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The role of mucus in particle processing by suspension-feeding marine bivalves: unifying principles

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Abstract Contemporary research on bivalve suspension-feeding has revealed a diversity of particle processing mechanisms depending on the anatomy and functioning of the pallial organs involved. On the biochemical level, however, some evidence of homogeneity has emerged concerning the types of mucopolysaccharide associated with particle processing. The present study uses both previous data and original research combining video endoscopy and mucocyte mapping to further explore the relationships between pallial organ topography, functional correlates, direction of current flow, and mucocyte secretion type. Five species representing five different families and all four major gill types are represented: Mytilus edulis, Placopecten magellanicus, Crassostrea virginica, Mya arenaria, and Spisula solidissima. Viscous acid or acid-dominant mucopolysaccharides are used when particle transport occurs on an exposed surface, or on a structure leading directly to such a surface, counter to the prevailing current flow. Associated functions are indiscriminate transport in gill ventral particle grooves and rejection of pseudofeces. Lower-viscosity mixed mucopolysaccharides are used when particle transport is on an enclosed or semi-enclosed surface, leading to other such surfaces, and with the current flow. Associated functions are transport of particles destined for ingestion, and ingestion itself. Low-viscosity neutral mucopolysaccharides are found in regions where reduction of mucus viscosity is important, such as the areas of the labial palps responsible for fluidization of the high-viscosity mucus-

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particle cord of the gill ventral particle groove prior to particle extraction. There thus appears to be a specialization of mucus type corresponding to functional specialization of the various pallial organs in suspensionfeeding marine bivalves.

Introduction

Recent advances in investigate techniques have led to a re-examination of previous paradigms concerning the mechanisms of suspension-feeding in bivalves (see Beninger et al. 1992, 1993; Ward et al. 1993, 1994). In particular, the role of mucus has been debated for decades, with protagonists sometimes adopting completely contradictory positions (MacGinitie 1941: Jørgensen 1966, 1990). Notwithstanding recent assertions to the contrary (Jørgensen 1996), it is now clear that mucus secreted by the epithelia of the pallial organs plays a key role in all aspects of particle processing (Beninger et al. 1991, 1993, 1997a, b; Beninger and Le Pennec 1993; Beninger and Dufour 1996; Beninger and St-Jean 1997). Reports that specific types of mucopolysaccharide are associated with certain anatomical configurations and mediate distinct functions on the bivalve gill (Beninger et al. 1993; Beninger and Dufour 1996) have been extended to the labial palps (Beninger and St-Jean 1997). It is thus pertinent to examine the role of mucus in bivalve particle processing to determine whether a uniform set of principles applies regardless of species, gill type, or pallial organ involved. Indeed, behavioral observations to date have revealed a multiplicity of transport characteristics according to gill and palp type; the eventual existence of a common underlying thread would greatly facilitate the understanding of bivalve particle processing.

The present study draws upon both original research using video endoscopy and mucocyte mapping, as well as previous data from various sources, to outline common features of mucus involvement in particle processing on the pallial surfaces of five species of sus-

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pension-feeding bivalves, representing five different families and all four major gill types.

Materials and methods

Data base

The data for the present study were derived from three sources: (1) previous reports of organ function and/or mucocyte distribution for the gills and labial palps of *Mytilus edulis*, *Placopecten magellanicus*, *Mya arenaria*, and *Spisula solidissima* (Beninger et al. 1993, 1997a; Beninger and St-Jean 1997), (2) current investigations of these characteristics for the mantle and lips of *M. edulis* and *P. magellanicus* and (3) previous reports of mucocyte distribution and function of the gills of *Crassostrea virginica* (Ward et al. 1994; Beninger and Dufour 1996). Details of data type and sources are summarized in Table 1.

Sampling of Mytilus edulis and Placopecten magellanicus

A total of 11 Mytilus edulis were sampled from lease sites in Mahone Bay (44°32'N; 64°13'W) and from the intertidal in Passamaquoddy Bay (45°00'N; 67°05'W) in May and November 1992. Three specimens were used for histological preparation and eight for in vivo observations. A total of seven adult Placopecten magellanicus were taken from drag samples at Digby Bank (44°40'N; 65°50'W) and Passamaquoddy Bay in May and June 1992. Three of these were used for histological preparation, while four were used for in vivo observations. All specimens destined for histological preparation were maintained in a recirculating seawater system (5 °C, 31‰S) for 30 d prior to dissection. They were fed weekly with 2 g of spray-dried *Tetraselmis* sp. (Algal 161, Celsys Inc.) suspended in 1 litre of seawater. Individuals destined for in vivo observations were maintained for several days either in an open ambient seawater circulating system at the Huntsman Marine Laboratory or in a recirculating seawater system at University of New Brunswick (St. John, New Brunswick, Canada), where they were fed a mixture of the unicellular algae Isochrysis (Tahitian strain), Chaetoceros muelleri, and Dunaliella tertiolecta at 10000 cells ml^{-1} final volume (15 to 16 °C, 33‰ S).

In vivo observations

The technique of video endoscopy (Ward et al. 1991) was used to observe pseudofeces transport from the labial palps to the external medium in two *Mytilus edulis*. The mussels were presented with the unicellular algal mixture, to which was added dropwise a suspension of reflective paint particles. Using a dissecting microscope this was compared to transport of carmine particles in six specimens, in which one valve had been removed. As no differences were observed in particle treatment, only the latter technique was used for *Placopecten magellanicus*.

Mucocyte staining and counting

The combined periodic acid–Schiff (PAS) and Alcian-blue technique was used for staining mucocytes (Beninger et al. 1993). Because of the nature of the tissues (thickness, mucocyte density), the mantle was stained in toto, while the lips and anterior part of the oesophagus were embedded in paraffin and sectioned at 10 μ m (mean mucocyte diameter) prior to counting.

The lips of Mytilus edulis and Placopecten magellanicus were divided into counting zones representing regions from the distal to the proximal extremities, as shown in Fig. 1.2 and 1.4. The mucocytes of *M. edulis* were situated both within and beneath the epithelium. All mucocytes within a 100 µm length of epithelium were counted first in one direction, then in the opposite direction. In order to maintain an error of less than 5%, a third count was performed if a difference of > 2 was found for counts between 30 and 50, or if a difference of > 3 was found for counts between 51 and 100. The means of these counts all then had ranges of 5% or less. Above 100, a 5% range was maintained in a similar manner. This procedure was repeated for each of three individuals. Since the subepithelial mucocytes were often arranged in poorly defined, dense clusters, a semiquantitative visual scale of their density was adopted, with the following values: 4 = cells indistinguishable;3 = cells barely distinguishable; 2 = cells easily distinguished; 1 = few mucocytes. Nine histological sections were counted as described for the lips of each of three specimens. Mean values and standard deviations were calculated for each region. The lip mucocytes of P. magellanicus were exclusively epithelial, and were counted as for the epithelial mucocytes of M. edulis.

The pallial surface of *Mytilus edulis* was divided by three transects perpendicular to the dorsal margin, across the anterior, median, and posterior regions, as shown in Fig. 2.2; the pallial surface of the mantle of *Placopecten magellanicus* was similarly divided by seven transects as shown in Fig. 2.4, because of its more circular shape compared to the mussel. Counts were performed in toto at intervals corresponding to 1/3 of each transect length, to the ventral margin (Fig. 2.4). All mucocytes within a $100\times$ field (170 µm diameter) were counted once in a clockwise direction and once in a counter-clockwise direction using a hand-held haemocyte counter. The counts were standardized to a 1-mm-diameter circle. An error of 5% was maintained as described above.

Table 1 Data sources for particle transport observations and mucocyte mapping of pallial organs in representatives of five bivalve families [1 Beninger et al. (1992); 2 Beninger et al. (1993); 3 Ward et al. (1993); 4 Ward et al. (1994); 5 Beninger and Dufour (1996); 6 Beninger and St-Jean (1997); 7 Beninger et al. (1997a); 8 Present study; – no known study]

| Family, species | Gill | | Palp | | Lips | | Mantle | |
|--|-----------|---------------------|-------------------------------|------------------|-----------|------------------|-----------|---------------------|
| | Endoscopy | Mucocyte mapping | Endoscopy (E)/ Carmine (C) | Mucocyte mapping | Endoscopy | Mucocyte mapping | Endoscopy | Mucocyte mapping |
| Mytilidae Mytilus edulis | 3 | 2 | 6 (E+C) | 6 | 8 | 8 | 8 | 8 |
| Pectinidae Placopecten magellanicus | 1 | 2 | 6 (E+C) | 6 | _ | 8 | 8 | 8 |
| Myacidae Mya arenaria | 3, 7 | 7 ^a | 6 (E+C) | _ | 7 | _ | 7 | _ |
| Mactridae Spisula solidissima | 7 | 7 ^a | 7 (E+C) | _ | _ | _ | - | _ |
| Ostreidae Crassostrea virginica | 4 | 5 | 4 (E) | _ | 4 | _ | _ | _ |

^a Quantitative mapping not possible

Fig. 1 Mytilus edulis and Placopecten magellanicus. Mucocyte densities on lips. 1.1 Mean semi-quantitative scale values for M. edulis subepithelial mucocytes in counting zones represented in 1.2 (L lip; O oesophagus). 1.3 Mean counts \pm standard deviation for epithelial mucocytes in three counting zones indicated in 1.4 (D distal; M median; Pproximal to mouth). Standard deviation for proximal zone negligible



Results and discussion

Current flow

The direction of current flow in the bivalve pallial cavity should be kept in mind when considering the pathways and mechanisms of particle transport. Since current flow in the various regions of the pallial organs can only be observed in intact animals, such data is derived from the endoscopic studies cited in Table 1, to which the reader is referred for detailed information. The general features of current flow are summarized in Fig. 3. These movements reflect the direction of water flow above the zone of influence of the transporting cilia beats (Sleigh 1989).

Composition and characteristics of mucus

Mucus is the hydrated form of secretions produced by mucocytes within epithelia, notably those of the pallial organs. The secretions consist of polysaccharide units with associated protein moieties. Classically (i.e. in vertebrates) these secretions are termed either mucopolysaccharides (synonym glycosaminoglycans) when the polysaccharide component consists of long, usually linear chains and the protein component is very small, whereas they are termed glycoproteins when the polysaccharide component consists of short, mainly branched chains and the protein component is relatively high. In invertebrates, these distinctions are not clearcut, and mucocytes may contain a mixture of secretions (Denny 1983). In the present work, we adopt the standard histochemical classification of mucus secretions (Vacca 1985; Cook 1990), wherein all mucocyte secretions are termed mucopolysaccharides (MPS), and are classed according to their degree of acidity (a result of acid groups on the saccharide units). Neutral mucopolysaccharides (NMPS) are PAS positive, Alcianblue negative, and present low viscosity. Acid mucopolysaccharides (AMPS) are PAS negative, Alcianblue positive, and present high viscosity. Between these two endpoints we distinguish mixed mucopolysaccharides (MMPS), containing roughly equal proportions of NMPS and AMPS, and acid-dominant

Fig. 2 Mytilus edulis and Placopecten magellanicus. Mean mucocyte densities for the pallial surface. 2.1 Mean mucocyte densities for *M. edulis* counting zones depicted in 2.2 (*A* anterior; *P* posterior; *D* dorsal; *M* median; *V* ventral zones).
2.3 Mean mucocyte densities for *P. magellanicus* counting zones depicted in 2.4. Only AMPS mucocytes were found on this surface in this species



mucopolysaccharides (ADMPS), which contain a majority of AMPS (see Table 5).

Participation of mucus in particle processing

In recent years, a considerable body of data using a variety of different techniques has both confirmed the participation of mucus in all aspects of particle processing and elucidated which types of mucus are used for which functions on which pallial organs (Beninger et al. 1991, 1992, 1993, 1997b; Ward et al. 1991, 1993, 1994; Tankersley and Dimock 1993; Beninger and Dufour 1996; Silverman et al. 1996; Beninger and St-Jean 1997). This body of data has recently been peremptorily dis-

missed (Jørgensen 1996), ostensibly on the grounds that in those studies using endoscopy, prolonged observation results in enhanced mucus production. This interpretation of the endoscopic work is fundamentally incorrect, as it confuses increased mucus production following prolonged exposure to low or medium particle concentrations (i.e. normal ingestion volume control, see Beninger et al. 1992) with artefactual mucus production due to the presence of the optical insertion tube (OIT). Such artefactual mucus production has never been observed with a properly positioned OIT.

To our knowledge, the only datum which has been interpreted to support the contention that mucus is not involved in normal feeding of marine suspension-feeding bivalves is the single visual observation of apparent lack



Fig. 3 Principal pathways of current flow and particle transport in a representative bivalve, Mytilus edulis. Water arrives via the incurrent siphon (IS), and the frontal surface of the gill (G) is exposed to a postero-anterior flow at the ventral margin, while the rest of the frontal surface is swept by a ventro-dorsal flow, with a progressive through component to the abfrontal region and out the exhalent siphon (ES). Solid arrows indicate current flow after crossing through to abfrontal region and on to exhalent siphon. Arrowheads represent particle transport typical of the many species which possess homorhabdic gills and a ventral particle groove (VG) (P labial palp)

of particle cohesiveness in alimentary tract aspirates (Kiørboe and Møhlenberg 1981). Using a similar technique, Bernard (1974) concluded the exact opposite; moreover, we now know that the mucus which accompanies ingestion is of relatively low viscosity (Beninger et al. 1992; Ward et al. 1994, Beninger and St-Jean 1997; present paper), such that it may not engender particle cohesiveness. Observations of this type are thus not appropriate for determining the presence of mucus. The overwhelming weight of data therefore supports the participation of mucus in all aspects of particle processing.

Lips and buccal region – *Mytilus edulis* and *Placopecten magellanicus*

Mytilus edulis possesses small, simple lips in contrast to the hypertrophied arborescent lips of *Placopecten mag*- ellanicus. In the M. edulis lips, only ADMPS were found in the epithelial mucocytes, whereas in the much more extensive subepithelial network, both ADMPS and NMPS mucocytes were present. A clear preponderance of subepithelial ADMPS (relatively high viscosity) was observed in the distalmost regions of the lips, compared to NMPS (Fig. 1.1, 1.2). This difference tended to even out toward the mouth and oesophagus, where the secretions would thus present a lower viscosity. In P. magellanicus, no mucocytes were found in the distalmost regions of the lips, but the middle region was characterized by a preponderance of AMPS and ADMPS (Fig. 1.3, 1.4), which would result in a relatively high-viscosity mucus on these epithelial surfaces. One of the proposed functions of the arborescent pectinid lips is to prevent the removal of material to be ingested by the strong currents produced by valve clapping, a characteristic defense response and the routine method of clearing the pallial cavity of detritus and pseudofeces. Food particles lifted out of the buccal region would thus tend to adhere to the lip epithelia and be more readily returned to the mouth via the amply

Mantle – Mytilus edulis and Placopecten magellanicus

ciliated lip tracts (Beninger et al. 1990).

The dorsal region of the mantle in Mytilus edulis was characterized by a marked dominance of AMPS, which declined sharply in the median region and disappeared altogether in the ventral region. This corresponded to the observed behavior of particles transported as pseudofeces from the gill particle groove to the palp (Fig. 4.1), then ventrally and posteriorly along the palp ventral margin (Fig. 4.2) to the mantle. Pseudofeces followed a very discrete pathway along the mantle just ventral to the dorsal bend of the gill, through to the inhalent siphon, where they were ejected (Fig. 4.3 to 4.6). This is in fact the only possible trajectory, since pseudofeces must be (and is) deposited in the infrabranchial chamber of the pallial cavity. Access to the exhalent siphon is blocked by the gill lamella itself, so the only possible exit is the inhalent siphon. It should be noted that the transport of pseudofeces on the *M. edulis* mantle is therefore almost entirely counter to the strong pallial current; the prevalence of AMPS along the rejection pathway thus corresponds to the requirement to anchor pseudofeces to the epithelium and prevent its resuspension in the pallial cavity, where it would again impinge upon the frontal surface of the gill filaments. The dorsal route of the rejection pathway corresponds to the location of the posteriormost tips of the labial palps and the very dorsal position of the gill dorsal bends (Beninger et al. 1995).

In contrast to the siphonal voidance of pseudofeces seen in *Mytilus edulis*, the Pectinidae reject pseudofeces via valve clapping, which periodically flushes out the mantle cavity (Yonge 1967; Morton 1979). There is thus no specific rejection pathway on the mantle, and this is 394

Fig. 4 Mytilus edulis. Still images from Hi-8 video endoscopic recordings of particle processing. Arrows indicate direction of movement. **4.1** Mucus-particle cord (*MC*) exiting gill particle groove (GPG) and being drawn onto palp dorsal fold (DF), and then onto the palp crests (C). Note that mucus-particle cord is transported parallel to palp ridges. 4.2 Pseudofeces (PF) exiting palp ventral margin (VM) and being transported as a mucus-particle bridge to the mantle (M). 4.3, 4.4, 4.5 Pseudofeces transported on a distinct mantle tract toward inhalent siphon. 4.6 Pseudofeces exiting inhalent siphon (IS) (Tsiphon tentacles). Scales are not indicated due to large depth of field



reflected in the mucocyte distribution (Fig. 2.3, 2.4). The greatest AMPS densities are found in the region where the palps are situated (Zone 1a) and near the centre of the mantle (Zones 3b, c), indicating that this may be where pseudofeces collect prior to expulsion radially during valve clapping.

Although in the case of both species, AMPS are used almost exclusively in the transport of particles (pseudofeces) on the mantle, the presence of siphons in *Mytilus edulis* is correlated with a very precise rejection pathway, which is not seen in the siphonless *Placopecten magellanicus*. Studies underway in our laboratory show that a discrete mantle rejection pathway appears typical of siphon-bearing bivalves. Patterns in bivalve particle transport and associated mucus types

Despite considerable differences in pallial organ structure and particle processing, certain common features characterize the involvement of mucus in the representatives of all five families examined here. In all cases where particle transport occurs on an exposed surface, or on an enclosed surface leading directly to an exposed surface, the accompanying mucus secretion is either ADMPS or AMPS, i.e. very viscous (Tables 2 to 4). In all these cases, particle transport is counter to the current flow (here counter is defined as an angle approximately 0 to 90° with the current flow), indicating that viscous MPS is essential for particle transport under

| Site | Surface type | Current flow | Function | Dominant secretion | Source |
|-----------------------------------|-----------------------|--------------|-----------------------------|--------------------|-----------------------------|
| Gill (frontal) | Open | Counter | Indiscriminate transport | Acid-dominant | Beninger et al. (1993) |
| Palp (dorsal fold) | Open | Counter | Indiscriminate transport | Acid-dominant | Beninger and St-Jean (1997) |
| Mantle (dorsal) | Open | Counter | Rejection (pseudofeces) | Acid | Present study |
| Palp (trough rejection tracts) | Enclosed ^a | Forward | Rejection (pseudofeces) | Acid | |
| Palp (postero-ventral margin) | Open | Counter | Rejection (pseudofeces) | Acid | (1997) |
| Palp (oral tract) | Semi-enclosed | Forward | Ingestion | Mixed | |
| Oesophagus | Enclosed | Forward | Ingestion | Mixed | Present study |

Table 2 Mytilus edulis. Mucus secretion type in relation to organ topography, function and current flow relative to particle movement

^a Enclosed surface leading directly to open surface (labial palp ventral margin)

 Table 3 Placopecten magellanicus. Mucus secretion type in relation to organ topography, function and current flow relative to particle movement

| Site | Surface type | Current flow | Function | Dominant secretion | Source | |
|---------------------------------|---------------|--------------|----------------------------|--------------------|--------------------------------|--|
| Gill (ordinary filament) | Open | Counter | Rejection (pseudofeces) | Acid | Beninger et al. (1993) | |
| Palp (postero-vental margin) | Open | Counter | Rejection (pseudofeces) | Acid | Beninger and St-Jean (1997) | |
| Gill (principal filament) | Semi-enclosed | Forward | Transport for ingestion | Mixed | Beninger et al. (1993) | |
| Palp (oral tract) | Semi-enclosed | Forward | Transport for ingestion | Mixed | Beninger and St-Jean | |
| Palp (ridged surface) | Semi-enclosed | Uncertain | Uncertain | Mixed) | (1997) | |
| Lips (proximal) | Enclosed | Forward | Transport for ingestion | Mixed } | Present study | |
| Oesophagus | Enclosed | Forward | Transport for ingestion | Mixed J | | |

 Table 4 Crassostrea virginica, Spisula solidissima, Mya arenaria. Mucus secretion type in relation to gill topography, function and current flow relative to particle movement

| Species, site | Surface type | Current flow | Function | Dominant secretion | Source |
|--------------------------------|----------------------------|--------------|-----------------------------|--------------------|-----------------|
| Crassostrea virginica | | | | | |
| Gill ordinary filament crest | Open | Counter | Indiscriminate transport | Acid-dominant | Beninger and |
| Gill principal filament trough | Semi-enclosed | Forward | Transport for ingestion | Mixed) | Dufour (1996) |
| Spisula solidissima | | | • | | |
| Homorhabdic filaments | Open | Counter | Indiscriminate transport | Acid | |
| Gill particle groove | Semi-enclosed ^a | Forward | Indiscriminate transport | Acid | Beninger et al. |
| Mva arenaria | | | | ~ ~ ~ | (1997a) |
| Homorhabdic filaments | Open | Counter | Indiscriminate transport | Acid | () |
| Gill particle groove | Semi-enclosed ^a | Forward | Indiscriminate transport | Acid | |

^a Semi-enclosed surface leading directly to exposed surface (palp ridged surface)

such conditions. The functions associated with particle transport accompanied by viscous MPS are rejection, as well as initial particle transport on the gill for a later "decision" (i.e. selection, ingestion volume regulation –

see Beninger and Dufour 1996; Beninger and St-Jean 1997) by the other pallial organs. This latter function is found only in bivalves possessing a single ventral gill particle groove (Mytilidae, Myacidae, Mactridae, etc.).

Table 5Summary of mucussecretion types used in particleprocessing on bivalve pallialcavity organs (NMPS neutralmucopolysaccharides; MMPSmixed mucopolysaccharides;ADMPS acid-dominantmucopolysaccharides; AMPSacid mucopolysaccharides;

| | Viscosity: | | | | | |
|----------------|--|----------------------------|--|---------|--|--|
| | Low | \rightarrow | \rightarrow | High | | |
| Secretion type | NMPS | MMPS | ADMPS | AMPS | | |
| Surface type | Various | Enclosed/ semi-enclosed | Open | Open | | |
| Current flow | Forward | Forward | Counter | Counter | | |
| Function | Dilution of AMPS and ADMPS to reduce viscosity | Ingestion | Indiscriminate transport for later "decision"Rejection of pseudofeces | | | |

Bivalves which possess both a ventral and a dorsal tract use the ventral tract mainly for rejection (Atkins 1937; Beninger et al. 1992).

In all cases where particle transport is on an enclosed or semi-enclosed surface, leading to other such surfaces, the accompanying mucus is a lower-viscosity MMPS (Tables 2, 3). This is also associated with forward current flow. Although the mucus-particle cord in the gill ventral particle groove is composed of either AMPS (among Mya, Spisula species) or ADMPS (among Mytilus, Crassostrea species), this is not really an exception to the rule, since the preceding surface (gill frontal surface) is exposed, and transfer of the cord to the palps usually occurs in an extremely exposed context; the subsequent surfaces are also exposed, and current flow is counter or perpendicular. The data to this point thus indicates a set of constants in the participation of mucus in particle processing, regardless of the pallial organ or the species considered (Table 5). Hence, in spite of the diversity of structure and processing function of bivalve pallial organs, an underlying common thread now appears evident.

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References

- Atkins D (1937) On the ciliary mechanisms and interrelationships of lamellibranchs. Part III: types of lamellibranch gills and their food currents. Q Jl microsc Sci 79: 375–420
- Beninger PG, Dufour SC (1996) Mucocyte distribution and relationship to particle transport on the pseudolamellibranch gill of *Crassostrea virginica* (Bivalvia: Ostreidae). Mar Ecol Prog Ser 137: 133–138
- Beninger PG, Dufour SC, Bourque J (1997a) Particle processing mechanisms of the eulamellibranch bivalves *Spisula solidissima* and *Mya arenaria*. Mar Ecol Prog Ser 150: 157–169
- Beninger PG, Le Pennec M (1993) A histochemical study of the bucco-oesophageal glands of the blue mussel *Mytilus edulis* L.: the importance of mucus in ingestion. J mar biol Ass UK 73: 237–240
- Beninger PG, Le Pennec M, Auffret M (1990) Peribuccal organs of Placopecten magellanicus and Chlamys varia (Mollusca: Bival-

via): structure, ultrastructure, and implications for feeding. II. The lips. Mar Biol 107: 225–233

- Beninger PG, Le Pennec M, Donval A (1991) Mode of particle ingestion in five species of suspension-feeding bivalve molluscs. Mar Biol 108: 255–261
- Beninger PG, Lynn JW, Dietz TH, Silverman H (1997b) Mucociliary transport in living tissue: the two-layer model confirmed. Biol Bull mar biol Lab, Woods Hole (in press)
- Beninger PG, St-Jean SD (1997) Particle processing on the labial palps of *Mytilus edulis* and *Placopecten magellanicus* (Mollusca: Bivalvia). Mar Ecol Prog Ser 147: 117–127
- Beninger PG, St-Jean SD, Poussart Y (1995) Labial palps of the blue mussel *Mytilus edulis* (Bivalvia: Mytilidae). Mar Biol 123: 293–303
- Beninger PG, St-Jean S, Poussart Y, Ward JE (1993) Gill function and mucocyte distribution in *Placopecten magellanicus* and *Mytilus edulis* (Mollusca: Bivalvia): the role of mucus in particle transport. Mar Ecol Prog Ser 98: 275–282
- Beninger PG, Ward JE, MacDonald BA, Thompson RJ (1992) Gill function and particle transport in *Placopecten magellanicus* (Mollusca: Bivalvia) as revealed using video endoscopy. Mar Biol 114: 281–288
- Bernard FR (1974) Particle sorting and labial palp function in the Pacific oyster *Crassostrea gigas* (Thunberg, 1795). Biol Bull mar biol Lab, Woods Hole 146: 1–10
- Cook HC (1990) Carbohydrates. In: Bancroft JD, Stevens A (eds) Theory and practice of histological techniques. Churchill Livingstone, Edinburgh, pp 177–215
- Denny M (1983) Molecular biomechanics of molluscan mucus secretions. In: Hochachka PW (ed) The Mollusca. Vol. 1. Metabolic biochemistry and molecular biomechanics. Academic Press, New York, pp 431–465
- Jørgensen CB (1966) The biology of suspension feeding. Pergamon Press, Oxford
- Jørgensen CB (1990) Bivalve filter-feeding: hydrodynamics, bioenergetics, physiology and ecology. Olsen and Olsen, Fredensborg
- Jørgensen CB (1996) Bivalve filter feeding revisited. Mar Ecol Prog Ser 142: 287–302
- Kiørboe T, Møhlenberg F (1981) Particle selection in suspensionfeeding bivalves. Mar Ecol Prog Ser 5: 291–296
- MacGinitie GE (1941) On the method of feeding of four pelecypods. Biol Bull mar biol Lab, Woods Hole 80: 18–25
- Morton B (1979) A comparison of lip structure and function correlated with other aspects of the functional morphology of *Lima lima*, *Limaria* (*Platilimaria*) *fragilis*, and *Limaria* (*Platilimaria*) *hongkongensis* sp. nov. (Bivalvia: Limacea). Can J Zool 57: 728– 742
- Silverman H, Lynn JW, Dietz TH (1996) Particle capture by the gills of *Dreissena polymorpha*: structure and function of laterofrontal cirri. Biol Bull mar biol Lab, Woods Hole 191: 42–54
- Sleigh MA (1989) Adaptations of ciliary systems for the propulsion of water and mucus. Comp Biochem Physiol 94A: 359–364
- Tankersley RA, Dimock RV Jr (1993) Endoscopic visualization of the functional morphology of the ctenidia of the unionid mussel *Pyganodon cataracta*. Can J Zool 71: 811–819

- Ward JE, Beninger PG, MacDonald BA, Thompson RJ (1991) Direction observations of feeding structures and mechanisms in bivalve molluscs using endoscopic examination and video image analysis. Mar Biol 111: 287–291
- Ward JE, MacDonald BA, Thompson RJ, Beninger PG (1993) Mechanisms of suspension-feeding in bivalves: resolution of current controversies by means of endoscopy. Limnol Oceanogr 38: 265–272
- Ward JE, Newell RIE, Thompson RJ, MacDonald BA (1994) In vivo studies of suspension-feeding processes in the eastern oyster, *Crassostrea virginica* (Gmelin). Biol Bull mar biol Lab, Woods Hole 186: 221–240
- Yonge CM (1967) Observations on *Pedum spondyloideum* (Chemnitz) Gmelin, a scallop associated with reef-building corals. Proc malac Soc Lond 37: 311–323