Evaluation of the Deuterium Dilution Method to Estimate Body Composition in the Barnacle Goose: Accuracy and Minimum Equilibration Time

Götz Eichhorn1,*, G. Henk Visser1,2
1Centre for Ecological and Evolutionary Studies, University of Groningen, P.O. Box 14, 9750 AA Haren, The Netherlands; 2Centre for Isotope Research, University of Groningen, 9747 AG Groningen, The Netherlands

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ABSTRACT

We examined body composition in barnacle geese (Branta leucopsis) by proximate carcass analysis and by deuterium isotope dilution. We studied the effect of isotope equilibration time on the accuracy of total body water (TBW) estimates and evaluated models to predict fat-free mass (FFM) and fat mass (FM) from different measurements varying in their level of invasiveness. Deuterium enrichment determined at 45, 90, and 180 min after isotope injection did not differ significantly. At all sampling intervals, isotope dilution spaces (TBWd) consistently overestimated body water determined by carcass analysis (TBWc). However, variance in the deviation from actual TBW was higher at the 45-min sampling interval, whereas variability was the same at 90 and 180 min, indicating that 90 min is sufficient time to allow for adequate equilibration. At 90 min equilibration time, deuterium isotope dilution overestimated TBWc by 7.1% ± 2.6% (P < 0.001, paired t-test, n = 20). This overestimate was consistent over the range of TBW studied, and TBWc could thus be predicted from TBWd (r² = 0.976, P < 0.001). Variation in TBWc and TBWd explained, respectively, 99% and 98% of the variation in FFM. FM could be predicted with a relative error of ca. 10% from TBW estimates in combination with body mass (BM). In contrast, BM and external body measurements allowed only poor prediction. Abdominal fat fresh mass was highly correlated to total FM and, if the carcass is available, allows simple means of fat prediction without dissecting the entire specimen.

Introduction

Somatic stores are a key factor in defining an animal’s body condition, and much of the variation in survival and reproduction has been attributed to the optimization of body reserves (Blem 1990; Carey 1996; Houston et al. 2007). Because of the central importance of energy and nutrient storage, their assessment has become an increasingly important aspect in current research (Brown 1996; Stevenson and Woods 2006). Birds in particular face the energetic dilemma of high energy expenditure for activity and maintenance and limitations of storage abilities due to their aerial lifestyle (McNab 2002).

A variety of methods have been applied to assess fat mass (FM) and fat-free mass (FFM) in vivo and in vitro (overviews in Blem 1990; Brown 1996; Gessaman 1999; Speakman 2001; Stevenson and Woods 2006). These methods differ in the accuracy of measuring the variable of interest and in the degree of invasiveness for the study animal. Although proximate body composition analysis is regarded as the most accurate method and the standard against which all other methods are evaluated, it obviously represents the most invasive, that is, lethal, avenue. Additionally, it is a labor- and time-intensive method. From such carcass analyses, researchers have recognized that an animal’s body water content represents a fairly stable proportion of the FFM because lipids are stored nearly free of water (Pace and Rathbun 1945; Odum et al. 1964; Ellis and Jehl 1991). While some animals can experience large changes in body mass and composition, particularly during extensive periods of fasting, the relative water content in the FFM (H₂OFMM) seems not to be significantly affected (Groscolas et al. 1991; Cherel et al. 1992). Consequently, estimating total body water (TBW) content enables prediction of FFM and subsequently, in combination with total body mass (BM), FM. Indeed, many studies used body water content estimates to successfully predict body stores (Campbell and Leatherland 1980; Miller 1989; overview Table VI in Blem 1990; Boos et al. 2000; but see Jamieson et al. 2006). Using calibrated regressions with dissectible fat depots, such as abdominal and leg fat pads, represents another method to estimate total FM in carcasses without analyzing the entire specimen (Thomas et al. 1983; Piersma 1984; Boos et al. 2000; Jamieson et al. 2006).
The refinement of isotope analysis techniques has yielded nondestructive means to estimate TBW by the principle of isotope dilution, allowing longitudinal studies and work where killing the animal is not an option. This method relies on releasing isotopically labeled water molecules into the body water pool and, after mixing, determining isotope concentrations in body fluids (usually blood) of a single timed sample (plateau approach) or a series of samples (intercept approach) to derive an estimate of TBW (Speakman et al. 2001). Because it involves only one sampling event and a shorter experimental period, the plateau approach is less invasive for the animal. Additionally, a single sample reduces costs for material and analysis. However, with the plateau approach, appropriate timing of the single sample is important, that is, after mixing of the marker with the body water is completed and before wash-out of the marker becomes effective. To reduce the latter, captive animals are deprived of food and water during the equilibrium period. The time a wild animal is held in captivity can crucially affect its performance, in particular during the breeding period when parental care for a clutch or brood has to be provided. Our general research goal is to employ isotope dilution to assess body composition of incubating barnacle geese (Branta leucopsis) in the field. To this end, we want to reduce the equilibrium time without affecting data quality. Furthermore, it is known that the isotope dilution method generally overestimates the actual TBW volume because some of the labeled atoms exchange with nonaqueous body constituents (Culebras and Moore 1977; Schoeller et al. 1980). A review of studies employing hydrogen isotopes in four bird species revealed a considerable variation by which the dilution space differed from actual TBW, ranging from underestimates of 2.3% to overestimates of 18% (Table 3 in Speakman et al. 2001). As has been noted frequently (Gessaman 1999; Mata et al. 2006; Shaffer et al. 2006), there is clearly a need for more bird studies evaluating estimates of TBW and other body components measured by isotope dilution against standard proximate body composition analysis.

In this study on captive barnacle geese, we compared estimates of TBW, FFM, and FM measured by proximate carcass analysis and by deuterium isotope dilution. Our specific objectives were to (1) assess how equilibration time may compromise accuracy of the dilution method; (2) determine the level of overestimation of TBW by deuterium dilution; and (3) evaluate the accuracy of predictions of FFM and FM from TBW and/or other predictor variables (BM, external morphological measurements, abdominal fat), depending on lethal and non-lethal approaches.

**Methods**

**Animals and Experimental Setup**

Animals were taken from a stock of barnacle geese Branta leucopsis kept at the Biological Centre of the University of Groningen in Haren, The Netherlands. Geese were kept on grassland while receiving ad lib. supplementary food (a mixture of grain and pellets). A total of 21 adult birds (≥2 yr old), consisting of 10 females and 11 males, were selected to achieve maximal range in body condition; the body condition criterion was residual BM from a regression of BM on the first principal component from a factor analysis including tarsus and total head length. To further increase the variation in body condition, we kept a subsample of two females and two males separate from the stock on grassland of lower food quality and with only limited supplementary food for 2 wk before the experiment. Their average mass loss during this period was 210 ± 72.5 g. All animals were used in the carcass analysis but only 10 of the 11 males for the isotope dilution space measurements because of leakage of the isotope mixture through the puncture hole of the thin skin in one male. Birds were sexed by cloacal inspection and confirmed by examination of gonads during dissection. Most birds (16) were collected from February to mid-March 2006 and the others (four males, one female) on April 21, 2005.

To standardize treatment, animals were put in bird cages with no access to food but access to drinking water on the evening before the isotope dilution experiment. The next morning (ca. 15 h later), 2 h before administration of the isotope solution, drinking water was removed until the end of the experiment, 4–6 h later. The birds were intraperitoneally injected with a 99.9% deuterium isotope solution (Sigma Chemicals) using 1.0-mL insulin syringes. The actual mass of each volume injected was determined by weighing the syringe before and after injection on an analytical balance (Mettler AG204) to the nearest 0.1 mg. Average dose mass was 1.1153 ± 0.0021 g (n = 20). Blood samples were collected from nine females and seven males at each of the following sampling times: 45, 90, and 180 min after injection. Additionally, one female and three male birds were sampled at 90 min. To estimate deuterium background levels, blood samples before isotope administration were taken from three female and three male birds. Blood was collected from the brachial and intertarsal veins and stored in flame-sealed microcapillaries. After the last blood sample was taken, birds were fully anesthetized with 3 mL intraperitoneally injected Nembutal (natriumtoaborbit 60 mg/mL), followed by cervical dislocation 10–15 min later. BM was then measured to the nearest 1 g, and carcasses were placed in plastic bags and refrigerated until being further processed on the next day or were double-packed and frozen at −20°C until dissection and body composition analysis. Daily care and management of the animals and the experimental protocol were approved by the Animal Experiments Committee of the University of Groningen (license DEC 4081B).

**Isotope Analyses**

The blood in the capillary tube was distilled in a vacuum line, where water vapor was cryogenically trapped in a quartz tube using liquid nitrogen. After complete transfer, the vacuum system was brought to room pressure by admitting dry nitrogen.
The insert was then quickly brought into a standard vial for automatic injection and sealed with a septum. During the sample preparation, internal water standards (gravimetrically prepared from pure deuterated water and also stored in flame-sealed capillaries) covering the entire enrichment range of the blood samples were distilled alongside the samples. This way, possible systematic effects on isotope enrichment due to the distillation process were accounted for. Such effects were also monitored in every batch by comparing the distilled standard waters with the same waters that were introduced into the vials directly. The actual δ^2H measurements were performed in automatic batches using a Hekatech high-temperature pyrolysis unit (Gehre et al. 2004) in which the injected water reacted with the glassy carbon available in the reactor according to H_2O + C → H_2 + CO. The H_2 and CO gas, emerging into a continuous He flow through the system, were then led through a gas chromatography (GC) column to separate the two gases in time and fed into a GVI IsoPrime isotope ratio mass spectrometer for the actual isotope analysis. For the analysis of δ^2H (from the H_2 gas emerging first from the GC column), every sample was injected typically six times from the same vial into the furnace in 0.2-μL quantities. Memory effects of the high-temperature pyrolysis oven were corrected for using a memory correction algorithm similar to the one described by Olsen et al. (2006). In the complete analysis scheme, several quality checks were incorporated. The isotope scales were calibrated using multiple distilled samples of two of the standard waters (being at the lower and the higher end of the sample range, respectively), whereas the measured δ^2H values for a third standard, representing the expected midrange of blood samples, were used as quality “targets” and had to be measured within 1% of their assigned values in order to meet the quality criterion for the batch. All sample analyses were run at least in duplicate (more times if values differed by more than 2.5%), and we used the average of values differing from each other by less than 2.5%.

Measurement of the Hydrogen Dilution Space (TBW_d)

Using the plateau approach (Speakman et al. 2001) and employing equation (1), we calculated the hydrogen dilution space (TBW_d) by taking into account the quantity of the dose (Q_d, mol), the δ^2H concentration of the dose (C_d, atom%), the δ^2H background concentration (C_b, atom%), and the δ^2H concentration of individual blood samples (C_i, atom%) taken at the various sampling intervals:

$$\text{TBW}_d = 18.02 \times Q_d \times \frac{C_d - C_b}{C_i - C_b}. \quad (1)$$

Background levels of δ^2H measured in six birds before dose administration averaged 4.11%, with a range of 33.01%, which represented only 0.60% of the average measured deuterium enrichment after dose injection (5.532%, n = 52). Therefore, we applied this average background value for all birds.

Dissection and Body Composition Analysis

Fresh or thawed carcasses were weighed; then, all feathers were plucked and the birds were reweighed, with the difference being plumage fresh mass. All skin was removed as was associated subcutaneous fat. The following organs were dissected out, weighed, and analyzed for water and fat content: left flight muscle complex (pectoralis and supracoracoideus), left leg musculature (attached to the tibiotarsus and femur), abdominal fat (a discrete deposit in the abdominal cavity, excluding mesenteric fat adhering to the intestines), gizzard, intestines (including mesenteric fat and ceca), heart, liver, spleen, and kidneys. Before analysis, the esophagus, gizzard, and intestines were emptied and reweighed. The total wet content excised from these organs was 20 ± 9 g. The right flight and right leg musculatures were excised and retained for other work, and their contributions to dry mass and FFM were estimated via the equivalent masses of their left counterparts. Organs were cut into small pieces of ca. 1 cm³, and bones of the skeleton were broken to expose marrow and brain before being oven-dried at 60°C until constant mass (7–15 d). Total body water from the carcass analyses (TBW_c) was calculated as the carcass fresh mass after plucking minus the sum of all dried tissues, thereby accounting for general water loss during dissection. Thus, water absorbed by feathers is intentionally not included in TBW_c because (external) plumage water is not part of the body water pool measured by isotope dilution. Lipids were extracted from the tissues with a soxhlet apparatus using petroleum ether as a solvent. We refer to whole-body FFM as total wet lean mass, including feathers and skeleton, calculated from fresh BM minus extractable FM.

Calculations and Statistics

Statistical analyses were performed with SPSS, version 14. All results are reported as mean ± 1 SD unless stated otherwise and were considered to be significant at P < 0.05. For all parametric tests, assumption of normality and homogeneity of variances were evaluated using the Kolmogorov-Smirnov test and Levene’s test, respectively (Zar 1999). As a measure of structural size, we derived scores of the first principal component (PC1) from a factor analysis based on four external measurements: the lengths of tarsus, total head (i.e., including bill), maximum wing chord, and keel (measured from the anterior tip of the carina to the end of the sternum, at the transition with the abdominal cavity). Tarsus was measured with calipers to the nearest 0.1 mm; all other variables were measured with a ruler at 1-mm accuracy. Variables had a similar factor loading (0.72–0.94) on PC1, which explained 73% of the total variance. We used ANOVA to test for mean differences among sex and GLM, with PC1 included as covariate.

Differences in estimated dilution space with time after isotope injection were compared by a repeated-measures ANOVA. Two-tailed paired t-tests were used to compare means of TBW_d and TBW_c. We used estimated TBW and other predictor var-
The coefficient of determination increased slightly from
be more appropriate than a linear fit (Piersma 1984). Although
cluded abdominal fat in multiple regressions, we tested whether
dissected fat pad, not chemically extracted fat). Before we in-
ally, we took the set of predictor variables from the basic model
to investigate whether FM predictions could be improved. Fi-

variables (see below) to predict FM and FFM by two approaches:
(1) multiple regression analyses and (2) assuming a constant
FFM hydration.

We applied a stepwise backward elimination procedure in
the multiple regression analyses. Starting with the nondestruc-
tively obtained variables tarsus, head, wing, keel, BM, and sex
as a basic model, we extended the set of predictor variables
and included either TBWd and dry BM or TBWc and dry BM
to investigate whether FM predictions could be improved. Fi-
ally, we took the set of predictor variables from the basic model
and included additionally abdominal fat fresh mass (i.e., of the
dissected fat pad, not chemically extracted fat). Before we in-
cluded abdominal fat in multiple regressions, we tested whether
a curvilinear relationship between FM and abdominal fat would
be more appropriate than a linear fit (Piersma 1984). Although
the coefficient of determination increased slightly from \( r^2 = 0.90 \)
to \( r^2 = 0.92 \), adding a quadratic term did not significantly
improve a linear fit of FM to abdominal fat \( (P = 0.06) \).

We used double cross-validation to evaluate the robustness
and replicability of regression equations following the proce-
dure described by Guan et al. (2004). Briefly, subjects from the
original data set were randomly assigned to two groups, equal
or similar in number and sex ratio of subjects. The statistically
significant predictor variables derived from the original full data
set were applied in both subgroups to develop predictive equa-
tions and derive coefficients of determination \( (r_{12}^2 \) and \( r_{22}^2, \) where
the first subscript number refers to subsample’s data and the
second subscript number to subsample’s regression coeffi-
cients). Standardized regression coefficients and \( Z \) scores of
predictor variables and of the response variable were used in
all cross-validation procedures. The predictors’ regression co-
efficients were crossed over the two subsamples to produce equations and
coefficients of determination \( (r_{12}^2 \) and \( r_{22}^2, \) from actual group data using the regression coefficients from the
other group for the predictions. Using this double-cross pro-
cedure, we calculated two shrinkage values: \( r_{12}^2 - r_{22}^2 \) and \( r_{22}^2 - r_{12}^2. \) The more closely the shrinkage estimate approaches 0, the
greater the degree of stability is across subsamples. Further-
more, two invariance coefficients were derived by correlating
the predicted values of subsample 1 with the predicted values
of subsample 1 using the regression coefficients of subsample
2 \( (r_{11, 12}) \) and vice versa \( (r_{21, 22}) \). As these invariance coefficients
approach 1, more confidence can be obtained in the replicability
of the results.

2. As an alternative to calibrated regression equations and
assuming a constant water content in the FFM \( (H_{2}O_{FFM} = \)
TBW : FFM = constant), we can infer the individual FFM and
FM from the following equations:

\[
\text{FFM} = \frac{\text{TBW}}{H_{2}O_{FFM}},
\]

\[
\text{FM} = \text{BM} - \text{FFM}.
\]

We will refer to this approach as the Pace and Rathbun (1945)
approach (sensu Mata et al. 2006).

Results

Body Composition by Carcass Analysis

Carcass analyses are listed in Table 1. Animals in this study
covered a broad scale of BM and composition, ranging twofold
in BM and from 2% to 25% in lipid content. Males were larger
than females according to PC1, the first principal component
from a factor analysis including tarsus, wing, skull, and keel
length \( (F_{1, n} = 31.18, P < 0.001) \). Significant differences between
sexes were also found for BM, TBW, and FFM. However, these
were caused by the sex-related differences in structural size.
Thus, when sex was tested together with PC1 in one analysis
to explain differences in BM, TBW, and FFM, only PC1 ex-
plained a significant part \( (P < 0.05 \) in all models), whereas var-
ation due to sex became nonsignificant. Females in our sample
tended to have higher fat loads \( (FM : BM, F_{1, 19} = 3.39, P = 0.08) \).

<table>
<thead>
<tr>
<th>Table 1: Whole-body composition by carcass analysis</th>
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<tr>
<td>Variables</td>
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<tr>
<td>BM ( (g) )</td>
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<tr>
<td>TBWc ( (g) )</td>
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<tr>
<td>FFM ( (g) )</td>
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<tr>
<td>FM ( (g) )</td>
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<tr>
<td>TBWc : BM (%)</td>
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<tr>
<td>TBWc : FFM (%)</td>
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<td>FM : BM (%)</td>
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</table>

Note. Given are body mass \( (BM) \), total body water from carcass analysis \( (TBW_c) \), fat-free mass \( (FFM) \), fat mass \( (FM) \), and fractions
(\%) of water and fat content for sexes combined and males and females separately.

\* Including fresh plumage mass in BM; if plumage is included from BM, the fractions are TBW : BM = 60.6% ± 4.1% and
TBWc : FFM = 68.8% ± 0.8%.

\* Significant differences between sexes \( (P < 0.05) \).
**TBW Measured by Deuterium Dilution and the Effect of Equilibration Time**

Within-individual variation in deuterium isotope enrichment occurred mainly between the first two sampling events and was independent of the size of the individual body water pool (Fig. 1). However, a pronounced increase over this period was restricted to a few individuals only, and overall changes between sampling times were too low to be significant (repeated-measures ANOVA, $F_{2,30} = 1.894, P = 0.168$).

At all sampling intervals, carcass body water was significantly overestimated by the deuterium dilution space (paired $t$-test, $P < 0.001$), ranging from average values of 9.2% at 45 min to 7.1% at 90 min (Table 2). This overestimate was consistent over the range of TBW, studied ($P > 0.05$ for all regression models at the various sampling times). The accuracy in predicting TBW from deuterium dilution was, on average, slightly better at 90 and 180 min than at 45 min, but this difference was not significant (repeated-measures ANOVA, $F_{2,30} = 2.130, P = 0.136$). Variance of the ratio of TBW$_d$ : TBW$_c$ was significantly different between sampling events (Levene’s test, $F_{2,49} = 4.473, P = 0.016$). The error in the deviation of TBW$_d$ from TBW$_c$ was twice as high at 45 min compared to 90 and 180 min after dose administration (Table 2), leading to a reduced precision of TBW predictions from isotope measurements at this early sampling stage. The following regression equations were derived to predict TBW$_c$ from TBW$_d$ at 90 min ($n = 20$) and 180 min ($n = 16$), respectively:

\[
\begin{align*}
\text{TBW}_c &= 96.034 + 0.852 \times \text{TBW}_d, \\
\text{TBW}_c &= 94.544 + 0.848 \times \text{TBW}_d.
\end{align*}
\]

When using deuterium dilution space as a predictor variable to estimate body composition, we employed TBW$_c$ values measured at 90 min.

**Estimates of Body Composition by Lethal and Nonlethal Methods**

TBW determined from either carcass analysis or isotope dilution was a very strong single predictor for FFM, explaining 98%–99% of the variation in FFM (Fig. 2). Table 3 compares the outcome of multiple regression analyses using TBW measured by deuterium isotope dilution and carcass analysis and further predictors related to different levels of invasiveness. The errors in the prediction of FM and FFM (calculated as deviations of predicted from observed values) following the Pace and Rathbun (1945) approach are also listed in Table 3. For the latter approach, we applied individual estimates of TBW from the established relationship between TBW$_c$ and TBW$_d$ (measured at 90 min equilibrium time) and a $\text{H}_2\text{O}_{\text{FMM}}$ of 63.2% (i.e., the average from our sample). We presented only predictive equations for FM because the absolute error of prediction was the same for FM and FFM, regardless of which component was taken as a response variable. This was due to the same significant predictors (models 1–3) for both FM and FFM and the fact that they add up exactly to BM. Thus, FFM was calculated as $\text{FFM} = \text{BM} - \text{FM}$. Also, we were mainly interested in the variation of FM (the relatively smaller of both components) and to what extent it can be accounted for by the various regression models.

Much of the variation in FM was unaccounted for by model 1 based on BM and external body measurements. Furthermore, relatively strong $r^2$ shrinkage and low invariance coefficients indicate lower replicability of the prediction when applied to different subsamples compared to the other models. FM was not significantly related to any of the potential predictors offered to model 1 separately. Only the combination of BM and a structural measurement (keel) revealed a significant relationship with FM. The combined variables dry BM (i.e., BM – TBW) and TBW determined from isotope dilution (model 2) or carcass analysis (model 3) both explained a large part of the variation in FM (92% and 97%, respectively). Moreover, the results from a cross-validation showed good replicability of the respective equations. When dry BM was replaced by BM in models 2 and 3, errors of prediction and $r^2$ were the same. However, such alternative models suffered notably from multicollinearity of the predictors (see VIF in Table 3), making them less robust. Abdominal fat fresh mass was highly correlated with total FM, and if it was the single predictor in the model, it accounted for 90% of the variation in FM. The variation increased slightly to 93% when head length was added to this model.
Table 2: Deuterium dilution space (TBWₙ) measured at different sampling intervals and in relation to total body water from carcass analysis (TBWₑ).

<table>
<thead>
<tr>
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<th>45 min</th>
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<th>90 min</th>
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<th>180 min</th>
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<tr>
<td></td>
<td>n = 16</td>
<td></td>
<td>n = 16</td>
<td></td>
<td>n = 20</td>
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<tr>
<td>TBW₈ₙ (g)</td>
<td></td>
<td></td>
<td>TBWₑ (g)</td>
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<tr>
<td></td>
<td>1,198 ± 185</td>
<td></td>
<td>1,179 ± 187</td>
<td></td>
<td>1,190 ± 169</td>
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<tr>
<td>TBWₙ : TBWₑ</td>
<td>1.092 ± 0.54</td>
<td></td>
<td>1.073 ± 0.27</td>
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<td>1.071 ± 0.26</td>
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<tr>
<td>r²TBWₑ-TBWₙ (SEE in g)</td>
<td>.90 (51)</td>
<td></td>
<td>.98 (22)</td>
<td></td>
<td>.98 (23)</td>
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</table>

Note. Data represent mean ± SD. Also given are coefficients of determination (r²) and the standard errors of the estimates (SEEs; i.e., the root mean square errors) from linear regressions of TBWₑ versus TBWₙ (P < 0.001 for all regressions).

* Includes only birds that were also measured at 45 and 180 min.

Discussion

We found a strong relationship between TBW measured by carcass desiccation and by deuterium isotope dilution in barnacle geese, which was consistent over a large range of body composition. With knowledge of this relationship and of the minimum equilibrium time needed for the isotope dose to mix completely with the body water pool, TBW can be very accurately predicted by deuterium isotope dilution as a nonlethal method. Furthermore, an accurate estimate of TBW was a strong predictor of FFM and, in combination with BM, FM. However, the accuracy of the estimates of FFM and FM for additional samples on the basis of TBW measured isotopically or by desiccation largely depends on the variation of the relative water content in the FFM.

Body Composition by Carcass Analysis

The dissected barnacle geese showed pronounced sexual differences in structural size and, related to this, differences in BM, FFM, and TBW. Higher fat loads in females may be expected in geese, in particular during the prebreeding phase, in anticipation of egg production and incubation (Raveling 1979).

FFM Hydration in Birds: How Variable Is It?

Whenever TBW, or an estimate thereof, is used to predict FFM, the variation of H₂O=label{_{FFM}} is of crucial relevance because it comprises the other major error source in addition to the error of the TBW estimate. Wang et al. (1999) reviewed this issue for adult mammals and concluded that species share a relatively constant H₂O=label{_{FFM}} in spite of differences in BM ranging by a factor of 10⁴. The relatively higher water content in the lean body component of the young growing organism until it reaches chemical maturity is well known (e.g., for mammals, Arnould et al. 1996; for birds, Bech and Østnes 1999), and thus we will restrict ourselves to a discussion of mature birds.

When we compare results of H₂O=label{_{FFM}} among studies, first of all, our attention has to be on possible differences in methodologies and definitions applied in these studies. Researchers often include water adsorbed to the feathers in the amount of TBW, which may amount to ca. 1%–2.5% of TBW (Crum et al. 1985; Mata et al. 2006). We intentionally did not do so because this “external water” is not in exchange with the body water pool estimated by isotope dilution (Crum et al. 1985). Further, FFM may be differently defined among studies, excluding, for instance, plumage (e.g., Boos et al. 2000; Mata et al. 2006) or bones and plumage (e.g., Groscolas et al. 1991) from the FFM component. Consequently, estimates of H₂O=label{_{FFM}} from those studies (71%–73%) are necessarily considerably higher than the value reported here.

In Table 4, we compiled data on H₂O=label{_{FFM}} reported or calculated from studies of waterfowl carcass analyses using comparable methods and definitions as applied in this study. H₂O=label{_{FFM}} can be reasonably stable over different seasonal and/or physiological stages within a given study. As in our study, no sex-related differences are indicated. Variation between studies can, however, be considerable.

Compared to those of other studies, our values (mean = 63.2%) are at the lower range of H₂O=label{_{FFM}} values reported so far. Although the geese had access to water during the captive period until 4–6 h before termination of the experiment, due to...
general capture stress they may have not made sufficient use of it and may have experienced a certain degree of dehydration. Birds can tolerate notable water losses under restrained conditions. For instance, Davidson (1984) noted a decrease of H$_2$O$_{FM}$ of 0.8%–1% per hour during the first 4 h after capture in knots (Calidris canutus) and dunlins (Calidris alpina), that is, from 66.7% to 63.5% and from 65.8% to 61.8%, respectively. Interestingly, after the first 4 h, dunlins appeared fully to compensate further water loss by metabolically produced water, whereas in knots, dehydration continued (at a lower rate) until 60.9% at 24 h after capture.

We conclude that both methodological factors and tolerance for a (temporarily) negative water balance in birds may contribute to the considerable variation in H$_2$O$_{FM}$, reported among studies. Nevertheless, homeostasis is crucial for organismal functioning. Therefore, under unrestrained circumstances, birds will probably strive to avoid dehydration and keep their water balance within small margins. Currently, it is not possible to rule out whether and to what extent variation in H$_2$O$_{FM}$ reflects truly species-specific diversity or varying sampling procedures and other methodological differences. Future applications of the isotope dilution method to estimate body composition would benefit from a better understanding of general patterns of variation in H$_2$O$_{FM}$ and how it may be related to species and/or physiological state.

Table 3: Predictive equations for total fat mass (FM; g) based on nonlethal and lethal measurements in 21 barnacle geese

<table>
<thead>
<tr>
<th>Multiple regression models</th>
<th>r$^2$</th>
<th>SEE</th>
<th>Absolute Error FM and FFM (g)</th>
<th>Error FFM (%)</th>
<th>Error FM (%)</th>
<th>VIF</th>
<th>Shrinkage $r_{11} - r_{12}$</th>
<th>$r_{21} - r_{22}$</th>
<th>$r_{31} - r_{32}$</th>
<th>$r_{41} - r_{42}$</th>
<th>$r_{51} - r_{52}$</th>
<th>$r_{61} - r_{62}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonlethal:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. FM = 368.7 + 0.329 ×</td>
<td></td>
<td></td>
<td>1.2 ± 3.4</td>
<td>13.7 ± 24.9</td>
<td>1.42</td>
<td>.19</td>
<td>.11</td>
<td>.84</td>
<td>.74</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BM – 6.997 × keel</td>
<td>.418</td>
<td>89.1</td>
<td>69.9 ± 44.8</td>
<td>4.0 ± 2.8</td>
<td>37.3</td>
<td>.12</td>
<td>.01</td>
<td>.99</td>
<td>.99</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. FM = –200.6 – 2.66 ×</td>
<td></td>
<td></td>
<td>1.2 ± 1.4</td>
<td>10.1 ± 10.1</td>
<td>1.15</td>
<td>.02</td>
<td>.01</td>
<td>.99</td>
<td>.99</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TBW$_a$ + 0.936 × dry BM</td>
<td>.918</td>
<td>33.7</td>
<td>21.3 ± 23.3</td>
<td>1.2 ± 1.4</td>
<td>10.1</td>
<td>.10</td>
<td>.01</td>
<td>.99</td>
<td>.99</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lethal:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. FM = –57.48 – 0.418 ×</td>
<td></td>
<td></td>
<td>1.2 ± 1.3</td>
<td>10.6 ± 11.4</td>
<td>1.18</td>
<td>.14</td>
<td>.12</td>
<td>.93</td>
<td>.94</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TBW$_a$ + .856 × dry BM</td>
<td>.966</td>
<td>21.6</td>
<td>16.0 ± 12.3</td>
<td>0.9 ± 0.8</td>
<td>10.6</td>
<td>.10</td>
<td>.01</td>
<td>.99</td>
<td>.99</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. FM = –382.7 + 4.492 ×</td>
<td></td>
<td></td>
<td>1.2 ± 1.5</td>
<td>16.0 ± 18.4</td>
<td>1.01</td>
<td>.01</td>
<td>.01</td>
<td>1.00</td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$F_{茴}$ + 5.223 × head</td>
<td>.931</td>
<td>30.1</td>
<td>24.9 ± 13.8</td>
<td>1.4 ± 0.8</td>
<td>16.0</td>
<td>.10</td>
<td>.01</td>
<td>.99</td>
<td>.99</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note. TBW$_a$ = deuterium dilution space; TBW$_c$ = total body water from the carcass analyses. n = 20 if TBW$_c$ is used in the equation. Modeling started with noninvasively measurable predictors in model 1: tarsus, head, wing, keel, body mass (BM), and sex. This set of potential predictors was expanded by TBW$_a$ and dry BM (i.e., BM – TBW$_a$) in model 2 and by TBW$_a$ and dry BM (i.e., BM – TBW$_c$) in model 3. For model 4, the same predictors were applied as for model 1, with the addition of abdominal fat fresh mass ($F_{茴}$). Shown are the final models containing only significant predictor variables derived by stepwise backward multiple regressions. For all models, *P* < 0.001 (except for model 1, *P* = 0.008). Summary statistics include coefficients of determination ($r^2$), standard error of the estimate (SEE), absolute and relative error of predicted FM and fat-free mass (FFM) calculated as deviations of predicted from observed values (mean ± SD), variance inflation factor of predictors (VIF), and statistics from a cross-validation procedure (see "Methods" for details). FFM was calculated as BM – FM (see text). Outcomes of the Pace and Rathbun (1945) approach are given in the last row (n = 20).

1. When dry BM is replaced by BM in the model, VIF = 4.51.
2. When dry BM is replaced by BM in the model, VIF = 3.68.

**TBW Measured by Deuterium Dilution and the Effect of Equilibration Time**

In this study, TBW$_a$ overestimated TBW$_c$ by 7.1% (at 90 min equilibrium time) consistently over a large range of TBW$_c$ and can thus be corrected according to the values given here. Speakman et al. (2001; Table 3) calculated an average of 4.7% by which actual TBW was overestimated by hydrogen isotope dilution in birds, based on nine studies on four species. Additional studies using hydrogen isotopes and the plateau approach reported 8.1% for the chicken Gallus gallus (Mata et al. 2006), 8.4% for glaucous-winged gull Larus glaucescens nestlings (Hughes et al. 1987), and 3.3% for the glaucous gull Larus hyperboreus (Shaffer et al. 2006). However, at least part of the variation among studies is due to methodological factors. For instance, the study by Degen et al. (1981), which was included in the average calculated by Speakman et al. (2001), as well as those by Hughes et al. (1987) and Shaffer et al. (2006) included water adsorbed to feathers in TBW$_c$. This plumage water cannot be accounted for by isotope dilution. As a consequence, the difference between TBW$_a$ and TBW$_c$ becomes smaller. Thus, an average value by which carcass TBW (excluding plumage water) is overestimated by the hydrogen dilution space in birds certainly exceeds the 4.7% given by Speakman et al. (2001).

Our results indicate that mixing of marker solution and body water was not completed after 45 min. Average levels and var-
Table 4: Data on the relative water content in the fat-free mass (H$_2$OFFM) in adult waterfowl

<table>
<thead>
<tr>
<th>Species and Period of Collection</th>
<th>Females (H$_2$OFFM)</th>
<th>Males (H$_2$OFFM)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Barnacle goose (Branta leucopsis)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Winter–spring</td>
<td>62.8 (.2)</td>
<td>63.5 (.4)</td>
<td>This study</td>
</tr>
<tr>
<td>Black duck (Anas rubripes)</td>
<td></td>
<td></td>
<td>Reinecke et al. 1982</td>
</tr>
<tr>
<td>Prelay</td>
<td>64.5</td>
<td>64.3</td>
<td></td>
</tr>
<tr>
<td>Laying</td>
<td>65.0</td>
<td>63.8</td>
<td></td>
</tr>
<tr>
<td>Postlay</td>
<td>63.2</td>
<td>63.8</td>
<td></td>
</tr>
<tr>
<td>Moult</td>
<td>63.2</td>
<td>63.2</td>
<td></td>
</tr>
<tr>
<td>Autumn</td>
<td>63.2</td>
<td>63.2</td>
<td></td>
</tr>
<tr>
<td>Winter</td>
<td>63.2</td>
<td>63.2</td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>63.5</td>
<td>63.5</td>
<td></td>
</tr>
<tr>
<td>Brent goose (Branta bernicla)</td>
<td></td>
<td></td>
<td>Korte 1988</td>
</tr>
<tr>
<td>Winter–spring</td>
<td>69.0 (.6)</td>
<td>68.4 (.5)</td>
<td></td>
</tr>
<tr>
<td>Canada goose (Branta canadensis)</td>
<td></td>
<td></td>
<td>Raveling 1979</td>
</tr>
<tr>
<td>Autumn migration</td>
<td>65.2</td>
<td>65.8</td>
<td></td>
</tr>
<tr>
<td>Midwinter</td>
<td>65.7</td>
<td>65.4</td>
<td></td>
</tr>
<tr>
<td>Spring migration</td>
<td>65.4</td>
<td>66.5</td>
<td></td>
</tr>
<tr>
<td>Prelay</td>
<td>65.4</td>
<td>65.4</td>
<td></td>
</tr>
<tr>
<td>Midincubation</td>
<td>66.1</td>
<td>65.2</td>
<td></td>
</tr>
<tr>
<td>Hatch day</td>
<td>68.7</td>
<td>68.1</td>
<td></td>
</tr>
<tr>
<td>Midmoult</td>
<td>67.5</td>
<td>65.3</td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>66.6</td>
<td>65.3</td>
<td></td>
</tr>
<tr>
<td>Common eider (Somateria mollissima)</td>
<td></td>
<td></td>
<td>Parker and Holm 1990</td>
</tr>
<tr>
<td>2–3 wk prelay</td>
<td>63.6</td>
<td>63.5</td>
<td></td>
</tr>
<tr>
<td>Prelay breeders</td>
<td>64.4</td>
<td>62.9</td>
<td></td>
</tr>
<tr>
<td>Prelay nonbreeders</td>
<td>66.5</td>
<td>65.8</td>
<td></td>
</tr>
<tr>
<td>Postlay</td>
<td>63.9</td>
<td>65.4</td>
<td></td>
</tr>
<tr>
<td>Hatch day</td>
<td>64.6</td>
<td>65.2</td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>64.6</td>
<td>65.3</td>
<td></td>
</tr>
<tr>
<td>Lesser scaup (Aythya affinis)</td>
<td></td>
<td></td>
<td>Austin and Fredrickson 1987</td>
</tr>
<tr>
<td>Premoult</td>
<td>73.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Moult</td>
<td>70.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Postmoult</td>
<td>71.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Migratory</td>
<td>70.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>71.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lesser snow goose (Anser c. caerulescens)</td>
<td></td>
<td></td>
<td>Campbell and Leatherland 1980</td>
</tr>
<tr>
<td>Through the year</td>
<td>69.3 (.06)$^a$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean of all studies</td>
<td>66.6 (1.2)$^a$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note. In most cases, H$_2$OFFM was calculated from mean values of body mass, fat mass, and total body water (TBW; except from individual values in Korte 1988; Campbell and Leatherland [1980] gave mean values directly). We used only samples (means) where the same birds were analyzed for all of those body components. n = sample size. Standard errors are shown in parentheses.

$^a$ TBW component included water adsorbed to feathers. Reference was unclear about the actual sample size related to the reported mean and standard error of 0.693 ± 0.06. Translated into percentage, this standard error would equal a questionable 0.0006%. We suspect that 0.06 is the correct value.

$^b$ Sexes were pooled.

$^c$ Mean of studies’ means; for studies where data for both sexes were available, the average was used.

Estimates of Body Composition by Lethal and Nonlethal Methods

Regression models to predict body composition based on BM in combination with other external morphological parameters performed poorly in this study of the barnacle goose (Table 3;
FFM includes virtually all structural mass and, at the same time, nearly all water, a measure of TBW better accounts for differences due to structural mass than do most morphological measurements. Indeed, variation in TBW determined from carcass analysis or isotope dilution both explained a large part of the variation in FFM and, consequently, FM.

Abdominal fat fresh mass was highly correlated with total FM and, if the carcass is available, allows simple means of fat prediction without dissecting the entire specimen. Model 1 represented the least invasive of the methods tested in Table 3. However, the error of prediction was highest, equating to 37% and 4.0% of actual FM and FFM, respectively. Using estimates of TBW from deuterium isotope dilution and either a regression or the Pace and Rathbun (1945) approach as alternative non-destructive methods reduced the relative error of the FM and FFM estimate to 10%–13% and 1.2%–1.3%, respectively.

Differences in the accuracy of a regression and the Pace and Rathbun (1945) approach depend on how much individual animals in the study sample deviate from the assumption of a constant FFM hydration. Variation in $H_2O_{FFM}$ was low in our sample (Table 1), resulting in a similar accuracy of both approaches (Table 3; Fig. 3). Irrespective of which approach is applied, prediction accuracy for a new sample will be reduced if $H_2O_{FFM}$ differs between the calibration sample and a new sample. A direct measurement of $H_2O_{FFM}$ in a subsample of the study population is always advisable to ensure confidence about resulting predictions. If this is not possible, we suggest applying an average value of 66.6% for studies on adult waterfowl, as calculated from the species-specific studies listed in Table 4, when using the Pace and Rathbun (1945) approach.

**Acknowledgments**

We want to thank Astrid Tijdens and René Adelerhof for their assistance in experimental work and carcass analysis. Bertke Verstappen-Dumoulin determined the $^2$H enrichments, and Harro Meijer provided a description of the isotope analysis procedure. Marcel Klaassen, Sarah Jamieson, Joost Tinbergen, and Rudi Drent are thanked for their valuable comments on earlier versions of the manuscript. G.E. was supported by scholarships from Marianne und Dr. Fritz Walter-Fischer Stiftung, Germany, and the Ubbo Emmius Programme at the University of Groningen.

**Literature Cited**


Bech C. and J.E. Østnes. 1999. Influence of body composition...


