 2. A) Total lung capacity (TLC) estimated using two methods. TLC was estimated as the volume of air in the lung between the lower and upper asymptote of the sigmoidal function published (Eq. 1) and the average minimum air volume (MAV). TLC<sub>30</sub> was estimated as the air volume injected for a transpulmonary pressure of 30 cmH<sub>2</sub>O plus the MAV. The solid line is the line of identity. B) Data showing the difference between TLC and TLC<sub>30</sub> plotted against body mass ( $M_b$ , kg).

Figure 3. Animal body mass versus model-predicted total lung capacity (TLC). TLC was estimated as  $TLC = a \cdot (1-0.07)^{-1}$ , where a is the parameter for the difference in inflated volume between the lower and upper asymptote in equation 1, and 0.07 the average estimated residual volume as determined by water displacement experiment. The broken regression line is the best fit to the data ( $TLC = 0.225 \cdot M_b^{0.81}$ ), and the solid regression line is the equation that predicts TLC in marine mammals ( $TLC_{est} = 0.135 \cdot M_b^{0.92}$ , Kooyman and Sinnett, 1979). Pg = Pagophilus groenlandicus, Pv = Phoca vitulina, Hg= Halichoerus grypus, La = Lagenorhynchus acutus, Dd = Delphinus delphis, Gm = Globicephala melas, Pp = Phocoena phocoena, Gg = Grampus griseus. Closed symbols are phocids and open symbols odontocetes.

Figure 4. The inflation and deflation curves from fitting Eq. 1 to the data for the phocids and odontocetes. The inflation volume data were expressed as a proportion of the TLC<sub>est</sub>. The deflation data were expressed as a proportion of the maximum inflation volume (including the estimated residual volume of 7%) before each deflation.







