Chemical defenses in the sea hare *Aplysia parvula*: importance of diet and sequestration of algal secondary metabolites

David W. Ginsburg*, Valerie J. Paul**

ABSTRACT: Marine algae produce a variety of secondary metabolites that function as herbivore deterrents. Algal metabolites, however, often fail to deter damage by some herbivores such as mesograzers that both live and feed on their host alga. In addition, the degree to which intraspecific chemical variation in an alga affects a mesograzers's feeding behavior and its ability to deter predators is poorly understood. The red alga *Portieria hornemannii* contains the secondary metabolites apakaochtodene A and B, which have been shown to vary in concentration among sites on Guam and act as significant deterrents to fish feeding. On Guam, the sea hare *Aplysia parvula* preferred and grew best when fed its algal host *P. hornemannii*. However, high concentrations of *P. hornemannii* crude extract and the pure compounds apakaochtodene A and B acted as feeding deterrents to *A. parvula*. Despite differences among sites in the levels of apakaochtodenes A and B, *A. parvula* showed no significant preference for *P. hornemannii* from any one location. *Aplysia parvula* found on *P. hornemannii* sequestered apakaochtodenes, and both whole animals and body parts were unpalatable to reef fishes. Sea hares found on the red alga *Acanthophora spicifera*, which contains no unpalatable secondary metabolites, had no apakaochtodene compounds and were eaten by fishes. This observation is consistent with the hypothesis that diet-derived algal metabolites in sea hares play a role in deterring predation.

KEY WORDS: Sea hare · Tri-trophic interaction · Antipredator chemical defense · Apakaochtodene A and B · *Aplysia* · *Portieria* · Guam

INTRODUCTION

Marine algae produce a variety of secondary metabolites that function as herbivore deterrents (Hay & Fenical 1988, Paul 1992, Hay 1996). Under natural conditions, these compounds provide a selective advantage to the algae (Hay 1992, 1996, Paul 1992). Algal secondary metabolites may also have other roles as defenses against pathogens and fouling organisms, thus increasing the adaptive value of these metabolites (Paul 1992, Schmitt et al. 1995).

Algal metabolites fail to deter damage by some herbivores and may stimulate feeding by small, relatively sedentary herbivores such as amphipods, crabs, polychaetes, and some gastropods. These herbivores, collectively termed mesograzers, both live and feed on their algal host (Hay et al. 1989, 1990a,b, Brawley 1992, Hay 1992, Trowbridge 1992, Di Marzo et al. 1993). It has been hypothesized that the preference of some mesograzers for chemically-defended algae has been driven primarily by the advantages of decreased predation that mesograzers might experience while living and feeding on a host plant that is a deterrent to reef fishes (Hay et al. 1987, 1989, 1990a,b, Hay 1992, 1996).

The degree to which intraspecific chemical variation in an alga affects a mesograzers's feeding behavior and...
its ability to deter predators is poorly understood. While several studies have examined quantitative variation in algal secondary chemistry (Carlton et al. 1989, Meyer & Paul 1992, 1995, de Nys et al. 1996, 1998, Puglisi & Paul 1997, Matlock et al. 1999), few have examined the effects of within-algal variation on the feeding preferences of mesograzers (Van Alstyne 1989, Poore 1994, Cronin & Hay 1996a,b). Such studies are necessary not only to understand the factors affecting the production of chemical defenses, but also to provide insight into the ecological consequences of such variation (Hay 1996, Becerro et al. 1998).

How intraspecific variation of algal secondary metabolites influences the vulnerability of mesograzers to predation has not been examined. These types of interactions can best be studied by examining the chemistry as well as the palatability of organisms within an alga-mesograzerpredator, or tri-trophic, system. In this study, we chose to investigate the red alga Portieria hornemannii (Lyngbye) Silva (Gigartinales: Rhizophyllidaceae), the mesograzer Aplysia parvula (Opisthobranchia: Anaspidea) and generalist, reef fish predators to examine how chemical variation in the primary producer affects these tri-trophic interactions.

The red alga Portieria hornemannii was selected for this study because it exhibits notable variation among sites in the production of secondary metabolites (Paul et al. 1987, Fuller et al. 1992, 1994, Puglisi & Paul 1997, Matlock et al. 1999) and is host to the sea hare Aplysia parvula (Switzer-Dunlap & Hadfield 1977, Carefoot 1987). On Guam, the major secondary metabolite of P. hornemannii is apakaochtodenede B (Fig. 1) which is an effective feeding deterrent against herbivores (Paul et al. 1987, 1990, 1992, Meyer et al. 1994). Several other halogenated monoterpenes including apakaochtodene A (Fig. 1), the double-bond isomer of apakaochtodene B, are minor metabolites (Paul et al. 1987, Puglisi & Paul 1997, Gunatilaka et al. 1999, Matlock et al. 1999).

The sea hare Aplysia parvula is an oligophagous herbivore that feeds primarily on red algae (Carefoot 1987, Rogers et al. 1995) and is able to sequester algal metabolites from its diet that have been hypothesized to function in defense against predators (Faulkner 1992, de Nys et al. 1996). Switzer-Dunlap (1978) reported that, in laboratory cultures, larvae of A. parvula preferentially metamorphose in the presence of Portieria hornemannii (= Chondrococcus hornemannii) relative to other algae. On Guam, we have observed A. parvula to primarily live on and to preferentially graze P. hornemannii, from which animals sequester the algal metabolites apakaochtodenede A and B (this study).

To investigate how seaweed chemical defenses mediate ecological interactions among the sea hare Aplysia parvula, its algal host Portieria hornemannii, and predatory reef fishes, we asked the following questions: (1) Does A. parvula prefer to consume and grow best on P. hornemannii compared to other algae? (2) Do among site differences in the concentrations of apakaochtodenedes found in P. hornemannii affect the feeding behavior of A. parvula? (3) Are A. parvula (whole animals and body parts) unpalatable to reef fish predators? (4) Do the varying concentrations of apakaochtodenedes found in P. hornemannii from different sites, which are sequestered by A. parvula, differentially affect predation on sea hares by reef fishes?

**MATERIALS AND METHODS**

**Study sites and organisms.** All experiments were conducted from June 1995 to June 1999 on Guam, the largest and southernmost island in the Mariana Islands archipelago located in the western North Pacific Ocean. Portieria hornemannii and Aplysia parvula were collected from 5 sites on Guam: 4 sites on the leeward side, Double Reef (13° 36’ N, 144° 48’ E), Gun Beach (13° 31’ N, 144° 48’ E), Apaca Point (13° 24’ N, 144° 40’ E), and Anae Island (13° 23’ N, 144° 38’ E) and 1 site on the windward side of the island, Pago Bay (13° 25’ N, 144° 48’ E) (Fig. 2). Double Reef and Gun Beach are located at the northern end of Guam. Sea hares and algae were collected from Double Reef's northeastern reef and from the northern end of Gun Beach. Pago Bay, Apaca Point and Anae Island are located further south on Guam. Samples from Pago Bay were collected behind the University of Guam Marine Laboratory and collections from Apaca Point were made from the southern end of the point. Specimens collected from Anae Island, located ~450 m offshore, were collected from the fringing reef on its eastern side.

The alga Portieria hornemannii grows as tufts attached to rocks or dead corals in subtidal areas with...
heavy current (Trono 1969, 1997, Puglisi & Paul 1997). Tufts are comprised of small individual thalli 3 to 9 cm tall with primary branches 0.5 to 1 mm in diameter (Trono 1969, 1997). On Guam, tufts are found 0.5 to 3 m apart (Puglisi and Paul 1997) and were collected haphazardly from the reef at depths of 0.5 to 8 m.

_Aplysia parvula_ is a small (0.4 to 10 cm max. length), oligophagous herbivore that feeds primarily on red algae (Carefoot 1987, Rogers et al. 1995, de Nys et al. 1996). These animals are camouflaged with coloration that matches their host algae (Rogers et al. 1995). _A. parvula_ were obtained by collecting whole tufts of _Portieria hornemannii_. Bulk collections of algae, from their respective sites, were brought back to the laboratory and placed in flow-through, outdoor aquaria, under natural light for ~2 wk. The handling of animals was kept to a minimum. Sea hares were gleaned from _P. hornemannii_ as they were needed for each experiment.

**Chemical extraction.** Organic compounds were extracted from individual _Portieria hornemannii_ thalli. Fresh or frozen whole individual thalli were rinsed quickly in freshwater to remove surface salt, cleaned of epiphytes, spun in a plastic salad spinner for 20 revolutions and blotted dry. After recording fresh weights (to the nearest 0.1 mg), thalli were ground with a Virtis high speed homogenizer in an equal volume solvent mixture of dichloromethane:methanol. The solvent and solid material was transferred to a beaker and the volume brought to ~50 ml. After 8 h, the extract was decanted through weighed filter paper (Whatmann GF/A Glass Microfibre filter), sealed and stored in a freezer at −20°C. Algal material left in the flask was resuspended in ~50 ml of fresh solvent and extracted for an additional 8 h. Three successive extractions were performed on each sample, after which all remaining solids were dried on the filters and weighed. The combined extract from each thallus was concentrated by rotary evaporation and weighed. These crude extracts were then dissolved in hexanes, filtered through glass wool, dried, weighed, and stored at −20°C prior to quantitative chemical analysis of apakaochtodene compounds. The extract yields of individual thalli were calculated using the dry weights from filtered, crude extracts.

Monoterpenes also were extracted from _Aplysia parvula_. Sea hares sequester algal metabolites in their digestive gland, which is located internally and supplies enzymes to the gut. A layer of tissue called the mantle, which, in this study, includes the parapodial folds and the foot, covers the animal. Individual sea hares were frozen and either left intact, or dissected and their digestive gland and mantle removed. Because of the small size of _A. parvula_, it was necessary to combine replicate digestive gland and mantle parts (independent of one another) in order to obtain sufficient material for chemical analysis. Pooled samples were freeze dried while whole animals were stored frozen. Both pooled and whole animals were chemically extracted with acetone and prepared for analysis of apakaochtodenes as described above.

**Quantitative analysis.** Dried extracts for gas chromatography-mass spectrometry (gc-ms) were re-dissolved in hexanes with naphthalene as an internal standard at a concentration of 50 µg ml⁻¹. Samples were diluted to a concentration of 1 ml of hexanes (with the internal standard naphthalene) per 1 mg crude extract (1 mg ml⁻¹). Gas chromatography was conducted using a Hewlett-Packard 5980 Series II Plus gas chromatograph with a cross-linked 5% methyl silicone column (HP-5, 30 m × 0.25 mm). Injections were made in the splitless mode with an inlet pressure of 13.9 kPa at 70°C. The injection port was held at 250°C with a 70 to 290°C temperature ramp at 10°C min⁻¹. The carrier gas was helium at a flow rate of 0.5 ml min⁻¹.

Mass spectrometry was conducted with an HP 5972 Mass Selective Detector (MSD). Ions characteristic of the internal standard naphthalene and apakaochtodenes A and B were monitored in the selected ion monitoring (SIM) mode and were quantitatively analyzed using purified standards. Quantification was performed with a multiple point (6 dilutions) external method using the HP ChemStation (1993) software.
The peak areas of apakaochtodenes A, B and the internal standard naphthalene were measured and their ratio (compound:internal standard) calculated and converted to concentration by reference to standard curves specific to each pure compound. For both apakaochtodenes A and B, the amount of pure compound was converted to % yield based on total dry mass of the individual thallus or sea hare.

**Choice assays.** The dietary preferences of *Aplysia parvula* were determined by conducting choice assays in laboratory aquaria (Paul & Pennings 1991). *A. parvula* used in each set of choice assays were collected from either Apaca Point or Gun Beach. Animals (n = 8 to 15) were placed individually in separate 1 l flow-through aquaria (diam. = 16 cm, max. depth = 8 cm). Each sea hare was offered a choice between 2 species of algae that were weighted with clothespins at opposite ends of the aquarium. Individual sea hares were placed in the middle of each aquarium at the beginning of the experiment and their presence or absence on either of the 2 algae was recorded hourly beginning on the 2nd day. Following the methods of Paul & Pennings (1991), individual animals were excluded from the experiment if they were not recorded on algae at least 3 times. Individual sea hares were used only once during the experiment. Five separate choice tests were conducted: *Portieria hornemannii* vs *Acanthopora spicifera*; *P. hornemannii* vs *Asparagopsis taxiformis*; *P. hornemannii* vs *Dictyota cervicornis*; *P. hornemannii* vs *Galaxaura marginata*; *P. hornemannii* vs *Gracilaria tsudae*; and *P. hornemannii* vs *Laurencia papillosa*. Sea hares were recorded as choosing the alga upon which they were observed the most (ties were excluded). For each choice assay, the total number of animals that chose 1 seaweed compared to another was analyzed using a 2-tailed binomial test. These particular seaweeds, all of which are red algal species except the brown alga *Dictyota cervicornis*, were used in choice and no-choice (below) assays because they are among the most common seaweeds in habitats where *A. parvula* is found on Guam.

**No-choice assays.** No-choice assays were based on *Aplysia parvula* growth and consumption on 5 separate, single species, algal diets: *Laurencia papillosa, Dictyota cervicornis, Gracilaria tsudae, Acanthopora spicifera,* or *Portieria hornemannii* (Paul & Pennings 1991).

Sea hares were blotted dry and weighed (to the nearest 0.1 mg). After confirming that there were no significant differences among the initial weights of animals (1-way ANOVA, F_4, 42 = 0.36, p = 0.83), each algal species was provided *ad libitum* to individual sea hares (n = 10) kept in separate 250 ml flow-through aquaria (diam. = 7 cm, max. depth = 9 cm). After 6 d, each animal was weighed again and its relative change in mass was calculated using the following equation: \[ \frac{(T_f - T_o)}{T_o} \], where \( T_o \) and \( T_f \) are the weight of *A. parvula* before and after the assay. Growth rates were log(x + 2) transformed to meet parametric criteria, slopes were compared by ANCOVA with the initial weights of *A. parvula* held as the covariate, and means were compared using Tukey’s HSD pairwise comparisons test.

During the growth assays described above, measurements of the consumption of algae by *Aplysia parvula* were also conducted. Following the protocol suggested by Peterson & Renaud (1989) and modified by Cronin & Hay (1996b), each algal species (see above) was divided into replicate treatments and no-herbivore controls. Each replicate sample was blotted dry and weighed on an electronic balance (to the nearest 0.1 mg). No-herbivore controls were used to control for autogenous changes in mass in the absence of herbivores. After 6 d, each alga was reweighed and the amount of algal tissue consumed by sea hares was calculated after correcting for autogenous weight changes in seaweeds using the following equation: \[ \frac{(T_f \times C_o/C_i) - T_o}{T_o} \], where \( T_o \) and \( T_f \) are the weight of the algal pieces exposed to herbivory before and after the assay, and \( C_o \) and \( C_i \) are the weight of the controls for autogenic changes before and after the assay (Cronin & Hay 1996b). Consumption data could not be transformed to meet parametric criteria. Consequently, the non-parametric Kruskal-Wallis test was used to test for differences in mean consumption rates among the 5 seaweed diets. The non-parametric equivalent of Tukey’s HSD test was used to assess differences among means.

Whole samples of the seaweeds *Acanthopora spicifera, Dictyota cervicornis, Gracilaria tsudae, Laurencia papillosa,* and *Portieria hornemannii* were analyzed for total Kjeldahl nitrogen (TKN) (Lachat Instruments 1992) and organic carbon (Nelson & Sommers 1975) using a Lachat QuickChem Automated Ion analyzer. Portions of each algal thallus (n = 3) were rinsed in seawater to remove extraneous material, oven dried at 62°C for 48 h and stored at –20°C in tightly capped vials. TKN and organic carbon data were calculated as the % yield of nitrogen and carbon from total dry mass of the seaweed. TKN values present in different species of seaweed were arcsin-square-root transformed to meet parametric criteria. Organic Carbon and TKN data were analyzed by 1-way ANOVA and means were compared using Tukey’s HSD multiple comparisons test.

**Chemical bioassays.** Crude *Portieria hornemannii* extract and pure apakaochtodenes were tested at varying concentrations to determine their effects on feeding by *Aplysia parvula*. Mean natural wet mass concentrations of *P. hornemannii* crude extract were...
between 0.5 to 4%. Conversely, pure apakaochtodenes were approximately 10% of the crude extract and natural wet mass concentrations ranged from 0.01 to 0.6%. Thalli of the palatable red seaweed *Acanthophora spicifera* (5 to 7 mg wet mass) were trimmed with a razor blade to a length of 5 cm and coated with a solution of either crude *P. hornemannii* extract or pure apakaochtodene A or B dissolved in hexanes (Hay et al. 1998). Treated thalli were tested over a range of concentrations that approximate their natural wet mass concentrations. Controls consisted of *A. spicifera* thalli that were coated with hexanes only. Since metabolites extracted from *P. hornemannii* are lipid-soluble, they adhere to the surface of the *A. spicifera* thalli after the hexanes evaporate. Hay et al. (1998) found that lipid soluble metabolites from other sea-weeds were not lost after 24 h underwater in similar experiments. For each assay, individual sea hares (n = 10) were given a choice between treated and control *A. spicifera* thalli offered simultaneously. Bioassays were run as described previously (Choice assays) over an interval of 24 to 48 h. The amount (in cm) of treated and control *A. spicifera* thalli eaten by each sea hare was recorded and analyzed using a paired t-test. Controls for autogenous changes in mass in the absence of herbivores were not necessary for this experiment because the length of *A. spicifera* thalli did not change without herbivory.

**Chemical variation among sites.** The preference of *Aplysia parvula* for Portieria hornemannii collected from different locations was determined using choice assays in laboratory aquaria. *P. hornemannii* was collected from different sites around Guam, and each population was held in a separate tank. Individual sea hares (n = 10 to 15), collected at Gun Beach, were placed in 1 l flow-through aquaria and offered a choice between individual *P. hornemannii* thalli from 2 different sites. Choice assays using *A. parvula* were run as described previously (Choice assays) and were recorded hourly for 5 to 7 h. Three separate choice experiments were conducted for the *P. hornemannii* (different populations are listed by site): Apaca Point vs Double Reef; Pago Bay vs Apaca Point; and Pago Bay vs Double Reef. Assays testing algae from Apaca Point vs Pago Bay were conducted on 2 separate d using different sea hares on each day. The assay results from these 2 d were combined and used as replicates. For each choice assay, the total number of animals choosing one population of *P. hornemannii* compared to another was analyzed using a 2-tailed binomial test. Additionally, individual *P. hornemannii* were collected separately from Apaca Point (n = 7), Double Reef (n = 6) and Pago Bay (n = 7), and chemically analyzed by gc-ms for their concentrations of apakaochtodenes A and B. Apakaochtodene A data could not be transformed to meet parametric criteria. Thus, the non-parametric Kruskal-Wallis test was used to test for differences in the levels of apakaochtodene A among the *P. hornemannii* populations. The non-parametric equivalent of Tukey’s HSD test was used to assess for differences among means. Differences in the levels of apakaochtodene B among the populations of *P. hornemannii* were analyzed by 1-way ANOVA followed by Tukey’s HSD multiple comparisons test. Sea hares were also collected at Anae Island (n = 11) and Double Reef (n = 10) and were analyzed chemically by gc-ms. These data, however, could not be transformed to meet parametric criteria. The apakaochtodene A and B contents of sea hares were compared between the 2 sites using the non-parametric Mann-Whitney test.

**Palatability to predators.** Sea hares and other opisthobranch molluscs can be preyed upon by fish, crustaceans, and even other opisthobranchs (Carefoot 1987, Pennings 1990a, Avila & Paul 1997, Gochfeld & Aeby 1997, Johnson & Willows 1999). In tropical habitats, coral reef fishes may be particularly important at controlling populations of opisthobranch molluscs (Gochfeld & Aeby 1997). We tested the palatability of *Aplysia parvula* to reef fish predators because these assays could be done in the field with natural assemblages of reef fishes.

The palatability of *Aplysia parvula* to reef fishes was determined at 2 sites on Guam: Gun Beach and Western Shoals (Paul & Pennings 1991). At both sites, frozen, whole *A. parvula* collected from Anae Island (n = 10 to 12) and Double Reef (n = 10 to 11) were released in random order into the water column by a scuba diver ~4 m below the surface. The fate of each animal (rejected or eaten) after ~30 s exposure to fish predation was recorded by a second diver underwater. At Western Shoals, live *A. parvula* from Anae Island (n = 10) were also tested to determine if ink and/or mucus secretions affected feeding by fish. Chunks of squid (n = 10), cut into pieces comparable to whole sea hares, served as controls to ensure that fish were readily feeding. A G-test of independence was used to analyze results via a 3 × 2 contingency table for Gun Beach and a 4 × 2 contingency table for Western Shoals.

*Aplysia parvula* from Apaca Point (n = 10) and Double Reef (n = 10) were frozen, dissected, and their body parts (digestive gland and mantle) offered to reef fishes at Gun Beach. Digestive gland (n = 10) and mantle (n = 10) parts were removed from each animal and treated as independent replicates. Body parts and squid chunks (n = 10, fish feeding controls) were cut into equal-sized pieces and released in random order into the water column as described above. Assays were analyzed using a 3-way log-linear model, classifying responses based on site of collection (Apaca Point or
Double Reef), type of body part (digestive gland or mantle), and number of body parts eaten or rejected. Selective feeding on digestive gland and mantle from Apaca Point and Double Reef was analyzed for each site using Fisher’s Exact Test. It was not possible, however, to collect a sufficient number of *A. parvula* from either Apaca Point or Double Reef for chemical analysis. Thus, sea hares were collected from Cocos Lagoon. The chemical content of animals collected from this site is similar to animals from Apaca Point and Double Reef. Sea hares (n = 10) were frozen, dissected and their digestive gland and mantle parts chemically analyzed by gc-ms for their apakaochtodene A and B content.

Individual sea hares maintained in laboratory aquaria on *Portieria hornemannii* (n = 15, from Double Reef) and the red alga *Acanthophora spicifera* (n = 15, from Pago Bay) were offered to reef fishes at Western Shoals. While *Aplysia parvula* is not commonly found on *A. spicifera* on Guam, animals were found once on this alga, which is known to contain no unpalatable secondary metabolites. We have no data to suggest that the *A. parvula* found on *A. spicifera* had ever eaten *P. hornemannii*. Sea hares and squid chunks (n = 15, fish feeding controls) were released in random order into the water column as described previously, and the fate of each animal was recorded underwater. Assays were analyzed using a G-test of independence via 2 × 2 contingency table; results were based on the sea hare’s algal diet (*A. spicifera* or *P. hornemannii*) and whether sea hares were eaten or rejected. *A. parvula* that were rejected by fishes in the field were analyzed chemically by gc-ms. The apakaochtodene A and B contents of sea hares maintained on *P. hornemannii* were compared to those maintained on *A. spicifera* using the non-parametric Mann-Whitney test.

### Results

#### Choice assays

*Aplysia parvula* chose *Portieria hornemannii* over all other tested seaweeds including the red algae *Acanthophora spicifera*, *Asparagopsis taxiformis*, *Galaxaura marginata*, *Gracilaria tsudae*, *Laurencia papillosa*, and the brown alga *Dictyota cervicornis* (2-tailed binomial, p < 0.05; Table 1). Of all choice tests performed, only one individual *A. parvula* preferred an alga other than *P. hornemannii*, and it chose *G. tsudae*.

#### No-choice assays

The relative change in mass of *Aplysia parvula* maintained on the 5 seaweed diets differed significantly over the 7-d period (ANCOVA, \( F_{4,41} = 12.07, p < 0.001 \); Fig. 3a). During these assays, 3 sea hares died

<table>
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<th>Alga</th>
<th>Sample size</th>
<th>No. of preferring</th>
<th>p</th>
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<td></td>
</tr>
<tr>
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<td></td>
</tr>
<tr>
<td><em>P. hornemannii</em></td>
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<td>8</td>
<td>0.007</td>
</tr>
<tr>
<td><em>Dictyota cervicornis</em></td>
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<td></td>
<td></td>
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<tr>
<td><em>P. hornemannii</em></td>
<td>11</td>
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<tr>
<td><em>Laurencia papillosa</em></td>
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Table 1. Dietary preferences of *Aplysia parvula* for different algal species. p values determined with binomial test (2-tailed)
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and were excluded from the experiment. Individual sea hares grew best on *Portieria hornemannii* compared with all other tested seaweeds.

The amount of algae consumed by *A. parvula* also was significantly different among treatments (Kruskal-Wallis, $H = 17.30$, df = 4, $p < 0.001$; Fig. 3b). Sea hares consumed significantly greater amounts of *Gracilaria tsudae*, *Acanthophora spicifera* and *P. hornemannii* than *L. papillosa* or *D. cervicornis*.

The mean amounts of organic carbon and total Kjeldahl nitrogen (TKN) differed significantly within the 5 seaweeds analyzed (ANOVA, $F_{4,10} = 5.90$, $p = 0.01$, and $F_{4,10} = 11.67$, $p = 0.001$, respectively; Fig. 4a,b). Despite significant differences among algal treatments, high levels of organic carbon were present in all of the seaweeds analyzed (Fig. 4a). The amounts of TKN in the different algae were more variable and reached their highest levels in *Gracilaria tsudae*, *Acanthophora spicifera* and *Portieria hornemannii* and their lowest levels in *Laurencia papillosa* and *Dictyota cervicornis* (Fig. 4b).

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**Chemical bioassays**

While crude *Portieria hornemannii* extract did not significantly deter feeding by *A. parvula* at concentrations ≤2% of algal wet mass (paired *t*-test, $p > 0.05$), feeding was deterred at concentrations of 4% and 6% (paired *t*-test, $p < 0.05$) (Fig. 5a). In a direct comparison, sea hares did not significantly distinguish (paired *t*-test, $p = 0.39$) between the crude *P. hornemannii* extract (0.9% wet mass) from Anae Island and Gun Beach (Fig. 5b). Both apakaochotodenes A and B, at all concentrations tested, significantly deterred (paired *t*-test, $p < 0.05$) feeding by sea hares (Fig. 6, b).

**Among site chemical variation**

In choice assays, individual *Aplysia parvula* did not show a significant preference for *Portieria hornemannii* collected from Apaca Point, Double Reef or Pago Bay (2-tailed binomial, $p > 0.05$; Table 2). Chemical

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**Fig. 4.** Organic carbon and TKN (total Kjeldahl nitrogen) content of 5 algal diets. (A) Mean % organic carbon and (B) mean % TKN (+1 SE; n = 3 for each algal treatment). TKN values arcsin-square-root transformed. Organic carbon and TKN data converted to % yield based on total dry mass of seaweed. Identical letters above bars indicate means are not different ($p > 0.05$). Pairwise comparisons made using Tukey’s HSD. Untransformed means presented for clarity.

**Fig. 5.** *Aplysia parvula*. Chemical bioassays using crude extracts from red alga *Portieria hornemannii*. (A) Crude extracts tested at wet mass concentrations that span the natural range of concentrations found in alga and (B) in direct comparison between *P. hornemannii* from 2 different locations: Anae Island; Gun Beach. Data represent mean amount of *Acanthophora spicifera* eaten (+1 SE; n = 10 paired replicates for each comparison). *values differed significantly ($p < 0.05$) between controls and treatments for a given extract concentration. ns: means did not differ significantly ($p > 0.05$)
analysis by gc-ms indicated significant differences in the levels of apakaochtodene A and B within individual algae among sites (Kruskal-Wallis test, H = 14.03, df = 2, p < 0.001, and ANOVA, F\_2, 17 = 6.60, p = 0.007, respectively; Fig. 7a). Mean levels of apakaochtodenes A and B were significantly higher in thalli from Apaca Point compared to Double Reef and Pago Bay, while there was no difference between thalli from Double Reef or Pago Bay.

Table 2. Dietary preferences of Aplysia parvula for Portieria hornemannii collected from different sites. p values were determined with a binomial test (2-tailed)

<table>
<thead>
<tr>
<th>Site</th>
<th>Sample size</th>
<th>No. of preferring</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apaca Point</td>
<td>13</td>
<td>10</td>
<td>0.092</td>
</tr>
<tr>
<td>Double Reef</td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pago Bay</td>
<td>10</td>
<td>6</td>
<td>0.507</td>
</tr>
<tr>
<td>Double Reef</td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Apaca Point</td>
<td>15</td>
<td>8</td>
<td>0.387</td>
</tr>
<tr>
<td>Pago Bay</td>
<td>4</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Chemical analysis of Aplysia parvula showed that animals from Anae Island sequestered significantly greater levels of apakaochtodenes A and B than those from Double Reef (Mann-Whitney test, U = 110.0, p < 0.001, and U = 84.0, p = 0.044, respectively; Fig. 7b).

Palatability to predators

In field assays at Gun Beach and Western Shoals, neither frozen sea hares from Anae Island nor Double Reef were palatable to reef fishes compared with squid pieces (G-test, G = 31.60, df = 2, p < 0.001, and G = 9.21, df = 3, p < 0.01, respectively; Fig. 8a). At Gun Beach, the fishes Thalassoma lutescens, Halichores hortulanus, and Balistapus undulatus ate frozen sea hares. Conversely, at Western Shoals, the fishes Scarus sp. and Naso vlamingii ate frozen sea hares, however, no
live sea hares were eaten. For nearly all cases, sea hares were eaten only after being mouthed and released by numerous fishes.

Field assays at Gun Beach also indicated that digestive gland and mantle parts from Aplysia parvula were not palatable to reef fishes (Fig. 8b). Reef fishes readily consumed squid parts. However, so as not to confound the specific question of the palatability of A. parvula body parts from different sites, squid data were excluded from statistical analyses. Since a log-linear model including all possible 2-way interactions did not significantly fit the observed preference data ($G = 4.54, df = 1, p = 0.033$), a 3-way interaction must be used to account for this variation. Fishes ate similar amounts of digestive gland parts from Apaca Point (1 eaten: 9 rejected) and Double Reef (0:10), whereas, fish ate different amounts of mantle parts from Apaca Point (6:4) and Double Reef (1:9). The fishes Abudeful saxatilis, Thalassoma lutescens and Arothron manilensis consumed both types of sea hare body parts. Chemical analysis by gc-ms indicated that levels of apakaochtodenes A and B were higher within the digestive gland of A. parvula compared with its mantle (Fig. 9).

Reef fishes significantly preferred whole Aplysia parvula found on Acanthophora spicifera compared with sea hares found on Portieria hornemannii ($G$-test, $G = 9.19, df = 1, p = 0.002$; Fig. 10a). Squid data were excluded from statistical analyses, as described above. Chemical analysis of A. parvula not eaten by fishes in the field showed that the apakaochtodene A and B content of sea hares found on P. hornemannii was significantly higher than animals found on A. spicifera (Mann-Whitney test, $U = 67.5, p = 0.003$ and $U = 70.0, p = 0.001$, respectively; Fig. 10b).

**DISCUSSION**

**Dietary specialization**

On Guam, Aplysia parvula chose and exhibited better growth on Portieria hornemannii over all other seaweed diets tested. While sea hares consumed algae that contained similar amounts of organic carbon and TKN, the relative growth of animals was greatest when they consumed P. hornemannii. Similarly, in other areas of the Pacific, A. parvula consumes a variety of red algae that are rich in secondary metabolites (Carefoot 1987, Rogers et al. 1995), many of which contain...
halogenated secondary metabolites (Hay & Fenical 1988). For example, in New South Wales, Australia, Rogers et al. (1995) reported that *A. parvula* preferentially consumes the nutritionally and chemically rich red algae *Laurencia obtusa* and *Delisea pulchra*, which are its primary algal hosts. Surprisingly, however, sea hares did not grow when fed *D. pulchra*, from which animals sequester secondary metabolites that are unpalatable to reef fishes (de Nys et al. 1996), but grew best on *L. obtusa*. The dietary preferences of herbivores and the nutritional content of the seaweed species that they consume are often unrelated (Carefoot 1987, Duffy & Hay 1991, Rogers et al. 1995, Wakefield & Murray 1998). Indeed, feeding specialization by *A. parvula* on chemically defended algae may reflect their need for algal hosts to serve as refuges against predation rather than some unique dietary requirement (Futuyma & Moreno 1988, Pennings 1990a, Rogers et al. 1995, Wakefield & Murray 1998).

*Aplysia parvula* may become less susceptible to predation by selectively living and feeding on *Portieria hornemannii*. The major secondary metabolite produced by this alga is apakaochtodene B which is an effective feeding deterrent to herbivorous fishes (Paul et al. 1987, 1990, 1992, Meyer et al. 1994). Therefore, *P. hornemannii* may offer *A. parvula* ‘enemy-free space’ (Price et al. 1980, Jeffries & Lawton 1984); that is, sea hares may avoid being eaten by inhabiting an alga that its predators (e.g., reef fishes) avoid (Rogers et al. 1995). These results are consistent with other studies of alga-mesograzers-predator interactions and support the hypothesis that mesograzers can minimize predation by specializing on chemically defended seaweeds that are unpalatable, and thus approached less by fish predators (Hay et al. 1987, 1989, 1990a, Hay & Fenical 1988, Duffy & Hay 1991, 1994, Paul & Pennings 1991, Hay & Steinberg 1992, de Nys et al. 1996).

Dietary specialization by *Aplysia parvula* may also be influenced by the types and concentrations of algal secondary metabolites found in its host algae. As previously mentioned, the work of Rogers et al. (1995) with *A. parvula* in Australia suggested that sequestration of secondary metabolites from the red alga *Delisea pulchra* might restrict sea hare’s growth. On Guam, *A. parvula* encountered individual *Portieria hornemannii* thalli that contain crude extract concentrations ranging from 0.5 to 4% of algal wet mass. At low concentrations, crude extract had little effect on sea hare grazing, but it deterred feeding at higher concentrations. Crude extract was a deterrent at 4%, a concentration found to be present in ≤2% of individual *P. hornemannii* thalli collected from several different locations on Guam (Matlock et al. unpubl. data). These findings are consistent with previous studies reporting that it is not uncommon for mesograzers, who by definition both live and feed on their host alga, to be indifferent to low concentrations of seaweed chemical defenses, yet be deterred at higher concentrations of these same metabolites (Hay et al. 1989, 1990, Pennings & Paul 1993, Steinberg 1995, Hay 1996, Nagle et al. 1998).

In contrast, the pure apakaochtodene A and B compounds produced by *Portieria hornemannii*, significantly deterred feeding by *Aplysia parvula* at all concentrations tested (Fig. 6); a surprise, considering that sea hares were indifferent to equivalent proportions of crude *P. hornemannii* extract. The underlying causes of this disparity between the ability of crude extract and apakaochtodenes to deter feeding by sea hares are not known. It is possible that compounds such as

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**Fig. 10. Aplysia parvula.** (A) Palatability and (B) apakaochtodene content of sea hares found on either red alga *Portieria hornemannii* or *Acanthophora spicifera*. Palatability data represent number of animals eaten by reef fishes at Western Shoals and were analyzed using a 2 × 2 contingency table. Chunks of squid served as fish feeding controls. Chemical data indicate mean % apakaochtodenes (+1 SE). Amounts of apakaochtodenes A and B converted to % yield based on total dry mass of sea hares. Numbers above bars represent number of samples analyzed. **Values differed significantly (U = 67.5, p = 0.003 and U = 70.0, p = 0.001, respectively) between sea hares found on *P. hornemannii* compared to *A. spicifera*.
triglycerides or sterols within the crude extract of *P. hornemannii* mask the potency of apakaochtodenes A and B. Another factor may be that pure compounds were applied to the surface of *Acanthophora spicifera* thalli, which is probably not where the monoterpenes are located in live *P. hornemannii*.

*Portieria hornemannii* shows significant variation in apakaochtodene content among thalli within the same site (Matlock et al. 1999). It is possible that such differences in the chemical makeup of *P. hornemannii* may present a problem to *Aplysia parvula* in that its algal host may vary in its suitability as a resource both temporally and spatially (Nagle et al. 1998, Matlock et al. 1999).

Despite among-site differences in levels of apakaochtodenes A and B, *Aplysia parvula* showed no significant preference for *Portieria hornemannii* from any location (Fig. 7, Table 2). These results are consistent with previous studies that have demonstrated significant variation in the production of apakaochtodenes by *P. hornemannii* among different locations on Guam (Puglisi & Paul 1997, Matlock et al. 1999). Our data suggest that at the site level, the feeding behavior of *A. parvula* was not influenced by the composition or concentration of apakaochtodenes produced by *P. hornemannii*, and are consistent with findings by Steinberg (1995), where feeding by the herbivorous sea urchin *Holopneustes purpurescens* was not affected by variation in levels of phlorotannins in its host kelp *Ecklonia radiata*.

**Predation on Aplysia parvula**

Similar to other sea hares (Faulkner 1992, Pennings 1994, Johnson & Willows 1999), *Aplysia parvula* sequesters diet-derived chemical defenses from its host algae (Rogers et al. 1995, de Nys et al. 1996), which on Guam is *Portieria hornemannii*. Our field experiments indicated that sea hares found on *P. hornemannii*, regardless of where they were collected, were unpalatable to reef fish. Thus, despite differences in the composition and concentrations of apakaochtodenes in *P. hornemannii* among sites, *A. parvula* that fed on *P. hornemannii* from Apaca Point were no more vulnerable to predation than when they had fed on *P. hornemannii* from Double Reef. The fish species found to consume whole *A. parvula* in this study were similar to fishes reported (most notably, wrasses) to eat the sea hare *Stylocheilus longicauda* (Paul & Pennings 1991) and other opistobranchs (Gochfeld & Aeby 1997, Avila & Paul 1997). The susceptibility of *A. parvula* to predators was slightly higher at Gun Beach than at Western Shoals. Generally, there were more parrotfish and damselfish (*Scarus* spp. and *Amblyglyphidodon* spp., respectively), and fewer wrasses (*Thalassoma* spp.) present at Western Shoals compared with Gun Beach. This may be because Western Shoals is a popular tourist site where recreational divers feed fish several times a day. It is possible that diet-derived secondary metabolites in opistobranchs may deter some fish predators but are ineffective against others, such as tolerant (e.g., wrasses) or opportunistic (e.g., damselfish) predators (Pennings & Paul 1993, Avila & Paul 1997). While fishes are not the only predators on opistobranch molluscs, they will readily consume nudibranchs (Avila & Paul 1997, Gochfeld & Aeby 1997) and ascoglossans (Trowbridge 1994) when conspicuous and available to them. Crabs (Trowbridge 1994) and other invertebrate predators (Carefoot 1987, Pennings 1990a, b, Johnson & Willows 1999) also have been shown to consume opistobranchs and may act as important predators in limiting natural assemblages of these animals.

In field assays, reef fishes consumed frozen *Aplysia parvula*, however, no live animals were eaten. *Aplysia parvula* releases ink when physically disturbed, similar to that reported for other sea hares (Carefoot 1987, Johnson & Willows 1999). While Dimatteo (1981, 1982) has suggested that the inking behavior of sea hares acts as a means of defense against predators, sea hare ink is not known to contain any algal secondary metabolites nor is its natural function(s) well known (Carefoot 1987, Pennings 1994, Johnson & Willows 1999). Carefoot et al. (1999) showed ink to act as a sensory irritant to a number of types of invertebrates and fishes. Whether the ink of live *A. parvula* was a basis for its unpalatability to reef fishes is not clear, but is a possible factor in affecting the vulnerability of these animals to predation on Guam.

Sea hares found on *Portieria hornemannii* were unpalatable to reef fishes, but animals found on *Acanthophora spicifera* were readily eaten. These results are consistent with chemical data (Fig. 10b) which showed that *A. parvula* sequester apakaochtodenes from a *P. hornemannii* diet. Concentrations of apakaochtodenes in animals were nearly an order of magnitude higher than those found in the alga. Sea hares found living on *A. spicifera*, which is not known to contain any unpalatable secondary metabolites, had no apakaochtodene compounds or any other detectable chemical defenses. Similarly, Pennings (1990a) showed that when the sea hare *Aplysia californica* was grown on an algal diet rich in secondary metabolites animals were less palatable to fish than when they had been grown on an algal diet lacking secondary metabolites altogether.

The body parts (digestive gland and mantle) of *Aplysia parvula* were also unpalatable to reef fishes. However, while reef fishes clearly showed no preference for individual body parts from Double Reef, fishes tended to more often consume mantle compared with
digestive gland parts from Apaca Point. Concentrations of apakaochtodenes in *A. parvula* varied between different body parts with algal compounds significantly present in the digestive gland compared to mantle. These data are consistent with previous findings of quantitative (Pennings & Paul 1993, de Nys et al. 1996) and qualitative variation (Winkler 1969, Gerwick & Whatley 1989, Faulkner 1992) of algal secondary metabolites in sea hares. Since sea hares sequester algal metabolites in their digestive gland (where they are not optimally located for defense) rather than mantle tissues, the anti-predatory role of these compounds has been disputed (Pennings & Paul 1993, Pennings 1994, de Nys et al. 1996). However, several studies have demonstrated that sea hare mantle is unpalatable to predators (Ambrose et al. 1979, Pennings 1990a, 1994, Pennings et al. 1999). It is possible that sea hares may produce an unpalatable compound which is of greater defensive value than diet-derived metabolites (Pennings & Paul 1993, Pennings 1994, Pennings et al. 1999). While this latter argument cannot be ignored, our data indicate that *A. parvula* body parts are unpalatable to most reef fish and are consistent with the hypothesis that algal secondary metabolites acquired by sea hares function as predator deterrents (Pennings 1990a, Paul & Pennings 1991, Faulkner 1992, de Nys et al. 1996).

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LITERATURE CITED


Hay ME, Duffy JE, Paul VJ, Renaud PE, Fenical W (1990b) Specialist herbivores reduce their susceptibility to predation by feeding on the chemically defended seaweed A vrainvillea longicaulis Limnol Oceanogr 35:1734–1743


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