Testing angular velocity as a new metric for metabolic demands of slow-moving marine fauna: a case study with Giant spider conchs *Lambis truncata*

Lloyd W. Hopkins1*, Nathan R. Geraldi2, Edward C. Pope1, Mark D. Holton3, Miguel Lurgi1, Carlos M. Duarte2 and Rory P. Wilson1

Abstract

**Background:** Quantifying metabolic rate in free-living animals is invaluable in understanding the costs of behaviour and movement for individuals and communities. Dynamic body acceleration (DBA) metrics, such as vectoral DBA (VeDBA), are commonly used as proxies for the energy expenditure of movement but are of limited applicability for slow-moving species. It has recently been suggested that metrics based on angular velocity might be better suited to characterise their energetics. We investigated whether a novel metric—the ‘Rate of change of Rotational Movement (RocRM)’, calculated from the vectoral sum of change in the pitch, roll and yaw/heading axes over a given length of time, is a suitable proxy for energy expenditure.

**Results:** We found that RocRM can be used as an alternative energy expenditure proxy in a slow-moving benthic invertebrate. Eleven Giant spider conchs *Lambis truncata* (collected in the Red Sea) were instrumented with multiple channel (Daily Diary) tags and kept in sealed chambers for 5 h while their oxygen consumption, $\dot{V}O_2$, was measured. We found RocRM to be positively correlated with $\dot{V}O_2$, this relationship being affected by the time-step (i.e. the range of the calculated differential) of the RocRM. Time steps of 1, 5, 10 and 60 s yielded an explained variability of between 15 and 31%. The relationship between $\dot{V}O_2$ and VeDBA was not statistically significant, suggesting RocRM to provide more accurate estimations of metabolic rates in *L. truncata*.

**Conclusions:** RocRM proved to be a statistically significant predictor of $\dot{V}O_2$ where VeDBA did not, validating the approach of using angular-based metrics over dynamic movement-based ones for slower moving animals. Further work is required to validate the use of RocRM for other species, particularly in animals with minimally dynamic movement, to better understand energetic costs of whole ecosystems. Unexplained variability in the models might be a consequence of the methodology used, but also likely a result of conch activity that does not manifest in movement of the shell. Additionally, density plots of mean RocRM at each time-step suggest differences in movement scales, which may collectively be useful as a species fingerprint of movement going forward.

**Keywords:** Angular velocity, Dynamic body acceleration (DBA), Energy expenditure, Movement costs, RocRM, Rotational movement

*Correspondence: lloyd.william.hopkins@gmail.com
1 Department of Biosciences, Singleton Campus, Swansea University, Wallace Building, Swansea SA2 8PP, UK
Full list of author information is available at the end of the article
**Background**

Animal movements, behaviours and life processes account for much of an organism’s energy demands, with the process-linked metabolism of an individual determining fitness, which in turn can affect ecosystem structure and functioning [1–5]. Accurate determination of energy expenditure in animal species is therefore pivotal to understanding the value and costs of behaviours and how these relate to the ecology of species within ecosystems [6–8].

However, directly measuring the rate at which an animal expends energy in the field, where it is unrestrained and allowed to exhibit a full suite of behaviours, is challenging [9, 10]. Doubly labelled water (DLW)—measuring the rate of CO$_2$ production, VCO$_2$ [10–12]—is the only direct measure, while other methods, such as recording heart rate, use proxies for energy expended [10, 13, 14]. All such methods have limitations. For example, heart rate monitors are generally intrusive, often requiring implantation, and are less suitable for smaller animals [15]. The doubly labelled water method, on the other hand, is logistically difficult to conduct and has limited temporal resolution [16]. A fairly recent approach has used tri-axial accelerometers in externally attached tags to provide a less invasive and high-resolution alternative deriving ‘dynamic body acceleration’ (DBA) as a proxy for movement-based energy expenditure [17, 18]. DBA is calculated from the dynamic acceleration (i.e. the acceleration signal that remains following the subtraction of static acceleration, or the gravitational component [19]) summed over all three dimension axes (x, y, z; surge, heave, sway) [20, 21].

DBA has been repeatedly shown to correlate strongly with the rate of oxygen consumption (MO$_2$ and VO$_2$) in a diverse range of taxa including, inter alia, birds [20], cephalopods [22], bivalves [23], fish [24, 25] and mammals [26]. These laboratory studies have produced statistical relationships between DBA and VO$_2$ that can be used to calculate energy expenditure from DBA extracted from accelerometer tags used on wild animals.

Despite the strength of this relationship and its applicability across many taxa, DBA is less likely to be a useful proxy for energy expended for species that move slowly because of their inherently weak dynamic acceleration signals. This limitation is indeed significant, because many marine benthic invertebrates, such as gastropods and crustaceans move slowly [27–29]. Indeed, where dynamism occurs, activity and energy expenditure within these periods may be measured by DBA metrics [28], but these may be a small percentage of the overall time budget for these animals. Lyons et al. [30], for example, showed that DBA correlated with oxygen consumption in the American lobster (*Homarus americanus*) during the more active periods of movement, but had limited capacity to resolve metabolic rates in periods when movement was slower.

Recently, a comprehensive review of DBA highlighted the need for alternative tag-derived metrics to address the problem posed by slow-moving species [27]. In particular, the authors suggested that rates of change of rotational axes (pitch, roll and yaw) might offer better insights into the degree of movement—and by proxy energy expenditure [17]—for such species compared to DBA (cf. [31]). In other words, the movement of, for example, a benthic invertebrate, where dynamic motion is consistently negligible and often only one locomotion type is used, may be manifest most notably in the speed at which it changes its whole-body orientation. Movement along these rotational axes (assuming no drift or external factors are acting upon the animal), regardless of the speed at which this happens, still requires the exertion of force and, therefore, the expenditure of energy. Indeed, Wilson et al. [27] presented preliminary data showing clearer changes in body angular velocity during movement bouts compared to DBA in the Giant spider conch *Lambis truncata*. Variability in the body roll angle has since been shown to describe slow-moving (walking) activity patterns of the European spiny lobster (*Palinurus elephas*) much better than DBA metrics, due to very smooth acceleration signals [29].

We investigate the validity of a new metric, derived from the rates of change in rotational movement (which we term the ‘Rate of Change of Rotational Movement’, or ‘RocRM’), as an alternative to DBA for estimating energy expenditure in a slow-moving benthic invertebrate, the Giant spider conch, *L. truncata*. *Lambis truncata* is a gastropod of the *Strombidae* family, a family that is commercially important and heavily overexploited in many countries [32–34]. Like many other conch species, it moves mostly via ‘jumps’ and rotational (about the yaw axis) ‘drifts’ [28, 35, 36]. In other species of conch, jumps have been shown to accompany increased oxygen consumption [37]. These jumps do produce a notable acceleration signal and the metric of mean DBA is able to differentiate between leaps and drifts in a movement recognition model [28]. Whether DBA is an accurate proxy for oxygen consumption in conchs, however, is unknown. For example, ODBA during a drift is typically much lower than during a leap [28], but might actually demand a considerable energetic commitment in practice [31, 38]. With enough drift movements, therefore, it is possible that a large proportion of energetic expenditure would simply be missed by DBA metrics. An alternative metric that can capture the energetic expenditure involved in either mode of movement would therefore be preferable. The major focus of this paper is accurately capturing
these drift movements. We used laboratory-based static respirometry to test the hypothesis that RocRM correlates with the rate of oxygen consumption in a slow-moving animal. We further examined how RocRM compared to DBA as a predictor of $V\dot{O}_2$ in this species. We hypothesised that RocRM would correlate more strongly with $V\dot{O}_2$ than DBA.

**Methods**

**Animal collection and holding facilities**
Eleven Giant spider conchs, *Lambis truncata sebae* (Kiener 1843), were collected by snorkellers from areas of rubble and coarse sand on a shallow (< 1 m) reef bed near the KAUST campus, Saudi Arabia, in the Red Sea during February 2019. Similarly sized conchs were selected (maximum width and length of shells, measured to the nearest cm, are shown in Table 1). A GPS fix (using a handheld GPS 73 unit; Garmin, Schaffhausen, Switzerland) was taken of what was deemed to be the most representative of the area where the specimens were collected from (22°17’22.7” N, 39°03’25.5” E), with all individuals being found within a 50 m radius.

Specimens were housed in two separate holding tanks (dimensions 118 × 56 × 46 cm L × W × H) supplied with a continuous flow of seawater, pumped directly from the neighbouring Red Sea and then micro-filtered. The water level in the holding tanks was 40 cm (ca. 264 L). Tanks of the same size were used during experiments.

Animals were kept in their holding tanks for 3 days prior to experiments and fed throughout the trial on the algal film that was allowed to coat the tank walls. The shells of the conchs were cleaned with a rigid brush to remove as much biological growth as possible and further cleaning was carried out when deemed necessary throughout the experimental period to minimise the effect of non-conch oxygen consumption during respirometry. However, conchs were not cleaned within 24 h of being experimentally tested, to reduce the potential stress of handling.

**Respirometry experiments**

**Overview**
The tanks used in the respirometry experiments (see above) were chosen to encourage ‘normal’ conch movement during the experiments; the tanks were large enough to allow short periods of traversal. Before each trial, the water flow into the tank was stopped and the tanks drained to a water level of 20 cm. This water height was more than sufficient to allow ‘jumping’ movement without the conch touching the plastic top covering. Flow was restored between experiments and left long enough to completely replace the water. Air was bubbled through the water prior to the start of the experiment to fully oxygenate the water. A water pump was used to circulate the water within the tank to reduce both microbial build-up on the sensor foil and oxygen stratification inside the respirometer (tank), providing more consistent estimates of both background respiration and the $V\dot{O}_2$ of the study animal [39]. MiniDOT $O_2$ sensor loggers (PME, California, USA) were anchored to the bottom of the tank at the end opposite to the pump (Fig. 1), with the sensing face pointing towards the tank centre.

Respirometry trials were conducted using closed chamber/static respirometer techniques [40]. For the control/blank runs, initial trials (i.e. when calibrating equipment) and in the experiments themselves, aquaria were sealed with plastic sheeting (similar to those described in [25] and [41]) so as to be gas tight. Four layers of 1-mm-thick

| Conch | Max. shell width (cm) | Max. shell length (cm) | Calculated approximate volume (cm$^3$) |
|-------|-----------------------|------------------------|---------------------------------------|
| 1     | 11                    | 17                     | 1047                                  |
| 2     | 12                    | 19                     | 1277                                  |
| 3     | 11                    | 14                     | 739                                   |
| 4     | 10                    | 14                     | 1008                                  |
| 5     | 11                    | 15                     | 1188                                  |
| 6     | 9                     | 14                     | 806                                   |
| 7     | 12                    | 16                     | 1382                                  |
| 8     | 11                    | 16                     | 1126                                  |
| 9     | 11                    | 16                     | 1126                                  |
| 10    | 12                    | 17                     | 1469                                  |
| 11    | 13                    | 17                     | 1591                                  |
| Mean  | 11.2 (± 1.1 sd)       | 15.9 (± 1.6 sd)        | 1159.9 (± 261.9)                      |

Measurements, to the nearest whole cm, of maximum length and width of conchs used in this study and their approximate volumes
Sheeting were bonded together with duct tape (taking care that the complete material was watertight) to ensure that the final cover had enough weight to rest against the water and push out trapped air. The sheet was sealed to the tank walls as close as possible to the water level using duct tape and the seal inspected for any obvious gaps or unwanted gas exchange. Air bubbles between the water and plastic were pushed out by smoothing the plastic towards the edges until none remained. Care was taken to seal around the water pump cable in such a way as to also be airtight. The salinity of the water was measured following each experiment and control using a Pro Plus Quatro salinity probe (YSI, Ohio, USA).

**Data loggers**

Water-dissolved oxygen concentration (mg/L DO) and water temperature were measured with the miniDOT loggers, with temperature corrections applied automatically by the sensor. Salinity corrections of DO required manual input of salinity using provided ‘miniDOT concatenate’ software. MiniDOT loggers took one reading of each variable every 30 s—the minimum time that O₂ was predicted to take to diffuse through the sensing foil (PME, pers. comm.) and therefore the shortest interval that could be confidently used. The sensing foil was cleaned between each trial.

Conch movement was recorded using an ‘elongated’ model Daily Diary tag (Wildbyte Technologies Ltd., UK), recording tri-axial acceleration (surge, heave and sway) and tri-axial magnetometry at 20 Hz. Each Daily Diary was powered by a 750 mAh, 3.6 V single-use battery (EVE Energy Co., China). The complete package was enclosed within a vacuform polystyrene plastic housing and sealed using Poly Cement (Humbrol, Hornby Hobbies, UK). The overall maximum package dimensions were 38 × 20 × 13 mm L × W × H with a weight in air of 6 g. Velcro (Velcro BVBA, UK) was glued to the Daily Diary package. All conchs had a 1.5 × 1.5 cm patch of complementary Velcro fixed to their shell using super glue (Loctite Power Flex, Loctite, Germany), at an approximately 25°–30° angle on the anterior shell spine (as it was an area conserved between individuals, close to the centre of mass, and relatively flat for ease of attachment). The Daily Diary package was attached to the animal during transfer of the conch from holding to experiment tanks and removed when conchs were returned to holding tanks.

**Respirometry experiments**

The Daily Diary-equipped conchs were allowed to settle within their testing tanks for at least 30 min prior to experimentation, with the miniDOT loggers placed in the tank at the same time. Tanks were sealed for 5 h and the DO and conch movement (via the Daily Diaries) recorded throughout this time. To increase the number of tests, conchs were tested four times across two general time periods (twice each): (1) ‘afternoon’ (average start and end times = 11:04 and 15:59, respectively—Saudi Arabia local time = GMT + 3) and ‘evening’ (average start and end times = 17:36 and 22:30, respectively). These times were used as preliminary work had shown *L. truncata* collected from this area appear to slowly increase their movement through the late afternoon and evening (unpublished data), therefore giving the best chance of capturing different ‘levels’ of movement frequency. Conchs were not reused in an experiment for at least 24 h.

All conchs were released following respirometry experiments and returned as close to their collection location as possible. The Velcro patch and glue were easily removed, leaving only superficial markings.

**Calibrations and controls**

A control/blank tank, without conchs, was included for both time periods mentioned (3 in the afternoon and 4 in the evening), lasting at least 5 h each. Blank runs were used to assess microbial (‘background’) VO₂. The average conch respirometry experiment start and end times informed the times used in calculations of corrections. Two mean control slopes, for afternoon and evening periods, were calculated from linear regression of each control run. Each conch VO₂ calculation was subsequently corrected for microbial VO₂ through subtraction of control rates (see [39]). The lowest recorded DO concentration in any experiment was 5.75 mg/L (87.54% O₂ saturation).

**Data analysis**

**VO₂**

The strength of the relationship of VO₂ as a function of time (as a linear model) was evaluated immediately after each experiment run, to check that the system was sealed sufficiently and that there was a generally consistent negative correlation. Origin 2019 (Origin Lab Corp., Massachusetts, USA) was used to compute $R^2$ values for this. Experiment runs were repeated if plots suggested erratic DO readings (potentially due to poor chamber sealing) or a low $R^2$ value comparing oxygen concentration and time (in the final analysis, all slopes had an $R^2$ of > 0.87). As a result of the corrections discussed in 2.2.3, three experiments were discounted due to background-corrected slopes turning positive (suggesting breaches in the tank sealing), all of which were in the afternoon period, leaving 41 slopes across 11 conchs (3 conchs with 3 slopes, all others with 4 slopes) to
be used in statistical analysis. Of these, five experiments did not record for the complete 5 hours (240, 267, 271, 286, 295 min, respectively), but were included in all analyses.

\( \dot{V}O_2 \) for a single organism in a static system was calculated as in [40]:

\[
\dot{V}O_2 = \Delta[O_2] \cdot V_{ind},
\]

where \( \Delta[O_2] \) is the regression slope of \( O_2 \) in milligrams per litre per hour, \( V_{ind} \) is the water volume in litres specific to the tested individual and \( V_{O_2} \) is the rate of \( O_2 \) consumption for that individual in the same time units as the slope (mg ind\(^{-1}\) h\(^{-1}\)). \( V_{ind} \) was calculated as the effective tank volume (1,321,600 cm\(^3\)) minus the volume of the miniDOT logger (314.16 cm\(^3\)), the water units as the slope (mg ind\(^{-1}\) h\(^{-1}\)).

**Movement**

Daily Diary data were preliminarily analysed and data extracted using DDMT (Daily Diary Multiple Trace) software (Wildbyte Technologies Ltd, UK). Acceleration (1 g = 9.81 m/s\(^2\)) was separated into dynamic and static (postural) components using a smoothing window of 3 s [the window length informed by Shepard et al. [19] and previously collected data on the movement durations of this species (unpublished data)].

Daily Diary data were used to calculate two movement metrics for each conch experiment run: RocRM and VeDBA.

The RocRM metric was calculated from the vectoral sum of the change (differential) in pitch, roll and heading over a specified time-step given by:

\[
RocRM = \sqrt{Roc_p^2 + Roc_r^2 + Roc_h^2},
\]

where \( Roc_p \), \( Roc_r \) and \( Roc_h \) refer to the rate of change in the angles of pitch, roll and heading, over the given time-step, respectively. The rate of change over a time-step is calculated within a moving window, therefore RocRM is presented as a continuous variable. Pitch and roll angle were derived from postural acceleration data [42] while heading is derived from the body orientation data in tandem with the geomagnetic field strength measured in three axes [43]. Specifically, pitch and roll were calculated as:

\[
Pitch(p) = \text{atan2}(S_x, \sqrt{S_y^2 + S_z^2}) \cdot \frac{180}{\pi},
\]

\[
Roll (r) = \text{atan2}(S_y, \sqrt{S_x^2 + S_z^2}) \cdot \frac{180}{\pi},
\]

where \( S_x \), \( S_y \) and \( S_z \) are the static acceleration as recorded on the heave, surge and sway axes, respectively.

Heading was derived as:

\[
\text{Heading} (h) = \text{mod}\left\{\frac{\text{atan2}(-m_y, m_x) \cdot \frac{180}{\pi}}{360}\right\},
\]

where \( m_x \) and \( m_y \) refer to the x and y magnetometer channels following normalisation (to bring the data onto a sphere) and rotational correction (using pitch and roll to account for device tilt) and mod refers to the modulo operator [44].

There was some magnetic noise within the tanks due to the pump, so heading was smoothed over 5 s. For consistency, pitch and roll were also smoothed over 5 s. RocRM was subsequently calculated for four different time-steps (i.e. the vectoral sum of the change in these smoothed channels); over 1 s (RocRM\(_1\)), 5 s (RocRM\(_5\)), 10 s (RocRM\(_{10}\)) and 60 s (RocRM\(_{60}\)).

The dynamic component of acceleration, used in calculating VeDBA, was calculated as:

\[
\text{Dynamic acceleration} = \text{total acceleration} - \text{static acceleration}.
\]

VeDBA (measured in g) was calculated using the formulation proposed by Qasem et al. [21]:

\[
\text{VeDBA} = \sqrt{A_x^2 + A_y^2 + A_z^2},
\]

where \( A_x \), \( A_y \) and \( A_z \) are the dynamic accelerations from each of the tri-axial channels at any given time. To be consistent with pitch, roll and heading, VeDBA was also subsequently smoothed over 5 s prior to statistical analysis.

Overall means for VeDBA and each RocRM measure were calculated for each experiment run and matched to their corresponding \( V_{O_2} \) measurement.

Kernel density distributions of the four different RocRM time-steps were calculated to provide an overview of the differences between timescales of conch movement.

**Statistical analyses**

Statistical analyses were conducted in R [45] version 3.6.0. Five linear mixed models (LMMs) were constructed to investigate the relationship between \( V_{O_2} \) and each one of the movement metrics independently, using the lme4 package [46] version 1.1.21. Thus, in each model, \( V_{O_2} \) was the response variable and log(VeDBA), log(mean RocRM\(_1\)), log(mean RocRM\(_5\)), log(mean RocRM\(_{10}\)) and log(mean RocRM\(_{60}\)) were included as predictor variables in their respective models (subscript refers to the differential time-step period in seconds). To account for
potential dependency of observations obtained from the same individual, ‘conch individual’ was included as a random-effect term in all models. All models included random intercepts only (i.e. only the by-individual intercept, and not the slope, were allowed to vary), as attempts to fit random slopes for the conch ID term had issues with singularity. Mean experiment temperature was initially included as a fixed effect in each model, but was removed from all of them through stepwise elimination of terms through ANOVA comparisons of models (see Additional file 1), as well as judged on Akaike Information Criterion (AIC) values, starting with the full model [47]. Likelihood ratio tests (R function ANOVA, with Maximum Likelihood) were used to determine the validity, effect size and significance of full versus null models. Graphical procedures (Q–Q plot, histograms, and residual vs fitted value plots) were used to visually assess the fit of the model and adherence to assumptions of normality and homoscedasticity of residuals Zuur et al. [48]. RocRM was log-transformed to better meet these assumptions. VeDBA was also log-transformed, to make comparisons between models fairer and clearer, although in practice the VeDBA models were near identical. Pseudo-\( R^2 \) values in the form of marginal (\( R^2_m \)) and conditional (\( R^2_c \)) values—where the conditional value considers the full model and the marginal value considers the fixed model effects alone relative to the conditional value—were calculated according to Nakagawa and Schielzeth [49]. 95% confidence intervals were computed via a bootstrapping method using the ‘confint’ (type = ‘percentile’, \( n = 500 \)) function within lme4 [46]. The 95% prediction interval was also calculated (using the calculator provided in [50]) for each reported \( p \) value, allowing comment on the \( p \) value uncertainty and replicability [51].

Where \( p \leq 0.37 \), \( p \) values were additionally converted into Bayes factor upper bounds (\( \overline{B} \)) values, giving the odds of the alternative hypothesis being true over the null hypothesis [51, 52]. \( \overline{B} \) values were calculated following the equation in [51]:

\[
\overline{B} \leq \frac{-1}{e \times p \times \ln(p)}, \tag{8}
\]

where ‘\( p \)’ is the model \( p \) value.

AIC evidence ratios, comparing the relative likelihood of one model over another [51], were calculated according to:

\[
ER = \frac{1}{e^{-0.5 \times \Delta AIC}}, \tag{9}
\]

where \( \Delta AIC \) is the difference in AIC between the two models of the interest. Note that the equation denominator calculates the relative evidence for a given model.

**Results**

**Controls**

The mean background \( V02 \) slopes of the ‘afternoon’ and ‘evening’ controls were \(-2.98\) and \(-4.03\) mg h\(^{-1}\), respectively. Despite the use of micro-filtered water in these experiments, there remained a clear presence of background/microbial respiration, particularly in the evening period, requiring the use of these corrections.

**Differences in movement traces between VeDBA and RocRM**

Movement data traces of rotational movement axes, VeDBA and RocRM reveal appreciable differences in the RocRM signal with different movement types (Fig. 2). RocRM10 movement peaks differed between movements predominantly in the pitch and roll axes, exhibiting split peaks, and those predominantly in the heading/yaw axis, exhibiting peaks that are more singular. VeDBA, in contrast, produced relatively consistent peaks.

All RocRM time-steps (1, 5, 10 or 60 s) had a positive skew of mean values (Fig. 3), with increasing values of RocRM with increasing time-step as well as increasing interquartile range (IQR) relative to the median value (RocRM1: median = 0.13, IQR = 0.07; RocRM5: median = 0.37, IQR = 0.26; RocRM10: median = 0.49, IQR = 0.43, RocRM60: median = 1.43, IQR = 1.76).

**RocRM as a proxy for oxygen consumption**

Mean temperature was removed from all five models due following ANOVA tests between models with and without the term (VeDBA model: \( \chi^2 (1) = 0.05, p = 0.83 \); RocRM1: \( \chi^2 (1) = 0.001, p = 0.97 \); RocRM5: \( \chi^2 (1) = 0.22, p = 0.63 \); RocRM10: \( \chi^2 (1) = 0.01, p = 0.91 \); RocRM60: \( \chi^2 (1) = 0.09, p = 0.77 \)). See Additional file 1 for further details.

Log-transformed mean VeDBA was not a statistically significant predictor of \( V02 \) mg ind\(^{-1}\) h\(^{-1}\) with large confidence interval bands (which include 0) and an \( R^2_m \) of 0.01 (Table 2; Fig. 4). In contrast, log-transformed mean RocRM increased with \( V02 \) (Table 2; Fig. 4) at all time-steps, with smaller (statistically significant) \( p \) values for larger time-steps. \( R^2_m \) values increased with increasing RocRM time-step (0.15, 0.18, 0.23 and 0.31). 95% prediction interval bands for these \( p \) values indicated increasing replicability of the model comparison outcome with increasing time-step of RocRM.
$B$ values for RocRM models also increased with increasing time-step. For RocRM time-steps of 1, 5, 10 and 60 s, the alternative hypothesis (that there is a difference between the full and null models) was at most 16.66, 53.26, 83.86 and 685.31 times as likely as the null hypothesis (that there is no difference between the full and null models). BFUB could not be reported for the VeDBA model as the $p$ value was $> 0.37$.

AIC-derived evidence ratios showed similar trends, with increasing evidence for the full model over the null model with increasing RocRM time-step. ER for RocRM time-steps of 10 and 60 s were particularly large. By contrast, ER for the VeDBA showed that the evidence for the full model was just 2.66 times as strong as for the null model.

**Discussion**

Dynamic body acceleration is the prevalent tag-derived metric by which to quantify energetic expenditure of animal movement. This methodology appears flawed in slow-moving organisms. Novel ways of assessing movement and their effects on metabolic rates are needed to better understand slow-moving species movements. Here we have shown that RocRM, a movement metric based on angular velocity about the three rotational axes (pitch, roll, yaw), is a good proxy for estimating...
energy expenditure in free-living spider conchs. The evidence is most convincing for time-steps of 10 and 60 s. Given the lack of a relationship between $\dot{V}O_2$ and VeDBA in this study, it appears that rotational-based metrics may provide a critical movement-linked energy metric where dynamism-based ones do not. These results support the suggested advantage of rotational metrics from preliminary data on conch movement presented in Wilson et al. [27]. They also build upon previous studies that show how slow-moving data in the European spiny lobster, $P$. elephas, is best described in terms of pitch and roll metrics [29] and that fine-scale behaviours in diving loggerhead turtles are manifest most noticeably in the angular velocity about the yaw axis [44].

Clearly, however, there is a sizeable source of variation around the predicted relationship not accounted for by RocRM, indicated by the confidence intervals and modest marginal $R^2$ values for shorter RocRM time-steps (particularly 1 and 5 s). Firstly, the need to place Daily Diaries on the hard shell of the conch inevitably means that conch activity that does not manifest in the movement of the shell is unaccounted for. For example, conchs feed using their proboscis whilst grazing [35, 53], which does not result in the movement of the shell [28]. The proportion of time, and energetic expenditure, that such movements account for is unknown, but might explain the variance in $\dot{V}O_2$ at the lowest RocRM values. We also note that we did not consider the non-movement energetic contributions of specific dynamic action (SDA) [54].

As conchs were allowed to graze whilst in their holding tanks on the growing algal film, the metabolic demands of digestion could well be relevant here. Indeed, other studies on gastropod species have found a postprandial peak following algal film feeding (see Table 2 of [54] and the references therein). Disentangling this effect, whether by explicitly studying the SDA of $L$. truncata in this context or by eliminating any food source in studies to validate our results, would certainly help clarify the extent of this potential variation.

Table 2  Linear mixed model results

| Model | Predictor | Est | CI        | t    | p (PI) | $\chi^2$ (df) | $\Delta$AIC (ER) | $R^2_m$/$R^2_c$ |
|-------|-----------|-----|-----------|------|--------|---------------|------------------|-----------------|
| 1     | Intercept | 4.20| 19.24–27.64| 2.49 | .      |               | 1.96 (2.66)     | 0.01/0.10       |
|       | log(mean VeDBA) | –0.37| –6.74–5.99 | 0.51 | (0.0001–1). | 0.03 (1) | 4.4 (9.02) | 0.15/0.41 |
| 2     | Intercept | 10.54| 6.74–14.27 | 5.82 | ***    | 16.66         | 6.40 (1)        | 0.18/0.40       |
|       | log(mean RocRM) | 2.59| 0.73–4.28  | 2.87 | (7.11$^{-6}–0.56$)* | 53.26 | 8.46 (1) | 0.23/0.46 |
| 3     | Intercept | 7.61| 6.10–9.19  | 9.27 | ***    |               | 6.46 (25.28)    | 0.31/0.51       |
|       | log(mean RocRM) | 2.26| 1.01–3.55  | 3.19 | (1.1$^{-6}–0.34$)** | 53.26 | 8.46 (1) | 0.23/0.46 |
| 4     | Intercept | 7.06| 5.67–8.22  | 11.00| ***    |               | 6.46 (25.28)    | 0.23/0.46       |
|       | log(mean RocRM) | 2.32| 1.03–3.54  | 3.84 | (6.909$^{-8}–0.14$)*** | 83.86 | 11.79 (1) | 0.23/0.46 |
| 5     | Intercept | 5.25| 4.27–6.09  | 10.74| ***    |               | 9.79 (133.62)   | 0.31/0.51       |
|       | log(mean RocRM) | 1.68| 0.95–2.37  | 4.57 | (2.04$^{-6}–0.04$)*** | 685.31 | 16.28 (1) | 14.28 (1261.43) |

Results of the final (following removal of terms through stepwise elimination) linear mixed models where $\dot{V}O_2$ mg ind$^{-1}$ h$^{-1}$ is predicted by either log-transformed mean VeDBA or log-transformed mean RocRM (over either 1 s, 5 s, 10 s or 60 s stepping value)

PI refers to the prediction interval of the p value

$\Delta$AIC refers to the difference between the null and full models, as shown by ANOVA model comparison, and ER refers to the AIC evidence ratio

$B$ refers to Bayes factor upper bound. A $B$ was not calculated for Model 1 due to the p value being $> 0.37$

$R^2_m$ (marginal $R^2$) refers to the variance of the (final) linear mixed model explained by only the fixed-effect terms, whereas $R^2_c$ (conditional $R^2$) refers to the variance explained by both fixed- and random-effect model terms [49]

$*$, ** and *** refer to p values of $> 0.05$, $< 0.05$, $< 0.01$ and $< 0.001$, respectively

Fig. 3  Density distributions of the 5-h mean RocRM values for each time-step (where ‘time-step’ refers to the time interval of the differential that RocRM was calculated over- either 1, 5, 10 or 60 s)
We also suggest that the method used here, using an overall mean taken over approximately 5 h, misses the shorter, more intense bouts of movement that might produce higher RocRM. However, reducing the time period over which \( \dot{V}O_2 \) is measured is likely to increase the error around \( \dot{V}O_2 \) estimates derived from the limitations in detecting very small changes in \( O_2 \) in a relatively large volume of water, especially as ‘high’ activity periods are brief. Nonetheless, further tests using more sensitive oxygen sensors may refine our work further.

The sensor probes employed here could reliably measure \( O_2 \) concentration only to a minimum resolution of once every 30 s, due to the traversal time of \( O_2 \) across the sensing foil. Other studies regularly employ probes taking measurements once per second [55, 56] and their use here would certainly have allow a more finely resolved \( \dot{V}O_2 \)–RocRM relationship to be quantified.

Conversely, reduction of the water volume, which might mitigate some of this problem, will tend to constrain animal movement. Our experimental set-up...

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**Fig. 4** Oxygen consumption in *Lambis truncata* is well explained by RocRM but not VeDBA. Relationships between oxygen consumption (\( \dot{V}O_2 \) mg ind\(^{-1}\) h\(^{-1}\)) and either mean log-transformed VeDBA or four different time-steps of log-transformed mean RocRM (1, 5, 10 or 60 s). Trend lines show the fitted relationships calculated from linear mixed models (LMMs) with 95% confidence intervals.
differed from many ‘traditional’ respirometry studies that use fixed-volume, solid-walled, smaller (relative to the animal’s body size) chambers, which would constrain animal movement. The tank volume employed here, approximately 114 times the volume of the conchs, allowed enough room for the animals to move in concentrated bouts.\(^1\) We note previous studies [25, 41] on blacktip (*Carcharhinus limbatus*) and lemon (*Negaprion brevirostris*) sharks encouraged consistent movement by having an animal to chamber volume ratio approximately two times larger than ours, although these animals are considerably more mobile than the spider conchs tested here. Any tank setup will restrict normal movement, of course. When coming up against a barrier to movement, conchs will often continue to try to move up against it until they can move again [28]—if this happened here, then it is possible that conchs would have expended large amounts of energy for what would appear, from the Daily Diary data, to be limited movement of the shell.

As the respirometry chamber had a soft plastic top in place of a solid lid, it is feasible that any conchs climbing up the tank sides could break the airtight seal by pushing the plastic covering away from the water’s surface. Conchs were regularly observed throughout the actual trials, however, and were never seen to climb the experiment tank sides beyond propping themselves at an angle between the sides and tanks bottom. As well as their greater weight and size compared to many gastropods, the lack of tortuous substrate may have stopped the conchs, whose shells are bigger and more protruding than many gastropods, from being able to traverse a 90° angle.

Chamber volume may not be the only factor affecting results. An important consideration when applying this relationship to free-living conchs is how the animal moves in relation to, and as a consequence of, its environment. Specifically, the substrate across which the conch travels will alter the relative contributions of changes in pitch, roll and yaw/heading. Further, the substrate may also negate or emphasise the inclusion of one or more rotational axes as part of RocRM. In simple respirometry chambers with flat bottoms as used here, pitch and roll changes are likely to make a smaller contribution towards the overall mean RocRM than if the animal was traversing rocky, uneven substrate such as the coral rubble that many conch species prefer [36]. Simulating such environments in respirometry conditions poses difficulties in, for example, the proper mixing of water and calculations of water volume when using fixed-volume chambers, as well as providing more substrate for microbes which may affect \(O_2\) measures. Nonetheless, comparison of different substrate rugosity/complexity in future studies would certainly improve the application of the RocRM–\(\dot{V}O_2\) relationship for a given subject species. The movement bout of the individual depicted in Fig. 2 suggests that changes in yaw/heading correspond closely to that of pitch, contrasting with a more negligible change in the roll. This is unsurprising, as the way in which the conch raises itself up onto its foot during a directional change will lead to a pitch change of the animal [28, 35]. As very few, if any, benthic animals move without any change whatsoever in at least two rotational axes, it would seem prudent to include all three axes as part of RocRM predictions of energy expenditure until further work indicates otherwise.

Apparent circadian rhythms were responsible for difference in movement levels, matching subsequently collected data on free-roaming *L. truncata* showing nocturnal preferences in movement (unpublished data; see also Additional file 1 on the difference in \(\dot{V}eDBA\) and RocRM between afternoon and evening periods). This was good from a perspective of finding a range of activities (i.e. data representative of both higher and lower movement levels). Interpretation of this would have been enhanced had we known more about the natural circadian behaviour of this study species, so that we might have been able to target periods of greater movement intensity.

In the frame of slow-moving animals that exhibit little-to-no dynamic acceleration signal (sensu Gleiss et al. [17]), the actual ‘jump’ of *L. truncata* is surprisingly dynamic—at least by comparison to the steadier motion of other invertebrates, such as crustaceans or urchins. These jumps do produce spikes in \(\dot{V}eDBA\), as seen in Fig. 2. However, the instances of these jumps were seemingly insufficiently large, or did not occur often enough, to make a difference in the relationship between \(\dot{V}eDBA\) and \(\dot{V}O_2\) over the 5-h time scale employed here—unsurprising when surge, heave and sway axes produce a vastly lower signal during these drifts [28]. In fact, in many instances, changes of heading were not even accompanied by a discernible change in acceleration, pitch or roll. The apparently greater contribution of drifts compared to jumps might also be a result of context—exploration of a small tank rather than an open benthic environment leading to the conch having less of a straight line path on which to jump continuously. It is also reasonable to assume that these jumps are at least partly anaerobic in

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\(^1\) ‘Bouts’ were considered to be groups of movements that interspersed non-movement periods (the non-movement periods lasting at least several minutes). Bouts lasted anywhere from < 1 min to > 15 min. Unpublished data on this species in the field have shown that 28.5% and 81.2% of movements occurred after an interval of < 1 min and < 10 min, respectively, with minimum and maximum between-movement times of 0.01 and 508.6 min, respectively.
nature, leading to a degree of post-exercise oxygen consumption (EPOC) [57]. Indeed, leaping bursts in other conch species lead to anaerobic-metabolism enzyme build-up [58] and EPOC up to 12 h post-burst [37]. We would suggest that our 5-h should account for at least part of this effect. However, more finely resolved VO2 measurements under different conditions should elucidate this (e.g., an experimental procedure more similar to [37]), as would further study specifically to quantify the length and size of EPOC. We suggest that researchers might consider a ‘scale’ of taxa to compare the competencies of VeDBA and RocRM according to the types of motion exhibited by the animals, whereby animal movement changes from being mostly described by its dynamic component of acceleration towards predominantly the change in the static component.

Beyond energy expenditure estimations, the use of different timescales (i.e. differentials over varying time-steps) in analysis of animal movement rotations may be a valuable tool in inferring, and in turn describing, scales over which an animal moves (Fig. 2). Smaller time-steps in RocRM, for example over just a few seconds, can pick up the instantaneous changes that occur during a jump, where larger time-steps (for example, in this instance 60 s) might best highlight longer-term changes in body posture (particularly if such body posture changes, relative to the plane of the sea bed, are a summation of slower, gradual, less dynamic movements such as drifts [28]). Indeed, different time scales of yaw-change have been shown to reveal fine-scale movement differences in Loggerhead turtles (Caretta caretta) that can be indicative of behaviour types better than DBA metrics alone [44].

Visualising these scales of movement, such as through density plots as shown here, may then show movement over different scales simultaneously to produce a ‘fingerprint’ for a species, individual or context and potentially be used as a means of comparing these against others. In Fig. 3, there is a clear bias towards scales of movement manifest at the largest time-step tested (RocRM60, i.e. 60 s). This is supported by the much larger Bayes factor upper bounds values and AIC evidence ratios for RocRM time-steps of 10 and 60 s compared to 1 or 5 s. Additionally, the p value prediction interval for RocRM60 has bounds entirely below 0.05, suggesting very high replicability and therefore that a time-step of 60 s does appear to be the most suitable for linking rotational movement to O2 consumption. We note, however, that increasing time intervals also allow the animal time to turn back on itself, affecting the angle traversed within the time interval and therefore the estimated angular costs. Researchers concerned about this miscalculation of the differentials might consider instead calculating the integral of each rotational axis over a given time-step; comparing the integral to the differentials may well identify instances of the animal turning back on itself.

A natural next step using L. truncata would be to derive estimates of energy expenditure through RocRM in free-living individuals to make ecologically relevant observations and inferences about this and related species. For example, ‘energy landscapes’ have typically been visualised and described for highly mobile species such as birds [59, 60] but, by using RocRM-derived estimates, spatial use and daily movement patterns could be framed more widely within the context of slow-moving benthic invertebrates. Additionally, RocRM may well convey much more information about an animal’s movement type than VeDBA when movement data are matched to behavioural observations; here, there are clear differences in the observed peak profiles depending on the relative sizes of change in the pitch, roll and heading axes.

Conclusions

The particular viability of RocRM to predict oxygen consumption (VO2) over different time intervals would indicate that it could work for a variety of differentially slow-moving species, even if their angular velocity is appreciably different from that of spider conchs. Further work should be conducted to understand the mechanisms behind the variability around the mean trend of RocRM vs VO2. Nonetheless, we have presented evidence suggesting that rotational movement metrics better correlate to energetic expenditure in slow-moving conchs than DBA. Using similar experiments to ours on species that will have fewer sources of movement not measured by attached tags would be an obvious first step to validating these results.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s40317-021-00255-x.

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Authors’ contributions

LWH and RPW conceived and designed the study with input from NRG, ECP and CMD. LWH and NRG conducted the static respirometry work and capture and release of specimens. LWH conducted data analysis with input from RPW,
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