The crown-filament pump of the suspension-feeding polychaete Sabella penicillus: filtration, effects of temperature, and energy cost

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ABSTRACT: The energetics of the ciliary crown-filament pump were studied for the suspension-feeding polychaete Sabella penicillus. Maximum filtration rate expressed as the clearance capacity \( F = 13.62W^{0.24} \) as a function of body size \( W, g \) dry wt was. Oxygen consumption \( (R, \text{ ml } O_2 \text{ h}^{-1} \text{ ind.}^{-1}) \) as a function of size was: \( R = 0.13W^{0.36} \). The water-processing capacity of a ‘standard’ 65 mg dry wt S. penicillus was estimated as 354 l of water filtered per ml of oxygen consumed. This suggests that the polychaete is adapted to live in waters with extremely low algal concentrations. Filtration rate as a function of temperature was measured in 2 size groups of worms. The relationship fitted straight lines and it was found that the viscosity effect may explain the whole correlation between filtration rate and temperatures between 5 and 20 °C. The operating point, \( O_p \), of the crown-filament pump was determined by equating pump characteristic and system characteristic, \( \Delta H_p = \Delta H_s \). The system characteristic was calculated as the sum of the 2 major contributions, namely the pressure drop across the pinnule-lattice of the crown-filaments, \( \Delta H_{pec} \) and the kinetic loss, \( \Delta H_{kex} \), in the water leaving each crown-filament, which was regarded as one of a series of parallel ‘pump units’. The calculated operating point and components for e.g. 15 °C were: \( O_p = \Delta H_{ipe} + \Delta H_{kex} = 0.0222865 + 0.000065 = 0.0224 \text{ mm } H_2O. \) The mechanical work done by the pump (pumping power) was 0.451 µW, compared to a total metabolic energy expenditure of \( R = 112 \text{ µW}. \)

INTRODUCTION

During recent years, several attempts have been made to characterize the pump systems of suspension-feeding animals in terms of pumping power output and energy expenditures of filtering water. Thus, the bivalve pump (Jørgensen et al. 1986, 1988; Jørgensen & Riisgård 1988), the ascidian pump (Riisgård 1988b) and the pump of the filter feeding polychaete Chaetopterus variopedatus (Riisgård 1989) have been analysed. The energetics of ciliary filter feeding in ciliates has been studied by Fenchel (1986). In a recent study the effects of temperature were studied in the mussel Mytilus edulis, which was modelled as a viscous, leaky, constant-force pump (Jørgensen et al. 1990). It was found that the filtration rate increased linearly with temperature and that 88 to 97 % of the temperature effect could be ascribed to changed viscosity of seawater.

The aim of the present work was to study the properties and energetics of the ciliary crown-filament pump of the suspension-feeding polychaete Sabella penicillus and to compare it with previous studies of biological pumps. Particular attention was paid to the effect of temperature.

Sabella penicillus lives in a tube, which it builds from suspended mud. The feeding mechanism of Sabella pavonina (= S. penicillus) has been described by Nicol (1931). The feeding organ is composed of 2 lateral lobes which are joined at the base on the dorsal side only. They curve around on either side of the mouth (Orrhage 1980), and each bears numerous filaments which stand out stiffly at regular intervals to form a crown (Fig. 1A). Each filament bears a double row of alternating pinnules (Fig. 1B, C). Water is drawn into the interpinnule canals from the outside by the beating of compound latero-frontal cilia consisting of 5 or 6 cilia (scanning electron micrographs made by Dr Claus Nielsen, University of Copenhagen; unpubl.) (Fig. 1D). Along with this water stream enter suspended food particles, mainly algal cells. By means of an unknown
Fig. 1. Sabella penicillus. (A) Glass holder with a worm within its natural tube. The water pumping and particle capturing filament-crown is extended. (B) Crown-filament (23 mm long) with 2 rows of pinnules (1.2 mm long, ca 210 pinnules in each row) branching off from the filament in an angle of 110° to each other. One filament represents one pump unit; the whole crown pump consists of 39 parallel pump units. (C)Idealized model of section of filament with pinnules. (D) Cross section of 2 pinnules with compound latero-frontal cilia in resting position, at end of recovery stroke, lfc(1), and end of active stroke, lfc(2); fc: frontal cilia; afc: abfrontal cilia. Dimensions refer to a 'standard' 65 mg dry wt worm. Dashed lines indicate water flow due to the pumping activity of the latero-frontal cilia. Drawings based on photographs of fresh preparations.

MATERIALS AND METHODS

Sabella penicillus were collected by SCUBA divers on vertical rock walls at about 30 m water depth in the Gullmarnfjord on the Swedish west coast. The worms were taken to the nearby Kristineberg Marine Biological Station and kept in running seawater (12 °C) before they were used for experiments at either the Marine Biological Station, at the Marine Pollution Laboratory, Charlottenlund, Denmark, or at the Institute of Biology, University of Odense, Denmark. The worms were transported to Danish laboratories by car in insulated seawater boxes, and then transferred to aquaria with 31 to 33 °C seawater (12 to 14 °C) from the Kattegat or the North Sea. The seawater in the aerated aquaria was changed several times a month, and fresh seawater was used in all experiments. The worms were fed monocultures of Phaeodactylum tricornutum or Dunaliella marina. During periods of up to 4 mo mortality was less than 2 %.

For filtration rate measurements, worms within their natural tubes were transferred to holders made of glass tubes (Fig. 1). Single worms or groups of worms could then be transferred to appropriate sized aquaria in which they were placed in a regular and standardized manner prior to clearance experiments.

Filtration rate measurements. The filtration rate was measured as the volume of water cleared of flagellate cells (Dunaliella marina or Rhodomonas sp., almost spherical cells, 5 to 6 μm in diameter) per unit time. The optimum particle size for retention efficiency is ca 3 to 8 μm (Jørgensen et al. 1984). Flagellate cells were added to a glass-beaker or an appropriate sized aquarium with a known volume of water (V) and a single worm (or groups of worms in ‘temperature effect experiments’, see below). The reduction in the number
of particles as a function of time was then followed by taking water samples and subsequently measuring the particle concentration by an electronic particle counter (Coulter Counter, Model TA II or Elzone 80XY). Clearance (F) was determined from the exponential reduction in algal cell concentration as a function of time (always verified as a straight line in a semi-log plot made by hand during the experiment) using the formula: 

\[ F = \frac{V}{n} \ln \left( \frac{C_0}{C_t} \right) \]

where \( C_0 \) and \( C_t \) are the algal concentrations at time 0 and time t, respectively, and \( n \) = number of worms.

**Temperature effect experiments.** Sabella penicillus were collected in the Gullmarfjord in the beginning of July. The clearance capacities of 2 size groups of worms (Group A = 10 'small' and Group B = 6 'large' individuals) were measured at different temperatures adjusted at intervals <5 °C per day (to prevent disturbance which causes the worm to produce mucus and/or retreat into its tube). Group A and B were kept in aerated aquaria holding 17.6 and 18.9 l of seawater, respectively. After addition of algal cells at an initial concentration of about \( 4 \times 10^3 \) cells ml\(^{-1}\), water samples (15 ml) were taken for determination of cell concentration every 5 min by means of a pipette, and the number of active worms was noted. Mean clearance (= filtration rate) was calculated and plotted as a function of temperature.

**Respiration experiments.** Worms were collected in September and the respiration measurements carried out at the Marine Pollution Laboratory during the following months. Oxygen consumption was measured using the experimental set-up shown in Fig. 2. The worms were placed in respiration chambers of an appropriate size (14, 26, 40, 66 and 162 ml) according to the size of the worms. The temperature of the respiration chamber was kept constant by submerging the chamber in a constant temperature aquarium (17.3 ± 0.4 °C). Adequate mixing of the water in the respiration chamber was ensured by the pumping activity of the specimen itself and by the aid of a peristaltic pump (Masterflex, Model 7567-10) recirculating water in 2 tubing systems (Tygon tube, Type 6408-41). One of the tubing systems was connected to a constant temperature cell (Radiometer, Type D 616) in which was mounted an oxygen electrode (Radiometer, Type E 5046). The oxygen electrode continuously recorded the oxygen tension, and the decrease was followed on a pen-recorder. A respiration experiment consisted of a 2 h measurement, during which the decrease in the oxygen tension was monitored. After 2 h fresh oxygen-saturated seawater was added and the oxygen tension was followed for another 2 h. All measurements were performed under oxygen tensions between 147 ± 6 and 129 ± 11 mm Hg (i.e. 94 to 83 % oxygen saturation). Oxygen uptake was corrected for each individual by following the oxygen decrease in a control experiment performed with the empty worm tube.

**Constants and conversion factors.** The following physical constants and conversion factors were used: density of seawater (35 %o S), \( \rho = 1.028 \text{ g cm}^{-3} \) at 5 °C; 1.027 at 10 °C; 1.026 at 15 °C; 1.025 at 20 °C; kinematic viscosity, \( \nu = 1.562 \times 10^{-6} \text{ m}^2 \text{s}^{-1} \) at 5 °C; 3.500 \times 10^{-6} \text{ at 10 °C}; 1.84 \times 10^{-6} \text{ at 15 °C}; 1.056 \times 10^{-6} \text{ at 20 °C}; acceleration due to gravity, \( g = 980.7 \text{ cm s}^{-2} \); 1 ml O\(_2\) = 5333 \( \mu \text{W} \). The expressions for density and kinematic viscosity were calculated from Wheaton (1977). When no references are given for formulae or concepts these matters may be found in general textbooks (e.g. Leyton 1975, Vogel 1981, Fox & McDonald 1985).

**RESULTS AND DISCUSSION**

**Filtration and respiration**

Two series of grazing experiments each with one individual of Sabella penicillus kept in an aerated aquarium, to which were added algal cells at different concentrations at different times, are shown in Figs. 3...
Reduction in the concentration of algal cells due to grazing by a single worm (56 mg dry wt) in an aerated aquarium. Three experiments on different days are shown. Arrows indicate additions of algal suspension.

Fig. 3. Sabella penicillus. Reduction in concentration of algal cells due to grazing by a single worm (56 mg dry wt) in an aerated aquarium. Three experiments on different days are shown. Arrows indicate additions of algal suspension.

Reduction in the concentration of algal cells due to grazing of the worm is rapid and constant (i.e. the clearance, expressed by the slope of the line fitted for the algal reduction in the semi-logarithmic plot, is high and appears constant) for concentrations ranging between about $2 \times 10^3$ and $4 \times 10^3$ cells ml$^{-1}$. At algal concentrations above about $10^4$ cells ml$^{-1}$ the filtration rate is reduced. At the 2 highest algal concentrations in Fig. 3 the filtration rate is relatively high after the first algal addition, but then becomes very low after the second addition. When the worm is allowed to graze the algal concentration down over a prolonged period (Fig. 4) the grazing rate gradually becomes higher. It may be concluded that the clearance is high and constant (up to at least 8 h) at algal concentrations below about $4 \times 10^3$ cells ml$^{-1}$, but at higher concentrations it may be suggested that the gut capacity is exceeded thus leading to a reduced filtration rate.

The maximum filtration rate expressed as the clearance capacity ($F$, l h$^{-1}$ ind$^{-1}$) measured in Sabella penicillus as a function of body size (W, g dry wt) in algal concentration intervals between $4 \times 10^3$ and $2 \times 10^5$ cells ml$^{-1}$ is shown in Fig. 5. The equation for the regression line is: $F = 13.62W^{0.24}$ ($r = 0.985$). The equation for the total filament length of all crown-filaments (L, cm) as a function of body size (W, g dry wt) is: $L = 437W^{0.32}$ ($r = 0.999$, $n = 15$, W interval: $5 \times 10^{-7}$ to 0.136 g). Because the total crown-filament length may be regarded as an indirect measure of the filtration rate, the exponent 0.32 for total filament length may be compared to the exponent 0.24 for filtration rate as a function of size. The agreement between the 2 slopes is reasonably good.

Clearance as a function of temperature in 2 size groups of worms is shown in Fig. 6. The relationship fits a straight line in both groups. The slopes of the regression lines are 2.17 and 1.68 in Group A and Group B, respectively.

Fig. 4. Sabella penicillus. Reduction in algal cell concentration due to grazing by a single worm (82 mg dry wt) in an aerated aquarium. Two experiments on different days are shown. Arrows indicate additions of algal suspension.

Fig. 5. Sabella penicillus. Clearance capacity as a function of body size at 13 °C. Regression line is shown.

Fig. 6. Sabella penicillus. Mean clearance as a function of temperature in 2 groups of worms (I) Group A: 10 individuals, 65 ± 30 mg dry wt; (II) Group B: 6 individuals, 120 ± 40 mg dry wt. Regression lines are shown.
Fig. 7 shows the oxygen consumption ($R, \text{ ml } O_2 \text{ h}^{-1}$ ind.$^{-1}$) in Sabella penicillus as a function of size (W, g dry wt). The equation for the regression line is: $R = 0.13W^{0.66}$ ($r = 0.985$). During several of the respiration experiments the worm frequently withdrew into its tube, but no difference in oxygen uptake was noted. This indicates that the crown of S. penicillus is not a respiratory organ, but may exclusively serve feeding purposes. A few observations were made with the use of a video system connected to an inverted microscope on which was placed a transparent plexiglass beaker with seawater and a small (ca 2 cm long) S. penicillus. The worm was in its natural tube, fixed in an inclined position (by means of a holder made of a piece of glass-tube). It was observed that the water-pumping latero-frontal cilia on the pinnules of the filaments did not beat when the crown was more or less withdrawn into the tube – or even when the crown was fully extended but not wide open. Since we could not detect any difference in oxygen consumption between the undisturbed, water-pumping worm and the disturbed worm with resting latero-frontal cilia, it seems reasonable to conclude that the beating of the latero-frontal cilia contributes only marginally to the total respiration rate.

The respiration rate of Sabella penicillus measured in this study (Fig. 7) may be compared to that of other suspension-feeding polychaetes. Kayar (1978) measured the oxygen uptake in S. melanostigma. By converting wet body weight to dry weight (5:1) Kayar’s data conform to the following equation for respiration ($R, \text{ ml } O_2 \text{ h}^{-1}$) as a function of dry weight (W, g): $R = 0.36W^{1.02}$ (W interval: 2 to 160 mg). Riisgård (1989) measured respiration ($R, \text{ ml } O_2 \text{ h}^{-1}$) as a function of dry body weight (W, mg) of Chaetopterus variopedatus and found the relationship: $R = 1.90W^{0.59}$ (W interval: 274 to 886 mg). The estimated respiration rate of S. melanostigma and C. variopedatus, each of 65 mg dry weight, is found to be 23.4 and 22.3 $\mu l$ $O_2$ h$^{-1}$, respectively. In the present study a ‘standard’ 65 mg dry weight worm has a respiration rate of 21 $\mu l$ $O_2$ h$^{-1}$. This value is close to those found for S. melanostigma and C. variopedatus.

The individually measured clearance capacities (Fig 5) are systematically higher than the mean clearances measured in the 2 groups with 10 and 6 individuals (Fig. 6). The lower values may be due to insufficient mixing in the aquaria. Instantaneous mixing of the total water volume is a prerequisite for the use of the clearance formula, and this condition is more easily approached with only one individual than with groups of worms. As the error of underestimation is systematic, the slopes of the lines in Fig. 6 are believed to describe the true relationship between clearance and temperature. An approximate slope of 2 for the ($t, \text{ °C}$) for the ‘standard’ worm was adopted and the relation expressed as:

$$F = 90 + 2t$$

The following discussion considers a ‘standard’ 65 mg dry weight Sabella penicillus with a clearance capacity, $F = 7.44$ $l$ $h^{-1}$ at 17 °C. The amount of water that a temperate zone, near-coastal, marine suspension feeder must filter to obtain enough food to cover the minimal energy requirements exceeds 10 l of water per ml of oxygen consumed (Jørgensen 1975). The water-processing capacity of the ‘standard’ Sabella penicillus is estimated as 354 l of water filtered per ml of oxygen consumed. This value may be compared to 50 l per ml of oxygen in Chaetopterus variopedatus (Riisgård 1989) which coexists with S. penicillus in the Gullmarnfjord. The 7-fold difference in water-processing capacity between C. variopedatus and S. penicillus may reflect differential particle capturing efficiency. C. variopedatus retains particles down to about 1.5 $\mu m$ with 100% efficiency, while the retention efficiency in the ciliary feeding S. penicillus rapidly declines below 3 $\mu m$ (Jørgensen et al. 1984). The high water-processing capacity of S. penicillus suggests that the measured clearances are representative of those applying in nature, and further, that the polychaete is adapted to live in waters with extremely low algal concentrations. Throughout its ontogenetic development the mussel Mytilus edulis, which retains particles somewhat less efficiently than S. penicillus (i.e. 4 um particles are retained with 100% efficiency), processes 15 to 50 l water per ml of oxygen consumed (Riisgård et al. 1980). Thus, the 2 species seem to be adapted to different regimes of suspended food particles, and the present data suggest that M. edulis may not be able to live in the same localities as S. penicillus. The rapid saturation of the digestive system (i.e. low clearance rates) at algal concentrations above $10^4$ cells ml$^{-1}$ in S. penicillus (Fig 3 and 4) supports this interpretation S. penicillus

$$\text{Fig. 7. Sabella penicillus. Respiration as a function of body size at 17.3 ± 0.4 °C. Regression line is shown}$$
seems to process water continuously and both the filter- and digestive systems are designed to operate optimally at low phytoplankton concentrations. The same adaptation may be found in other suspension-feeding polychaetes, and the use of unnaturally high particle concentrations in filtration experiments may explain the generally low values found for suspension-feeding polychaetes by other workers (Table 1). In a study of *Lanice conchilega*, Buhr (1976) used an algal concentration of $4 \times 10^4$ cells ml$^{-1}$. This is about 10 times the 'saturation' concentration found in the present work for *S. penicillus* (Figs. 3 and 4). The very low filtration rates measured by Dales (1969) and Brown (1977) in *C. variopedatus* have been discussed and partly explained by Rüsgård (1989). Presumably, the low filtration rates shown in Table 1 are not representative of the water processing capacities of the species.

A functional response similar to that seen in Figs. 3 and 4 has previously been found in early veliconchia of *Mytilus edulis* (Rüsgård et al. 1980) and veliger larvae of the hard clam *Mercenaria mercenaria* (Rüsgård 1988a) fed *Isochrysis galbana* cells. At low algal concentrations ($2 \times 10^3$ to $4 \times 10^3$ cells ml$^{-1}$) the clearance of *M. edulis* larvae was high and constant over 8 h. At high algal concentrations (above $2 \times 10^4$) the clearance capacity was only maintained until the stomach was full. The time at which this took place was indicated by a sharp shift in the relationship between clearance and algal concentration in a semi-logarithmic plot. Similar shifts can be seen in Fig. 3. Other experiments with *S. penicillus* exposed to algal concentrations between $4 \times 10^3$ and $10^4$ *Rhodomonas* cells ml$^{-1}$ have also shown such shifts. The reduction in cell concentration after the gut has been filled probably reflects simultaneous refilling of the gut. This may explain the curves for algal reduction in the experiments with high algal concentrations in Fig. 4.

The present work emphasizes the need for more knowledge about actual algal concentrations in the immediate vicinity of benthic suspension-feeding polychaetes. The food particle concentration in the boundary layer may be extremely low. Benthic suspension feeders seem to have developed filter-, pump- and digestive systems to cope with low algal concentrations. The absence of control of filtration rate as a means to control feeding rates in *M. edulis* (Jørgensen et al. 1988) stresses the importance of performing laboratory experiments at natural algal concentrations and of interpreting laboratory findings in a meaningful ecophysiological context. The apparently inverse rela-

### Table 1. Clearance (= filtration rate) in different suspension-feeding polychaetes. (Body size expressed as wet wt has been converted to dry wt by dividing by 5)

<table>
<thead>
<tr>
<th>Family</th>
<th>Species</th>
<th>Dry wt (mg)</th>
<th>Clearance (l h$^{-1}$)</th>
<th>Suspended particles</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terebellidae</td>
<td><em>Lanice conchilega</em></td>
<td>1.3</td>
<td>0.009</td>
<td>6.92</td>
<td>Dunaliella Buhr (1976)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3.6</td>
<td>0.011</td>
<td>3.13</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>8.1</td>
<td>0.013</td>
<td>1.63</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>9.5</td>
<td>0.013</td>
<td>1.41</td>
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<td></td>
<td>35.9</td>
<td>0.026</td>
<td>0.75</td>
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<tr>
<td>Serpulidae</td>
<td><em>Pomatoceros triqueter</em></td>
<td>2.2</td>
<td>0.011</td>
<td>5.0</td>
<td>Dunaliella Klockner (1978)</td>
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<tr>
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<td>7.3</td>
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<tr>
<td></td>
<td><em>Salmacina dysteri</em></td>
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<td>0.00029</td>
<td>10.4</td>
<td>Graphite Dales (1957)</td>
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<tr>
<td></td>
<td><em>Spirorbis borealis</em></td>
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<td>0.00023</td>
<td>4.8</td>
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<td><em>Schizobranchia insignis</em></td>
<td>200</td>
<td>0.35</td>
<td>1.75</td>
<td>Graphite Dales (1961)</td>
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<tr>
<td></td>
<td><em>Myxicola infundibulum</em></td>
<td>539</td>
<td>0.286</td>
<td>0.531</td>
<td>Graphite Dales (1957)</td>
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<tr>
<td></td>
<td>100-500</td>
<td>2.8</td>
<td>1.34</td>
<td>Thalassiosira</td>
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<tr>
<td></td>
<td></td>
<td>1.04</td>
<td></td>
<td>Chroomyces</td>
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<td></td>
<td><em>Sabella penicillus</em></td>
<td>37</td>
<td>0.073</td>
<td>1.973</td>
<td>Graphite <em>Duania</em></td>
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<tr>
<td></td>
<td>65</td>
<td>7.1</td>
<td>109</td>
<td><em>Duania</em></td>
<td>Present study</td>
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<td>Chaetopteridae</td>
<td><em>Chaetopterus variopedatus</em></td>
<td>50</td>
<td>1.08</td>
<td>21.6</td>
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<td></td>
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<td>864</td>
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<tr>
<td></td>
<td></td>
<td>904</td>
<td>1.94</td>
<td>2.15</td>
<td>? Brown (1977)</td>
</tr>
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</table>
tionship between algal concentration and filtration rate in, e.g., mussels (Winter 1973, Sprung & Rose 1988) should not be interpreted as a regulatory mechanism (Riisgård & Mohlenberg 1979, Jørgensen et al. 1988, 1990). The decrease in filtration rate may simply be due to an overloading of the feeding system when unnaturally high algal concentrations are used in laboratory experiments.

**Effects of temperature and energy cost**

In order to analyse the ciliary crown-filament pump of *Sabella penicillus*, dimensions referring to a ’standard’ 65 mg dry weight worm were obtained from video recordings and microscope photographs of fresh crown filament preparations. The following dimensions were measured: length of crown-filaments, LCF = 23 mm; number of crown-filaments, N = 39 (estimated from the expression: N = 18.94D0.48, where D = crown-diameter = 2 X LCF); length of pinnules = 1.2 mm; width of pinnules = 30 μm; height of pinnules, L = 35 μm; distance between 2 pinnules, l = 60 μm; angle between 2 rows of opposite rows of pinnules on the same filament = 110°. From these data and Eq. (1), the mean water velocity (v) between the pinnules at 5, 10, 15 and 20 °C is estimated to: v = 1.063, 1.169, 1.276 and 1.382 mm s⁻¹, respectively.

The operating point, Oₐ, of the crown-filament pump may be determined by equating pump characteristic and system characteristic (Jørgensen et al. 1986, 1988, Riisgård 1988b, 1989), ΔHₛ = ΔHₑ. The system characteristic in *Sabella penicillus* is calculated as the sum of only 2 major contributions, namely the pressure drop across the pinnule-lattice of the crown-filaments, ΔHₑ, and the kinetic loss, ΔHₑₓ, in the water leaving each crown-filament which may be regarded as one of 39 parallel 'pump units' (Fig. 1).

The structure of the crown-filaments approximates that of a filter consisting of parallel circular cylinders, the pinnules (diameter, d = 35 μm), with a distance, l = 110 μm, between centers of neighbouring cylinders. The model developed by Tamada & Fujikawa (1957) to predict pressure drop over such filters is used to calculate the pressure drop across the crown-filament:

\[ \Delta Hₑ = K₁vᵢu/gd \]  

(2)

where K₁ = 8r/(1 - 2ln r + r²/6), r = d/l, u is the undisturbed upstream velocity (0.773 mm s⁻¹ at 5 °C; 0.850 at 10 °C; 0.928 at 15 °C; 1.005 at 20 °C). The pressure drop has also been estimated from the experimentally obtained formula of Munson (1988) for estimating pressure drop across square-mesh screens at very low Reynolds number (Re ≤ 0.1):

\[ \Delta Hₑ = K₂(vₑ)²/2g \]  

(3)

where K₂ = 4.75(1 - α²)/2α², α = (1 - d/l)². The Reynolds number in the present case fulfills the prerequisite for using the formula (Re = ud/v = 0.04). The calculated pressure drops found by using both formulae are shown in Table 2. There is fairly good agreement between the 2 set of estimates, and a mean value is used for calculating the operating point.

The kinetic component is calculated from (cf. Jørgensen et al. 1990):

\[ \Delta Hₑₓ = (vₑ)²/2g \]  

(4)

where vₑ is the mean velocity in the exhalant split of the crown-filament estimated as: (filament volume flow)/(filament 'exhalant area' = 2 X 23 mm; see Fig. 1) = (51/46) = 1.11 mm s⁻¹. The calculated components and operating points, Oₑ, for 5, 10, 15 and 20 °C are

<table>
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<tr>
<th>Filtration rate (ml s⁻¹)</th>
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<th>10 °C</th>
<th>15 °C</th>
<th>20 °C</th>
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<td>Eq. (1)</td>
<td>1.67</td>
<td>1.88</td>
<td>2.00</td>
<td>2.17</td>
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<tr>
<td>Eq. (5)</td>
<td>1.67</td>
<td>1.93</td>
<td>2.20</td>
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</table>

<table>
<thead>
<tr>
<th>Head losses (mm H₂O)</th>
<th>5 °C</th>
<th>10 °C</th>
<th>15 °C</th>
<th>20 °C</th>
</tr>
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<tbody>
<tr>
<td>Interpinnule canals ΔHₑ</td>
<td>0.021663</td>
<td>0.020587</td>
<td>0.019713</td>
<td>0.019040</td>
</tr>
<tr>
<td>Mean</td>
<td>0.024491</td>
<td>0.023275</td>
<td>0.022865</td>
<td>0.021526</td>
</tr>
<tr>
<td>Kinetic loss ΔHₑₓ</td>
<td>0.000045</td>
<td>0.000055</td>
<td>0.000065</td>
<td>0.000077</td>
</tr>
<tr>
<td>Operating point Oₑ</td>
<td>0.0246</td>
<td>0.0234</td>
<td>0.0224</td>
<td>0.0217</td>
</tr>
<tr>
<td>Pumping power Pₑ (μW)</td>
<td>0.414</td>
<td>0.443</td>
<td>0.451</td>
<td>0.473</td>
</tr>
<tr>
<td>Pump work Pₑ/R (%)</td>
<td>0.370</td>
<td>0.396</td>
<td>0.402</td>
<td>0.423</td>
</tr>
</tbody>
</table>

Table 2. *Sabella penicillus*. Estimated filtration rates from empirically derived expression, Eq. (1), and from Eq. (5), in an optimally pumping 65 mg dry weight 'standard' at reference temperatures of 5, 10, 15 and 20 °C. The head losses, operating points (Op), power output and pump work (Pₑ/R) are shown.
shown in Table 2. There is a decrease in the operating point from 0.0246 to 0.0217 mm H2O when the temperature is increased from 5 to 20 °C, and it may be noted that the main contribution to the system resistance is the pressure drop across the pinnule-lattice.

Filtration rates calculated from Eq. (1) at 5, 10, 15 and 20 °C may be compared to filtration rates at these temperatures, Fe, estimated by using the expression:

$$F_e \approx F_0 \left( \frac{v_f}{v_e} \right) = 1.67 \left( \frac{v_f}{v_e} \right)$$

(5)

From the calculated filtration rates shown in Table 2 it is seen that the viscosity effect may explain the whole correlation between filtration rate and temperature.

The temperature effects on filtration rate in *Sabella penicillus* may be compared to a recent study on *Mytilus edulis* by Jørgensen et al. (1990). It was found that 88 to 97% of the increase in filtration rate in the temperature interval from 5 to 22 °C was due to change in viscosity. It is interesting to note that the slope of the linear regression of clearance versus temperature in *M. edulis* is lower (about 1.2) than that found in the present work (Fig. 6). The significant kinetic loss in the exhalant siphon of *M. edulis* is more likely to offset increased filtration rate due to reduced frictional loss at higher temperatures than is the case for *S. penicillus* in which the kinetic exit loss is negligible.

The power output, $P_p$, from the *Sabella penicillus* pump can be calculated as the product of pumping pressure ($\Delta P = \rho g \rho_0$) and filtration rate ($F$):

$$P_p = \rho g \rho_0 F$$

(6)

Pumping power at the 4 reference temperatures is shown in Table 2. The work done by the pump may be compared to the total metabolic energy expenditure of the 'standard' *Sabella penicillus*, as expressed by the oxygen consumption $R = 0.021 \text{ ml O}_2 \text{ h}^{-1}$ which corresponds to 112 μW. The pump work, $P_p/R$, which expresses the mechanical work done by the pump as a percentage of the total metabolic rate, is also shown for the four reference temperatures in Table 2.

The ratio between metabolic and mechanical efficiency of the latero-frontal cilia in *Sabella penicillus* is not known, but to arrive at an estimate another ciliary suspension-feeding animal may be used for comparison, namely *Mytilus edulis*. The mechanical pumping power as a percentage of the power generation of the latero frontal cilia in a 35 mm 'standard' *M. edulis* may be found by relating the pump power output of 10 μW (Jørgensen et al. 1988) to the metabolic rate of 78 μW of the cells carrying the water-pumping lateral cilia (Clemmesen & Jørgensen 1987), i.e. $10/78 \times 100 = 13\%$. By using 13% as a general estimate of metabolic to mechanical efficiency in ciliary suspension-feeders the percentage of total metabolic power used for water pumping may be found in suspension-feeders belong-
ing to various taxonomic groups (Table 3). Only 3.1 % of the total metabolic energy is used for water processing in S. penicillus. The generally low values in Table 3 suggests that the energetic cost of water pumping is not a restricting factor for the success of suspension-feeding animals. The pump- and filter systems in different species of suspension-feeders seem to be discretely dimensioned to the biotope to which the animals are adapted. The energetic cost of having a large filtering organ is not likely to be the limiting factor for a suspension feeder even in an extremely meagre environment.

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