# A Review of the Chemical Ecology of Antarctic Marine Invertebrates<sup>1</sup>

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SYNOPSIS. The interdisciplinary field of marine invertebrate chemical ecology is relatively young, and particularly so in polar marine environments. In this review we present evidence that the incidence of chemical defense in antarctic benthic marine invertebrate phyla is widespread. Mechanisms of chemical defense have been detected in antarctic representatives of the Porifera, Cnidaria, Brachiopoda, Tunicata, Nemertea, Mollusca and Echinodermata. This argues against earlier biogeographic theories that predicted a low incidence of chemical defense in polar waters where levels of fish predation are low. Selection for chemical defense in benthic sessile and sluggish marine invertebrates is likely a response to an environmentally stable community shown to be structured primarily by biotic factors such as predation and competition. Holoplankton and the eggs, embryos and larvae of both benthic and planktonic antarctic macroinvertebrates may also employ chemical defense to offset mortality during characteristically slow development and long life span where susceptibility to predation is seemingly high. While most research to date has focused on the role of secondary metabolites in mediating predation, it is likely that bioactive compounds in antarctic marine invertebrates also serve roles as antifoulants and allelochemics. The diversity of bioactive metabolites detected to date in antarctic marine invertebrates sets the stage both for continuing and for broadening efforts to evaluate their functional and ecological significance.

# Introduction

The role of chemical defense in mediating marine invertebrate predator-prey interactions has received considerable attention (reviewed by: Bakus et al., 1986; Paul, 1992; Pawlik, 1993; Hay, 1996). As is true of marine plants (Hay and Fenical, 1988; Hay, 1992; Hay and Steinberg, 1992), sessile and sluggish marine invertebrates are extremely vulnerable to mobile predators. Moreover, marine invertebrates are subject to having their tissues fouled by settling larvae, diatoms and other algal cells, and in the case of sponges, soft corals, bryozoans and other sessile marine invertebrates, to overgrowth by species competing for space. Ultimately, the ability of these organisms to chemically defend themselves can play a significant role in regulating predator-prey

interactions and structuring marine communities (Bakus *et al.*, 1986; Paul, 1992; Pawlik, 1993; Hay, 1996).

Most of the research on marine invertebrate chemical ecology has focused on tropical environments (Bakus et al., 1986). The coral reefs of the tropics are well known for their high species diversity and communities comprised of individuals and colonies competing intensely for space and food. Under such conditions, it is not surprising that many sessile or sluggish marine invertebrates have evolved chemical means of defense (Bakus et al., 1986; Paul, 1992, Pawlik 1993; Faulkner, 1996 and references within). Early studies examining geographic patterns of chemical defense (as measured by ichthyotoxicity) in marine invertebrates noted a decrease in chemically defended species as one moved from tropical to temperate marine habitats (Bakus and

<sup>&</sup>lt;sup>1</sup> Invited mini-review.

Green, 1974; Green, 1977; Bakus, 1981). Based on these comparisons, and the observation that fish predation on benthic marine invertebrates decreased with increasing latitude (Vermeij, 1978), an inverse correlation between chemical defense and latitude in marine invertebrates was proposed (Bakus and Green, 1974).

Although the antarctic benthic marine environment has been classically considered to be harsh environmentally, studies have shown that with the exception of organisms living in the most shallow water (less than 33 m depth) where anchor ice and ice scour are important structuring forces (Dayton et al., 1969, 1970), the benthic community is remarkably rich and stable (Dell, 1972; Dayton et al., 1974; Richardson and Hedgpeth, 1977; Dayton 1989, 1990; White, 1984). The assemblage of marine organisms comprising this community has evolved during the events preceding the cretaceous breakup of Gondwanaland and the movement of the antarctic continent southward. The establishment of a circumpolar current effectively isolated the continent, and its climate remained temperate to sub-tropical until 22 million years ago (Dayton et al., 1994). The marine biota, therefore, has its origins in warmer climates and is relatively old (Dell, 1972). According to Dayton et al. (1994) the shallow-water antarctic marine fauna is derived from 1) relict autochthonous fauna, 2) eurybathic fauna from deeper water, and 3) cool-temperate species, mostly from South America.

The factors responsible for structuring this complex sponge-dominated community appear to be primarily biological in nature. For example, competition for space is intense between certain species of sponges, several of which are capable of dominating space if released from predation pressure (Dayton et al., 1974; Dayton, 1989). Moreover, although fish predators that browse on sessile marine invertebrates such as sponges are indeed rare in high latitudes (Eastman, 1993), there is a very high incidence of predation in Antarctica by mobile macroinvertebrates, particularly echinoderms such as sea stars (Dearborn, 1977; McClintock, 1994). In the water column, there are a variety of abundant planktivorous fish that are known to ingest zooplankton (e.g., Foster et al., 1987; Eastman, 1991) as well as extremely dense swarms of predatory amphipods (e.g., Rakusa-Suszczewski, 1972). While overall grazing or predation pressure (per unit time) is arguably higher in tropical marine systems (Vermeij, 1978), some antarctic macroinvertebrate grazers and predators occur in extremely high densities and consequently are capable of exerting considerable predation pressure (Fig. 1A). Moreover, antarctic organisms often have very long lifespans (Clarke, 1983), the effect of which may be that predation intensity averages out over lifespan to be similar to that in temperate or even tropical marine systems (Fig. 1B). As one of the oldest, most environmentally stable and intensely biologically structured marine systems, the antarctic benthos is well suited for the evolution of chemical defense mechanisms.

There has been a plethora of bioactive secondary metabolites (defined as those compounds with no definitive role in primary metabolic pathways) isolated from marine invertebrates (Scheuer, 1990; Attaway and Zaborsky, 1993; Faulkner, 1996). These compounds have been isolated primarily from sponges and cnidaria (predominantly soft corals), although smaller numbers of other members of marine invertebrate groups have also yielded bioactive secondary metabolites (Baker, 1996; Faulkner, 1996 and references within). Although the descriptive chemical studies published on novel secondary metabolites from marine invertebrates number in the thousands, there are comparatively few studies which provide ecological information about the functional significance of these compounds. In many instances, biological activity has been measured simply as the ability of a compound to cause cell lysis or inhibit growth of a non-marine microbe. While such information may have utility in pharmaceutical studies (Munro et al., 1987; Scheuer, 1990; Hay and Fenical, 1996), much remains to be learned about the ecological significance of such bioactive compounds. This is particularly true in polar waters, where chemical ecological studies of marine invertebrates have only recently begun, with the vast majority of studies having been conducted in McMurdo Sound, Antarctica (77°S, 166°E).



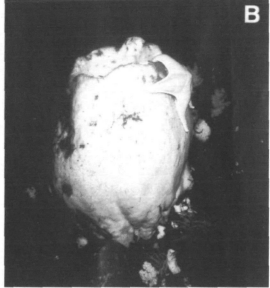


Fig. 1. In situ photographs of antarctic benthic predation events representative of different time scales. A. Mass feeding aggregation of the antarctic nemertean worm *Parborlasia corrugatus* and the antarctic sea star *Odontaster validus*. These common predation events may occur over a time scale of a few hours (McClintock, Baker, Heine pers. obs.; photo by J. Heine). B. The antarctic spongivorous sea star *Acodontaster conspicuus* preying on a large vase sponge. These predation events may last months to years (Dayton *et al.*, 1974; photo by J. Mastro).

While the data base for antarctic waters is consequently regional in its scope, it can be expected to have wide applicability as the shallow epi-macrofauna of McMurdo Sound is generally circumpolar in its distribution (Hedgpeth, 1969; Dell, 1972; Arnaud, 1974; White, 1984). The purpose of this paper is to provide an overview of what is currently known of the chemical ecology of antarctic marine invertebrates and suggest directions for further research. We include in this review both information on the broad bioactive nature of secondary metabolites and wherever possible information that is of ecological significance. To date, ecological studies in Antarctica have focused primarily on evaluations of how defensive chemistry may inhibit feeding by likely or known predators, similar to studies in temperate and tropical marine environments (Paul, 1992; Pawlik, 1993; Hay, 1996).

# **EMPIRICAL STUDIES**

# Sessile macroinvertebrates

Sponges.—The sponges of McMurdo Sound, Antarctica have been the focus of chemical studies to date (Table 1). On the

eastern side of McMurdo Sound, Antarctica, approximately twenty-one species of conspicuous marine sponges occur in nearshore waters accounting for 55% of the benthic surface area below 33 m depth (Koltun, 1970; Dayton et al., 1974). Dayton (et al., 1974) was the first to suggest chemical defenses occurred in antarctic sponges, having observed no predation on the sponges Leucetta leptorhapsis, Dendrilla membranosa and Isodictya erinacea, all of which apparently lacked physical defenses against spongivorous sea stars. In the first quantitative studies of chemical bioactivity, McClintock (1987) and McClintock and Gauthier (1992) found a number of antarctic sponge extracts caused mortality in goldfish or inhibited growth in allopatric microorganisms. Moreover, Blunt et al. (1990) found the incidence of antiviral activity similar in marine sponges from temperate waters of New Zealand (37%) and polar antarctic waters (33%). In addition, Battershill (1990) conducted immunological assays with three antarctic sponges from McMurdo Sound (Cinachyra antartica, Latrunculia sp., Polymastia sp.) and found

TABLE 1. Chemical and ecological bioactivity of antarctic shallow-water marine invertebrate.

	Anti- micro- bial	Anti- fou- lant	Cyto-	Tube- foot retrac- tion	Feeding deter- rence	Activity in:				
						Com-	Extract	Tiscue	Whole animal	Reference <sup>b</sup>
DANIFER !	- Ulai	Idill	IOXIC	Sea stars	1.1211	pound	LAHaci	113340		Kererenee
PORIFERA										_
Calyx acuarius	+			+			+			1
Cinachyra antarctica				+			+			1,2,3,4,5,6
Dendrilla membranosa	+			+		+	+			1,2
Gellius benedeni	+			+			+			1,2
Haliclona sp.				+			+			1
Haliclona dancoi	+			+			+			1,2
Homaxonella balfourensis	+			+			+			1,2
Inflatella belli				+			+			1
Isodictya erinacea	+			+		+	+		+¢	1,2,3
Kirkpatrickia variolosa	+			+		+	+			1,2,3,7,8,9
Latrunculia apicalis	+			+		+	+			1,2,3,6,10
Leucetta leptorhapsis	+		+	+		+	+			1,2,3
Polymastia invaginata	+			+			+			1,2
Rosella nuda				+			+			1
Rosella racovitzae				+			+			1
Sphaerotylus antarcticus				+			+			1
Tetilla leptoderma				+			+			1
CNIDARIA										
Alcyonium paessleri	+	+	+	+	+		+	+		11,12,13
Clavularia frankliniana	_	_	_	+	+		_	+		11,12,13
Gersemia antarctica	+	+	+	+	+		+	+		11,12,13
BRACHIOPODA										, ,
				+	+		+	+		14
Liothyrella uva				+	+		T	т		14
ASCIDACEA										
Cnemidocarpa verrucosa			_		+		+	+		15
NEMERTEA										
Parborlasia corrugatus			+		+	+	+	+	+	16,17
MOLLUSCA										
Austrodoris kerguelensis				+	+	+		+		18,19,20
Clione antarctica	_				+	+	_		+	2,21,22,23
Marseniopsis mollis				+	+	+		+	+	18,19,24
Tritoniella belli				+	+	+		•	+	18,19,25
ECHINODERMATA				•	•	•			•	- 0, ,
				,					+c	26
Perknaster fuscus			+	+					+c	26 27
Diplasterias brucei				+			+		+,	<i>21</i>

<sup>&</sup>lt;sup>a</sup> Data only presented for species tested against antarctic microbes, microalgae, macroinvertebrate gametes, macroinvertebrates or fish.

them all capable of rejecting tissues from congeners, concluding these sponges were capable of producing defensive chemicals that were toxic to genetically dissimilar tissues. The results of these studies suggested that a variety of antarctic sponges harbor toxic compounds. Nonetheless, while these bioassay approaches provide useful infor-

<sup>&</sup>lt;sup>b</sup> References: McClintock et al., 1993a<sup>1</sup>; Baker and McClintock unpublished<sup>2</sup>; Baker et al., 1993<sup>3</sup>, 1995<sup>4</sup>; Molinski and Faulkner, 1987<sup>5</sup>; Yang et al., 1995<sup>6</sup>; Jayatilake et al., 1995<sup>7</sup>; Perry et al., 1994<sup>8</sup>; Trimurtulu et al., 1994<sup>9</sup>; Blunt et al., 1990<sup>10</sup>; Slattery and McClintock, 1995<sup>11</sup>; Slattery et al., 1995<sup>12</sup>; Slattery, 1994<sup>13</sup>; McClintock et al., 1993b<sup>14</sup>; McClintock et al., 1991<sup>15</sup>; Heine et al., 1991<sup>16</sup>; W. Kem unpublished<sup>17</sup>; McClintock et al., 1992a<sup>18</sup>; McClintock et al., 1994d<sup>19</sup>; Davies-Coleman and Faulkner, 1991<sup>20</sup>; McClintock and Janssen, 1990<sup>21</sup>; Bryan et al., 1995<sup>22</sup>; Yoshida et al., 1995<sup>23</sup>; McClintock et al., 1994<sup>24</sup>; McClintock et al., 1994c<sup>25</sup>; McClintock and Baker, 1997<sup>27</sup>.

<sup>&</sup>lt;sup>c</sup> Whole animal activity detected in developing embryos.

Fig. 2. Chemical structures of bioactive compounds isolated from antarctic organisms (clockwise from upper left): The bacterium *Pseudomonas aeruginosa* isolated from the surface of the sponge *Isodictya setifera*, the pteropod *Clione antarctica*, and the sponges *Isodictya erinacea*, *Dendrilla membranosa*, *Latrunculia apicalis* and *Kirkpatrickia variolosa*.

mation which may be of pharmacological or agrochemical importance, they provide little information of demonstrative ecological significance. Subsequent investigations by McClintock et al. (1994a) focused on examinations of the feeding deterrent characteristics of hexane, chloroform and methanol (non-polar to polar) extracts of antarctic sponges employing a tube-foot assay using an ecologically relevant predator, the common antarctic spongivorous sea star Perknaster fuscus. Significant tube-foot retraction activity was detected in the chloroform and methanol extracts of 75% of eighteen species tested. These levels of deterrence are similar to levels measured in reef fish presented feeding pellets containing extracts of Caribbean sponges in the laboratory (Pawlik et al., 1995). The bioactive compounds likely responsible for feeding deterrence or additional biological activity have been elucidated in five of these antarctic species.

A variety of bioactive metabolites have been identified from antarctic sponges (Baker et al., 1993, 1994; Fig. 2). Diterpene metabolites, including dendrillin, 9,11-dihydrogracilin A, membranolide and picolinic acid have been extracted from the tissues of *Dendrilla membranosa* (Molinski and Faulkner, 1987, 1988; Baker et al., 1993, 1994, 1995). The latter compound causes sea star tube-foot retraction (Mc-

Clintock et al., 1994b). Kirkpatrickia variolosa has been found to produce variolins, a novel group of alkaloids, which have been demonstrated to have antitumor and antiviral activity (Perry et al., 1994; Trimurtulu et al., 1994). A purple pigment in K. variolosa causes tube-foot retraction in the antarctic sea star Perknaster fuscus (Baker and McClintock, unpublished). Moreover, a bioactive stilbene derivative has been isolated from K. variolosa, the first such observation of a stilbene from a marine invertebrate (Jayatilake et al., 1995). The antarctic sponge Latrunculia apicalis has been found to contain a variety of discorhabdin pigments (Yang et al., 1995; Baker et al., 1993, 1994) which are cytotoxic (Blunt et al., 1990) and cause tube-foot retraction in P. fuscus (Baker and McClintock, unpublished). Recent work with the sponge Leucetta leptorhapsis has indicated the presence of highly cytotoxic lipids (Baker et al., 1993) that cause feeding deterrence in antarctic seastars (McClintock et al., 1993a). The antarctic sponge Isodictya erinacea contains the secondary metabolites eribusenone, 7-methyladinine, and p-hydroxybenzoic acid (Baker et al., 1994; Moon et al., in preparation). Bioactive metabolites have also been isolated from microbes associated with antarctic sponges. Jayatilake et al. (1996) recently isolated a series of diketopiperazines and phenazine alkaloids from the bacterium *Pseudomonas aeruginosa* associated with the antarctic sponge *Isodictya setifera* (Fig 2.). Finally, the antarctic sponge *Artemisina apollinis* has been found to possess 28 steroidal derivatives including a new group of natural products (Seldes *et al.*, 1990). The ecological roles of bioactive compounds isolated from *L. leptorhapsis*, *I. erinacea*, and *I. setifera* are currently unknown.

Although it is possible that some of the bioactive metabolites in antarctic sponges are "evolutionary baggage" with no current adaptive significance, it is more likely that given the extreme age of the marine benthos, there has been ample time for defensive metabolites to have evolved in response to sympatric predators, fouling organisms, or competitors. In this respect, it is of interest to consider whether congeners from other latitudes have similar or different chemical defenses, or lack chemical defenses entirely. The answer appears to vary with species, as might be expected of sponges under different selective pressures. For example, the antarctic congeners of Haliclona and Mycale are devoid of the types of secondary metabolites found in their tropical counterparts (alkaloids in Haliclona (Schmitz et al., 1978; Sakai et al., 1986; Baker et al., 1988; Crews et al., 1994) and terpenes or mycalamides from Mycale (Perry et al., 1988; Fusetani et al., 1991; Tanaka et al., 1993)). Alternately, antarctic Latrunculia and Dendrilla both produce derivatives (discorhabdin G and dendrillin, respectively [Yang et al., 1995; Baker et al., 1995]) of secondary metabolites found in temperate and tropical congeners (Karuso et al., 1984; Mayol et al., 1985; Molinski and Faulkner, 1987; Blunt et al., 1990) and the tropical sponge Leucetta microraphis harbors leucettamols (Kong and Faulkner, 1993) that are similar in structure to cytotoxic lipids produced by the antarctic congener Leucetta leptorhapsis (Baker et al., 1993). More information is needed to provide a more comprehensive evaluation of phylogenetic influences on patterns of chemical defense in antarctic sponges.

In many cases it is those antarctic sponges with color that have proven to be bioactive to date (Baker and McClintock, per-

sonal observation). At first examination these would appear to be examples of warning coloration or aposematism (Guilford and Cuthill, 1991; Rosenberg, 1991). However, Pawlik et al. (1995) found no evidence of aposematism in tropical marine sponges. Moreover, it is unlikely aposematism occurs in antarctic sponges, as the antarctic marine benthos is depauperate in visually oriented browsing predators such as fish which feed on sessile invertebrates (most antarctic benthic fish eat crustaceans; Eastman, 1991, 1993). In this regard, even if aposematism occurred in sponges (but see Pawlik et al., 1995), there should be little evolutionary selection for warning coloration in antarctic sponges when their primary predators are sea stars (Dearborn, 1977; McClintock, 1994), which orient to prey chemically (Sloan, 1980). Nonetheless, it is intriguing that antarctic sponges employ their colored pigments as a chemical defense. For example, pigments from the colored sponges Latrunculia apicalis and Kirkpatrickia variolosa are bioactive and cause sea star tube-foot retraction (Baker and McClintock, unpublished).

Soft corals.—Chemical studies of antarctic soft corals have focused on three antarctic species that occur in McMurdo Sound (Table 1). Two of these species, Alcyonium paessleri and Clavularia frankliniana are extremely abundant along the east side of McMurdo Sound (Dayton et al., 1974; Slattery, 1994). The third species, Gersemia antarctica, is much rarer, and within Mc-Murdo Sound is confined to its western regions (Slattery and McClintock, 1995). All three of these soft corals are chemically defended. Organic extracts of A. paessleri and G. antarctica inhibit the attachment of marine antarctic bacteria (Slattery et al., 1995). In addition, organic extracts exhibit antifoulant activity in field experiments causing inhibition of diatom settlement. This is important as diatoms are one of the primary fouling organisms in antarctic benthos (El-Sayed and Fryxell, 1993). Two potential soft coral predators, the antarctic sea stars Odontaster validus and Perknaster fuscus, show tube-foot retractions to organic and aqueous extracts of all three soft coral species (Slattery and McClintock, 1995).

Aqueous homogenates of two of the soft corals, *A. paessleri* and *G. antarctica*, are toxic to antarctic sea urchin larvae, suggesting that similar to tropical marine invertebrates toxicity and feeding deterrence are not predictably related (Pawlik *et al.*, 1995). Distinct secondary metabolites responsible for toxicity or feeding deterrence may be present in these soft corals (Slattery *et al.*, 1995). Chimyl alcohol has been detected in the tissues of *C. frankliniana* and causes rejection of treated shrimp-discs in the omnivorous sea star *Odontaster validus* (McClintock *et al.*, 1994c).

Bryozoans.—While antarctic bryozoans can be dominant members of antarctic benthic communities (Foster, 1969), only one study has investigated their biological activity. Winston and Bernheimer (1986) investigated the extracts of five species of antarctic bryozoans collected from the Antarctic Peninsula and examined hemolytic activity against mammalian erythrocytes. Extracts of one species, Carbasea curva, caused significant lysis of mammalian cells, suggesting the presence of bioactive secondary metabolites. No investigations have yet been conducted to determine the nature of the secondary metabolite(s) nor to examine their ecological significance. Winston and Bernheimer (1986) indicate that C. curva is one of the most abundant bryozoans found in benthic samples from the Antarctic Peninsula or the Ross Sea, and that the toxic substance may be employed in chemical defense, explaining the success of a weakly calcified body plan.

Brachiopods.—One of the few Antarctic Peninsular organisms which has been studied for its chemical defense is the common antarctic brachiopod Liothvrella uva. This punctate terebratulid species is common at depths of 5-300 m (Voss, 1988). Ecological assays indicate that extracts of whole soft body tissues cause significant retraction of the sensory tube-feet of six species of antarctic sea stars (Table 1; McClintock et al., 1993b), several of which are likely predators of brachiopods (McClintock, 1994). Ground body tissues embedded in agar pellets containing krill meal resulted in significant feeding deterrence in an allopatric fish, the Sheepshead Minnow Cyprinidon

variegatus (McClintock et al., 1993b). Studies are needed to determine if ecologically relevant fish are also deterred. The nature of the bioactive compound(s) responsible for feeding deterrence are unknown. The results of this study further support the hypothesis that unpalatability occurs in brachiopods in temperate, tropical, and polar environments (Thayer and Allmon, 1991). A chemical means of defense may be particularly important from an evolutionary perspective due to the slow growth and late reproduction which characterizes the life history of brachiopods (Thayer, 1985).

Ascidians.—Although there are many species of antarctic ascidians, only one species has been chemically investigated. The common solitary ascidian Cnemidocarpa verrucosa is a large conspicuous species which is found in McMurdo Sound, and has a circumpolar distribution (Kott, 1969; Dayton et al., 1974). McClintock et al. (1991) conducted an analysis of the palatability and chemical defense of the tunic, ovitestes, branchial basket, body wall, endocarps and intestines. They found that the tunic was deterrent to sympatric pelagic and benthic fish, in addition to an allopatric fish model (Table 1). Nonetheless, homogenates of the tunic were not cytotoxic to sea urchin gametes, suggesting that the bioactive compound(s) are not toxic. The heavy fouling of the tunic surface by bryozoans and hydroids indicates that bioactive chemicals are not effective antifoulants. Mature ovitestes are rejected by antarctic fish and alginate krill pellets containing lipophilic extracts of ovitestes are rejected by the antarctic sea star Odontaster validus, a likely predator of settling larvae (Dayton et al., 1974), indicating that the eggs and larvae may possess a chemical defense (McClintock et al., 1991; McClintock and Baker, unpublished).

# Mobile macroinvertebrates

Nemerteans.—One of the most conspicuous benthic invertebrates in shallow antarctic waters is the large (up to 1 m extended body length) nemertean worm Parborlasia corrugatus (Fig. 1A; Knox, 1970, Gibson, 1983). Despite their high abundance and rich energy content, and obvious

lack of skeletal defenses, individuals are rarely if ever preyed upon (Dayton et al., 1974; Heine et al., 1991). Cytotoxicity assays indicate homogenates of the whole body tissues cause mortality in gametes of the antarctic sea urchin Sterechinus neumayeri. In addition, ecological bioassays indicate two common species of antarctic fish show significant rejection of body tissues (Table 1; Heine et al., 1991). Both toxicity and feeding-deterrent characteristics of this species may result from the production of a copious acidic (pH = 3.5) mucus (Heine et al., 1991). The defensive properties of this acidic mucus may be diminished through neutralization in seawater. Nonetheless, the mucus of P. corrugatus also harbors a potent toxic neuropeptide (William Kem, unpublished).

Molluscs.—Investigations of chemical defense in antarctic molluscs have been limited to species occurring in McMurdo Sound (Table 1). Nonetheless, many of these species have circumpolar distributions, and their defensive chemistry is likely similar across a wide geographic range. Ecological bioassays indicate the mantle tissues of the antarctic nudibranchs Austrodoris kerguelensis and Tritoniella belli are rejected by two species of sympatric antarctic fish, and cause significant tube-foot retractions in four species of sympatric antarctic sea stars (McClintock et al., 1992a). Austrodoris kerguelensis produces a series of bioactive acid glycerides (Davies-Coleman and Faulkner, 1991) which remain to be evaluated for ecological function, while T. belli contains chimyl alcohol which causes feeding deterrence in the sea star Odontaster validus (McClintock et al., 1994b). This proven feeding deterrent may be derived from its diet of the soft coral Clavularia frankliniana (McClintock et al., 1994b). The prosobranch Marseniopsis mollis, which has an internalized vestigial shell, has mantle tissues which are rejected by antarctic fish and sea stars (McClintock et al., 1992a) and the compound homarine has been isolated from the mantle, foot and viscera (McClintock et al., 1994c). Homarine causes feeding deterrence in the omnivorous O. validus at levels spanning those detected in the body tissues of M. mollis (McClintock et al., 1994c). Homarine was not found in the tunic of the antarctic ascidian Cnemidocarpa verrucosa, the presumed primary prey of this gastropod (McClintock et al., 1994c). Nonetheless, homarine has been isolated from the dense assemblage of epizooites which foul the tunic of this ascidian (McClintock et al., 1994c), suggesting that these fouling organisms (primarily bryozoans and hydroids) may be grazed by M. mollis and be the dietary source of this defensive compound.

The common pelagic pteropod Clione antarctica occurs in vast swarms seasonally in antarctic continental shelf waters (Foster, 1987). Intact live sea butterflies and whole body homogenates imbedded in pellets containing feeding stimulants are consistently rejected by the antarctic zooplanktivorous fish Pagothenia borchgrevinki, a significant pteropod predator (Foster et al., 1987), indicating the presence of chemical defense (McClintock and Janssen, 1990). Interestingly, the abundant antarctic hyperiid amphipod Hyperiella dilatata abducts and carries a single live individual of C. antarctica (Fig. 3), providing itself with a means of chemical defense from fish predators (McClintock and Janssen, 1990). The nature of the compound responsible for chemical deterrence in C. antarctica has recently been elucidated to be a novel bioactive linear hydroxyketone, named pteroenone (Fig. 2; Bryan et al., 1995; Yoshida et al., 1995). This is one of the first defensive compounds isolated from a planktonic macroinvertebrate, and supports recent evidence that feeding deterrent properties of oceanic holoplankton may be widespread (McClintock et al., 1996). As C. antarctica is extremely abundant, the trophic implications of chemical defense in modeling energy flow in the antarctic water column must be considered. Moreover, if additional organisms receive defensive characteristics through their association with chemically defended species (e.g., McClintock and Jansson, 1990), this furthers the complexity of planktonic ecosystems (McClintock et al., 1996).

Echinoderms.—Research conducted with temperate and tropical echinoderms has indicated that a large number of species may possess bioactive compounds, particularly those which are saponin-related (Verbist, 1993). Bioactivity appears to occur most fre-

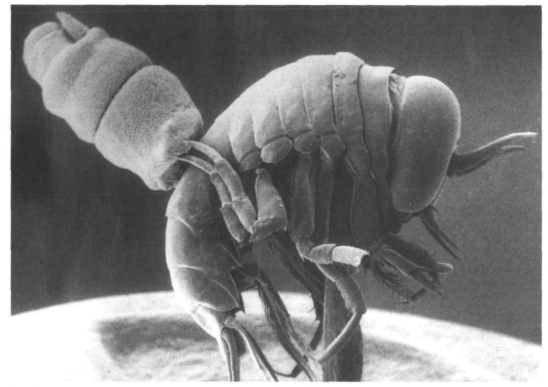


Fig. 3. The antarctic hyperiid amphipod *Hyperiella dilatata* shown grasping the chemically defended pteropod *Clione antarctica* (McClintock and Janssen, 1990). This unique association provides a mechanism for amphipods to defend themselves against their primary predators, antarctic fish. Magnification is approximately ×500.

quently in the body tissues of the holothurians and asteroids (Mackie et al., 1977). McClintock (1989) examined the toxicity of the body wall of 23 species of antarctic echinoderms to the mosquitofish Gambusia affinis. Patterns of toxicity were class specific, and similar to patterns seen in temperate and tropical species, with the highest levels of toxicity occurring in the body tissues of holothuroids and asteroids. Two of three species of holothuroids tested were toxic, while seven of thirteen species of asteroids tested caused mortality in fish. Toxicity was not found to be related to the amount of skeletal material present in the echinoderm body wall, although the degree of skeletal armament was generally low among antarctic species (McClintock, 1989).

The common spongivorous sea star *Perknaster fuscus* possesses the bioactive tetrahydroisoquinoline alkaloid fuscusine (Kong *et al.*, 1992). This is only the second report of an alkaloid in the body wall tissues of a

sea star, the first being imbricatine, a benzyltetrahydroisoquinoline alkaloid which occurs in the sea star Dermasterias imbricata on the west coast of North America. This compound is known to induce a flight response in a sea anemone (Patherina and Anderson, 1986). Crude homogenates of the body wall tissues of P. fuscus are toxic to the sperm of the antarctic sea urchin Sterechinus neumayeri, cause an inhibition of the righting response of the antarctic sea star Odontaster validus, and elicit significant tube-foot retractions in the sympatric antarctic sea stars O. validus, O. meridionalis, Acodontaster conspicuus and Diplasterias brucei (Table 1; McClintock et al., 1992b). At least one of these sea stars, Odontaster validus, is an opportunistic echinovore (McClintock, 1994). It is of some significance that while a number of antarctic sea stars are included in the diets of echinoderms and sea anemones (Dearborn, 1977), P. fuscus has never been ob-

served as a prey item (Dayton et al., 1974). This may be attributable to its chemical defense. Possessing such an effective means of defense has important ramifications in maintaining populations of this spongivorous sea star, as it is responsible for keeping one of the primary space-dominating antarctic sponges (Mycale acerata) at levels which allow increased species diversity (Dayton et al., 1974). Steroidal glycosides and polyhydroxylated steroids have recently been isolated from the antarctic sea star Acodontaster conspicuus and the ophiuroids Ophiosparta gigas and Ophionotus victoriae (D'Auria et al., 1993, 1995; Iorizzi et al., 1996; De Marino et al., 1997). Bioassays indicate that steroidal glycosides and polyhydroxylated steroids isolated from A. conspicuus cause significant growth inhibition in three species of antarctic marine bacteria isolated from the surfaces of antarctic sponges and echinoderms (De Marino et al., in press). This suggests one ecological role of these compounds may be in the prevention of microbial fouling.

In one of only two studies to focus on the chemical defense of early life stages of antarctic marine invertebrates, McClintock and Vernon (1990) examined the icthyonoxic characteristics of the eggs and embryos of fifteen species of antarctic echinoderms, primarily sea stars. Using the common killifish Fundulus grandis as a model predator, they offered pellets impregnated with echinoderm tissues and a feeding stimulant to fish. Chemical feeding deterrents occurred in the large yolky eggs of the pelagic lecithotrophic sea stars Perknaster fuscus and Porania antarctica. Brooded embryos of the sea stars Notasterias armata and Diplasterias brucei were also noxious. The chemical deterrents were effective at ecologically relevant concentrations. In a more recent study, Mc-Clintock and Baker (1997) demonstrated that the lecithotrophic eggs, embryos and larvae of five species of antarctic marine invertebrates (including three echinoderms) were unpalatable or chemically defended from ecologically relevant benthic and water column invertebrate predators (sea stars, sea anemones and amphipods). Two echinoderms with planktotrophic modes of reproduction (the sea urchin Sterechinus neumayeri and the sea star *Odontaster validus*) lacked chemical defense in their eggs or larvae. Therefore, similar to tropical marine invertebrates (Lindquist and Hay, 1996), large yolky lecithotrophic embryos and larvae are more likely to be chemically defended. This may be especially true in antarctic embryos or larvae where very slow development results in substantial periods of time spent in the plankton or on the benthos (Pearse *et al.*, 1991).

### SUMMARY AND FUTURE DIRECTIONS

The unique environmental and biotic factors that characterize shallow antarctic shelf waters have resulted in communities structured by biotic factors such as predation and competition (Dayton et al., 1974). The present review indicates that selection for ecologically relevant secondary metabolites has occurred in a wide variety of representative organisms (reviewed in Table 1). Nonetheless, while much of the research conducted to date on chemical defense has been conducted in the laboratory using ecologically relevant predators (as essentially has been the case for temperate and tropical marine invertebrates), there is a need to extend studies to the field, as has been done to a limited extent in tropical marine environments (Paul, 1992; Pawlik, 1993; Hay, 1996). Much also remains to be learned about additional novel bioactive compounds, and there is a need for further evaluation of the functional roles of known compounds. In addition to defense from predation, factors most likely to be mediated by defensive metabolites include fouling and allelochemical interactions. For example, sessile antarctic benthic organisms are subject to particularly intense fouling pressure by benthic diatoms. Allelochemistry could be important because in contrast to the generalized perception of reduced growth rates in polar species, there are species that grow rapidly and crowd out other species, eventually monopolizing space if released from predation (Dayton et al., 1974, Dayton, 1989).

To date, few investigations have extended studies of defensive metabolites to antarctic macroalgae (Amsler *et al.*, submitted). Such studies will be of particular interest given current theories relating defensives

sive chemistry in macroalgae to biogeographic patterns of herbivory (Estes and Steinberg, 1988; Steinberg and Paul, 1990; Hay, 1992). Initial investigation suggests that antarctic brown algae produce defensive polyphenolics, similar to their temperate and tropical counterparts, perhaps explaining their seemingly low levels of herbivory (Amsler, McClintock and Baker, unpublished).

As in all planktonic systems, there is a need to extend the evaluation of chemical defense to marine invertebrate holoplanktonic organisms in antarctic waters. Of particular interest will be those holoplanktonic organisms that comprise significant biomass in pelagic foodwebs, such as salps and pteropods. Chemically mediated predation may have profound effects on modeling energy flow in such systems (McClintock et al., 1996). Moreover, more information on the role of secondary metabolites in mediating patterns of predation on the pelagic embryos and larvae of macroinvertebrates is needed, along with more quantitative information on abundances and diets of likely planktonic predators.

Finally, studies are needed to determine if antarctic microorganisms (e.g., bacteria, microalgae) employ defensive chemistry (e.g., Jayatilake et al., 1996), if macroinvertebrates utilize microbial defensive metabolites, and if defensive chemistry in antarctic organisms is static or can be induced in response to a variety of ecological constraints.

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### REFERENCES

- Arnaud, P. M. 1974. Contribution a la bionomie marine benthique des regions antarctiques et subantarctique. Tethys 6:645-656.
- Attaway, D. H. and O. R. Zaborsky. (eds.) 1993. *Marine biotechnology*. Plenum Press, New York.
- Baker, B. J. 1996. Beta-carboline and isoquinoline alkaloids from marine organisms. In S. W. Pelletier (ed.), Alkaloids: Chemical and biological perspectives, Vol. 10, pp. 357–407. Permagon Press, London.
- Baker, B. J., P. J. Scheuer, and J. N. Schoolery. 1988. An antifungal pentacyclic alkaloid from a marine sponge *Haticlona* sp. J. Am. Chem. Soc. 110:965– 966.
- Baker, B. J., W. Kopitzke, M. Hamann, and J. B. McClintock. 1993. Chemical ecology of antarctic sponges from McMurdo Sound, Antarctica: Chemical aspects. Antarctic J. U.S. 28:132–133.
- Baker, B. J., W. Y. Yoshida, and J. B. McClintock. 1994. Chemical constituents of four antarctic sponges in McMurdo Sound, Antarctica. Antarctic J. U.S. 24:153–155.
- Baker, B. J., R. W. Kopitzke, W. Y. Yoshida, and J. B. McClintock. 1995. Chemical and ecological studies of the antarctic sponge *Dendrilla membranosa*. J. Nat. Prod. 58:1459–1462.
- Bakus, G. J. 1981. Chemical defense mechanisms in the Great Barrier Reef, Australia. Science 211: 497-498.
- Bakus, G. J. and G. Green. 1974. Toxicity in sponges and holothurians: A geographic pattern. Science 185:951-953.
- Bakus, B. J., N. M. Targett, and B. Schulte. 1986. Chemical ecology of marine organisms: An overview. J. Chem. Ecol. 12:951-987.
- Battershill, C. N. 1990. The chemical ecology of Antarctic benthic marine invertebrates—initial observations. New Zealand Antarctic Rec. 10:9-21.
- Blunt, J. W., M. H. G. Munro, C. N. Battershill, B. R. Copp, J. D. McCombs, N. B. Perry, M. Prinsep, and A. M. Thompson. 1990. From the antarctic

- to the antipodes:  $45^{\circ}$  of marine chemistry. New. J. Chem. 14:751-775.
- Bryan, P. J., W. Yoshida, J. B. McClintock, and B. J. Baker. 1995. Ecological role for pteroenone, a novel antifeedant produced by the conspicuous antarctic pteropod *Clione antarctica* (Gymnosomata: Gastropoda). Mar. Biol. 122:271–278.
- Clarke, A. 1983. Life in cold water: The physiological ecology of polar marine ectotherms. Mar. Biol. Ann. Rev. 21:341-453.
- Crews, P., X.-C. Cheng, M. Adamczeski, J. Rodrigues, M. Jaspars, F. J. Schmitz, S. C. Traeger, and E. O. Pordesino. 1994. 1,2,3,4-Tetrahydro-8-hydroxymanzamines, alkaloids from two different Haplosclerid sponges. Tetrahedron 50, 13567–13574.
- D'Auria, M. V., L. G. Paloma, L. Minale, R. Riccio, A. Zampella, and M. Morbidoni. 1993. Isolation and structure characterization of two novel bioactive sulphated polyhydroxysteroids from the antarctic ophiuroid *Ophiosparte gigas*. Nat. Prod. Lett. 3:197-201.
- D'Auria, M. V., L. G. Paloma, L. Minale, R. Riccio, and A. Zampella. 1995. On the composition of sulfated polyhydroxysteroids in some ophiuroids and the structure determination of six new constituents. J. Nat. Prod. 58:189-196.
- Davies-Coleman, M. T. and D. J. Faulkner. 1991. New diterpenoic acid glycerides from the antarctic nudibranch Austrodoris kerguelensis. Tetrahedron 47:9743–9750.
- Dayton, P. K. 1989. Interdecadal variation in an antarctic sponge and its predators from oceanographic climate shifts. Science 245:1484-1486.
- Dayton, P. K. 1990. Polar benthos. In W. O. Smith (ed.), Polar oceanography, Part B, Chemistry, biology and geology, pp. 631-685. Academic Press, New York.
- Dayton, P. K., G. A, Robilliard, and A. L. DeVries. 1969. Anchor ice formation in McMurdo Sound, Antarctica and its biological effects. Science 163: 273-274
- Dayton, P. K., G. A. Robilliard, and R. T. Paine. 1970. Benthic faunal zonation as a result of anchor ice at McMurdo Sound, Antarctica. In M. W. Holgate (ed.), Antarctic ecology, pp. 244–258. Academic Press, London.
- Dayton, P. K., B. J. Mordida, and F. Bacon. 1994. Polar marine communities. Amer. Zool. 34:90–99.
- Dayton, P. K., G. A. Robilliard, R. T. Paine, and L. B. Dayton. 1974. Biological accommodation in the benthic community at McMurdo Sound, Antarctica. Ecol. Monogr. 44:105-128.
- Dearborn, J. H. 1977. Food and feeding characteristics of antarctic asteroids and ophiuroids. In G. A. Llano (ed.), Adaptations within antarctic ecosystems, pp. 293–326. Gulf Publ. Co., Houston, Texas.
- Dell, R. K. 1972. Antarctic benthos. Advances in Marine Biology 10:1-216.
- De Marino, S., M. Iorizzi, F. Zollo, C. D. Amsler, B. J. Baker, J. B. McClintock, and L. Minale. 1997. Isolation, structure elucidation and biological activity of the steroid glycosides and polyhydroxysteroids from the antarctic starfish Acodontaster conspicuus. J. Nat. Prod. (In press)

- Eastman, J. T. 1991. Evolution and diversification of antarctic notothenioid fishes. Amer. Zool. 31:93–109.
- Eastman, J. T. 1993. Antarctic fish biology: Evolution in a unique environment. Academic Press, Inc., New York.
- El-Sayed, S. Z. and G. A. Fryxell. 1993. Phytoplankton. In E. I. Friedman (ed.), Antarctic microbiology, pp. 65–122. John Wiley and Sons, Inc. New York.
- Estes, J. A. and P. D. Steinberg. 1988. Predation, herbivory, and kelp evolution. Paleobiology 14:19–36.
- Faulkner, D. J. 1996. Marine natural products. Nat. Prod. Repts. 13:75–125.
- Foster, M. W. 1969. Brachiopoda. *In* V. C. Bushnell and J. W. Hedgpeth (eds.), *Antarctic map folio series* 11, p. 9. American Geographical Soc. New York.
- Foster, B. A. 1987. Composition and abundance of zooplankton under the spring sea-ice of McMurdo Sound, Antarctica. Polar Biol. 8:41-48.
- Foster, B. A., J. M. Cargill, and J. C. Montgomery. 1987. Plantivory in *Pagothenia borchgrevinki* (Pisces: Nototheniidae) in McMurdo Sound. Polar. Biol. 8:49-54.
- Fusetani, N., T. Sugawara, S. Matsunaga, and H. Hirota. 1991. Cytotoxic metabolites of the marine sponge *Mycale adhaerens* Lambe. J. Org. Chem. 56:4971–4974.
- Gibson, R. 1983. Antarctic nemerteans: The anatomy, distribution, and biology of *Parborlasia corru*gatus (McIntosh, 1976) (Heteronemertea: Lineidae). Antarct. Res. Ser. 39:289-316.
- Green, G. 1977. Ecology of toxicity of marine sponges. Mar. Biol. 40:207-215.
- Guilford, T. and I. Cuthill. 1991. The evolution of aposematism in marine gastropods. Evolution 45: 449-451.
- Hay, M. E. 1992. The role of seaweed chemical defenses in the evolution of feeding specialization and in the mediation of complex interactions. *In V. J. Paul (ed.)*, *Ecological roles of marine natural products*, pp. 93– 118. Cornell University Press, Ithaca.
- Hay, M. E. 1996. Marine chemical ecology: What's known and what's next. J. Exp. Mar. Biol. Ecol. 200:103-134.
- Hay, M. E. and W. Fenical. 1988. Marine plant-herbivore-predator interactions: The ecology of chemical defense. A. Rev. Ecol. Syst. 19:111-145.
- Hay, M. E. and W. Fenical. 1996. Chemical ecology and marine biodiversity: Insights and products from the sea. Oceanography 9:10-20.
- Hay, M. E. and P. D. Steinberg. 1992. The chemical ecology of plant-herbivore interactions in marine versus terrestrial communities. In G. A. Rosenthal and M. R. Berenbaum (eds.), Herbivores: their interactions with secondary plant metabolites, 2E, Vol. II: Evolutionary and ecological processes, pp. 371-413. Academic Press, New York.
- Hedgpeth, J. W. 1969. Introduction to antarctic zoogeography. Antarctic Map Folio Ser. 11, 109.
- Heine, J. N., J. B. McClintock, M. Slattery, and J. Weston. 1991. Energetic composition, biomass and chemical defense in the common antarctic nemertean *Parborlasia corrugatus* McIntosh. J. Exp. Mar. Biol. Ecol. 153:15-25.
- Iorizzi, M., S. De Marino, L. Minale, F. Zollo, V. Le

- Bert, and C. Roussakis. 1996. Investigation of the polar steroids from an Antarctic starfish of the family Echinasteridae: Isolation of twenty seven polyhydroxysteroids and steroidal glycosides, structures and biological activities. Tetrahedron 57:10997–11012.
- Jayatilake, G. S., B. J. Baker, and J. B. McClintock. 1995. Isolation and identification of a stilbene derivative from the antarctic sponge Kirkpatrickia variolosa. J. Nat. Prod. 58:1958–1960.
- Jayatilake, G. S., M. P. Thronton, A. C. Leonard, J. E. Grimwade, and B. J. Baker. 1996. Metabolites from an antarctic sponge-associated bacterium, *Pseudomonas aeruginosa*. J. Nat. Prod. 59:293–296.
- Karuso, P., B. W. Skelton, W. C. Taylor, and A. H. White. 1984. The constituents of marine sponges. I. The isolation from *Aplysina sulphurea* (Dendroceratida) of 1R, 1'S, 1"R, 3R-1-acetoxy-4-ethyl-5(1,3,3-trimethycyclohexyl)-1,3,dihydroisobenzofuran-1'(4),3carbolactone and the determination of its crystal structure. Australian J. Chem. 37:1081–1093.
- Knox, G. A. 1970. Antarctic marine ecosystems. *In*, M. W. Holgate (ed.), *Antarctic ecology*, Vol. 1., pp. 69-96. Academic Press, New York.
- Koltun, V. M. 1970. Sponges of the arctic and antarctic: A faunistic review. Symp. zool. Soc. Lond. 25:285–297.
- Kong, F. and D. J. Faulkner. 1993. Leucettamols A and B, two antimicrobial lipids from the calcareous sponge *Leucetta microraphis*. J. Org. Chem. 58:970-971.
- Kong, F., M. K. Harper, and D. J. Faulkner. 1992. Fuscusine, a tetrahydroisoquinoline alkaloid from the sea star *Perknaster fuscus antarcticus*. Nat. Prod. Lett. 1:71-74.
- Kott, P. 1969. Antarctic ascidians. Antarctic Res. Ser. 13:1–239.
- Linqvist, N. and M. E. Hay. 1996. Palatability and chemical defense of marine invertebrate larvae. Ecol Monogr. 66:431–450.
- Mackie, A. M., H. T. Singh, and J. M. Owen. 1977. Studies on the distribution, biosynthesis and function of steroid saponins in echinoderms. Comp. Biochem. Physiol. 56B:9-14.
- Mayol, L., V. Piccialli, and D. Sica. 1985. Gracilin A, a unique nor-diterpene metabolite from the marine sponge *Spongionella gracilis*. Tetrahedron 26: 1357–1360.
- McClintock, J. B. 1987. An investigation of the relationship between invertebrate predation and the biochemical composition, energy content, spicule armament and toxicity of sponges in the benthic community at McMurdo Sound, Antarctica. Mar. Biol. 94:479–487.
- McClintock, J. B. 1989. Toxicity of shallow-water antarctic echinoderms. Polar Biol. 9:461-465.
- McClintock, J. B. 1994. Trophic biology of shallowwater antarctic echinoderms. Mar. Ecol. Prog. Ser. 111:191–202.
- McClintock, J. B. and B. J. Baker 1997. Palatability and chemical defense in the eggs, embryos and larvae of shallow-water antarctic marine invertebrates. Mar. Ecol. Prog. Ser. (In press)
- McClintock, J. B., B. J. Baker, M. Hamann, M. Slat-

- tery, R. Kopitzke, and J. Heine. 1994a. Tube-foot chemotactic responses of the spongivorous sea star *Perknaster fuscus* to organic extracts of antarctic sponges. J. Chem. Ecol. 20:859–870.
- McClintock, J. B., B. J. Baker, M. Hamann, W. Yoshida, P. J. Bryan, J. Heine, G. S. Jayatilake, and B. H. Moon. 1994c. Homarine as a feeding deterrent in the common shallow-water antarctic lamellarian gastropod *Marseniopsis mollis*: A rare example of chemical defense in a marine prosobranch. J. Chem. Ecol. 20:2539-2549.
- McClintock, J. B., B. J. Baker, M. Slattery, J. N. Heine, P. J. Bryan, W. Yoshida, M. T. Davies-Coleman, and D. J. Faulkner. 1994b. Chemical defense of the common antarctic shallow-water nudibranch *Tritoniella belli* Eliot (Mollusca: Gastropoda) and its prey, *Clavularia frankliniana* Rouel (Cnidaria: Octocorallia). J. Chem. Ecol. 20:3361–3372.
- McClintock, J. B. and J. J. Gauthier. 1992. Antimicrobial activities of antarctic sponges. Antarctic Science 4:179–183.
- McClintock, J. B., J. Heine, M. Slattery, and J. Weston. 1991. Biochemical and energetic composition, population biology and chemical defense of the antarctic ascidian *Cnemidocarpa verrucosa*. J. Exp. Mar. Biol. Ecol. 147:163-175.
- McClintock, J. B. and J. Janssen. 1990. Pteropod abduction as a chemical defense in a pelagic antarctic amphipod. Nature 346:462-464.
- McClintock, J. B., M. Slattery, B. J. Baker, and J. Heine. 1993a. Chemical ecology of antarctic sponges from McMurdo Sound, Antarctica: Ecological aspects. Antarctic J. U.S. 20:134–135.
- McClintock, J. B., M. Slattery, P. Bryan, W. Yoshida, M. Hamann, and J. N. Heine. 1994d. Chemical ecology of three antarctic gastropods. Antarctic J. U.S. 24:151-152.
- McClintock, J. B., M. Slattery, J. Heine, and J. Weston. 1992a. Chemical defense, biochemical composition and energy content of three shallow-water antarctic gastropods. Polar. Biol. 11:623–629.
- McClintock, J. B., M. Slattery, J. Heine, and J. Weston. 1992b. The chemical ecology of the antarctic spongivorous sea star *Perknaster fuscus*. Antarctic J. U.S. 27:129-130.
- McClintock, J. B., M. Slattery, and C. W. Thayer. 1993b. Energy content and chemical defense of the articulate brachiopod *Liothyrella uva* (Jackson, 1912) from the Antarctic Peninsula. J. Exp. Mar. Biol. Ecol. 169:103-116.
- McClintock, J. B., D. P. Swenson, D. K. Steinberg, and A. A. Michaels. 1996. Feeding deterrent properties of common oceanic holoplankton from Bermudian waters. Limnol. Oceanogr. 41:798–801.
- McClintock, J. B. and J. Vernon. 1990. Chemical defense in the eggs and embryos of antarctic sea stars. Mar. Biol. 105:491–495.
- Molinski, T. F. and D. J. Faulkner. 1987. Metabolites of the antarctic sponge *Dendrilla membranosa*. J. Org. Chem. 52:296–298.
- Molinski, T. F. and D. J. Faulkner. 1988. An antibacterial pigment from the sponge *Dendrilla membranosa*. Tetrahedron Lett. 29:2137–2138.
- Munro, M. H. G., R. T. Ludibrand and J. W. Blunt.

- 1987. The search for antiviral and anticancer compounds from marine organisms. *In* P. J. Scheuer (ed.), *Bioorganic marine chemistry*, pp. 93–176. Springer-Verlag, Berlin.
- Pathirana, C. and R. J. Anderson. 1986. Imbricatine, an unusual benzyltetrahydroisoquinoline alkaloid from the starfish *Dermasterias imbricata*. J. Am. Chem. Soc. 108:8288–8289.
- Paul, V. J. (ed.) 1992. Ecological roles of marine natural products. Comstock, Ithaca, New York.
- Pawlik, J. R. 1993. Marine invertebrate chemical defense. Chem. Rev. 93:1911-1922.
- Pawlik, J. R., B. Chanas, R. T. Toonen, and W. Fenical. 1995. Defenses of Caribbean sponges against predatory reef fish. I. Chemical deterrency. Mar. Ecol. Prog. Ser. 127:183–194.
- Pearse, J. S., J. B. McClintock, and I. Bosch. 1991. Reproduction of antarctic benthic marine invertebrates: Tempos, modes and timing. Amer. Zool. 31:65–80.
- Perry, N. B., J. W. Blunt, M. H. G. Munro, and L. K. Pannel. 1988. Mycalamide A, an antiviral compound from a New Zealand sponge of the genus Mycale. J. Am. Chem. Soc. 110:4850~4851.
- Perry, N. B., L. Ettouati, M. Litaudon, J. W. Blunt, M. H. G. Munro, and H. Hope. 1994. Alkaloids from the antarctic sponge *Kirkpatrickia variolosa*. Part 1. Variolin B, a new antitumour and antiviral compound. Tetrahedron 50:3987–3992.
- Rakusa-Suszczewski, S. 1972. The biology of *Paramoera walkeri* Stebbing (Amphipoda) and the antarctic sub-fast ice community. Polski Archiwum Hydrobiologii 19:11–36.
- Richardson, M. D. and J. W. Hedgpeth. 1977. Antarctic soft-bottom, macrobenthic community adaptations to a cold, highly productive, glacially affected environment. In G. A. Llano (ed.), Adaptations within Antarctic ecosystems, pp. 181-195. The Smithsonian Institution, Washington D. C.
- Rosenberg, G. 1991. Aposematism and synergistic selection in marine gastropods. Evolution 45:451– 454.
- Sakai, R., T. Higa, C. W. Jefford, and G. Bernardinelli. 1986. Manzamine A, a novel antitumor alkaloid from a sponge. J. Am. Chem. Soc. 108:6404–6405.
- Scheuer, P. J. 1990. Some marine ecological phenomena: Chemical basis and biochemical potential. Science 248:173-177.
- Schmitz, F. J., K. H. Hollenbeak, and D. C. Campbell. 1978. Marine natural products: Halitoxin, toxic complex of several marine sponges of the genus *Haliclona*. J. Org. Chem. 43:3916–3922.
- Seldes, A. M., M. E. DeLuca, E. G. Gros, J. Rivirosa, A. San-Martin, and J. Darias. 1990. Steroids from aquatic organisms, XIX. New sterols from the an-

- tarctic sponge Artemisina apollinis. Zeitschrift für Naturforschung 45:83–86.
- Slattery, M. 1994. The chemical ecology of three antarctic soft corals. Ph.D. Diss., Univ. Alabama, Birmingham.
- Slattery, M. and J. B. McClintock. 1995. Population structure and feeding deterrence in three shallow-water antarctic soft corals. Mar. Biol. 122:461–470.
- Slattery, M., J. B. McClintock, and J. N. Heine. 1995. Chemical defenses in antarctic soft corals: Evidence for antifouling compounds. J. Exp. Mar. Biol. Ecol. 190:61-77.
- Sloan, N. A. 1980. Aspects of the feeding biology of asteroids. Oceanogr. Mar. Biol. Ann. Rev. 18:57– 124.
- Steinberg, P. D. and V. Paul. 1990. Fish feeding and chemical defense of tropical brown algae in Western Australia. Mar. Ecol. Prog. Ser. 58:253-259.
- Tanaka, J., T. Higa, K. Suwanborirux, U. Kokpol, G. Bernardinelli, and C. W. Jefford. 1993. Bioactive norsesterterpene 1,2-dioxanes from a Thai sponge Mycale sp. J. Org. Chem. 58:2999-3002.
- Thayer, C. W. 1985. Brachiopods versus mussels: Competition, predation, and palatability. Science 228:1527-1528.
- Thayer, C. W. and R. A. Allmon. 1991. Unpalatable thecideid brachiopods from Palua: Ecological and evolutionary implications. In D. J. MacKinnon, D. E. Lee, and J. D. Campbell (eds.), Brachiopods through time, pp. 253–260. Balkema Press, Rotterdam.
- Trimurtulu, G., D. J. Faulkner, N. B. Perry, L. Ettouati, M. Litaudon, J. W. Blunt, M. H. G. Munro, and G. B. Jamieson. 1994. Alkaloids from the antarctic sponge Kirkpatrickia variolosa, Part 2, Variolin A and N(3')-methyltetrahydrovariolin B. Tetrahedron 50:3993-4000.
- Verbist, J. F. 1993. Pharmacological effects of compounds from echinoderms. In M. Jangoux and J. M. Lawrence (eds.), Echinoderm Studies 4, pp. 111-186. Balkema Press, Rotterdam.
- Vermeij, G. J. 1978. Biogeography and adaptation. Harvard University Press, Boston.
- Voss, J. 1988. Zoogeography and community structure analysis of macrozoobenthos of the Weddell Sea (Antarctica). Rep. Polar Res. 45:1-145.
- White, M. G. 1984. Marine benthos. In R. M. Laws (ed.), Antarctic ecology, Vol 2. pp. 421-461. Academic Press, London.
- Winston, J. E. and A. W. Bernheimer. 1986. Hemolytic activity in an antarctic bryozoan. J. Nat. Hist. 20:369-374.
- Yang, A., B. J. Baker, J. E. Grimwade, A. C. Leonard, and J. B. McClintock. 1995. Discorhabdin alkaloids from the Antarctic sponge *Dendrilla membranosa*. J. Nat. Prod. 58:1459–1462.
- Yoshida, W. Y., P. J. Bryan, B. J. Baker, and J. B. McClintock. 1995. Pteroenone: A defensive metabolite of the abducted pteropod *Clione antarctica*. J. Org. Chem. 60:780-782.