Effect of Swimming Activity on Relative Weight and Body Composition of Juvenile Rainbow Trout

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Abstract.—Fisheries managers often assess body condition using relative weight (W_r) because it provides a comparative measure of fish plumpness among individuals and populations. However, it is not known whether the morphological information that W_r summarizes reflects physiological measures, such as relative lipid reserves, in rainbow trout Oncorhynchus mykiss. The purpose of this study was to determine whether swimming activity affects either the W_r or proximate body composition of juvenile (total length, 170–260 mm) rainbow trout. When rainbow trout from a hatchery were fed ad libitum for 147 d, inactive (no current) and active (15 cm/s current velocity) fish did not differ in W_r. However, inactive rainbow trout maintained relatively constant lipid levels, whereas active fish declined in lipid content. Relative weight may provide a comparable measure of body form, but it is not an accurate index of lipid content between active and inactive rainbow trout fed an excess ration. For assessing the physiological condition of rainbow trout, measurement of proximate body composition appears to be more accurate than indices based on length and weight.

Fisheries surveys frequently include the collection of length and weight data to describe various characteristics of fish populations, such as growth, length structure, and body condition. Assessments of body condition are important to biologists who manage fish populations, and various body condition indices have been developed to aid in these assessments. Condition indices measure the “plumpness” of fish and have been used to assess the effects of environmental variation on fish populations, such as changes in habitat and prey availability (Austen et al. 1994; Filbert and Hawkins 1995; Lio et al. 1995; Simpkins and Hubert 2000). Condition indices have frequently been used as monitoring tools and to assess the status of fish populations for management purposes (Gabelhouse 1991; Johnson et al. 1992; Marwitz and Hubert 1995; Weiland and Hayward 1997).

Debate has centered on methodological and statistical issues of using body condition indices (see Bolger and Connolly 1989; Cone 1989; Springer et al. 1990). From these discussions, relative weight (W_r) has become the most accepted index for assessing body condition (Murphy et al. 1990; Ney 1999). Relative weight compares the weight of an individual fish to a standard weight derived from a standard length–weight relationship for the species and the measured total length of the fish. A characteristic of W_r is that it provides a comparative measure of fish plumpness (Murphy et al. 1990). However, little is known whether the morphological information that W_r summarizes reflects possible physiological differences among fish, such as lipid or protein reserves.

The assumption in the use of W_r is that the weight of an individual fish reflects its physiological condition (Gutreuter and Childress 1990; Murphy et al. 1990). Physiological condition of fish has often been described in terms of reserve nutrients, particularly lipids, present in the body (Love 1970; Gershanovich et al. 1984; Mommersen 1998). Physiological condition determines the ability of fish to compete for foraging and spawning sites, maintain and repair tissues, and cope

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with stress caused by environmental change (Brown and Murphy 1991). Changes in endogenous energy reserves are associated with metabolic processes that either store or deplete lipid or protein (Wedemeyer et al. 1990; Navarro and Gutiérrez 1995). Rose (1989) found that Wc could be used to assess body composition of immature walleyes Stizostedion vitreum during feeding experiments. Brown and Murphy (1991) observed that Wc was correlated with body composition of juvenile striped bass Morone saxatilis and palmetto bass (striped bass × white bass M. chrysops) that were fed different rations. Studying adult white crappie Pomoxis annularis, Neumann and Murphy (1992) found that seasonal variation in Wc corresponded to changes in percent whole body lipid. Jonas et al. (1996) found significant relationships between seasonal changes in Wc and energy density of age-0 and age-1 muskellunge Esox masquinongy but suggested that Wc was not a robust indicator of seasonal changes in energy density.

Allocation of consumed energy can differ among active and sedentary fish (Warren and Davis 1967; Adams and Breck 1990). Fish that swim actively tend to partition more consumed energy into metabolic processes and less into storage than do fish that are relatively sedentary (Kitchell 1983; Mømmersen 1998). Consequently, at similar feeding rates, actively swimming fish should have lower lipid reserves than fish that are inactive (Busacker et al. 1990; Moyes and West 1995; Mømmersen 1998). Relative weight is thought to serve as a surrogate for estimating proximate body composition (Blackwell et al. 2000). So, actively swimming fish should have lower Wc values than fish that are relatively sedentary if they are eating similar amounts. However, research on lipid reserves of fish has been limited to monitoring rates of depletion associated with periods of starvation and spawning migrations (Idler and Bitner 1959; Jöbling 1980; Black and Love 1986). Little attention has been devoted to studying the effects of swimming activity on the allocation of consumed energy into protein and lipid reserves (Moyes and West 1995; Mømmersen 1998). To our knowledge, no studies have evaluated whether Wc and measures of proximate body composition are different between actively swimming and sedentary populations of fed fish.

The purpose of this study was to determine whether swimming activity affects the Wc and proximate body composition of juvenile (169–233 mm total length [TL]) rainbow trout Oncorhynchus mykiss. Our objectives were to describe the changes in Wc and measures of proximate body composition (i.e., whole body estimates of lipid, water, and protein) and evaluate differences in Wc and measures of proximate body composition between actively swimming and sedentary juvenile rainbow trout held under controlled laboratory conditions.

Methods

Juvenile rainbow trout (130–150 mm TL; 35–45 g total weight) were obtained from the Wyoming Game and Fish Department’s Como Bluffs Fish Hatchery and transported to the University of Wyoming’s Red Buttes Environmental Research Laboratory near Laramie, Wyoming. Seventy-five fish were marked with soft visual-implant tags (Northwest Marine Technologies, Inc.) and placed in each of two 240-L circular tanks (height = 0.85 m; radius = 0.30 m) that received a continuous flow (10 L/min) of aerated well water maintained at 7.5°C. An automated timer and fluorescent lights were used to produce a photoperiod of 12 h light: 12 h dark. Fish were allowed to acclimate for 21 d and were fed an excess ration of Silver Cup pelleted trout chow (Nelson and Sons, Inc.).

After the acclimation period, rainbow trout were sedated with MS-222 (3-aminobenzoic acid ethyl ester methanesulfonate), individually identified, weighed to the nearest gram, and measured to the nearest millimeter TL. Two experimental treatments were used to affect Wc and proximate body composition. Fish continued to receive an excess ration of trout chow but were held either in a current (approximately 15 cm/s) produced by circulating water pumps (March Manufacturing, Inc., Model 2U) or in standing water.

All of the rainbow trout from each tank were sedated, individually identified, weighed, and measured every 21 d. Six fish were randomly selected from each treatment and killed to allow assessment of proximate body composition on each sampling date. The dead fish were quartered and dried at 60°C to constant mass (approximately 48 h). Whole body water content was determined as the difference between wet and dry tissue weight. Dry tissues of individual fish were homogenized and stored in air-tight plastic containers at −20°C away from light. Total lipid content of dry tissue was assessed by quantifying total fatty acid concentration. Duplicate 200-mg samples of dried tissue were subjected to direct transesterification with boron trifluoride in methanol to obtain fatty acid methyl esters (Rule 1997). Fatty acid methyl esters were quantified by gas–liquid chromatography (Hewlett-Packard 5890) as described by
Table 1.—Mean length, weight, and relative weight ($W_i$) did not differ significantly ($P < 0.05$) between sedentary and actively swimming juvenile rainbow trout at the onset (day 0) or conclusion (day 147) of the experiment.

<table>
<thead>
<tr>
<th>Measure</th>
<th>Day 0</th>
<th>Day 147</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>SE</td>
</tr>
<tr>
<td>Sedentary</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Length (mm)</td>
<td>169</td>
<td>0.8</td>
</tr>
<tr>
<td>Weight (g)</td>
<td>40</td>
<td>0.7</td>
</tr>
<tr>
<td>$W_r$</td>
<td>94</td>
<td>0.6</td>
</tr>
<tr>
<td>Active</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Length (mm)</td>
<td>170</td>
<td>0.7</td>
</tr>
<tr>
<td>Weight (g)</td>
<td>49</td>
<td>0.7</td>
</tr>
<tr>
<td>$W_r$</td>
<td>94</td>
<td>0.7</td>
</tr>
</tbody>
</table>

Rule (1997). Nitrogen content of dry tissue was determined by using a CHNS analyzer (Fisons Instrument EA 1108) with 5–10-mg samples; results were converted to total crude protein (percent crude protein = percent nitrogen $\times 6.25$). Total lipid and crude protein were expressed as dry-weight percentages.

Relative weight ($W_r$; Wege and Anderson 1978) was calculated as

$$W_r = (W/W_i) \times 100,$$

where $W$ was the wet weight (g) of the individual fish and $W_i$ was the length-specific standard weight. The logit $W_r$ equation for rainbow trout ($\log_{10} W_r = -5.023 + 3.024 \log_{10} TL$) was from Simpkins and Hubert (1996).

Two-sample $t$-tests were used to test for differences in the distributions of initial lengths, weights, and $W_r$ values between treatments. This test was also used to evaluate differences in final measurements between treatments. Simple linear regression was used to evaluate relationships between dependent variables ($W_r$, percent total body water, percent lipid, and percent protein) and duration of swimming activity or inactivity by rainbow trout. A $t$-test was used to test for differences in intercepts of the relationships between treatments (Zar 1996). Coefficients of determination, residual plots, and an $F$ lack-of-fit test were used to evaluate the suitability of applying linear models to assess trends in $W_r$ and measures of proximate body composition over time (Neter et al. 1996). For relationships where initial estimates (i.e., intercepts) were not significantly different, the general linear model (GLM; Neter et al. 1996) was used to evaluate whether slopes of the relationships between dependent variables and time differed among treatments. The GLM took the form:

$$Y = \beta_0 + \beta_1 X_1 X_2 + \varepsilon,$$

where $Y$ was the dependent variable of interest, $X_1$ was a nominal variable equal to 0 if fish were inactive and 1 if fish were active, and $X_2$ was a continuous variable equal to the time (duration) of treatment ($d$). The value of $\beta_0$ was a derived constant equal to the common intercept of the relationship between the dependent variable and time. Statistical computations were performed by using SuperANOVA (Abacus Concepts 1991). Significance was determined at $P < 0.05$ for all tests.

**Results**

No significant differences were observed in the length ($t_{148} = 0.60; P = 0.55$), weight ($t_{148} = 0.41; P = 0.61$), or $W_r$ ($t_{148} = 0.78; P = 0.43$) of fish among treatments at the onset of the experiment (Table 1). Relative weight significantly increased over time for both sedentary ($t_{1239} = 11.13; P < 0.001$; $r^2 = 0.23$) and active ($t_{1239} = 8.74; P < 0.001$; $r^2 = 0.14$) fish, and relationships did not significantly differ from a linear pattern for active ($F_{6,423} = 2.01; P = 0.06$) or inactive ($F_{6,423} = 1.87; P = 0.08$) fish. Moreover, rates of change in $W_r$ were not significantly different between active and inactive treatments ($F_{1,369} = 1.29; P = 0.26$; $\beta_1 = 0.065 \pm 0.001$; $r^2 = 0.19$: Figure 1). By the end of the experiment, active and sedentary fish did not differ significantly in mean length ($t_{63} = 1.83; P = 0.07$), weight ($t_{63} = 1.73; P = 0.09$), or $W_r$ ($t_{63} = 0.46; P = 0.65$; Table 1).

Estimates of initial percent lipid content did not differ significantly between active and inactive fish ($t_{63} = 0.99; P = 0.32$; $\beta_0 = 16.27 \pm 0.04$), and relationships between percent lipid and time did not significantly differ from a linear pattern for active ($F_{5,36} = 0.26; P = 0.93$) or inactive ($F_{5,35} = 1.36; P = 0.26$) fish. However, slopes of the
relationships between percent lipid and time were significantly different between treatments ($F_{1,84} = 5.33; P < 0.05$). Percent crude lipid did not change significantly over time for inactive fish ($t_{40} = 0.37; \beta = 0.72$) but did decrease significantly in active fish ($t_{49} = 3.49; \beta = 0.005; \beta = -0.027 \pm 0.001; r^2 = 0.23$; Figure 1).

The estimated initial percent water content in fish was not significantly different between active and inactive treatments ($t_{63} = 1.22; \beta = 0.23; \beta = 0.005$), and relationships between percent water and time did not differ significantly from a linear model for active ($F_{1,36} = 0.31; P = 0.80$) or inactive ($F_{1,35} = 0.50; P = 0.77$) fish. However, the slopes of the relationships between percent total body water and time were significantly different between treatments ($F_{1,84} = 6.12; P = 0.05$). Percent total body water did not change significantly over time for inactive fish ($t_{40} = 0.68; P = 0.50$) but did increase significantly for active fish ($t_{49} = 2.92; P < 0.01; \beta = -0.015 \pm 0.001; r^2 = 0.19$; Figure 1).

Initial percent protein estimates were not significantly different between active and inactive treatments ($t_{63} = 1.14; \beta = 0.026; \beta = 0.07$). Percent protein increased significantly over time for both active ($t_{49} = 4.74; P < 0.001; r^2 = 0.35$) and inactive ($t_{40} = 2.49; P < 0.05; r^2 = 0.13$) fish, and the relationships did not differ significantly from a linear model for active ($F_{1,36} = 2.21; P = 0.07$) or inactive ($F_{1,35} = 1.72; P = 0.16$) fish. The rate of change in percent protein did not differ between active and inactive treatments ($F_{1,84} = 0.84; P = 0.36; \beta = 0.056 \pm 0.002; r^2 = 0.25$; Figure 1).

**Discussion**

Juvenile rainbow trout that were actively swimming differed from sedentary fish in proximate body composition. Fish that were swimming at 15 cm/s had substantially lower lipid reserves and higher water content than sedentary fish after 147 d (Figure 1). Despite differences in proximate body composition between actively swimming and sedentary fish, swimming activity did not affect $W_r$. Fish were equally plump in both treatments but differed significantly in lipid reserves. Consequently, our results do not support the hypothesis that $W_r$ serves as a surrogate for proximate body composition as a measure of fish health (Blackwell et al. 2000).

Swimming activity substantially influenced the allocation of consumed energy in juvenile rainbow trout. Fish that were sedentary maintained relatively constant lipid levels, whereas fish that were swimming decreased in lipid content. These results
suggest that active fish allocated more consumed energy to metabolic demands and less to storage than did inactive fish. Percent protein increased similarly among active and inactive fish, suggesting that both active and inactive fish consumed enough energy for the synthesis of muscle tissue. Despite differences in lipid content, mean TL and mean weight were not substantially different between active and inactive fish at the end of the experiment.

Concern has developed regarding the use of indices computed from measurements of length and weight to assess changes in body condition because assessments based on these indices do not account for the replacement of proximate body constituents by water (Navarro and Gutiérrez 1995; Moomsen 1998). Inverse relationships have been found between percent lipid and percent water content in sockeye salmon *O. nerka* (Idler and Bitners 1959; Brett et al. 1969), brown trout *Salmo trutta* (Elliott 1976), striped bass, palmetto bass (Brown and Murphy 1991), and several other species (Love 1970). Idler and Bitners (1959) found that the percent water content of sockeye salmon increased during the spawning migration, but the sum of percent water and percent lipid remained constant such that total body weight did not change. Similarly, we found that percent water increased as percent lipid decreased in actively swimming juvenile rainbow trout. However, we interpret these data cautiously because the percentages of water, lipid, and protein sum to 100% (Anderson and Neumann 1996). A decrease in the mass of one constituent would result in a corresponding increase in the percentage of another constituent with a constant mass. It is likely that *Wₚ* for rainbow trout was not sensitive enough to account for a 4% decline in lipid content. However, Brown and Murphy (1991) found that *Wₚ* was more strongly related to percent lipid in striped bass and palmetto bass over a wider range of fat content (20–30%).

A possible shortcoming of *Wₚ* is that statistical properties rather than physiological measures have been used to evaluate the suitability of the condition index (Murphy et al. 1990; Hyatt and Hubert 2001). The suitability of samples from fish populations for inclusion when developing *Wₚ* equations have been based on statistical properties of relationships between length and weight. Slopes, intercepts, and coefficients of determination for relationships between length and weight vary because of small sample sizes, narrow ranges of length and weights of fish within samples from populations, and differences in morphology of sampled fish in populations used in developing *Wₚ* equations (Murphy et al. 1990; Pope et al. 1995; Kruse and Hubert 1997). A more sensitive basis for evaluating samples to include when developing body condition indices may be the use of relationships between length and measures of proximate body composition. Anderson and Neumann (1996) suggested that the ratio of lipid to protein should be similar across lengths, when assessments of body condition remain relatively constant. Therefore, a change in lipid content could be reflected by a change in body condition.

We conclude that activity had a measurable effect on the lipid content of juvenile rainbow trout but that *Wₚ* failed to identify differences. Relative weight may provide a comparable measure of body form, but it is not an accurate index of lipid content between active and inactive rainbow trout fed an excess ration. Further research is needed to evaluate relationships between *Wₚ* and body composition of rainbow trout in lotic or lentic habitats that differ in food availability. Nevertheless, measurement of proximate body composition appears to be more accurate than indices based on length and weight when assessing physiological condition of rainbow trout.

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