Evidence of seawater drinking in fasting subadult hooded seals (*Cystophora cristata*)

Fernando Alvira-Iraizoz • Erling Sverre Nordøy

Abstract The purpose of this study was to investigate the total water turnover rate of fasting subadult hooded seals in order to elucidate to what extent these animals rely in seawater drinking/mariposia at this life stage. Considering mariposia is important for later accurate estimations of food consumption using water turnover rate as a proxy. Five subadult hooded seals were kept fasting for 4 days in a seawater pool. Total body water (39.6 ± 2.5 % of total body mass) decreased by 3.1 ± 0.4 % of initial body water over the experimental period. Turnover rates were 16.7 ± 3.9 (influx) and 24.6 ± 4.6 (efflux) ml · day⁻¹ · kg⁻¹ with a net water loss of 710 ± 51 ml · day⁻¹.

It was estimated that the seals drank approximately 947 ml of seawater per day, which corresponds to 61 % of total daily water influx. Initial body water was relatively low as a result of the high body fat (46.9 ± 3.2 % of initial body mass) shown in the animals. It is concluded that subadult fasting hooded seals drink significant amounts of seawater during fasting. Although mariposia stands out as the major source of free water in fasting hooded seals, the amount of seawater ingested is unlikely to provide a net gain of free water as it is provided by metabolic water. However, it may contribute to excrete the excess of urea produced during early phase 1 of fasting.

Keywords: marine mammals, mariposia, total body water, turnover rate, water homeostasis

Introduction

Water balance in pinnipeds has been previously studied by use of the labelled-water method in order to understand the basic physiology underlying water homeostasis (Depocas et al 1971; How and Nordøy 2007; Lea et al 2002; Nordøy et al 1992; Ortiz et al 1978; Ortiz et al 2002; Skalstad and Nordøy 2000). The labelled-water method has also been used to estimate body composition and milk and food intake (Arnould et al 1996; Beauplet et al 2003; Carlini et al 2001; Crocker et al 2012; Donohue et al 2002; Lydersen and Hammill 1993; Lydersen and Kovacs 1996; Lydersen et al 1997; Noren et al 2003; Noren et al 2008; Reilly and Fedak 1990) in different stages of life of pinnipeds. Therefore, accurate estimates of water turnover rate (rH₂O) are important to acquire a better insight about how pinnipeds maintain water homeostasis and to perform precise estimates of food/milk intake at different life stages.

Water homeostasis of a number of pinnipeds, such as the hooded seal (*Cystophora cristata*) for instance, is challenged throughout the entire life since such seal species spend most of their lifetime either swimming/diving in open waters or fasting. Folkow et al (2010), using satellite telemetry, determined the overall time spent in open seawater by young of the year hooded seals to be 76 %, covering a vast area of the north Atlantic. Subadult and adult (2-10 years old) hooded seals (Jan Mayen stock) also disperse after the moulting/breeding periods and their positions have been recorded by satellite-linked transmitters as far as southwest of Iceland, British Isles, Faeroe Islands, south of Bear Island, the Norwegian Sea and Svalbard. The trips performed by those animals lasted on average 47 ± 22 days (n = 46) (Folkow et al 1996). The northwest Atlantic subpopulation (Newfoundland and Labrador) have also been shown to travel large distances (Andersen et al 2009). During these migratory movements, seals have no access to fresh water. Moreover, hooded seals accomplish long post-weaning, moulting and breeding fasts (Folkow et al 1996; Lydersen et al 1997). In this case, they neither consume any food. The major effects of fasting have been described as profound mass loss, reduced metabolic rate, increased plasma osmolality, reduction in urine production and, ultimately, dehydration (Beauplet et al 2003; Nordøy et al 1993; Nordøy et al 1990; Nordøy et al 1992; Rosen and Trites 2002).

A number of studies have shown that pinnipeds effectively restrict water loss by a number of different adaptations. The efflux routes are respiratory evaporative water loss (REWL), urine, cutaneous evaporative water loss (CEWL) and water in faeces. Studies of the rate of REWL in
grey seals (*Halichoerus grypus*) have shown that water loss through respiration is relatively low due to a very well developed nasal heat and water exchange mechanism (Folkow and Blix 1987; Skog and Folkow 1994). In addition, Lester and Costa (2006) showed that apnoeic breathing, together with the complex nasal turbinates, allows northern elephant seals (*Mirounga angustirostris*) to reduce REWL by 41 %. Urine production, moreover, may also be effectively reduced when water is scarce due to a very high concentrating ability of the kidneys (Hong et al 1982; Storeheier and Nordøy 2001). Various studies of different species have shown that urine production is considerably reduced during fasting periods (Nordøy et al 1993; Nordøy et al 1990). CEWL has been generally considered very low in pinnipeds living in cold environments. In such conditions, seals increase or reduce body heat loss by adjusting perfusion to the flippers and heat transfer through the blubber (Kvadsheim and Folkow 1997). Nordøy et al (1992) estimated CEWL in fasting grey seal pups resting on land to be only about 3 ml · day$^{-1}$ · kg$^{-1}$ and active sweating has only been reported in California sea lions (*Zalophus californianus*) (Matsuura and Whittow 1974). Water in faeces in fasting seals is negligible since no food is ingested.

As mentioned before, during long migratory periods the seals have no access to fresh water in form of ice/snow. Snow and ice may be a source of water when hauling out on pack ice as described by Renouf et al (1990) in harp seals (*Phoca groenlandica*). Thus, most of water influx during these prolonged journeys in open waters must be obtained from free and metabolic water from food, inspired water and seawater drinking (mariposia) (Skalstad and Nordøy 2000). Depocas et al (1971) estimated inspired water rate to be 93-108 ml · day$^{-1}$ in harbor seals (*Phoca vitulina*) while Skalstad and Nordøy (2000) calculated respiratory influx to be about 114 ml · day$^{-1}$ or 5 % of total water influx in hooded seals fed capelin. Regarding seawater drinking, early studies concluded that marine mammals do not voluntarily drink seawater (Depocas et al 1971; Fetcher and Fetcher 1942; Irving et al 1935) and that mariposia is not an effective mechanism to obtain water (Tarasoff and Toews 1972). However, studies by Gentry (1981) and Skalstad and Nordøy (2000) showed evidences of spontaneous mariposia in a number of pinnipeds. Moreover, experimentally dehydrated harp seals were able to restore water balance with ad libitum access to seawater (How and Nordøy 2007). Early studies of ringed (*Pusa hispida*) and Baikal (*Pusa sibirica*) seals showed that these species are able to concentrate urine up to 2000 mOsm · kg$^{-1}$, well above seawater concentration, which serves as a remarkable mechanism for water conservation (Hong et al 1982) and reinforces the potential of mariposia. Unfortunately, direct measurement of the amount of water entering the body by each of those routes in wild subadult hooded seals is nearly impossible. When food is deprived (fasting), catabolism of subcutaneous fat has proven to be the main source of energy and water (Arnould et al 2001; Castellini et al 1987; Nordøy et al 1993; Ortiz et al 1978; Verrier et al 2009). Up to 1.07 grams of free water are produced per 1 gram of fat metabolized (Schmidt-Nielsen 1964).

The purpose of this study was, by use of a bolus injection of tritiated water ($^3$H$_2$O), to estimate the total water influx ($^3$H$_2$O) in order to elucidate to what extent fasting subadult hooded seals depend on seawater drinking/mariposia at this life stage. An accurate estimation of the level of mariposia will improve our understanding of the water balance of young hooded seals, which may result in more accurate estimations of food and milk consumption when using the labelled water method.

**Materials and Methods**

**Animals**

Five subadult hooded seals, 3 females (no. 1, 3 and 5) and 2 males (no. 2 and 4), aged 1.5 years were used in this study. The animals were captured as weanlings in the Greenland Sea (70°N, 16°W) and transported on board RV “Helmer Hanssen” to the research facility of the Arctic Chronobiology and Physiology research group at UiT-the Arctic University of Norway. The animals were kept in the facilities in a 42000 l water tank with continuous exchange of seawater and exposed to simulated (70°N) seasonal variation in day length prior the experiment onset. This study was carried out in strict accordance with the recommendations of the Animal Welfare Act of Norway and the Norwegian Regulation on Animal Experimentation. All applicable institutional, national and/or international guidelines for the care and use of animals were followed. All procedures performed involving animals were in accordance with the ethical standards of the Norwegian Ethical Committee for Animal Research (Permit Number: FOTS ID: 5422). During an experimental period of four days, the animals were fasted in order to investigate this status. Initial body mass (BM) ranged from 83.4 to 103.5 kg (Table 1) and seals appeared to be in excellent health before, during and after the experimental period.

**Table 1 Seal identification number (ID #), sex (F: female, M: male) and initial body mass (BM) of the 5 seals used in the study.**

<table>
<thead>
<tr>
<th>ID #</th>
<th>Sex</th>
<th>BM (kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Seal 1</td>
<td>F</td>
<td>103.5</td>
</tr>
<tr>
<td>Seal 2</td>
<td>M</td>
<td>87.6</td>
</tr>
<tr>
<td>Seal 3</td>
<td>F</td>
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<tr>
<td>Seal 4</td>
<td>M</td>
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<tr>
<td>Seal 5</td>
<td>F</td>
<td>87.5</td>
</tr>
</tbody>
</table>

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Experimental protocol

The seals were fasted for 24 h in the seawater tank prior to onset of the experiment. Then, a 1.0 ml intramuscular injection of a sedative (50 mg · ml⁻¹ tiletamine-50 mg · ml⁻¹ zolazepam, Zoetil Forte Vet, Reading, L ́Hay-Les-Roses, France) was administered and the animals were weighted (Dini Argeo MCWNT 1RF-1; 10-150 kg, accuracy ±0.5 kg; Spezzano di Fiorano, Italy). The animals were then restrained on an especially designed board and two catheters (Selacon-TM 16G/1.70x160 mm, The Hague, the Netherlands) inserted into the extradural intravertebral vein (EDV) at the level of the 1st and 4th lumbar vertebra. After that, a bolus dose of ²H₂O (1 - 2.5 µCi · kg⁻¹) was injected, as described by Reilly and Fedak (1990), through the catheter located at the 1st lumbar vertebra (EDV1). A volume of 10 ml of saline (9 g · ml⁻¹ NaCl, Melsungen, Germany) was gently flushed through the catheter after the injection. Whereafter an additional 5 ml of saline were used to rinse the outer part of the catheter. Blood samples were collected from the catheter inserted at the level of the 4th lumbar vertebra (EDV2) 30, 60 and 90 minutes after ²H₂O injection. After every blood sampling the catheter was gently flushed with 10 ml of saline in order to avoid contamination of subsequent blood samples and blood clotting. After the initial ²H₂O experiment, the seals were reintroduced into the seawater tank. After 96 h of fasting with ad libitum access to seawater, seals were restrained on the same board and blood samples collected from the EDV in order to estimate rH₂O. All blood samples were immediately centrifuged for 10 min at 2000 rpm (Kubota KS-8000, Tokyo, Japan) and plasma was transferred to cryovials (VWR International, Norway) and stored at -80ºC for later analysis.

In addition, blood was sampled from the femoral vein to determine whether sampling site might have an effect when determining rH₂O using labelled water. Nordgarden et al (2000) measured the directional flow in the extradural intravertebral vein (EDV) by use of Doppler flowmeter from non-diving to the diving situation in harp seals (Phoca groenlandica) and demonstrated changes in flow direction, although not correlated with diving. Pinnipeds have also proven to reduce their circulation to the flippers during exposure to low ambient temperatures (Hammelel al 1977). The major veins of the hind flippers are another important site for sampling blood. One may speculate that such circulatory changes may affect the rate of dilution of a radiolaubelled substrate when using veins in these locations for sampling. However, no statistical differences between the specific activity (SA) of plasma samples from any of the sampling sites were found, therefore, this issue was not further investigated.

All plasma samples were deproteinised with 70 % perchloric acid (HClO₄) (200 µl · ml⁻¹ plasma) for protein precipitation prior scintillation counting (Bray 1960). An aliquot of 0.5 ml deproteinized plasma was added to 9.5 ml of scintillation fluid Ultima GoldTM (LSC cocktail, Sigma-Aldrich Química S.L., Tres Cantos, Madrid) and analysed in triplicates to determine radioactivity of ³H₂O in a liquid scintillation counter (Packard 1900 TR Beta-Teller Liquid Scintillation Analyser, Oslo, Norway) (How and Nordøy 2007; Reilly and Fedak 1990; Skalsstad and Nordøy 2000). A set of standards was run prior every counting. The resulting data was adjusted to the real water content of deproteinized plasma samples. Plasma water content was calculated by desiccation of 1.0 ml plasma aliquots at 60 ºC for 24 hours and HClO₄ water content was assumed 30%. Background level of tritium in plasma, 69 ± 54 cpm · ml⁻¹ (n = 5), was subtracted from SA at equilibrium (SA₀).

Body composition

Initial total body fat (TBF₁) was calculated as described by Reilly and Fedak (1990) for grey seals: %TBF₁ = 105.1 – 1.47 (%TBF) where, %TBF is the initial body fat as a percentage of body mass and %TBF is the total body water as percentage of body mass. Thus, initial lean body weight (LBW₀) was calculated as %LBW₀ = 100 - %TBF₁.

Total body water (TBW) and rH₂O

TBW was calculated as described by Lifson and McClintock (1966): TBW = i.d. / SA₀ where; i.d. is the injected dose in counts per minute (cpm) and SA₀ is the specific activity of ³H₂O in plasma at equilibrium in cpm · ml⁻¹. rH₂O, total water influx and efflux, was calculated using equations 4 and 6 provided by Nagy and Costa (1980) assuming a linear change in TBW. The amount of water entering and leaving through different routes from the main TBW pool was then estimated. According to Nordøy and Blix (1985), Nordøy et al (1990) and Reilly (1991), 94 % of the energy expended during the fasting period is derived from subcutaneous fat in grey seal pups. Moreover, Nordøy et al (1993) showed that only 2 – 4 % of the energy expended by harp seal pups was derived from protein catabolism after day 3 of fasting. To simplify calculations, it was assumed that fasting subadult hooded seals cover the energy expenditure by subcutaneous fat loss. Metabolic water was calculated as a function of grams of fat metabolized, thus, 1 g of fat metabolized produced 1.07 g of water. Respiratory water influx was calculated as follows. Assuming 100 % relative humidity in air (Skalsstad and Nordøy 2000), the partial pressure of water vapour (Pᵥ) in inspired air was calculated for the given room temperature (6.5 ºC). Respiratory day volume was calculated from Folkow and Blix's equation (Folkow and Blix 1987): RMV = 0.042 · ADMR + 0.119; where RMV is respiratory minute volume in l · min⁻¹ and ADMR is the average daily metabolic rate.
expressed in Watts. Then, daily water vapour influx \( W_i \) was calculated using: 
\[ W_i = RDV \cdot P_{H_2O} / 760; \]
where RDV is respiratory day volume and 760 is the standard pressure in air in mmHg. \( W_i \) was converted to ml · day\(^{-1} \) of absorbed water using molar volume at the specific temperature and molar mass of water. Finally, the extent of mariposa was determined by subtracting metabolic water and respiratory influx from the total water influx determined by the \(^3\)H\(_2\)O method.

REW\(_L\) was calculated using a similar approach as when calculating respiratory influx. Exhaled air temperature was estimated in grey seals, according to Folkow and Blix (1987), to be 29º C at the given room temperature. \( P_{H_2O} \) was adjusted to this temperature and introduced in the calculations previously described to determine REW\(_L\). CEW\(_L\) was considered to be negligible in actively swimming hooded seals at a low ambient temperature since submersion and blood flow adjustments to the appendages are considered the ideal way to decrease body heat (Kvadsheim and Folkow 1997) rather than evaporation. Water loss in faeces was also assumed negligible in fasting seals (Nordøy et al 1992; Storeheier and Nordøy 2001). Urine production was determined by subtracting REW\(_L\) from total water efflux.

\( rH_2O \) was corrected for exchange using equation 39 provided by Lifson and McClintock (1966). Fractionation was corrected using a transformed version of equation 31 presented by Lifson and McClintock (1966): 
\[ r_{H_2O}(calc) = r_{H_2O}(x_f + 1 - x); \]
where \( r_{H_2O}(calc) \) is the corrected turnover rate, \( r_{H_2O} \) is the efflux rate, \( x \) is the fraction of turnover subjected to fractionation and \( x_f \) is the fractionation factor \( (x_f = 0.93, \) according to Sepall and Mason (1960)). Exchange and fractionation combined amounted in average \((n = 5)\) to an overall 6 % or 104 ml · day\(^{-1} \) overestimation of rH\(_2\)O which were subtracted from total water efflux.

**Statistical analysis**

Means for BM and radioactivity of each animal after isotope equilibration were compared to those at the end of the experimental period by T-test for paired samples. Data sets of each seal were previously analysed for normality using the Shapiro-Wilk test (for small sample sizes). In all cases, the data sets followed a normal distribution \((p > 0.05)\). Pearson correlation tests were run to investigate associations between body composition and rH\(_2\)O. Correlation was considered significant when \( p < 0.05 \). Pearson correlation coefficient was considered high when \( r < 0.5 \) and assumed to be the most representative parameter due to the reduced sample size. All statistical analyses were carried out using SPSS-software.

**Results**

BM showed a significant \((p < 0.05)\) linear decrease down to an average \((n = 5)\) of 96.9 ± 0.4 % of BM\(_i\) during the 96 h of experimental fasting. This amounted to 710 ± 58 g · day\(^{-1} \). Assuming all energy to be derived from subcutaneous fat reserves the amount of BM loss corresponded to an average \((n = 5)\) loss of 2.8 ± 0.2 kg (Table 2) of fat/water over four days of fasting. TB\(_F\) was in average \((n = 5)\) 43.1 ± 5.4 kg or 46.9 ± 3.2 % of BM\(_i\) while LB\(_W\) amounted in average \((n = 5)\) to 48.5 ± 3.9 kg or 53.1 ± 3.2 % of BM\(_i\) (Table 2).

<table>
<thead>
<tr>
<th>BM(_i) (kg)</th>
<th>BM(_f) (kg)</th>
<th>ΔBM (kg)</th>
<th>TBF(_i) (kg)</th>
<th>%TBF(_i)</th>
<th>LBW(_i) (kg)</th>
<th>%LBW(_i)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Seal 1</td>
<td>103.5</td>
<td>100.6</td>
<td>2.9</td>
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<td>51.1</td>
<td>50.6</td>
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<td>43.6</td>
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<td>42.5</td>
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<td>44.0</td>
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<td>87.5</td>
<td>84.8</td>
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<td>42.7</td>
<td>50.2</td>
</tr>
<tr>
<td>Av±SD</td>
<td>91.68±8.1</td>
<td>88.84±8.2</td>
<td>2.8±0.2</td>
<td>43.1±5.4</td>
<td>46.9±3.2</td>
<td>48.5±3.9</td>
</tr>
</tbody>
</table>

Av: Average; SD: Standard deviation

Tritium turnover rate was assumed to follow a linear decrease (Figure 1) and, thus, \( rH_2O \) was calculated assuming linear changes in TB\(_W\). Tritium activity in TB\(_W\) decreased in average \((n = 5)\) 16 ± 3 % in 4 days of fasting.

TB\(_W\) was on average \((n = 5)\) 39.6 ± 2.5 % of BM\(_i\). Total water influx and efflux were calculated to be on average \((n = 5)\) 1491 ± 294 ml · day\(^{-1} \) or 16.7 ± 3.9 ml · day\(^{-1} \) · kg\(^{-1} \) and 2200 ± 333 ml · day\(^{-1} \) or 24.6 ± 4.6 ml · day\(^{-1} \) · kg\(^{-1} \), respectively. Overall daily water loss amounted to 710 ± 51 ml · day\(^{-1} \) (Table 3). Net water loss matched BM loss as the result of assuming fat to be the only metabolic substrate during fasting. There was no correlation between BM\(_i\) and influx and/or efflux rates \((0 < r < 0.5)\). However, a high negative correlation \((r < -0.5)\) was shown between TBF\(_i\) and efflux or influx rates and all water pathways. In other words, a high TBF\(_i\) may contribute to reduce overall rH\(_2\)O. Lastly, no correlation was found \((r ≈ 0)\) between LB\(_W\) and efflux and/or influx rates.

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Respiratory influx, metabolic water and mariposia amounted to 6 % (1.0 ± 0.0 ml · day⁻¹ · kg⁻¹), 33 % (5.3 ± 0.1 ml · day⁻¹ · kg⁻¹) and 61 % (10.5 ± 3.8 ml · day⁻¹ · kg⁻¹) of total water influx, respectively. REWL and urine production amounted to 16 % (3.8 ± 0.1 ml · day⁻¹ · kg⁻¹) and 84 % (20.8 ± 4.6 ml · day⁻¹ · kg⁻¹) of total water efflux, respectively.

**Discussion**

On average, the hooded seals in this study lost 710 ± 58 g · day⁻¹ or 7.7 g · day⁻¹ · kg⁻¹ while fasting. Grey seal pups during the prolonged post-weaning fast (over 7 weeks) with no access to water showed a BM loss of 10.6 g · day⁻¹ · kg⁻¹ (Nordøy et al 1992). TBFᵢ was on average (n = 5) 46.9 ± 3.2 % of BMᵢ. Body composition compared well with recently weaned pups of other species. TBFᵢ has been reported to be 48.5 ± 0.9 % in sub Antarctic fur seal pups (Verrier et al 2009), 50 % in elephant seal pups (Ortiz et al 1978), 48.5 ± 0.9 % in subantarctic fur seal pups (Verrier et al 2009) and 40 % in grey seal pups (Sparling et al 2006), all just prior to weaning. These pups have all accumulated body fat for an extended period during lactation and, therefore, have a very high fat content and a very low TBW since subcutaneous adipose tissue contains only 10 % of free water (Ortiz et al 1978).

By using equations provided by Nagy and Costa (1980) total water influx and efflux were calculated to be 16.7 ± 3.9 ml · day⁻¹ · kg⁻¹ and 24.6 ± 4.6 ml · day⁻¹ · kg⁻¹, respectively. No other studies have been done in subadult hooded seals, thus a direct comparison is difficult. On one hand, hooded seal pups during the post weaning fast have shown a daily water influx of 15-16 ml · day⁻¹ · kg⁻¹ (Lydersen et al 1997; Schots et al 2017) which is very similar to that measured in the present study. On the other hand, nursing hooded seal pups (Lydersen et al 1997) and actively feeding juvenile hooded seals (Skalstad and Nordøy 2000) have shown a total water influx of approximately 125 and 70 ml · day⁻¹ · kg⁻¹, respectively. This demonstrates that fasting subadult hooded seals with access to seawater show rH₂O clearly more similar to those of fasting (food and water deprivation) hooded seal pups than to those of nursing/feeding individuals. Other species, however, have shown different turnover rates under different conditions. Nordøy et al (1992) calculated water influx and efflux to be as low as 8 and 12 ml · day⁻¹ · kg⁻¹, respectively, in fasting grey seal pups without access to any exogenous source of water. Elephant seal pups fasting under natural conditions showed, as well, an exceptionally low rH₂O of 5.4 ± 0.7 (influx) and 6.0 ± 0.8 (efflux) ml · day⁻¹ · kg⁻¹ (Ortiz et al 1978). Moreover, Verrier et al (2009) calculated influx and efflux rates in fasting subantarctic fur seal pups of 7.9 ± 0.9 and 8.8 ± 1.0 ml · day⁻¹ · kg⁻¹, respectively. In contrast, enclosed (no access to seawater) and free-ranging (access to seawater) Antarctic fur seal pups showed a water influx of 20.5 ± 0.3 and 33.0 ± 1.7 ml · day⁻¹ · kg⁻¹, respectively, during natural fasting (Lea et al 2002) having in both cases much higher rH₂O and percentage TBW than the hooded seals in the present study. This suggests ones more that different species of pinnipeds might be differently adapted with regard to water conservation when food and/or water are deprived as mentioned by Ortiz et al (1978). We could not find any correlation between BMᵢ and rH₂O suggesting that rates of influx and efflux are not dependant on the mass of the seal at the beginning of fasting. However, a negative correlation was shown between TBFᵢ and rH₂O. In this case, influx and efflux rates were lower in seals with higher TBFᵢ. This may suggest that seals with larger reserves of subcutaneous fat may better cope with situations of food/water deprivation not only by potentially producing more metabolic water but also by reducing overall rH₂O. It might be explained by the possibility of an even further decrease in metabolic rate and, consequently, the rH₂O (Reilly 1991; Richmond et al 1962), due to the larger isolative blubber layer.

![Figure 1](https://example.com/figure1.png)

Figure 1 Tritium activity (cpm) plotted as a function of time (Teq, equilibration time; Tf, end of the experiment) in five subadult hooded seals. Seal #1 (▲), seal #2 (♦), seal #3 (●), seal #4 (▼) and seal #5 (■).
With regard to influx rates, metabolic water production is likely to be overestimated since fat was considered the only source of energy/water and catabolized to its end-products. However, it was considered to be more accurate to calculate the metabolic rate as a function of fat catabolized than using a multiplier of BMR predicted by the Kleiber equation (Kleiber 1961). Rapid changes in metabolic rate at the beginning of a fasting period have been shown in different species (Boily and Lavigne 1995; Cherel et al 1992; Hudson 1973; Keys et al 1950; Markussen 1995; Markussen et al 1992; Mrosovsky and Sherry 1980; Rosen and Trites 2002). Thus, assuming a constant metabolic rate during early stages of fasting may lead to errors. For instance, hooded seal pups during the first days of the post-weaning fast have shown metabolic rates ranging 2.1 – 4.4 times the predicted BMR (Lydersen et al 1997). Besides, the calculated metabolic rate of the animals used in this study compared well with that of presumably less active fasting grey seal pups under similar conditions (Nordøy et al 1990). Respiratory influx and REWL were calculated as described by Folkow and Blix (1987) from empirical data obtaining similar results for REWL than in the present study. Moreover, this method has been validated in later studies of water balance (Skalstad and Nordøy 2000). Therefore, the rate of water influx through mariposia will be, if anything, underestimated.

Table 3 Total water efflux (rH\textsubscript{2}O efflux) and influx (rH\textsubscript{2}O influx), efflux and influx rates per kg of BM and overall water loss per day of five subadult hooded seals. Average (n = 5) values ± SD are given in the bottom line.

<table>
<thead>
<tr>
<th></th>
<th>rH\textsubscript{2}O efflux</th>
<th>rH\textsubscript{2}O influx</th>
<th>rH\textsubscript{2}O efflux</th>
<th>rH\textsubscript{2}O influx</th>
<th>Δ H\textsubscript{2}O loss</th>
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<tr>
<td></td>
<td>(ml · day\textsuperscript{-1})</td>
<td>(ml · day\textsuperscript{-1})</td>
<td>(ml · day\textsuperscript{-1} · kg\textsuperscript{-1})</td>
<td>(ml · day\textsuperscript{-1} · kg\textsuperscript{-1})</td>
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<td>Seal 4</td>
<td>2627</td>
<td>1827</td>
<td>32.1</td>
<td>22.3</td>
<td>800</td>
</tr>
<tr>
<td>Seal 5</td>
<td>2286</td>
<td>1612</td>
<td>26.5</td>
<td>18.7</td>
<td>674</td>
</tr>
<tr>
<td>Av±SD</td>
<td>2200±333</td>
<td>1491±294</td>
<td>24.6±4.6</td>
<td>16.7±3.9</td>
<td>710±51</td>
</tr>
</tbody>
</table>

Av: Average; SD: Standard deviation

Figure 2 Total water turnover (rH\textsubscript{2}O, ml · day\textsuperscript{-1} · kg\textsuperscript{-1}). Water influx rates via respiration, mariposia and metabolic water; and efflux rates via REWL and urine production of five subadult hooded seals. Percentage contribution of each different route of water influx and efflux to the total turnover rate is also given.

Mariposia contributed 10.5 ± 3.8 ml · day\textsuperscript{-1} · kg\textsuperscript{-1} or 61% to total water influx of short-term fasting subadult hooded seals. Despite of being the major contributor of free water during fasting, this amount of mariposia compares very well with that shown in actively feeding hooded seals (9 ml · day\textsuperscript{-1} · kg\textsuperscript{-1}) (Skalstad and Nordøy 2000). Although mariposia has been shown to restore water balance in dehydrated harp seals (How and Nordøy 2007) and potentially give a net gain of water (Storeheier and Nordøy 2001), ingested seawater shown in the present study was most probably used to create urinary osmotic space for urea, as described by Wolf et al (1959), rather than to provide a net gain of water. Storeheier and Nordøy (2001) showed that adult harp seals fasting for 2 days produced excess amount of urea during early phase I of fasting, which was more efficiently excreted when seawater was given as bolus administration. This voluntary seawater drinking may thus aid in eliminating the excess of urea produced in early stages of fasting due to protein catabolism. Therefore, this study supports the theory of “urea osmotic space” proposed by Wolf et al (1959) and further elucidated by Costa (1982) studying sea otters (Enhydra lutris).

It is concluded that also subadult fasting hooded seals drink significant amounts of seawater when available during early fasting. Although metabolic water remains as the most reliable source of free water for fasting subadult hooded seals, mariposia may play an important role in excreting the excess of urea produced during phase I of fasting.
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