Direct measurements of sugar uptake from seawater into molluscan larvae

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ABSTRACT: A method for the direct measurement of picomole amounts of mono-, di-, and trisaccharides in seawater was developed by combining high-performance liquid chromatography (HPLC) with amperometric detection. This new method is described and then applied to the study of sugar uptake from seawater into veliger larvae of a bivalve (Crassostrea gigas) and a gastropod (Haliotis rufescens). Bivalve veligers take up glucose, maltose, cellobiose, and cellotriose but not L-rhamnose, isomaltose, maltotriose, or isomaltotriose. Similarly, gastropod veligers take up glucose and maltose, but not L-rhamnose or maltotriose. Glucose and maltose are transported into the bivalve veligers by separate pathways and the influx of 14C-labeled glucose is a good measure of the net flux (measured by HPLC). The uptake of complex sugars, in addition to monosaccharides, permits molluscan larvae to utilize a greater part of the dissolved organic material in the sea as a source of nutrition.

INTRODUCTION

Most studies on the transport of dissolved organic matter (DOM) from seawater by soft-bodied marine invertebrates have focused primarily upon the flux of free amino acids (e.g. Stephens & Schinske 1961, Ferguson 1967, Davis & Stephens 1984, Manahan 1989). There is also a smaller body of literature that shows invertebrates can transport other classes of compounds such as sugars (e.g. Stephens 1962, Ahearn & Gomme 1975, Pajor et al. 1989), fatty acids (Testerman 1982), acetate (Holst & Zebe 1984), and nucleosides (Schneider & Whitten 1987). About 10% of the constituents in the total DOM pool have been chemically identified (Williams 1975); of this fraction only a small percentage can be quantified due to limitations in analytical chemistry. Thus, it is necessary to develop analytical methods that permit the measurement of other classes of compounds in seawater. Such advances are needed to better understand the role of DOM in the nutrition of marine invertebrates.

Two experimental approaches have been used to determine if organic compounds can be transported by marine invertebrates from seawater. The first uses radiolabeled substrates to measure either the increase in radioactivity within the organism, or the decrease of radioactivity (influx) in the medium (Stephens 1962). This approach is sensitive and relatively easy to perform but gives no information on net flux of the compound under study (Johannes et al. 1969, Manahan 1989). The second approach measures the compound of interest in seawater (Manahan et al. 1982) and monitors the subsequent changes in concentration with time (net flux). This approach has the advantage of yielding information on net flux, but is dependent upon advances in analytical chemistry. Both of these approaches have been successfully used in studies of amino acid transport, but only the former has been extensively employed for studies of sugar transport (Stephens 1962, Pajor et al. 1989).

The direct quantification of sugars in seawater by high-performance liquid chromatography (HPLC) has been confounded by analytical problems. Conventional detection methods for sugars, such as refractive index or ultraviolet detection, lack the sensitivity to measure these compounds in natural seawater at submicromolar concentrations. The low concentration of sugars in seawater has forced investigators to concentrate their samples either by rotary evaporation or air drying (Mopper et al. 1980, Ittekkot et al. 1981). However, quantification is difficult with all of these methods because pretreatment procedures can lead to unpredictable losses of material (Dawson & Mopper 1978). In the present study, we describe a series of modifications to an analytical procedure (cf. Watanabe 1985) that permits the quantification of individual sugars at sub-
micromolar concentrations in seawater. No desalting or concentrating of samples is required. In this paper we use this method to study sugar uptake by molluscan larvae.

**MATERIALS AND METHODS**

**Reagents and sugar standards.** Pure water for HPLC analysis was made by distilling deionized water and then removing organic material by intense UV oxidation (Organicpure, Sybron & Barnstead). HPLC-grade methanol and acetonitrile were purchased from Baxter Healthcare Corp. (Burdick & Jackson division). All chemicals and standards were purchased from either Sigma Chemical Co. or Spectrum Chemical Co.

**HPLC quantification of individual sugars in seawater.** For the analysis of sugars, a 500 µl sample of seawater was taken and gently passed through a 0.2 µm (pore size) filter housed in a 13 mm diameter 'Pop top' (Nuclepore) holder. A 400 µl sample of the filtrate was taken and 100 µl of water was added. To this mixture was added 500 µl of acetonitrile. This dilution factor of 2.5 was empirically determined to give the best peak symmetry while avoiding salt precipitation. A 100 µl aliquot of this final solution was injected onto the HPLC column. Thus this injection volume contained 40 µl of the original seawater sample. Sugars were separated isocratically using a mobile phase of 45% aqueous solvent and 55% acetonitrile. The aqueous solvent was a solution of 2.7% NaCl (w/v). The addition of salt to the mobile phase was necessary to match the ionic concentration of the mobile phase to that of the injected seawater sample. Failure to do so resulted in a large solvent-front peak that masked most of the sugar peaks. A Waters Model 5000A pump and a Rainin aminopropyl-bonded silica (particle size 5 µm) column (250 mm by 4.6 mm), with a guard column of the same packing, were used for the separation of the sugars. The sample injection valve was a Rheodyne Model 7010, fitted with a 100 µl sample loop. The post-column reagent pump was a Waters Model M45 equipped with a SSI (Scientific Systems) Model LP-21 diaphragm pulse dampener. The pulse dampener was connected to a stainless steel mixing tee with 6 m of stainless steel tubing of 1/16" (1.6 mm) outside diameter and 0.005" (0.13 mm) internal diameter. The combination of the pulse dampener and the small diameter of the stainless steel tubing provided the necessary back pressure to reduce pulsations from the reagent pump.

Sugars were detected with a post-column reaction technique (Watanabe & Inoue 1983) involving an electrochemical mediator, copper (II) bisphenanthroline (CBP). Phenanthroline (570.8 mg, Sigma Chemical Co.) was dissolved in 5 ml of methanol that was then added to a warm aqueous solution of copper acetate (191.7 mg in 300 ml water). This solution contained a 3:1 molar ratio of phenanthroline to copper. The excess phenanthroline ensured that all of the copper was complexed. The resulting solution was stirred on a hot plate until all the methanol had evaporated (ca 2 h). The CBP complex was then diluted to 960 ml with water to give a final concentration of 1 mM CBP, empirically determined to maximize sensitivity. Dibasic sodium phosphate (12.27 g, Spectrum Chemical) was added to the CBP solution and the pH was adjusted to 11.6 using a concentrated NaOH solution. The CBP reagent was filtered through a polycarbonate filter. Because acetonitrile partially dissolves polycarbonate, the mobile phase was filtered through a polyethylene filter. Both filter types were 0.2 µm pore size Nuclepore. The CBP reagent and the mobile phase were degassed (LKB Model 2156 Solvent Conditioner) with helium for at least 1 h immediately prior to use, and were kept under pressurized helium during all analyses.

Individual sugars were separated on the column and the CBP reagent was added by a mixing tee (post-column). A 2:1 ratio of eluent flow (0.8 ml min⁻¹) to reagent flow (0.4 ml min⁻¹) gave the best sensitivity and peak symmetry. Lowering this ratio resulted in an excessive dilution of the separated peaks by the post-column reagent, resulting in lower sensitivity and wider peaks. Increasing this ratio also resulted in decreased sensitivity, presumably because not enough CBP was present to react with the reducing sugars. The combined solutions were then pumped through an in-line teflon reaction coil of 6.2 m length [1/16" (1.6 mm)] outside diameter by 0.01" (0.25 mm) internal diameter. The entire length of the reaction coil was knitted through 1/16" holes in an aluminum board. Interweaving the reaction coil through an aluminum board helped to maintain peak symmetry. The reaction coil was immersed in a water bath maintained at 98°C ± 0.02 (Precision Model RDL 20), the maximum temperature that could be used without significant baseline shifts. A degassing period of at least 1 h was needed to avoid the formation of gas bubbles in the reaction coil at 98°C. Also, it was important that the length of the reaction coil be at least 6.2 m to maintain high back pressure needed to keep the eluent in solution at the high temperatures of the reaction. The sugars in the eluent, under the conditions of high temperature (98°C) and high pH (11.6), reduced the copper in the CBP complex from Cu²⁺ to Cu⁺. The resulting zones of reduced CBP in the eluent stream were then reoxidized as they passed across the working electrode (glassy carbon) in the cell of the electrochemical detector (Model LC-4B, Bioanalytical Systems). The low applied potential (40 mV) across the
working electrode ensured that only the CBP was oxidized because organic materials require much higher potentials for oxidation. The resulting current from the reaction at the electrode was converted to voltage by the detector and recorded with a computing integrator. Individual sugars were identified by retention times compared to known standards. Quantification was performed by integrating peak areas using a Waters Model 740 Data Module.

**Larvae.** Veliger larvae of the bivalve *Crassostrea gigas* were supplied by Coast Oyster Co. (Quilcene, WA, USA). Veliger larvae of the gastropod *Haliotis rufescens* were provided by the Ab Lab (Port Hueneme, CA, USA). Experiments with bivalve veligers were conducted at 20°C; experiments with gastropod veligers at 17°C.

**Net uptake of sugars from seawater.** Rates of net flux were determined by measuring the disappearance of substrates from seawater using HPLC. All seawater used for experiments was first passed through a 0.2 μm (pore size) polycarbonate filter and then autoclaved. Immediately prior to an experiment, all larvae were collected on a mesh screen, washed with sterile seawater, and then resuspended in a graduated cylinder. To determine the concentration of larvae, replicate aliquots were taken and the larvae were counted. Preliminary experiments revealed that concentrations of larvae ranging from 800 to 2800 ml⁻¹ were needed to produce a rate of sugar uptake that was measurable by HPLC (see figure legends for exact numbers). A mixture of sugars was added to a 50 ml flask (experiments shown in Fig. 2) or a 20 ml vial (experiments shown in Figs. 3 and 4) at the start of the experiment to give a final substrate concentration of ca 1 μM for each sugar. Samples of the seawater (500 μl) were taken at various time intervals for HPLC analysis. Each sample was gently passed through a 0.2 μm (pore size) filter and frozen at −70°C for later analysis. No differences were observed between sugar standards which had been filtered then frozen at −70°C for 4 mo and freshly prepared standards. Nonetheless, all analyses were completed within 14 d of taking the samples.

**Net flux and influx of glucose.** Isotopic techniques and HPLC analyses were combined to compare the net flux and influx of glucose into veliger larvae of *Crassostrea gigas*. Larvae were exposed to a concentration of 900 nM glucose which was comprised of 14C-glucose (353 μCi μmol⁻¹, Sigma Chemical Co.) and unlabeled glucose. A series of 500 μl samples was taken during a time course experiment, of which 400 μl was used for HPLC analysis and 20 μl for isotope counting. The sample of the filtrate used to determine the rate of depletion of 14C-glucose (influx) was placed in a scintillation vial with 50 μl of concentrated HCl. The acid converts all dissolved bicarbonate to CO2, thus removing any 14CO2. After 24 h, 5 ml of scintillation cocktail (Scintiverse, Fisher Scientific Co.) were added to each vial. The amount of radioactivity in each vial was determined by liquid scintillation counting (LKB model 1211). The decrease in radioactive in the seawater was used to determine glucose influx. The decrease in the total concentration of glucose (HPLC analyses) was used to determine net flux.

**Effect of maltose on the transport rate of 14C-glucose by larvae.** Veliger larvae of *Crassostrea gigas* were collected, washed, resuspended, and counted as described above. The rate of transport of 14C-glucose by larvae from a 1 μM solution was determined in the absence and presence of 200 μM maltose. A known number of larvae (636 ml⁻¹) was added to each of 3 vials containing 2 pCi of 14C-glucose and unlabeled glucose (10 ml of seawater, final glucose concentration was 1 μM). Identical experiments were set up in an additional 3 vials, except that maltose was also added (i.e. glucose: 1 μM, maltose: 200 μM). A time course experiment (duration 8 to 10 min) was performed with each vial during which 500 μl samples of the seawater containing larvae were taken at 1 min intervals. The larvae were separated from the seawater by centrifugation through silicone oil, and the radioactivity per larva was determined using procedures described elsewhere (Manahan 1989). The amount of radioactivity per larva was corrected for the change in specific activity due to the addition of unlabeled glucose and the transport rate was calculated as fmol glucose larva⁻¹ h⁻¹.

**RESULTS**

Fig. 1A shows an original chromatogram of the separation of 5 sugars added to natural seawater, each present at 1 μM (40 pmol injected). The sugars in seawater were measured directly with no pretreatment, other than matching the injected sample to the mobile phase as described above. The 5 sugars were separated and quantified in ca 3 min. The order of elution is L-rhamnose (monosaccharide), glucose (monosaccharide), maltose (disaccharide), melibiose (disaccharide), and maltotriose (trisaccharide). Sugars are separated by normal-phase chromatography and the retention times are related to the number and position of the hydroxyl groups. The detector response was directly proportional to the amount of glucose injected (Fig. 1B).

**Net uptake of sugars by larvae**

Fig. 2A shows the change in concentration of 4 sugars in the presence of veliger larvae of *Crassostrea gigas*. Maltotriose and rhamnose were not taken up,
while both glucose and maltose were completely removed (i.e. below the limit of detection) within 240 min. No significant changes in concentration were measured for these same 4 sugars in the control flask (no larvae) after 300 min (Fig. 2A, inset). A similar pattern of net uptake was observed for veliger larvae of *Haliotis rufescens* (Fig. 2B). Glucose and maltose were completely removed in 240 min, while there was no detectable uptake of maltotriose or rhamnose.

A range of other complex sugars was tested to see if larvae of *Crassostrea gigas* had the capacity to transport them. Larvae were exposed to the disaccharides cellobiose and isomaltose, and the trisaccharides cellobiose and isomaltotriose. Fig. 3A shows that larvae can take up cellobiose in addition to glucose but not isomaltose or maltotriose. In this experiment, glucose was completely removed to the limit of detection by 90 min. Cellobiose was taken up more slowly and was completely removed by 180 min. Larvae also took up the trisaccharide cellobiose, but at a lower rate than glucose (Fig. 3B). Larvae did not take up the trisaccharide isomaltotriose (Fig. 3B).

Fig. 4 illustrates the relationship for larvae of *Crassostrea gigas* between the rates of glucose net flux, as measured with HPLC, and the rates of influx, as measured with 14C-label. The rate of isotope influx was a good measure of the rate of glucose net flux into larvae.

**Effect of maltose on the rate of transport of 14C-glucose by larvae**

Fig. 2A shows that veligers of *Crassostrea gigas* can take up both glucose and maltose when both substrates were present in seawater. To resolve if maltose was being hydrolyzed and transported as glucose, an experiment was carried out to determine whether maltose and glucose were transported via the same carrier in larvae of *C. gigas*. The transport rate by larvae of 14C-glucose at 1 μM in seawater was measured in the presence and absence of 200 μM maltose. In 3 separate experiments, with no maltose present, 14C-glucose was transported at rates of 58, 63, and 76 fmol glucose larva⁻¹ h⁻¹ (Table 1). These rates of glucose transport were not significantly different from those measured in the presence of 200 μM maltose (see ANOVA, Table 1).
DISCUSSION

Measurement of sugars in seawater

The amperometric detection scheme of Watanabe (1985) allows for the measurement of individual sugars in pmol amounts. However, Watanabe's methods require that samples be diluted 50- to 20000-fold with mobile phase prior to HPLC analysis. Thus, his procedures, originally developed for analyzing fruit juices and human fluids, do not permit quantification of sugars at submicromolar concentrations in seawater. Our procedures overcame this limitation by using the following modifications: (1) adding NaCl to the mobile phase, (2) sparging the reagents with helium for at least 1 h, (3) diluting the seawater sample by a factor of 2.5 with distilled water and acetonitrile before injection, (4) utilizing a brand of aminopropyl-bonded column that can separate sugars even when using a relatively low percentage (volume to volume) of acetonitrile, and (5) using a knitted coil for the reaction at 98 °C. With these modifications, as little as 5 pmol of an individual reducing sugar can be detected in seawater without any sample concentration or desalting (Fig. 1). For the analyses presented here, with a 100 μl injection volume (i.e. 40 μl seawater), 5 pmol equates to a 125 nM concentration for each sugar. Lower sugar concentrations can be measured using larger injection vol-

Table 1. *Crassostrea gigas*. Effect of maltose on the transport rate of 14C-glucose by veliger larvae (207 μm valve length, 636 ml-l). Concentration of glucose, 1 μM; concentration of maltose, 200 μM. n: number of points used to calculate each value of r²

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ANOVA

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Uptake of sugars from seawater by larvae

Filtered seawater that had been sterilized by autoclaving was used as an incubation medium. Larvae were always thoroughly rinsed with this seawater prior to experiments. Thus, any bacteria in the experiments could only be those associated with the larvae. Bacteria-free (axenic) larvae of *Crassostrea gigas* (cf. Manahan 1989) were not used for the experiments presented here. However, no significant differences between axenic and xenic cultures have been observed for the transport rates of amino acids by larvae (Davis & Stephens 1984, Jaeckle & Manahan 1989). If bacteria were present in significant numbers in our experiments then the concentration of all sugars should decrease given the heterotrophic potential of bacterial populations (Williams 1975). However, of the monosaccharides tested (glucose and L-rhamnose) only glucose was taken up during our experiments. For the disaccharides, both maltose and cellulbiose were taken up, but not isomaltose. Similarly, only cellotriose was taken up of the 3 trisaccharides tested. Because of this selective uptake of sugars, we are confident that the larvae were responsible for the observed patterns of sugar uptake.

The influx rate of radiolabeled-glucose into *Crassostrea gigas* larvae was similar to the rate of glucose net flux when the latter was measured by direct chemical analysis (HPLC) (Fig. 4). This finding supports other reports for marine invertebrates that have used $^{14}$C-glucose to measure influx (e.g. Stephens 1962) or spectrophotometric assays to measure glucose net flux (Pajor et al. 1989). The application of HPLC to these kinds of measurements will be especially useful for those sugars for which radiolabeled substrates are not commercially available (e.g. many of the disaccharides and trisaccharides). HPLC measurements revealed that veliger larvae of *C. gigas* have the ability to take up the monosaccharide glucose but not L-rhamnose (Fig. 2A). The disaccharide maltose was taken up from seawater at a rate similar to glucose (Fig. 2A). However, the rate of energy input into the larvae from the transport of maltose would be approximately twice that from glucose transport. Glucose enters the larva by a transport system not shared with maltose: the rate of glucose transport from 1 μM is not affected by the presence of excess maltose (200 μM) (Table 1). This suggests that maltose is not being hydrolyzed prior to being transported as glucose monomers. Larvae of *C. gigas* also have the ability to take up the disaccharide cellubiose, the trisaccharide cellotriose, but not isomaltose, maltotriose nor isomaltotriose (Fig. 3A, B). Like bivalve larvae, veligers of the gastropod *Halosia rufescens* were also found to take up glucose and maltose, but not the monosaccharide L-rhamnose, nor the trisaccharide maltotriose (Fig. 2B). Cellulbiose and cellotriose were not tested with larvae of *H. rufescens*.

Glucose is believed to be the dominant monosaccharide in seawater (Itekkot et al. 1981, Sakugawa & Handa 1983) and is probably a result of oligosaccharide hydrolysis. Phytoplankton have been shown to produce α1-4 and β1-4 glucans as storage products (Craigie 1974, Lewin 1974). Maltose is also used by some phytoplankton as a storage product (Patil & Joshi 1970). The breakdown of these glucans produces smaller sugars (glucose, maltose, maltotriose) that are possibly released into the water column by exudation or upon cell lysis. Both β1-3 and β1-4 glucans are found in seawater and their presence has been attributed to algal blooms (Sakugawa & Handa 1983, Sakugawa & Nobukiko 1985). Some of the hydrolysis products of β1-4 glucans are glucose, cellulbiose, and cellotriose. Larvae can take up complex sugars that have 1-4 glycosidic linkages (maltose, cellulbiose, cellotriose). However, larvae cannot take up isomaltose or isomaltotriose both of which have 1-6 glycosidic linkages. This suggests that the 1-6 linkage might confer a steric hindrance for transport. By having the ability to take up more complex sugars, in addition to monomers, molluscan larvae are able to utilize a greater fraction of the DOM pool as a source of nutrition.

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LITERATURE CITED


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