Distribution and Efficiency of Bacteriolysis in the Gut of Arenicola Marina and Three Additional Deposit Feeders

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Distribution and efficiency of bacteriolysis in the gut of *Arenicola marina* and three additional deposit feeders

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ABSTRACT. A simple technique was developed to measure the bacteriolytic activities of the digestive fluids of the deposit-feeding polychaete *Arenicola marina*. Lysis of a cultured environmental isolate, incubated with extracts of gut luminal contents, was monitored spectrophotometrically. Concurrent direct counts were used to verify cell lysis. The ability of extracts from 8 longitudinal sections of the gut to lyse the bacterium was monitored. The digestive ceca, anterior stomach, and posterior stomach regions exhibited high lytic activities, whereas bacteriolytic activities in all other regions of the gut were negligible. Similarly, extracts of surface sediments and fecal castings showed negligible lytic capabilities. The sharply limited distribution of lytic activity implicates the ceca as the source of bacteriolytic agent and suggests a true plug-flow system, with little axial mixing. Questions regarding the fate of lytic agents, which disappear abruptly posterior to the stomach, remain unanswered. Localization of lysis in the gut coupled with estimates of gut residence time permit the calculation that ingested bacteria are exposed to strong lytic activity for approximately 20 min. Incubation of *in situ* sediment samples with gut fluids corroborates the distributional findings of the *in vitro* work although the efficiency of lysis is much reduced, possibly due to exopolymer capsules and slimes of natural sedimentary bacteria. Cross-phyletic comparisons of bacteriolytic activities reveal both qualitative and quantitative differences. Much less demarcation of lytic activity is observed in the guts of a holothuroid (*Caudina arenata*) and a hemichordate (*Stereobalanus canadensis*), with a pattern more similar to that of *A. marina* observed in another polychaete, *Amphitrite johnstonii*. Quantitatively, the polychaetes showed higher levels of activity with rates in *A. marina* exceeding those of the hemichordate and holothurian by more than 10-fold.

KEY WORDS: Bacteriolysis · Deposit feeding · Sedimentary bacteria · *Arenicola marina* · *Amphitrite johnstonii* · *Caudina arenata* · *Stereobalanus canadensis*

INTRODUCTION

The concept of the 'microbial loop' (Azam et al. 1983) has altered long-held views of pelagic food webs and trophic transfer. Knowledge of the benthic microbial loop is more fragmentary and extrapolations from the pelagic system are dangerous. Several important differences from the water column are evident. In pelagic systems the inability of larger zooplankton, with the possible exception of tunicates (Pomeroy et al. 1984, Diebel & Powell 1987), to effectively graze bacteria is an important constraint on the structure of the microbial loop. Benthic systems differ in that potential bacterivores include metazoans, both meio- and macrofauna, in addition to protozoans (Kemp 1990). This difference is largely because most bacteria are attached to mineral or organic particles (e.g. Dye 1983, Ellery & Schleyer 1984). The ubiquity and abundance of deposit-feeding macrofauna, coupled with their surprisingly high rates of sediment ingestion (e.g. Taghon 1988, Wheatcroft et al. 1990), suggest an important role as grazers of bacteria.

Ingestion by deposit-feeding macrofauna has been shown to affect biomass (e.g. Moriarty et al. 1985, Plante et al. 1989), growth (Deming & Colwell 1982, Duchene et al. 1988, Plante et al. 1989), activity (Juniper 1981, Aller & Yingst 1985) and community composition (Dobbs & Guckert 1988, Duchene et al. 1988) of microbes in sedimentary and detrital environments. Both experiments and field observations indi-
cate that invertebrate ingestion can strongly influence population dynamics of ingested bacteria, either by cell destruction (e.g. Deming & Colwell 1982, Austin & Baker 1988, Plante et al. 1989) or by stimulation of growth (e.g. Reyes & Tiedje 1976, Klug & Kotarski 1980, Deming & Colwell 1982, Plante et al. 1989). The net result, i.e. population increase or decrease, of one or both of these processes varies among ingested microbial strains (Wavre & Brinkhurst 1971, Baker & Bradnam 1976, Reyes & Tiedje 1976, Dobbs & Guckt 1988, Duchene et al. 1988, Prieur et al. 1990). Certain bacterial strains are efficiently eliminated due to gut passage whereas others simultaneously increase in number (e.g. Wavre & Brinkhurst 1971).

The nature of the interactions between bacteria associated with sediments and detritus and the animals ingesting these microbes is certainly diverse, and generally unclear. A central question is to what extent these ingested microorganisms contribute to, or compete for, resources with the animal. This simple question, however, masks numerous complexities. Different microbial populations will interact with the host in dissimilar ways — gut passage can be either mutually beneficial or mutually detrimental, or somewhere between these extremes. Also, the nature of association, within a given population, is likely to vary spatially within the animal's gut. Additionally, the effects of gut passage can be manifested well after the parties are no longer physically associated, e.g. stimulated bacterial growth can be observed in feces (Newell 1965, Hargrave 1970, Fenchel 1972, Juniper 1981).

We have recently presented predictions of association type and bacterial distributions within the guts of detritivores (Plante et al. 1990). The models giving rise to these predictions were based on 'optimal digestion theory' (Penry & Jumars 1986, 1987, Dade et al. 1990). Their theory adapts reactor theory, a body of theory from chemical engineering aimed at the design of chemical reactors, to analyze the process of digestion and absorption. We (Plante et al. 1990) adapted and extended this theory in order to predict type and location of animal-microbe interaction within the guts of detritivores. A crude optimal digestion model for microbes in a gut-like environment was formulated; costs and benefits of potential associations in terms of digestive model variables were then analyzed. Predictions of association type, e.g. mutualism, competition, or parasitism, and distribution of both attached and transient microbes were thus made possible. The major cost and benefit to the microbe were concluded, respectively, to be risk of enzymatic digestion and local supply of the products of animal digestion. Briefly, our analysis suggests that the midgut should be kept free of both resident and transiting bacteria via mechanical and chemical means because microbial invasion and competition in this region of absorption would be costly to the animal. Once the 'war of absorption' is over, however, hindguts and feces of detritivores are excellent environments for microbial growth — numbers and productivity of both endosymbiotic and transient bacteria should be highest here. Foregut associations might be extremely beneficial to both parties but are unlikely to arise in small animals because of the short residence times of material in their foreguts. In general, our predictions regarding site and type of association are supported by previous observations, made without benefit of theoretical guidance. However, few experimental data to more rigorously test hypotheses are yet available.

According to this model a non-resident, transiting bacterium will experience at least 2 contrasting environments and be subject to 2 distinct 'processes', digestion and regrowth. We focus in this work on the process experienced first, digestive removal. Digestion influences the bacterial communities of detritus in at least 2 direct ways: biomass is reduced and the bacterial species composition of the ingesta is altered.

The 2 specific goals of the present study are to assess the sites and strength of bacteriolysis in deposit-feeder guts. In this initial effort we focus on animals with simple tubular guts; therefore our results are unlikely to be applicable to mollusks and crustaceans. We will test whether the 'cost' of digestion to bacteria is confined to the midgut as predicted (Plante et al. 1990). Assuming plug-flow transport of gut materials, we can also calculate the length of time bacteria are exposed to lytic agents. Time is a key variable both from the animal's perspective in that it affects both the nature and amount of food digested and from the microbe's perspective in that digestive removal of bacteria appears to be directly proportional to gut residence time (Plante & Jumars 1993). Lytic profiles may also give hints as to the source, as well as the function, of the lytic agent. Our study also compares lysis among 4 species of deposit feeders representing 3 phyla. Similarity of results will give some indication of the extent to which we can generalize to other taxa with similar feeding modes and gut morphologies.

**METHODS**

**Specimen collection and preparation.** *In vitro* and *in vivo* bacteriolytic assays were performed using gut extracts of the lugworm *Arenicola marina* (Polychaeta: Arenicolidae). The guts of the tentaculate polychaete *Amphitrite johnstoni* (Polychaeta: Terebellidae), a subsurface-feeding enteropneust, *Stereothenus canadensis* (Enteropneusta: Harrimanaidae), and the apodous *Caudina arenata* (Holothuroidea: Molpadiidae)
have also been profiled for bacteriolytic activity to represent different modes of deposit feeding and to allow cross-phyletic comparisons. *Arenicola marina* and *Amphitrite johnstoni* were dug by shovel at low tide from the poorly-sorted sands surrounding Pratt's Island, Maine, USA in September and November of 1992, respectively. *S. canadensis* and *C. arenata* were collected from subtidal (~20 m) muds of the Damariscotta River estuary in mid-coastal Maine, via Van Veen grab. *S. canadensis* and *C. arenata* were collected in January and August of 1993, respectively. All animals were either dissected on site or immediately upon return to the laboratory, within 5 h of collection. For dissection of *Arenicola marina*, the body wall was cut longitudinally to reveal an intact alimentary tract from pharynx to anus, although the rectum at times could not be cleanly separated from the body wall. Coelomic fluids were then rinsed out of the body cavity with seawater and the gut was blotted dry. The contents of the pair of large ceca were collected by suspending each into a 1.5 ml microtube, followed by puncture with small dissecting scissors. Other gut regions, the esophagus (foregut, FG), anterior stomach (midgut, MG), posterior stomach (MG), anterior intestine (MG), posterior intestine (MG), anterior rectum (hindgut, HG), and posterior rectum (HG), were opened and sampled with a small scoopula, taking care to avoid collection of animal tissue or fluids of the body cavity. These samples were placed in 1.5 ml polypropylene tubes and frozen until further analysis. In preliminary tests, no effects of freezing on lytic activity were observed. Total (fluid plus sediment), dry sediment, and fluid (total minus sediment) weights were later obtained for normalizations.

In general, the sampling of the other 4 animals followed the procedures used for *Arenicola marina*. The discrete gut sections to be sampled for the other deposit feeders were determined by visual morphological distinctions of the gut and were generally fewer than those for *A. marina* (Fig. 1). The small size and nature of the enteropneust guts made dissection more difficult and a small degree of contamination of samples by coelomic fluid was likely. In addition, much of the trunk of these animals is regularly lost upon collection so the most posterior portions of the animals could not be sampled.

Egestion rates were estimated for *Arenicola marina* at the time of collection. During ebb tide, when the worms were still under approximately 0.5 m of water, fecal castings were gently brushed away and marked with numbered flags. After emersion, just prior to animal collection, new casts were collected by spatula for determination of dry weight. Subsequently the dry weights of the gut contents of *A. marina* could be obtained to estimate gut 'capacity'. (Gut fullness might be a more accurate term than capacity since there is no requirement that the gut be completely filled). Preliminary tests revealed that egestion rates remained constant for only about 0.5 h after emersion, after which drying of the sediment inhibited feeding and defecation (see also Kermack 1955), so all feces and worms were collected in this 0.5 h window following emersion. Gut residence time was then estimated by dividing gut capacity (weight) by egestion rate (weight time-1).

**In vitro lysis assays.** A traditional lysozyme assay, which follows the lysis of *Micrococcus lysodeikticus* (formerly *M. luteus*) turbidimetrically (e.g. McHenery & Birkbeck 1982, Morsky 1983), was initially tried to quantify the bacteriolytic activity of gut juices. *M. lysodeikticus* was so resistant to lysis by gut juices, however, that turbidity loss could hardly be differenti-
lated from that of the controls or the noise of the technique. Upon subsequent testing, numerous strains isolated from local sediments showed susceptibility to lysis under the same conditions. One such isolate, designated SS-1, was chosen to be the standard assay bacterium. This bacterium, likely a member of the Pseudomonadaceae (Table 1), was isolated from surface sediments of the *Arenicola marina* collection site.

SS-1 was cultured in filtered Marine Broth 2216 (Difco) overnight at 25°C on a shaker table. Mid- to late-log phase cells were pelleted (3000 rpm, −1200 × g, 10 min), washed in 0.066 M phosphate buffer (potassium phosphate, monobasic, pH = 6.2), and repelleted and resuspended in the same phosphate buffer. Preliminary tests showed that use of phosphate or tris-HCl buffers to wash and resuspend SS-1 enhanced bacteriolysis relative to those cells resuspended in seawater. Preliminary tests also revealed that lytic susceptibility was a function of culture age, with stationary phase cultures exhibiting markedly reduced sensitivity (unpubl. data; see also Stolp & Starr 1965); our cultures were always pelleted at between 14 and 16 h. Resuspensions were diluted to give an absorbance of approximately 0.600 at 450 nm (Spectronic 2000, Bausch & Lomb). This absorbance (A450) represented a concentration of about 1 × 10⁸ SS-1 cells ml⁻¹.

After obtaining total (sediment + fluid) weights of samples, gut juices were extracted from gut sediment with 0.5 ml phosphate buffer, then centrifuged (7000 rpm, −3000 × g, 10 min). Small volumes (between 10 and 100 μl) of extracted gut juices were added to 1.5 ml SS-1 suspensions in disposable 1 cm polystyrene cuvette tubes. The spectrophotometer was zeroed against phosphate buffer; addition of equal volumes of phosphate buffer to the SS-1 suspensions served as controls. Lysis was followed by optical clearing, with readings taken at 0, 0.25, 0.50, 1, 2, 3, 5, 10, and 30 min after addition of gut fluid or buffer. Lytic rates were calculated using the initial linear portions of absorbance vs time plots. All lysis experiments were performed at 20 to 23°C. The pellet from each centrifuged extract was then dried and weighed. Sample wet (i.e. fluid only) weights were obtained by subtraction so that values could be normalized to both dry and wet weight. Both normalizations were performed because, *a priori*, we were unsure whether lytic activity might be correlated with amount of ingesta. Samples of wet or dry weight below 7.5 mg were not used. Obtaining a signal (i.e. decrease in absorbance) in excess of the phosphate control was difficult below this sample weight on the other hand, a signal slightly above the control (but within the precision of the instrument) could result in anomalously high lytic activity after normalization.

From a small number of the total pool of gut sections analyzed, subsamples were fixed in 2% glutaraldehyde for microscopic enumeration. Direct counts were performed using epifluorescence microscopy. Samples were DAPI stained (20 ppm) for 6 min and viewed at 1000× using a Zeiss Axioskop.

Resuspension in phosphate buffer appeared to create artificial osmotic challenges for SS-1 which enhanced lysis relative to resuspension in seawater. On the other hand, the procedure outlined above undoubtedly presents an environment more dilute in gut fluids than would be experienced by sedimentary bacteria transiting the gut of *Arenicola marina*. To test whether SS-1 would be lysed under more realistic conditions, we washed and resuspended the bacterium in filtered seawater and added 30 μl undiluted (no phosphate buffer extraction) fluids from the ceca of *A. marina* to 120 μl of resuspended cells. The added gut fluids were filtered with a 0.2 μm micro-spin centrifuge unit (Lida, Kenosha, WI, USA) prior to addition to avoid addition of indigenous gut flora. Subsamples were taken at 0, 5, and 60 min for both viable and total counts. Serial dilutions were plated in duplicate on Marine Agar 2216 plates for viable counts, direct

<table>
<thead>
<tr>
<th>Characteristic Tested</th>
<th>Result</th>
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<tbody>
<tr>
<td>Gram stain</td>
<td>−</td>
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<tr>
<td>Morphology</td>
<td>Rods</td>
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<tr>
<td>O/F*</td>
<td>O</td>
</tr>
<tr>
<td>Anaerobic growth</td>
<td>−</td>
</tr>
<tr>
<td>Motility</td>
<td>+</td>
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<tr>
<td>Oxidase</td>
<td>+</td>
</tr>
<tr>
<td>Catalase</td>
<td>−</td>
</tr>
<tr>
<td>Production of indole</td>
<td>−</td>
</tr>
<tr>
<td>Nitrate reduction</td>
<td>−</td>
</tr>
<tr>
<td>Tryptophanase</td>
<td>−</td>
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<tr>
<td>Arginine dehydrogenase</td>
<td>−</td>
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<tr>
<td>Urease</td>
<td>−</td>
</tr>
<tr>
<td>Esculine hydrolysis</td>
<td>+</td>
</tr>
<tr>
<td>Gelatinase</td>
<td>+</td>
</tr>
<tr>
<td>Beta-galactosidase</td>
<td>−</td>
</tr>
<tr>
<td>Sodium required for growth</td>
<td>+</td>
</tr>
<tr>
<td>Utilization of D-glucose</td>
<td>−</td>
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<tr>
<td>Utilization of L-arabinose</td>
<td>−</td>
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<tr>
<td>Utilization of D-mannose</td>
<td>−</td>
</tr>
<tr>
<td>Utilization of N-acetyl-D-glucosamine</td>
<td>−</td>
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<tr>
<td>Utilization of maltose</td>
<td>−</td>
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<tr>
<td>Utilization of D-glucuronate</td>
<td>−</td>
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<tr>
<td>Utilization of caprate</td>
<td>−</td>
</tr>
<tr>
<td>Utilization of adipate</td>
<td>−</td>
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<tr>
<td>Utilization of L-malate</td>
<td>−</td>
</tr>
<tr>
<td>Utilization of citrate</td>
<td>−</td>
</tr>
<tr>
<td>Utilization of phenyl-acetate</td>
<td>−</td>
</tr>
</tbody>
</table>

*Oxidation/fermentation tests were done in MOF medium (Leifson 1963).*
counts of total bacteria were performed as described above. These experiments were performed in triplicate with seawater additions serving as controls.

**Lysis of sedimentary bacteria.** The digestive susceptibility of cultured bacteria in logarithmic growth may be much greater than that of natural sedimentary bacteria, most of which are attached to inorganic particles and surrounded by protective organic matrices. To assess whether extracted gut fluids would act on natural sedimentary bacteria, we added 100 µl of a 50% solution (in filtered seawater) of midgut (anterior stomach) fluid to 5 samples of approximately 0.2 g (wet wt) of surface sediments (top 3 mm from the Pratt’s Island site), vortexed, then removed subsamples at 5 and 60 min for viable counts. Sediment samples with equivalent volumes of seawater or hindgut fluid served as controls. To conserve gut fluids, log values were obtained with a separate treatment, with 5 separate surface sediment samples plus 100 µl added seawater, rather than sampling each of the above 3 treatments (MG, HG, and seawater additions) at logt. Colonies were counted after a 48 h incubation at room temperature (22 to 25°C) on 2216-agar plates. At logt, the treatments were fixed for total direct counts. Prior to filtration and DAPI staining, samples for epifluorescence enumeration were incubated for ≥30 min with 0.01 M sodium pyrophosphate, then sonicated at 60 W for 30 s (3 × 10 s). Finally, dry weights of each sediment sample were obtained for normalization to g dry wt⁻¹.

**Statistics.** One- or two-way ANOVAs were used to test for overall differences in means in all experiments performed. Generally, the Tukey HSD test was employed to make pairwise comparisons of means within the ANOVAs. Prior to performing these parametric statistical procedures, the assumption of homoscedasticity was checked using Bartlett’s test, and data were log-transformed if necessary. All statistical tests were performed using SYSTAT (version 5.2.1) for the Macintosh computer.

**RESULTS**

*In vitro* bacteriolysis

In each lugworm assayed, the rate of lysis of bacteria varied among gut sections but consistently decreased with time as an end-point was approached (Fig. 2); the speed at which this end-point was reached was proportional to overall lytic strength, suggesting substrate limitation in the experiments with the most active extracts. Lytic activity in the gut of *Arenicola marina* differed significantly among the various gut regions (1-way ANOVA, p < 0.001) (Fig. 3). Tukey’s multiple comparison test clearly showed that the gut extracts separated into 2 distinct groups: the anterior and posterior stomach extracts, as well as those from the ceca, resembled each other (p >> 0.10) but differed from the phosphate buffer control and all other gut samples (p < 0.001). Activity posterior to the stomach decreased slightly with distance but the levels were statistically indistinguishable from one another (p >> 0.10). In most animals, foregut activity was not observed; however, in a few specimens weak activity was noted. Activities of
surface sediment, fecal, and coelomic fluid extracts could not be distinguished from controls (ANOVA, p >> 0.10). A slight amount of optical clearing was observed with the phosphate buffer alone (Fig. 2).

The mean gut residence time of *Arenicola marina* in September was 1.09 ± 0.67 h (n = 10). On average, the length of the lytic portion of the gut (stomach) was 30% (± 4%, n = 8) of the total gut length. Assuming plug-flow, ingested bacteria therefore spent approximately 20 min in the lytic portion of the gut.

Direct counts showed that cell lysis was coincident with optical clearing. Significant reductions in number were observed relative to phosphate buffer controls (2-way ANOVA, p < 0.001) and with respect to time (p = 0.002). Qualitative changes in the staining intensity and cell integrity of SS-1 upon exposure to midgut fluids made counts more subjective with time (Fig. 4).

Addition of undiluted cecal fluid to SS-1 resuspended in seawater resulted in a large and rapid loss of cells. A significant treatment effect was observed (2-way ANOVA, p < 0.001). Time was also significant for the gut-fluid treatment (p < 0.001) but not for the controls (p >> 0.10). Cell number was reduced by 93% and 96% by 5 and 60 min, respectively, as compared with \( t_0 \). The reduction from \( t_5 \) to \( t_{60} \) was not significant (p >> 0.10). Qualitatively, cell integrity and staining intensity decreased with time in a fashion similar to that noted for phosphate-buffer-resuspended SS-1 cells (see Fig. 4). No loss was observed over 5 or 60 min in the controls (p >> 0.10 for both times).

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Fig. 4. Epifluorescence staining of SS-1 resuspensions (A) prior to addition of gut fluids, and (B) 5 min after midgut-fluid incubation 1000×
The decline in colony-forming units (CFUs) was even more dramatic with addition of cecal juice; numbers at both 5 and 60 min were at least 4 orders of magnitude lower than either initial numbers or controls (2-way ANOVA; \( p < 0.001 \)). Controls showed no significant numerical changes through time (\( p > 0.10 \)).

**Lysis of sedimentary bacteria**

A highly significant overall treatment effect (1-way ANOVA, \( p < 0.001 \)) was seen among bacterial numbers in sediments incubated for 60 min in seawater, midgut and hindgut fluids, and initial numbers \( t_0 \) treatment). The addition of midgut fluids to surface sediments resulted in a marginally significant (Tukey’s test, \( p = 0.063 \); Fisher’s LSD, \( p = 0.014 \)) decrease in total bacterial numbers over a 60 min incubation (Fig. 5a), although the decrease of 45% is much smaller than the reduction in numbers of the pure culture of SS-1 observed with added gut fluids. The controls, surface sediments incubated with seawater or hindgut fluids, did not exhibit significant numerical changes (Fig. 5a; \( p > 0.10 \)).

Similar trends were obtained for viable counts on Marine Agar 2216 plates, with significant numerical losses (\( p < 0.001 \)) observed with midgut fluid exposure but not in controls (Fig. 5b). The reduction in viability (or ability to culture on 2216 agar medium) was proportionally much higher, at >99%, than was the reduction in total cell counts (at 45%). The high variances in the hindgut-fluid incubations are due to anomalously high CFUs in 1 replicate (with high counts at both \( t_0 \) and \( t_{60} \)); if these values are deleted, both means and variance are nearly identical to those for the seawater controls.

**Cross-phyletic comparisons**

Quantitatively, the gut fluids among the 4 deposit feeders differed most obviously in fluid content and color. Arenicola marina represented one extreme; the midgut of this deposit feeder was more fluid-rich than the ingested sediments or other parts of the lugworm gut. Additionally, by far the greatest relative increase in fluid content was observed in A. marina as compared to the other animals studied (Table 2). Yellow color and an apparent soapiness not found in ingested fluids indicated that this excess fluid was secreted by the animal. Caudina arenata and Stereobalanus canadensis demonstrated the other extreme in that gut contents visually resembled ingested sediments, with no obvious soapiness, color change or changes in fluid content. Amphitrite johnstoni was more like A. marina but the evidence of secretory additions was somewhat less obvious.

The lytic activity in Amphitrite johnstoni was lower than that observed in Arenicola marina by roughly 50 to 70%; it was, however, more evenly distributed through the gut. Although activity was observed in both Stereobalanus canadensis and Caudina arenata gut sections, neither of the other deposit feeders exhibited activities comparable to that of the polychaetes (Table 2). Again, activity appeared more evenly distributed than in A. marina but some concentration in the middle and posterior regions was apparent.

**DISCUSSION**

We normalized our lytic activities both to fluid weight and to weight of dry sediment. Our observations convinced us that bacteriolytic activity can be found in portions of the gut devoid of particulate mat-
Table 2. Lytic activities [mean ± 1 SD (n)] normalized to fluid and dry sediment weights of ingesta, along with fluid content (%) for each gut region of 4 deposit feeders

<table>
<thead>
<tr>
<th>Gut section</th>
<th>Lytic activity</th>
<th>Wet wt. (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ΔA₄₅₀ min⁻¹ wet g⁻¹</td>
<td>ΔA₄₅₀ min⁻¹ dry g⁻¹</td>
</tr>
<tr>
<td><strong>Arenicola marina</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Esophagus</td>
<td>0.4 ± 0.9 (5)</td>
<td>0.8 ± 1.2 (4)</td>
</tr>
<tr>
<td>Ceca</td>
<td>22.9 ± 16.8 (5)</td>
<td></td>
</tr>
<tr>
<td>Anterior stomach</td>
<td>42.7 ± 31.2 (6)</td>
<td>44.0 ± 25.6 (6)</td>
</tr>
<tr>
<td>Posterior stomach</td>
<td>29.3 ± 28.4 (8)</td>
<td>14.1 ± 5.8 (7)</td>
</tr>
<tr>
<td>Anterior intestine</td>
<td>0.7 ± 1.0 (6)</td>
<td>0.2 ± 0.3 (7)</td>
</tr>
<tr>
<td>Posterior intestine</td>
<td>0.1 ± 0.2 (8)</td>
<td>0.4 ± 0.8 (9)</td>
</tr>
<tr>
<td>Anterior rectum</td>
<td>0.2 ± 0.2 (9)</td>
<td>0.1 ± 0.1 (9)</td>
</tr>
<tr>
<td>Posterior rectum</td>
<td>0.1 ± 0.2 (8)</td>
<td>0.1 ± 0.1 (7)</td>
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<tr>
<td><strong>Amphitrite johnstoni</strong></td>
<td></td>
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<tr>
<td>Esophagus</td>
<td>1.6 ± 1.3 (3)</td>
<td>0.3 ± 0.5 (2)</td>
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<tr>
<td>Anterior stomach</td>
<td>8.2 ± 3.4 (5)</td>
<td>33.4 ± 16.4 (4)</td>
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<tr>
<td>Posterior stomach</td>
<td>10.9 ± 1.1 (5)</td>
<td>29.8 ± 12.7 (5)</td>
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<tr>
<td>Anterior intestine</td>
<td>11.6 ± 3.0 (5)</td>
<td>21.8 ± 6.2 (4)</td>
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<tr>
<td>Posterior intestine</td>
<td>6.0 ± 2.0 (2)</td>
<td>11.7 ± 0.5 (2)</td>
</tr>
<tr>
<td><strong>Caudina arenata</strong></td>
<td></td>
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<tr>
<td>Esophagus</td>
<td>0.25 ± 0.44 (5)</td>
<td>0.60 ± 1.20 (5)</td>
</tr>
<tr>
<td>Anterior intestine-1</td>
<td>3.78 ± 6.44 (5)</td>
<td>30.67 ± 52.82 (3)</td>
</tr>
<tr>
<td>Anterior intestine-2</td>
<td>1.07 ± 2.03 (4)</td>
<td>2.10 ± 3.90 (4)</td>
</tr>
<tr>
<td>Posterior intestine</td>
<td>0.57 ± 0.78 (5)</td>
<td>1.30 ± 1.80 (5)</td>
</tr>
<tr>
<td>Cloaca</td>
<td>0.15 ± 0.18 (5)</td>
<td>0.50 ± 0.60 (5)</td>
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<tr>
<td><strong>Stereobalanus canadensis</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anterior intestine-1</td>
<td>0.30 ± 0.51 (3)</td>
<td>1.59 ± 2.75 (3)</td>
</tr>
<tr>
<td>Anterior intestine-2</td>
<td>3.69 ± 6.78 (5)</td>
<td>19.34 ± 36.93 (5)</td>
</tr>
<tr>
<td>Anterior intestine-3</td>
<td>3.19 ± 3.20 (5)</td>
<td>13.03 ± 13.24 (5)</td>
</tr>
<tr>
<td>Posterior intestine</td>
<td>3.71 ± 2.47 (5)</td>
<td>17.50 ± 14.69 (5)</td>
</tr>
</tbody>
</table>

We therefore regard as more significant the lytic values normalized to fluid weight, although trends for both are similar. Fluid-weight normalization gives an idea of the concentration of lytic agents in the fluids, a mixture of animal secretions and sediment porewater, to which ingested bacteria are exposed. Within a species, lytic activity was not correlated with amount of ingesta; gut sections nearly devoid of solid ingesta demonstrated lytic capabilities comparable to those regions filled with sediment. A high value of lytic activity normalized to dry weight, then, is more an indication of an empty gut than lytic strength. Dry-weight normalizations do, however, reveal the extent to which ingesta are bathed by digestive secretions in animals like Arenicola marina, in which changes in fluid content are obvious.

Bacteriolytic activity, expressed in terms of either dry sediment or fluid weight, in the gut of Arenicola marina is concentrated in the midgut. This observation supports our predictions based on cost-benefit models (Plante et al. 1990), and agrees with observed longitudinal patterns of bacterial numbers in various deposit feeder guts (e.g. Deming & Colwell 1982, Duchene et al. 1988, Plante et al. 1989). Digestive enzymes in A. marina are similarly distributed with proteases, carbohydrases and esterase/lipases largely confined to the ceca and stomach (Longbottom 1970, Mayer unpubl.). The ceca are indicated as the source of lytic agent as their contents exhibit strong lytic activity and visually resemble stomach contents in color and soapiness. That the stomach showed a still higher (although not significantly so) lytic capacity may implicate the stomach as an additional source of lytic agents.

New insights regarding the mechanisms of bacteriolyis are revealed in the present study. Filtered gut fluids added to sediments caused lysis, as did the supernatants of centrifuged extracts, suggesting that particulates are unimportant and intracellular digestion (e.g. by amoebocytes) of bacteria in the gut lumen is unlikely. Preliminary observations suggest that the lytic agent is not lysozyme as anticipated and likely is not an enzyme at all. Contrary to the results of Grossman & Reichardt (1991), who used whole Arenicola marina guts extracted in Tris-HCl buffer versus the phosphate-buffer-extracted gut contents employed in this study, we observed very little lysis of the Gram-
positive bacterium *Micrococcus lysodeikticus*, but rapid lysis of the Gram-negative SS-1. Further tests with 3 more Gram-negative and 2 more Gram-positive isolates resulted in similar trends (Plante unpubl.). Because enzymes often are associated with the cells of the gut wall (e.g. Feral 1989), the disparate results of Grossman & Reichardt (1991) and those of the present study may not be surprising. We made numerous efforts to replicate their results using whole-gut homogenates in Tris-HCl buffer, however, and were unable to obtain appreciable lysis of *M. lysodeikticus*. The complete explanation for this discrepancy is unclear at this time; closer examination of the data, however, reveals that our results are less contradictory than first appearances would suggest. Although not reported in their manuscript, their initial bacterial densities were approximately $3 \times 10^6$ ml$^{-1}$ (S. Grossman pers. comm.); these data combined with their rate data (their Fig. 6) reveal that the lytic rates that they observed for *M. lysodeikticus* (maximal values of $\sim 3\%$ min$^{-1}$) were much lower than what we observed for SS-1 (maximal values $> 70\%$ min$^{-1}$) and more comparable to our controls. The difference, then, may be that they were able to achieve non-lytic controls whereas our phosphate buffer or Tris-HCl controls always showed some small amount of lysis, roughly equal to what we observed in *M. lysodeikticus* resuspensions with added gut fluids. *M. lysodeikticus* is highly susceptible to lysozyme (and is used in standardized lysozyme assays) as are many Gram-positive bacteria, whereas Gram-negative bacteria are typically much more resistant. Additionally, preliminary experiments show that ca 90% of the lytic rate of SS-1 exposed to gut fluids is retained after these fluids are microwaved or boiled, indicating a non-enzymatic source of lysis, and further investigations have revealed high levels of surfactants in the gut which appear to correlate with lytic activity (Plante unpubl.). A surfactant lyasing agent is also consistent with the observed greater susceptibility of Gram-negative bacteria (e.g. Varra 1992, Johnson 1994). Although certain products of digestion, e.g. various fatty acids and peptides, can exhibit surfactant activity, the observation of bacteriolysis in the particle-free (Kermack 1955, Plante pers. obs.), and presumably food-free, ceca points to an actively secreted lytic agent.

Little activity was measured in the esophagus, indicating an absence of axial mixing of digestive secretions. In most cases bacteriolytic activity was much reduced or even negligible in all other portions of the gut, surface sediments, and fecal coils relative to the midgut. This observation allows the calculation that exposure time to the lytic portion of the gut on the dates of collection is approximately 20 min; this estimate is fairly rough, however, since relative lengths of gut sections were used as surrogate measures for gut volume. The diameter of the alimentary tract of *Arenicola marina* clearly is not constant from mouth to anus (e.g. Fig. 1); the stomach is generally wider but so too is the stomach less tightly packed with ingested materials and more filled with secretory fluids. In addition, our method of estimating mean gut residence time assumes that the weight of sediment in the gut upon dissection is similar to gut-sediment weight at defecation. The calculations of Kermack (1955), in which the weight of sediment in one defecation event exceeded the weight of sediment in the entire gut of *A. marina*, illustrate that a gross misrepresentation of residence time will result if gut fullness does vary. Given our observations that egestion rate remained constant up to 30 min after emersion, we are confident that assumptions regarding gut fullness were met.

The disappearance or inactivation of the lytic agent posterior to the stomach cannot be explained with conviction at this time. The apparent removal of fluids in posterior regions of the gut, especially apparent in *Arenicola marina*, however, may suggest the recycling of digestive secretions. Given the low quality of food ingested by deposit feeders, a strategy for efficient recycling seems especially appropriate, particularly in those animals which secrete relatively large quantities of digestive agents and exhibit short gut residence times. One function of the peritrophic membrane, found in the Terebellidae and other polychaetes (Dales & Pell 1970), may be to enable absorption and redistribution of secretions, potentially by a mechanism similar to that demonstrated in dipteran larvae (Espinoza-Fuentes & Terra 1987). One alternate hypothesis is that a countercurrent flow retains fluids within the midgut, resulting in a longer midgut residence time for liquids relative to the sediment plug. Such a scenario runs counter to predictions based on optimal digestion theory that deposit feeders should function as plug-flow reactors with negligible axial mixing of reactants (Penry & Jumars 1987). Plug-flow of sediment particles for some deposit feeders has been demonstrated (Penry 1989 Plante unpubl.); the absence of axial mixing in the liquid phase, however, remains undocumented.

Our experiments in which undiluted gut fluids were mixed with bacterial resuspensions in seawater demonstrate that gut fluids near their natural dilutions are sufficient to lyse bacterial cells under normal osmotic conditions. Although the addition of gut fluids enhanced the lysis of SS-1 over the lysis due to potassium phosphate buffer, resuspension in phosphate buffer alone caused some lysis and stimulated lysis in the presence of gut fluids relative to a seawater resuspension. Resuspension in phosphate buffer increased the sensitivity of the assay procedure, which proved...
useful given the small amounts of fluids encountered in some gut sections coupled with the dilutions necessary to extract these fluids from ingested sediments. The probable mechanism of enhanced lysis in the presence of phosphate buffer is osmotic disruption (Stolp & Starr 1965), although interference with stabilizing divalent cations (Ca$^{2+}$ and Mg$^{2+}$) on the lipopolysaccharide-covered cell surfaces (Varra 1992) could also contribute to greater lytic susceptibility. These same experiments demonstrated with parallel viable and total counts that cell viability (or cultivability) was lost prior to lysis.

Experiments in which fresh surface sediments were incubated with gut fluids of Arenicola marina showed that, although in situ bacteria were lysed by midgut fluids, the efficiency of lysis was substantially lower than that for a cultured single strain. Prominent explanations among many possibilities for this disparity are exopolymer secretions, varying growth stages, and differential susceptibility among in situ bacteria. Exopolymer secretions have been shown to inhibit enzymatic digestion of microbes, although mechanisms for this protection are still uncertain (Decho 1990). Preliminary portions of the present study offer some evidence for the latter 2 suggestions: stationary-phase cells were shown to be less susceptible to lysis and marked differences to susceptibility were observed between Micrococcus lysodeikticus and SS-1. That log-phase bacteria are more susceptible to digestion than stationary-phase cells may have implications for the concept of gardening (Hylleberg 1975); macrofaunal activity may not only stimulate microbial biomass production, but also simultaneously increase the microbes' susceptibility to digestion.

The experiments in which sediments were incubated with gut fluids also corroborated the idea that cell inactivation is more widespread than cell lysis, with viable counts decreasing much more than total counts. Although we suspect that bacteria in these experiments were killed, we conservatively use the term 'inactivation' to acknowledge the possibility that only cultivability was lost prior to lysis. These same experiments demonstrated with parallel viable and total counts that cell viability (or cultivability) was lost prior to lysis.

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