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Energetics, cost reduction and functional consequences of fish morphology

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Cost reduction strategies are often invoked as explanations when studies of adaptation fail to find predicted costs. This might seem discouraging, offering little opportunity for further investigation. In this paper, we demonstrate that cost reduction strategies can themselves be investigated by arguments from design. Recent work on inducible morphological defences has shown that hydrodynamical disadvantages (e.g. high drag) in fishes can be compensated for by standard metabolic rate (SMR) adjustments. Here, we theoretically investigate the possibilities and limitations for swimming cost compensation through SMR adjustment. We continue by modelling how intraspecific power curve variation affects the optimal swimming velocity between food patches. Our results show that, even though SMR modifications may compensate for hydrodynamical disadvantages, low-drag fishes will nevertheless have a marked advantage under high food abundance. The relative advantage will decrease with decreasing food levels. We also show that hydrodynamical properties of fishes can be used to predict their propensity to become foraging (or swimming) specialists. Low-drag fishes can use a broad range of swimming velocities without substantial increases in swimming cost, whereas the cost of deviating from the optimal swimming velocity increases markedly in high-drag fishes. The results have important implications for the evolution of morphological diversity in fishes.

Keywords: cost reduction; swimming; foraging; standard metabolic rate; fish morphology

1. INTRODUCTION

The link between morphology, physiology and ecological performance is central to the study of adaptation and has emerged as a main focus of modern evolutionary ecology (Lauder 1996). Functional analysis combined with optimality modelling has successfully explored this connection by relating design features to measures of performance, often with great predictive power (e.g. Pennycuick 1989; Webb 1993; Losos et al. 1997). However, one aspect has remained elusive. Phenotypes which deviate from performance optima are commonly predicted to suffer increased costs (e.g. Dodson 1984), but such costs have often proven negligible or absent (Spitze 1992; Tollrian 1995). This has led to the suggestion that disadvantages can be avoided through cost reduction strategies (Tollrian 1995; Moller 1996; DeWitt et al. 1998). However, if costs can disappear this could be a serious problem for the study of adaptation in general. In this paper, we demonstrate that cost reduction strategies can themselves be investigated by arguments from design. Our approach identifies the possibilities and limitations of such strategies and leads to specific predictions about their ecological and evolutionary consequences.

The study of animal adaptation often involves the analysis of energy budgets (e.g. Pennycuick 1989; Webb 1993). Whenever it does, the metabolic rate at rest plays an essential role. In mammals and birds, this is usually estimated as the basal metabolic rate (BMR), while the measure used for most other organisms is the standard metabolic rate (SMR). Both measures have traditionally been seen as more or less fixed, species-specific consequences of the animal’s body mass (e.g. Peters 1983). Recent work has fundamentally changed this view by demonstrating both considerable intraspecific variation and adaptive flexibility in the BMR and SMR (Ricklefs et al. 1996; Piersma & Lindström 1997; Secor & Diamond 1998). In fishes, downregulation of the SMR may enable increased growth (Wieser & Medgyesy 1990) and variation in the SMR can be used to explain fish growth strategies and life histories (Metcalfe et al. 1995; Metcalfe 1998). In the dimorphic crucian carp Carassius carassius, differences in the SMR correspond to differences in morphology (Pettersson & Brönmark 1999). Crucian carp increase in body depth in response to chemical cues from piscivorous fishes and the deeper body constitutes a morphological defence against gape-limited predators (reviewed in Brönmark et al. 1999). Brönmark & Miner (1992) suggested that the deep-bodied morphology should incur a fitness cost related to increased energy expenditure during swimming. However, deep-bodied fishes have a significantly lower SMR than shallow-bodied individuals (Pettersson & Brönmark 1999). As a result, the two morphs experience similar costs of transport despite marked differences in drag (Pettersson & Brönmark 1999). Could this be a strategy for reducing hydrodynamic disadvantages (Videler 1993)? If so, should we expect it to be widespread, also occurring in other types of polymorphisms (e.g. Smith & Skulason 1996)? Here, we address the problem by focusing on two idealized morphs which differ with respect to body drag and, hence, the cost of swimming. We then continue by addressing the functional consequences of SMR adjustment. Finally, we derive optimal swimming velocities and growth rates for these fishes when living in a patchy environment.
2. SWIMMING ENERGETICS AND COST REDUCTION

We begin our analysis by considering the shape of the swimming cost function and its consequences for characteristic velocities. When describing the metabolic cost versus swimming velocity in fishes, researchers typically fit one of two basic equations to data on metabolic rate (e.g. Webb 1993) (figure 1 and table 1). The exponential equation has the form

\[ P(U) = a \cdot e^{-\epsilon U}, \]

while the polynomial function can be written as

\[ P(U) = a + b \cdot U^c, \]

where \( a \) represents the SMR, \( b \) and \( c \) are constants obtained from the curve fitting procedure and \( U \) is the velocity relative to the water. Hydrodynamically, \( b \) and \( c \) represent the drag of the body, including efficiencies (cf. Webb 1993). Both equations have the dimensions work per unit of time. Two typical polynomial relationships are shown in figure 1, illustrating how power for one high-drag morph and one low-drag morph increases curvilinearly with increasing swimming velocity. For simplicity, we have assumed that these morphs differ only with respect to curve shape. Throughout this study, we will model cost reduction as a 40% reduction in the SMR, i.e. \( a \), of the high-drag morph. A similar SMR reduction has been found in deep-bodied crucian carp (Pettersson & Brönmark 1999).

To derive the velocity associated with the minimum cost of transport, i.e. the energy cost per unit distance, we set \( f(U) = P(U)/U \) and solve for the minimum (cf. Weih 1973; Alexander 1974). This velocity, \( U_{\text{inc}} \), can also be found by constructing a tangent from the origin to the power curve (figure 1) (Weih 1973). As a result, we find that the optimal swimming velocity \( U_{\text{inc}} \) may (equation (2)) or may not (equation (1)) be affected by SMR adjustments (see table 1). Under both equations, an SMR reduction leads to a decreased cost of transport (table 1) (flicker 1975), either through direct cost saving (equation (1)) or through a combination of direct and indirect effects as \( U_{\text{inc}} \) decreases (equation (2)). Assuming the swimming cost functions in figure 1, a 40% reduction in the SMR of the high-drag morph is sufficient to attain the same cost of transport as the low-drag morph, thus removing the effects of the hydrodynamical disadvantage.

The power curve shapes also have other implications. For instance, they provide information about the relative cost of deviating from the optimal swimming velocity \( U_{\text{inc}} \). If we denote this relative cost of transport \( \text{COT} \), the relationship can be described by

\[ \text{COT} = \frac{P(U_{\text{inc}} + \Delta U) \cdot U_{\text{inc}}}{P(U_{\text{inc}}) \cdot (U_{\text{inc}} + \Delta U)}, \]

where \( \Delta U \) is the deviation from \( U_{\text{inc}} \). This equation is dimensionless. The analysis shows that low-drag morphs are capable of adjusting their swimming velocities over broad intervals at the cost of only marginal increases in \( \text{COT} \) (figure 2). In contrast, high-drag individuals already suffer major costs at small deviations from \( U_{\text{inc}} \) (figure 2). Interestingly, this effect is even more marked for SMR-adjusting, high-drag fishes (figure 2).

3. MAXIMIZING GROWTH IN A PATCHY ENVIRONMENT

Foraging fishes commonly encounter patchily distributed food (e.g. Marschall et al. 1989). We continue our analysis by investigating how the two morphs should adjust their swimming velocities between such food concentrations in order to maximize growth (cf. Ware 1975), which is synonymous to maximizing their net rate of energy intake (Stephens & Krebs 1986). Alternatively, the surplus energy gained from foraging could be allocated to fuel reserves or energy used for reproduction. When the fish is in a food patch, we assume that the accumulated net energy gain \( E \) follows a function of diminishing return in relation to the time spent feeding in the patch (cf. Charnov 1976). When swimming between food patches the fish pay an energetic travel cost, which will be

\[ C = P(U_{\text{inc}}) \cdot U_{\text{inc}}^2 \cdot D, \]

where \( P(U) \) is the metabolic rate of swimming at velocity \( U \) (e.g. equation (2)) and \( D \) is the distance between food patches. The growth rate, measured as energy per unit of time, can be written as

\[ G = (E - C) \cdot (t_p + t_i)^{-1}, \]
Table 1. Two equations used to describe the relationship between metabolic cost and swimming velocity in fishes

(The predicted optimal swimming velocity ($U_{mc}$) associated with the minimum cost of transport, the cost of transport (COT) and the influence of a reduction in the SMR (parameter $a$) on $U_{mc}$ and COT. Parameters $b$ and $c$ represent the drag of the body, $m$ is the body mass, $g$ is the acceleration due to gravity and $U$ is the swimming velocity relative to the water. Please note that when using the exponential metabolic function (equation (1)), only $c$ will affect $U_{mc}$.)

<table>
<thead>
<tr>
<th>Power function</th>
<th>$U_{mc}$</th>
<th>COT</th>
<th>$U'_{mc}$</th>
<th>COT</th>
</tr>
</thead>
<tbody>
<tr>
<td>$P(U) = a \cdot e^{-U/m}$</td>
<td>$\frac{P(U)}{mg} \cdot U_{mc}$</td>
<td>$-$</td>
<td>$-$</td>
<td></td>
</tr>
<tr>
<td>$P(U) = a + b \cdot U^c$</td>
<td>$\left(\frac{a}{b(r-1)}\right)^{1/x}$</td>
<td>$\frac{P(U')}{mg} \cdot U_{mc}$</td>
<td>$-$</td>
<td></td>
</tr>
</tbody>
</table>

where $P'$ is the derivative of the power function with respect to the swimming velocity ($dP/dU$) (Hedenström & Alerstam 1995). The general solution to equation (6) for the optimal swimming velocity and patch residence time can easily be found graphically by constructing tangents as shown in figure 3. In this case we have assumed that the low- and high-drag morphs have the same energy gain function while foraging in a food patch (figure 3). The travel cost is for a constant distance between food patches, meaning that zero travel time will be associated with an infinite velocity. However, fishes will have a minimum travel time associated with the maximum sustainable (critical) swimming speed. The two curves denoted as $C_L$ and $C_H$ in the left quadrant of figure 3 represent the travel costs between food patches for the two fish morphs. Additionally, $C_H$ represents the travel cost for the high-drag morph after a 40% SMR reduction. The optimal swimming velocity is given implicitly from the travel time ($t = D \cdot U^{-1}$) by the mutual tangent between the gain curve and the transport cost curve (figure 3), which maximizes the overall net energy gain. Notice that the slope of this tangent also represents the overall net energy gain (equation (5)). Generally, the overall net energy gain and, therefore, the maximum growth rate is higher for the low-drag morph than for the high-drag morph. Figure 3 shows that the swimming velocity between food patches which is associated with the maximum growth rate ($G$) is higher in the low-drag morph (cheap swimming at relatively high velocities) than in the high-drag morph. The optimal swimming velocity for maximizing $G$ is higher than the velocity of the minimum cost of transport $U_{mc}$, which is the velocity where the travel cost curve has its minimum (figure 3). The associated optimal patch residence time is higher for the high-drag morph than for the low-drag morph (figure 3). When the SMR is reduced ($C_{H_{40}}$), the growth rate $G$ increases somewhat, but is still below that of the low-drag morph. The same graphical approach can also be used for analysing the effect of patch quality or foraging efficiency when in the patch (resulting in different net energy gain functions) and the effect of interpatch distances (Hedenström & Alerstam 1995). A decreased rate of foraging, e.g. as food resources decrease over the season, will yield lower optimal swimming velocities between food patches (cf. Ware 1978). The differences between morphs in net energy gain will then be lower. Furthermore, increased distances between patches will lead to reduced optimal swimming velocities between them.
If a fish is foraging just to balance its energy budget without any net energy reward, i.e. $G = 0$ in equation (7), we obtain the optimum swimming velocity condition as $dP/dU = P/|U|$. This velocity is identical to the velocity which gives the minimum cost of transport, i.e. $U_{mc}$.

4. DISCUSSION

Our work shows that cost reduction strategies can be successfully evaluated by arguments from design and biomechanics. The approach not only avoids the vagueness associated with invoking general cost avoidance but also provides entirely new predictions. First, we find that high hydrodynamical drag is not necessarily associated with a high cost of transport. By adjusting the SMR, fishes can potentially compensate for otherwise costly morphologies. Depending on the specific relationship between movement and energy consumption, i.e. the power curve, cost reductions may be the direct consequences of the decreased metabolic rate or combinations of direct and indirect cost savings. The latter is true when the power curve differs from a pure exponential shape, because reductions in the SMR then also cause $U_{mc}$, the swimming velocity associated with the minimum transport cost, to be shifted downwards, resulting in further cost reductions.

Furthermore, the foraging model shows that it is not possible to avoid the hydrodynamic disadvantage when trying to maximize the net energy gain from food patches. When the two morphs feed on the same food and patches, high-drag fishes will be further from their $U_{mc}$ than low-drag fishes will be and their resulting growth rate will be lower. Reductions in the SMR will not affect this outcome. The disadvantage will be particularly marked when patch quality is high. Thus, in nature, low-drag fishes will be able to use resources more efficiently at high patch quality. This also suggests that the difference between morphs in growth rate will decrease as resources are reduced. Interestingly, this pattern was found in a field study where deep- and shallow-bodied crucian carp competed for limited food resources (Pettersson & Bronmark 1997). Despite a lower SMR (Pettersson & Bronmark 1999) which could have released resources for growth (Wieser & Medgyes 1990; but see Friede 1983; this study), deep-bodied crucian carp grew slower than shallow-bodied competitors and the difference was most pronounced during high food abundance at the beginning of the experiment (Pettersson & Bronmark 1997). It should be noted that the present model assumes that the two morphs have similar energy gain curves. A complication occurs if the two morphs specialize in different food sources or foraging tactics (cf. Schluter 1993, 1998). If low-drag fishes are more efficient foragers while in a food patch, i.e. experience a steeper gain curve than high-drag fishes, this enhances the differences in growth rate.
between morphs. However, if the opposite is true, i.e. that a high-drag morph experiences a higher feeding rate than a low-drag morph, such as benthic and limnetic morphs of sympatric sticklebacks \((Gasterosteus\ \text{spp.})\) (Schluter 1995), differences in the growth function would reduce the effects from differential travel costs due to hydrodynamic drag. In the stickleback system the differences in growth rate among morphs may be due to secondary adaptations of the feeding apparatus, especially in the benthic form (cf. Schluter 1995). Our model might still apply to the initial state of morphological differentiation, before any other modifications of, for example, the mouth have occurred, and may explain the initial trajectory towards ecological specializations among polymorphic fishes.

We also analysed how deviations from the optimal swimming velocity affect high- and low-drag morphs. Our results established a significant but overlooked link between hydrodynamics and the evolution of trophic specialization in fishes. The cost of transport increases sharply in high-drag fishes when deviating from \(U_{mc}\). In contrast, the cost of transport for low-drag fishes is only marginally affected over a broad range of swimming velocities. This suggests that high-drag morphs should generally show less variable swimming velocities and should consequently have a propensity to specialize in these velocities and foraging strategies which involve low-velocity variance. Low-drag fishes may use a broad range of velocities without substantial cost increases and should thus be able to remain swimming and foraging generalists. It should be noted that these predictions are merely consequences of locomotion power curves.

Taken together, these results have important implications for the species which inspired this study (i.e. crucian carp), as well as for numerous other examples of dimorphisms and polymorphisms in fishes (Smith & Skúlason 1996) and, finally, for the general use of cost reduction as an evolutionary explanation (e.g. Möller 1996). First, we find that an SMR reduction may indeed allow high-drag fishes such as deep-bodied crucian carp to attain costs of transport similar to low-drag fishes. There is now abundant evidence for intraspecific variation in the metabolic rate (Ricklefs et al. 1996; Piersma & Lindström 1997; Metcalfe 1998) and SMR flexibility (Wieser & Medgyesy 1990; Piersma & Lindström 1997; Secor & Diamond 1998). However, the usefulness of this strategy is restricted by the narrow range of swimming velocities which can be used. If an SMR-reducing fish deviates from its \(U_{mc}\), the costs of transport will soon exceed those of normal fishes. In addition, the strategy most probably incurs other costs. For instance, a low-standard metabolism is normally associated with a low maximum workload, both in terms of locomotion, digestion and food conversion (Priede 1985). Nevertheless, given the large number of trophic polymorphisms in fishes (Smith & Skúlason 1996; Schluter 1998), modifications of the SMR could be a commonly occurring strategy for reducing the impact of the hydrodynamical disadvantages caused by, for example, gape-limited predation (Pettersson & Brönmark 1997, 1999) or sexual selection (Quinn & Foote 1994). Interestingly, our analysis of the relative cost of transport shows that the shapes of power curves will themselves be a factor strongly favouring ecological segregation. This aspect will definitely merit further study.

Finally, our work shows that cost-reducing strategies are not beyond the reach of functional analysis. Instead, the explicit modelling approach allows us to assess the possibilities and limitations of cost reduction. This design perspective should be taken into consideration in future studies of how morphology, physiology and ecological performance interact, as it is likely to have profound effects on how to interpret individual behaviour, trophic interactions and, ultimately, the evolution of morphological diversity.

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