Are the RNA:DNA ratio and dry-weight-at-length suitable growth proxies for brown shrimps (*Crangon crangon*)?

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Summary: Growth rates are of fundamental interest for studying population dynamics of species when no age information is available. In these cases in situ growth proxies must be identified and validated, and here we tested whether RNA-DNA ratio (RD) and dry weight condition (DWC) can act as such proxies for the shrimp *Crangon crangon*. Growth rates (mm d–1) were determined for male and female shrimps (20 to 67 mm) reared at five different temperatures (5°C to 25°C) and caught in two different months (May, July). Due to the potential interactive effects of the factors sex, size, temperature and time, different statistical analysis were applied to test for linear interactions (factorial analysis of mixed data, FAMDs), nonlinear additive effects (generalized additive models, GAMs) and nonlinear interactions (nonlinear regression combined with GAMs). The FAMD indicated positive correlations between length, month and growth rate. A positive correlation between DWC and sex and between RD and temperature was also indicated. RD was not significant in the GAM but a nonlinear link between length, temperature and growth was found. Finally, an extended Bertalanffy growth model was fitted to sex-specific growth rates and in a second step a GAM was fitted to the differences (residuals) between observed and predicted growth. RD again had no significant explanatory power for growth and, although it is a commonly used growth proxy, we concluded that it is not a suitable index for evaluating length growth of *C. crangon*.

Keywords: *Crangon crangon*; growth; dry weight; RNA; DNA; GAM; FAMD.

¿La relación ARN:ADN y el peso en seco a la longitud son indicadores adecuados para determinar el crecimiento de la quisquilla gris (*Crangon crangon*)?

Resumen: En el estudio de las dinámicas poblacionales de las especies, y en particular en las que no se dispone de información sobre la edad de los individuos, es fundamental conocer las tasas de crecimiento. Por lo tanto, es necesario identificar y validar las aproximaciones sobre estimación de crecimiento *in situ* adecuados. En este estudio se ha analizado si la relación ARN-ADN (RD) y la condición de peso seco (DWC) pueden actuar como indicador para la estimación de crecimiento en el camarón *Crangon crangon*. Se determinaron las tasas de crecimiento (mm d–1) para los camarones macho y hembra capturados (20 a 67 mm) en dos meses diferentes (mayo, julio) y cultivados a cinco temperaturas diferentes (5 a 25°C). Debido a los posibles efectos interactivos de los factores como sexo, tamaño, temperatura y tiempo, se aplicaron diferentes análisis estadísticos para probar las interacciones lineales (FAMD: análisis factorial de datos mixtos), efectos aditivos no lineales (modelos aditivos generalizados, GAMs) y interacciones no lineales (regresión no lineal combinada con GAMs). La FAMD indicó correlaciones positivas entre la duración, el mes y la tasa de crecimiento. También se observó una correlación positiva entre DWC y sexo, y entre RD y temperatura. La RD no fue significativa en el GAM, pero se encontró un vínculo no lineal entre la longitud, la temperatura y el crecimiento. Finalmente, se adaptó un modelo de crecimiento ampliado de Bertalanffy a las tasas de crecimiento específicas por sexo y, en un segundo paso, se adaptó un GAM a las diferencias (residuales) entre el crecimiento observado y el previsto. De nuevo RD no tenía un poder explicativo significativo para el crecimiento y aunque es un indicador de crecimiento utilizado comúnmente, concluimos que no es un índice adecuado para evaluar el crecimiento de longitud de *C. crangon*.

Palabras clave: *Crangon crangon*; crecimiento; peso seco; ARN-ADN; GAM; FAMD.


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INTRODUCTION

One of the fundamental variables in population analysis and ecology is age, as only with age factors such as maximum lifespan, productivity, sensitivity to the environment, reproduction rates and general life cycle can a population be described precisely. If no age information is available, age can be estimated from length and growth, but growth is influenced by factors such as energy availability, experienced temperatures, behaviour and physiological condition, and thus undergoes high variations. The aim of this study was to identify factors that explain this variability and that help us to determine growth rates accurately from field samples.

Crustaceans moult regularly, which leads to a loss of hard parts and all easily accessible age information (Hartnoll 2001), so ageing of crustaceans remains a challenging task. Although new methods for ageing shrimps have been developed recently (Kilada et al. 2012), these methods might, like the age pigment lipofuscin (Sheehy et al. 1999), not be suitable for all crustaceans and especially not the short-lived species. A solid management or a sound understanding of the population biology of these species is therefore only possible if good growth information is available to allow for a conversion of observed length to age. Here, the uncertainty in the age information gained from length increases significantly if individual growth is variable or not determinable in situ, e.g. due to common restrictions of cohort analysis (following the progression of length modes). Cross validations between field and laboratory studies are either few in number or not straightforward due to rearing biases and variable field conditions. A better length-age conversion could be obtained if robust proxies were available. Even a separation from slow and fast growing individuals would be of use.

For brown shrimp (*Crangon crangon*) such a proxy would be of interest because it is heavily exploited in the southern North Sea, it is not managed so far, partly due to missing age information, and, with a total exponential mortality of $Z \geq 5$ a$^{-1}$ (Hufnagl et al. 2010), the vast majority of the shrimps grow no older than two years, so age proxies such as lipofuscin are not applicable. More specifics on the general life cycle of the shrimp can be found elsewhere (Tiews 1970, Campos and van der Veen 2008, Siegel et al. 2008).

Reviewing the literature on *C. crangon* growth, we found a high growth variability within and between single studies (Hufnagl and Temming 2011b). Weight growth is seldom measured as it is unreliable, so we focused on total length growth. Growth rates for 20- to 30-mm-sized shrimps (total length, TL) at 20°C has been reported to be 0 mm d$^{-1}$ (Edwards 1978), about 0.2 mm d$^{-1}$ (van Lissa 1977) and 0.5 mm d$^{-1}$ (Dalley 1980). Even under ad libitum feeding conditions, growth variability has remained high for reasons that are so far unknown (Hufnagl and Temming 2011a, b). Individual differences in the genome, feeding behaviour or adaptation to winter conditions might be possible reasons but have not been tested systematically so far.

If we assume that the growth variability, observed between and within different studies, is related to different feeding behaviour (e.g. individual intake rates), preferences, activity, or different life histories (e.g. metabolic adaptation leading to lower activity and/or lower consumption and respiration rates), then this should be measurable. Protein synthesis or cell activity can, for example, be determined by the RNA content in a cell or by deriving other condition proxies such as dry weight in relation to length. Analysing and understanding the relation between growth and these proxies could help to better understand and describe the population structure, the seasonal cycle and the maximum age.

RNA-DNA$^{-1}$ (RD) is a commonly used proxy for individual growth (see references below). The amount of RNA in a cell is assumed to correlate with protein synthesis, and thus with nutrition and growth, whereas the amount of DNA, the carrier of genetic information, is assumed to be constant in the majority of cell types. Among a variety of species, significant correlations of RD and growth have been reported, for example, in larval fish (Hovenkamp and Witte 1991, Buckley et al. 1999, Peck et al. 2003), corals (Buckley and Szmań 2004), scallops (Lodeiros et al. 1996) and crustaceans such as *Artemia* (Dagg and Littlpage 1972), *Homarus* (Juinio and Cobb 1994), *Penaeus* (Moss 1994a, 1994b) and *Calanus* (Wagner et al. 1998). However, in a number of studies no correlations were determined (Mathers et al. 1994, Lee et al. 2006, Norkko et al. 2006), indicating that a priori testing and calibration of this proxy is required.

Hufnagl et al. (2010) found that, in starvation experiments and in field-caught animals, RD and DWC were meaningful proxies describing short-term starvation processes. Thus, RD and DWC have the potential to indicate nutrition history, leading to the question whether both measures can also be applied as growth proxies.

The aim of this study was therefore to determine whether RD and DWC are suitable in situ growth proxies for the brown shrimp *Crangon crangon* and whether or how temperature, shrimp size, sex and time of the year will influence these proxies.

MATERIALS AND METHODS

Growth experiments

Growth experiments were performed at the “AWI-Biological Institute Helgoland” between 15 May and 11 August 2006. Temperatures (T) of 5°C, 10°C, 15°C, 20°C and 25°C and shrimps of 20 to 67 mm TL were used. Conditions, preliminary handling, catch area, feeding and a detailed analysis of the observed growth, mortality and moult rates are described in detail in Hufnagl and Temming (2011a). A combined analysis of the data in relation to growth rates (GRs) obtained by other authors was given in Hufnagl and Temming (2011b). Only a short description of the experimental design will thus be given here.

Shrimps were reared in groups distributed over 73 aquaria. In 50 aquaria all individuals were marked with coloured plastic marks to track individual growth and
moult rates. No negative effects on growth or mortality were observed using this treatment. In the remaining 24 aquaria, the animals were sorted to one initial length, and growth was determined from the difference between start and end length divided by rearing time.

Shrimps were caught on 8 and 9 May on a cruise with the research vessel Uthörn in the Weser (58°49’N, 8°10’E), Elbe (54°02’N, 8°20’E) and Eider (54°17’N, 8°27’E) estuaries between 4 and 8 m water depth. Due to the large mesh sizes used on board, no small shrimps were collected during the first sampling. Furthermore, small shrimps prefer shallow water and were therefore obtained “on foot” using a push net (1.8 mm mesh size) off the coast of Büsum, Germany (54°07’N, 8°51’E) in about 1 m water depth on 12 May and 10 July.

All shrimps were fed daily with live polychaetes (Nereis spec., Lanice conchilega), green algae (Ulva lactuca) and live plankton. The plankton was obtained from regular samplings performed off the island of Helgoland every workday using a 280- and 500-µm mesh (Greve et al. 2004). Shrimps were reared at 5°C, 10°C, 15°C, 20°C and 25°C and 31.7±0.6 psu in closed recirculation systems with 50% daily water exchange. Only shrimps that spent at least one complete moult cycle plus ten days in the experiment tank at the rearing temperature were used for later analysis. This was done to make sure that no external uncontrolled factors from the time the animal had spent outside the laboratory would influence the results.

For marked animals, GR was obtained from the last observed moult increment (difference between pre- and post-moult length) divided by the inter-moult period. For all pre-sorted unmarked shrimps, the length was determined at the beginning and at the end of each experiment. Wet weight was not tracked during the experiment due to high measurement uncertainties and to avoid handling stress. Dry weight was determined at the beginning and at the end of each experiment due to high measurement uncertainties and to avoid handling stress. Dry weight was determined from the time the animal had spent outside the laboratory would influence the results.

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Determination of the RNA:DNA ratio

RD was based on fluorescence measurements following largely the procedure described by Caldarone et al. (2001). Deviations from their method, accuracy of the analysis and application of the method to C. crangon collected in the field and reared in the laboratory were described in Hufnagl et al. (2010). RD was determined from freeze-dried muscle samples (3rd segment). Samples were homogenized and dissolved. After adding of the fluorescence marker ethidium bromide, a first baseline fluorescence measurement was taken. After digestion of the RNA in the sample using RNase, the second fluorescence measurement was taken. After digestion of the DNA by DNase, the third measurement was taken. From the differences between the fluorescence values and by internal calibration (on the same 96-well plate) with a DNA and RNA standard, the amount of RNA and DNA and the RNA-DNA ratio were determined. Each sample was measured twice.

Statistical analysis

For each shrimp the following factors were available: RD, TL (mm), DWC, rearing temperature (T, °C), sex (juvenile, male, female) and catch month (May, July). In total 666 shrimps were analysed, 36 juveniles (sex could not be defined), 175 males, 389 females and 66 females with eggs (female+). Numbers per length class (ranges from 20 to 29 mm, 30 to 39 mm, etc.) were 12 (20 mm), 107 (30 mm), 261 (40 mm), 175 (50 mm), 111 (60 mm), respectively. At 5°C, 10°C, 15°C, 20°C and 25°C, 123, 189, 155, 137 and 62 animals were reared, respectively.

Due to the number of interacting factors and the complexity of the data set, we chose different approaches to identify whether RD and DWC are suitable growth proxies:

1) Data were split by sex and the level of RD or DWC and the observed GRs were compared. In this analysis temperature and shrimp length were not included. 2) A factorial analysis of mixed data (FAMD) was used to identify the importance of all available variables and to identify whether and which variables were linearly related. The FAMD is comparable to a principle component analysis but accounts for qualitative and quantitative variables at the same time. The FAMD will not account for direct nonlinear effects. 3) A generalized additive model (GAM) was used to identify nonlinear additive effects including all factors (T, TL, sex, month, RD, DWC). As GAMs account for additive but not for nonlinear interactive effects, 4) the first derivative of the von Bertalanffy growth function, expanded with a temperature term, was fit to the sex specific growth observations to allow for a description of growth by sex, length and temperature. We tried to explain the remaining variability by accounting for RD and DWC. Again, a GAM was used to identify whether the variability of the residuals (observed growth – predicted growth) can be explained by RD or DWC.

Analysis 1. Direct effect of RD and DWC

RD and DWC were rounded to the nearest 0.1 values and all female and male GRs were pooled by their corresponding RD and DWC values. All GRs in each bin were tested for normal distribution and variance homogeneity using Kolmogorov-Smirnov and Leven tests, respectively. If no normal distribution or variance homogeneity was determined, Mann-Whitney-U tests were applied to test for significant differences between groups.

Analysis 2. Linear interaction of variables: FAMD

The FAMD included all quantitative (RD, DWC, GR, TL) and qualitative (T, sex, month) variables. Temperature included five classes, 5°C, 10°C, 15°C, 20°C and 25°C; sex included female, male, female+...
(egg carrying) and juvenile; and month included May and July. FAMD was calculated using R 3.2.2 and the package FactoMineR (Lê et al. 2008).

**Analysis 3. Nonlinear additive effects of variables: GAMs**

GAMs were calculated using the R package "mgcv". No interactive effects were taken into account and qualitative and quantitative variables were used as described under Analysis 2, with one difference: here temperature was applied as quantitative variable.

The analysis was started using a full model including all variables:

\[
\text{GR} = \text{s}(\text{RD}) + \text{s}(\text{L}) + \text{s}(\text{DWC}) + \text{s}(\text{T}) + \text{gender} + \text{month}
\]

Starting from this full model, all significant parameters (p<0.05) were kept while all non-significant parameters and interactions were rejected and the model was refitted without them. Several error distributions ("Gaussian", "Poisson", "Negative binomial" and "Gamma") were tested and the model with the lowest Akaike information criterion (AIC) was finally used. The number of dimensions of all smooth terms was set to k=5.

**Analysis 4. Analysing residuals of observed and predicted GR**

Von Bertalanffy (1934) explained growth as the difference between anabolism and catabolism:

\[
\frac{dW}{dt} = H \cdot W^\frac{3}{2} - k \cdot W
\]

where \( W \) is the weight, \( k \) the catabolic constant, equivalent to % weight loss per time in starving individuals, and \( H \) the anabolic constant, related to food intake and synthesis of body mass. In Hufnagl and Temming (2011b) we expanded this growth model, after transformation, so length growth is represented with a temperature term:

\[
\frac{dL}{dt} = a + b \cdot T - c \cdot e^{d \cdot T} \cdot L
\]

The term \( c \cdot e^{d \cdot T} \) with constants \( c \) and \( d \) represents the von Bertalanffy length growth factor \( K \) (upper case as this is the length-based growth factor in relation to the \( k \) before, which is the weight-based factor), where \( a \) and \( b \) are constants. In Hufnagl and Temming (2011b) this model was fitted to growth data obtained from 25 different studies yielding an \( r^2 \) of 0.87 and a parameter “\( a^* \)” not significantly different from zero, which was therefore omitted. Owing to the good representation of the data, the same growth model was applied in this study, fitting \( b, c \) and \( d \) to observed sex specific GRs and to all GRs, including also juveniles. Residuals were determined as \( \text{Res} = \text{observed growth} - \text{predicted growth} \). For juvenile shrimps the general model was applied to predict growth.

Variability in growth residuals was analysed using GAMs as described above.

### RESULTS

#### Growth rates

Mean GRs were lowest at 5°C and 10°C, with values <0.1 mm d\(^{-1}\) for all size classes. In general, growth decreased with increasing body length and increased with increasing temperatures. Highest mean GRs (0.56 mm d\(^{-1}\)) were observed for animals that did not overwinter (those caught in July), which had a start length of 20 mm and were reared at 25°C. Lowest GRs (0.02 mm d\(^{-1}\)) were observed for overwintered adult shrimps (60 mm) reared at 10°C. Females grew on average 0.1 mm d\(^{-1}\) faster than males at comparable TL and T.

**Analysis 1. Direct effect of RD and DWC**

Higher median GRs were related to higher RD, but variability was generally high for male as well as for female shrimps (Fig. 1). Data were not normal distributed (KS, p>0.05), so Mann-Whitney-U tests were used to compare classes. Male shrimps with RD<0.5 displayed significantly lower GRs than those with RD>1. Female shrimps with RD<0.4 displayed significantly lower GRs than those with RD between 0.8 and 1.1. Variability was higher for female than for male shrimps. Shrimps with higher DWC also showed significantly higher GRs (Fig. 1).

**Analysis 2. Linear interaction of variables: FAMD**

Variables GR, month and TL were closely linked to axis 1, which explained 25% of the variance (Fig. 2). Variables DWC and sex were linked to axis 2, which explained 13% of the variance. Axes 3 to 5 (not shown) explained 11%, 10% and 9% of the variance. Variables RD and T were linked to both the first and the second axis. A high positive correlation was indicated for GR, TL, RD and month, while the variables representation indicated only a weak relation between month and DWC or month and sex.

In the individual factor map the variable temperature was represented by the first and second axis, while month was fully described by the first axis. Distributions of individuals, when described by axis 1 and axis 2, were also significantly different from zero on axis 1 for 25°C and on axis 2 for May and July as here \(-2<\text{v.test}<2\) (Table 1). The \( v \).test further indicated that shrimps from the 5°C experiment were significantly located on the negative parts of both axes (\( v \).test for dim 1 and dim 2 <−2), while those of the 15°C experiment were located on the positive sides (positively associated with axes 1 and 2), which can also be seen in Figure 2.
Crangon crangon growth proxies

Fig. 1. – Boxplots of growth vs. RNA-DNA$^{-1}$ (top panels) and dry weight condition (lower panels) for male (left) and female (right) shrimps. Lines above the boxes indicate groups which were not significantly different. Light grey boxes indicate groups which were not normally distributed.

Fig. 2. – Result of the factorial analysis of mixed data. Top left, variables representation; top right, factor map; lower panels, observations by first and second dimension coloured either by the temperatures the shrimps experienced (lower left) or the sex of the shrimp (lower right), where female+ are egg-carrying shrimps.
Analysis 3. Nonlinear additive effects of variables: GAMs

RD was not but all other variables were significant \((p<0.05, \text{Table 2})\), so the final GAM was

\[
GR = s(L) + s(DWC) + S(T) + \text{gender} + \text{month}
\]

\(\left(r^2=0.773, \text{deviance expl. 77.8}\%ight)\).

Excluding the factors sex or month neither decreased the AIC nor increased the explained deviance, so both factors were kept. Estimated degrees of freedom ranged between 3 and 4 and the k-index was 0.91, 0.69 and 0.53 for DWC, L and T, respectively.

Between 20 and 40 mm TL, the regression spline increased with size and decreased again, but with a shallower slope, for larger sizes (Fig. 3). The DWC regression spline increased with increasing values. The temperature smoother was on a comparable level for 5°C and 10°C and on a higher level for 15°C to 25°C.

Table 1. – Results of the factorial analysis of mixed data. Upper parts shows the results for qualitative variables, the lower part for the quantitative variables. For both sets of variables their contribution to dimension 1 and 2, and for the qualitative variables also the results of the v.test.

<table>
<thead>
<tr>
<th>Variable</th>
<th>ctr to Dim 1</th>
<th>v.test</th>
<th>ctr to Dim 2</th>
<th>v.test</th>
</tr>
</thead>
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<td>qualitative variables</td>
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<td></td>
<td></td>
<td></td>
</tr>
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<td>3.2</td>
<td>7.9</td>
<td>23.4</td>
<td>-15.6</td>
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<td>male</td>
<td>1.8</td>
<td>6.7</td>
<td>4.2</td>
<td>-7.5</td>
</tr>
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<td>female</td>
<td>0.3</td>
<td>-3.4</td>
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<td>11.4</td>
</tr>
<tr>
<td>female+</td>
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<td>-10.4</td>
<td>1.5</td>
<td>4.1</td>
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<td>-10.5</td>
<td>2.9</td>
<td>-6.0</td>
</tr>
<tr>
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<td>-6.2</td>
<td>9.4</td>
<td>-11.4</td>
</tr>
<tr>
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<td>8.1</td>
<td>5.1</td>
<td>8.1</td>
</tr>
<tr>
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<td>2.6</td>
<td>7.8</td>
<td>3.8</td>
<td>6.9</td>
</tr>
<tr>
<td>25°C</td>
<td>0.1</td>
<td>1.1</td>
<td>1.7</td>
<td>4.3</td>
</tr>
<tr>
<td>May</td>
<td>10.1</td>
<td>-22.7</td>
<td>0.0</td>
<td>0.1</td>
</tr>
<tr>
<td>July</td>
<td>17.2</td>
<td>22.7</td>
<td>0.0</td>
<td>-0.1</td>
</tr>
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<td>quantitative variables</td>
<td></td>
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<td></td>
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<td>RD</td>
<td>0.442</td>
<td>-0.293</td>
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<tr>
<td>L</td>
<td>-0.751</td>
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<td></td>
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<tr>
<td>GR</td>
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<td>0.389</td>
<td></td>
<td></td>
</tr>
<tr>
<td>DWC</td>
<td>0.187</td>
<td>0.539</td>
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</table>

Table 2. – Values obtained from the generalized additive model used in Analysis 2: Nonlinear additive effects of variables. Table includes the number of knots \(k'\), estimated degree of freedom \(\text{edf}\), k-index and the significance value \(p\).

<table>
<thead>
<tr>
<th>Variable</th>
<th>(k')</th>
<th>edf</th>
<th>k-index</th>
<th>(p)-value</th>
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<tbody>
<tr>
<td>s(L)</td>
<td>4</td>
<td>3.33</td>
<td>0.68</td>
<td>0</td>
</tr>
<tr>
<td>s(DWC)</td>
<td>4</td>
<td>3.197</td>
<td>0.91</td>
<td>0.02</td>
</tr>
<tr>
<td>s(T)</td>
<td>4</td>
<td>3.935</td>
<td>0.53</td>
<td>0</td>
</tr>
</tbody>
</table>

Fig. 3. – Results of the generalized additive model. Partial residuals for growth vs. length (top left), dry weight condition (top middle), temperature (top right), sex (middle left), sampling month (middle right), and distribution of the residuals (lower panels).
Analysis 4. Analysing residuals of observed and predicted GR

Fitting the extended von Bertalanffy growth function to sex-specific growth data, the following models were obtained:

\[
GR_{\text{pred}} = 0.04324 (\pm 0.00201) T - 0.00325 (\pm 0.00250) e^{0.07035 (\pm 0.00250) T} L
\]
\[
(r^2=0.44, n=663)
\]

\[
 fGR_{\text{pred}} = 0.05567 (\pm 0.00321) T - 0.00444 (\pm 0.00038) e^{0.06892 (\pm 0.00251) T} L
\]
\[
(r^2=0.49, n=386)
\]

\[
mGR_{\text{pred}} = 0.05494 (\pm 0.00369) T - 0.00463 (\pm 0.00055) e^{0.07551 (\pm 0.00289) T} L
\]
\[
(r^2=0.57, n=172)
\]

where \(GR_{\text{pred}}\) was based on all observations (juvenile, male, female, female+), \(fGR_{\text{pred}}\) on female/female+ and \(mGR_{\text{pred}}\) on male observations. Plus/minus deviations in brackets indicate the standard error of the parameter estimate.

Fitting a GAM to the growth residuals (GR–GR\(_{\text{pred}}\)) all variables except RD were significant and taking sex or month or both factors out did not increase the AIC or explained deviance. Thus, the final model was

\[
GR–GR_{\text{pred}} = s(L) + s(DWC) + s(T) + \text{gender} + \text{month}
\]

Estimated degrees of freedom were 3.9, 3.4 and 2.3 and k-indices were 0.62, 0.54 and 0.89 for T, TL and DWC, respectively (Table 3). The total deviance explained was 52.2% and the \(r^2 = 0.512\). The TL and DWC smoother increased with increasing TL and DWC, respectively (Fig. 4). July values were again higher than May values and no clear trend in sex or temperature was visible except for slightly higher values for 5°C.

Table 3. – Values obtained from the generalized additive model used in Analysis 4: Analysing residuals of observed and predicted growth. Table included the number of knots \(k'\), estimated degree of freedom (edf), k-index and the significance value \(p\).

<table>
<thead>
<tr>
<th>(k')</th>
<th>edf</th>
<th>k-index</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>s(L)</td>
<td>4</td>
<td>3.428</td>
<td>0.537</td>
</tr>
<tr>
<td>s(DWC)</td>
<td>4</td>
<td>2.267</td>
<td>0.885</td>
</tr>
<tr>
<td>s(T)</td>
<td>4</td>
<td>3.917</td>
<td>0.624</td>
</tr>
</tbody>
</table>
**RD measurement accuracy**

Corrected correlation coefficients of the standards, measured on the same plate as the samples, were always $r^2 > 0.98$. Repeatability was determined by the coefficient of variation between the calibration standards of all 294 measurements as 8% for RNA and 10% for DNA. These measurements were conducted on 21 dates distributed over one year and analysed together with the samples presented in Hufnagl et al. (2010). The variation between wells (each sample was prepared and analysed twice) was 2% before addition of RNase and DNase and 6% after addition. The slope ratio (DNA slope/RNA slope) for laboratory intercalibration (Caldarone et al. 2006) was 0.85.

**DISCUSSION**

The two proxies chosen here, RD and DWC, only explained a very low fraction of the variability observed in individual GRs of brown shrimps, despite the reports that RD explained growth in other species (see introduction) and despite the observation that a significant decrease of both proxies was observed in starvation experiments (Hufnagl et al. 2010). Surprisingly, nor was a difference observed between the faster growing summer (July) cohort and the slower growing overwintering (May), while in the latter a reduced protein synthesis could be expected. In the following section we will briefly summarize and discuss the findings of the statistical analysis and then focus on the reasons why RD might by a bad and DWC only a weak growth proxy for brown shrimp and perhaps shrimps in general.

Several statistical analyses were presented here to be able to tackle all linear and nonlinear as well as additive and interacting effects that might have masked the explanatory power of DWC and RD on GR. In the first analysis we tested whether in general faster-growing individuals showed higher DWC or RD ratios than slow-growing ones, which was the case, but variability in growth at similar RD and DWC was generally high. This first analysis did not account for shrimp length, temperature or sampling month. Taking these factors into account in a direct and linear way, as done in the FAMD analysis, indicated that all these factors are linked and that all of them influenced growth. However, RD and DWC had the least explanatory power and only explained a small part of the variability in the data set. Individual total length, sampling month and ambient temperature were significantly correlated with GR.

Temperature effects in biology are often nonlinear, as, for example, in physiology (e.g. Q10) growth vs. length (e.g. Gompertz equation) or enzymatic activity (e.g. Michaelis Menten). Trying to explain growth of brown shrimp by accounting only for linear effects, as assumed by the von Bertalanffy growth model. The maximum in the partial residuals at a shrimp length of about 40 mm. The probability of observing higher GRs increased with increasing temperature and was higher in July than in May. Sex effects were significant but weak and increased the explained deviation by only 1%, which is therefore negligible. As we found nonlinear interactions of length and temperature in an earlier literature study (Hufnagl and Temming 2011b), we took this into account in the last analysis. In that earlier study growth variability of shrimps of the same size and sex reared at the same temperature still displayed high growth variation, which we expected would be explainable by RD or DWC. In the GAM analysis of the residuals (difference of the observed and predicted growth), again RD was not significant, while DWC, T, L, month and sex were. Sex increased the explained deviance by 1.6% (from 50.6% to 52.2%) and can therefore again be neglected. This is mainly due to the fact that it was already accounted for in growth models (GRpred) which were sex-specific. A general sex effect is present as the growth models differed, which is in line with earlier studies (Campos et al. 2009, Labat 1977, Meixner 1969). The significant effect of T and L on the residuals is surprising, as T and L were also already included in the growth model. Mainly at the lowest temperatures (5°C), higher deviations between the observations and the growth model were observed. For the remaining temperatures no clear trend was visible. The length effect was more pronounced, indicating that for small length classes (<35 mm) the growth model predicted higher GRs in comparison with the observed ones. One reason might be that we fitted the general growth model also to the juvenile shrimps. No explicit growth model could be fitted due to the limited size range (20-30 mm). This could have biased the residuals if smaller length classes were dominated by slower growing males, as observed in the sampling area (Hufnagl et al. 2010). Another explanation could be that juvenile growth is generally not isometric, as assumed by the von Bertalanffy growth model. The GAM took this deviation into account but the observed remaining variation could still not be explained by RD or DWC; and also a strong effect of the sampling month remained.

One might argue that shrimp dry weight growth rather than length growth should have been used for a comparison with DWC or RD. However, weight growth (or carbon content) could not be tracked here as measurements were done on individuals. Getting a start and an end value from the same individual is simply not possible. As individual variability was high, so a reference sample from the start time (sampling) would not have helped either. Due to the high number of setae and extremities in relation to the relatively low body weight, the measurement error of wet weight would also have been high, covering the subtle growth differences (partly 0 mm per moult). Crustaceans also take up water to keep their gills wet and therefore an additional uncertain amount of water is trapped even after the animal has been blotted dry. Measurement errors of up to 10% can occur (unpublished data). Furthermore, water is permanently replaced by protein during the in...
ter-moult period (Passano 1960, Perger and Temming 2012), so wet weight will likely not change much over time although dry weight of course does.

**RNA:DNA**

From this study we can conclude that the RD is only of limited use for estimating in situ growth. RD was not significantly related to growth in both GAM analyses and showed only a weak correlation with growth in the FAMD. This is in contrast to species where RD significantly improved growth models (Hovenkamp and Witte 1991, Smith 2003, Peck et al. 2003) but in line with other species where authors reported that RD was not correlated with growth or that RD can only be used during special life stages and under special conditions. For example, for *Artemia salina* RNA∙µg⁻¹ depended on the growth phase and was higher in the exponential phase (Dagg and Littlepage 1972) or only juvenile GRs of the scallop *Euvola ziczac* were correlated with RD (Lodeiros et al. 1996). Although RD is generally a good proxy for growth determination of larval fish (Lee et al. 2006, Mathers et al. 1994), in *Solea solea* RD could only be used to compare larvae of the same ontogenetic stage (Richard et al. 1991). Anger and Hörche (1990) detected no correspondence of RD to instantaneous GR in carbon and nitrogen of the spider crab, *Hyas araneus*. RD of *Sardina pilchardus* varied only with length and food (zooplankton biomass, Chicharo et al. 1998a) and even in well-fed fish larvae a high variability of RD was observed (Bergeron and Boulhic 1994), so high RD values do not necessarily have to correspond to high GR.

There are several factors that might influence RD, but not growth. For shrimps these factors are most likely moulting, maturation and feeding (see more detailed discussion in DWC section). These factors influence a variety of biochemical processes e.g. the RNA level will likely increase to increase the level of moult hormones or steer gonad maturation. Both processes will increase RD although growth is not or even negatively influenced. RD responds to feeding events within a very short time span of the order of days or even hours (see below) but to determine length and weight GR appropriately it takes days to weeks.

RD was determined from *Crangon crangon* muscle samples which undergo changes during the moult cycle (Chang, 1995; Sánchez-Paz et al. 2003). Mitochondria (also containing RNA) are delocated during the moult cycle (Miyawaki and Tsuruda 1984) and activity and protein degradation is influenced (Boddeke 1976, Regnault 1979, Regnault and Lagardère 1983). Not surprisingly, RD changes related to the moult cycle have been observed e.g. for *Homarus americanus* post larvae (Juinio et al. 1992). Several hormones are included in the moult process (Hartnoll 2001) and Nott and Mavin (1986) described that metabolic activity of *C. crangon* peaks at premoult but decreases during the moult phase. RNA regulates all processes mentioned (enzyme and hormone productions) and therefore changes in the RNA concentration most likely occur during moult.

In *C. crangon* and other crustaceans RNA concentrations are therefore most likely not only related to feeding processes but also strongly dependent on the moult process and therefore most probably an unreliable indicator of growth.

Haefner and Spaargaren (1994) showed that biochemical parameters change significantly during vitellogenesis of *C. crangon*. Chicharo et al. (2007) observed higher RNA-DW⁻¹ in female than in male *C. crangon* and concluded that a possible reason might be maturation or sex-specific behaviour. Here, range and variability between male and female shrimps was comparable, which might indicate that maturation will not be the main factor affecting RD.

In addition to moulting and maturation, mainly feeding, short-term starvation during moult and the quick response of RD to feeding events will cause a mismatch between RD and GR (see also Hufnagl et al. 2010). In bivalves (Norkko et al. 2006), fish larvae (Bergeron 2000), fish juveniles (Malloy and Targett 1994, Richard et al. 1991) and decapods (Rosa and Nunes 2004) RD is influenced by feeding and starving within days in bivalves or hours (Lee et al. 2006, Chicharo et al. 1998b, Lough et al. 1996). This stands in conflict with the experimental time span (weeks) necessary to determine GR in *C. crangon*. Nevertheless, it was worthwhile testing and analysing it, as temperature, feeding and light conditions in the experiment were kept as constant as possible and all shrimps were fed the same diet. Assuming that RD mainly reflects metabolic and enzymatic processes, a link between constantly high growth and high RD could have been visible.

Injuries might also have influenced the results, as regeneration might reduce growth of *C. crangon* (Tiews 1970). Regeneration under ad libitum feeding could therefore lead to high RD ratios but low GRs. Only healthy (all extremities no black spots) animals were used in our experiments to minimize this effect, but animals were reared in groups to allow a larger sample size and cannibalism is well known for brown shrimp (Plagmann 1939).

**Accuracy**

Measurement errors cannot be fully excluded due to the determination method. Samples were freeze-dried, the muscle was dissected out on ice to prevent DNA and especially RNA degradation, and every sample was analysed twice. As in our previous study (Hufnagl et al. 2010) significant declining trends in RD in starvation experiments were determined and samples were measured alongside the samples presented here, it can be assumed that the observed variation in RD and the missing significant link to growth is due to other factors than measurement error.

**Dry weight condition index**

Our results indicate that DWC is, like RD, only of limited use as a growth proxy, but that the probability was high that an animal with high DWC also displayed
higher GR. Nevertheless, owing to the existing variability (Fig. 1) any growth prediction based on DWC will still be erroneous. Since TL and dry weight can be determined with sufficient accuracy, it can be assumed that growth variability must largely result from other sources than measurement error. These might be differences in the stage of the moult cycle, the stage of maturation of female shrimp and individual feeding behaviour, as described above.

Inter-moult periods vary between individuals, and the timespan to the next moult event could not be determined here. Individual moult events were only tracked for marked shrimps, and although this provides the time span since the last moult, it does not provide information on when the next moult will take place. Since crustaceans exhibit continuous weight growth but incremental (discontinuous) length growth, which is hindered by the hard structure of the exoskeleton (Blaxter and Outhward 1991), DWC can vary according to the moult stage (Perger and Temming 2012). Hence, growth is a steady replacement of water by protein (Rosa and Nunes 2004), and two similar sized shrimps can display different dry weights only because they are in different stages of the moult cycle. Additionally, in calculating the DWC, the whole animal is considered and differences in the thickness or calcification of the carapace might influence this proxy. Up to 17% of the whole dry weight can be made up by the exuvia (Regnault and Luquet 1978).

Furthermore, DWC is effected by weight loss due to higher energy demands or food deprivation. Several days before ecysis, crustaceans stop feeding (Blaxter and Outhward 1991). Effects of this pre-moult starvation and the energy loss during that phase are discussed differently. In most crustaceans, glycogen, glucose and lipids derived from the hepatopancreas and haemolymph are used to fuel the moultling process (Sánchez-Paz et al. 2007, Clifford and Brick 1983, Barclay et al. 1983). These substances have a high energy density and the influence should therefore be small on DWC. On the other hand, dry weight is mainly a measure of muscle protein, which is expected to decrease when growth is low (negative), which would be captured by the DWC (e.g. Hufnagl et al. 2010). During the moult process, Regnault and Lagardère (1983) assumed that *C. crangon* loses 10% of its protein biomass due to additional energy demands.

The second factor influencing dry weight is maturational ovary development of female shrimps. At about 55 mm TL, 50% of females carry eggs (Henderson and Holmes 1987, Boddeke 1961). Energy required for gonadogenesis might influence somatic GR (Taylor and Peck 2004). According to Haefner and Spargaen (1993) the weight of the ovary can increase the body weight of *C. crangon* by 5.5% and, according to Henderson and Holmes (1987), the smallest egg-bearing females were ~38 mm. In our experiments, less variability in GR per DWC (Fig. 1) was determined for male in comparison to female shrimps. This might be due to the influence of maturation.

*C. crangon* feeds mainly at night (Feller 2006), and Pihl and Rosenberg (1984) determined that the stomach content is about 1% to 2% of the body weight. This fact might add to the variability. In this study shrimps were fed and sampled at comparable times, but differences in individual feeding behaviour could have contributed to the variability.

**PERSPECTIVE**

Although the hypothesis that RD or DWC can be applied as growth proxies had to be rejected here, it cannot be rejected per se due to factors not included here. Future work should therefore focus on changes of DWC and RD over the moult cycle and maturation in relation to growth.

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