Identification of bacteria associated with a disease affecting the marine sponge *Ianthella basta* in New Britain, Papua New Guinea

James M. Cervino1,*, Kathryn Winiarski-Cervino2, Shawn W. Polson3, Thomas Goreau4, Garriet W. Smith5

1Department of Biological Sciences, Pace University, 1 Pace Plaza, New York, 10038, USA
2The New York Academy of Medicine, New York, 10029–5293, USA
3Medical University of South Carolina, Charleston, South Carolina, 29425, USA
4Global Coral Reef Alliance, 37 Pleasant St., Cambridge, Massachusetts, 02139, USA
5Department of Biology and Geology, University of South Carolina, Aiken, South Carolina, 29801, USA

**ABSTRACT:** *Ianthella basta* marine sponges in Kimbe Bay, west New Britain, Papua New Guinea were affected by a disease, and exhibited high mortality, between 1996 and 2000. These fan-shaped sponges were mottled with brown lesions, rotted tissue and large holes. The decayed tissue was surrounded by brown biofilm that smothered the ostia openings. Since 1996, *I. basta* suffered its highest mortality at 3 sites within 16–20 km of the shore of west New Britain. No mortality was observed at 3 other locations further from shore (between 27–41 km), nor at 10 sites located more than 41 km from shore outside of Kimbe Bay in deeper waters, nor at the site nearest to shore. Comparison of the carbon source utilization patterns of 99 bacterial isolates from all healthy and diseased sponges revealed 5 species of bacteria specifically present in diseased *I. basta*. These bacteria were not present in healthy sponge samples. Bacteria isolated from affected sponges and inoculated onto healthy sponges caused disease signs similar to those in field specimens. The 16S rRNA genes from these bacteria were sequenced and found to correspond with 2 species of *Bacillus* and 3 species of *Pseudomonas*. The closest relatives of these bacteria based on BLAST searches included many terrestrial pathogens and species that are used as pathogens against insects and fungi in integrated pest control management. The bacteria causing disease in *I. basta* may thus originate from pesticides applied to agricultural land, predominantly oil palm plantations, in west New Britain. The possibility that these bacteria can pass virulence factors to marine bacteria through horizontal gene transfer needs to be investigated, as this may have unexpected impacts on marine ecosystems.

**KEY WORDS:** Sponge disease · 16S rRNA · *Pseudomonas* · *Bacillus* · Pesticide · *Ianthella basta*
Webster et al. (2002). Rützler et al. (1988) reported that sponge mortality has been attributed to cyanobacterial infection. Recent reports indicate that temperature stress (Cervino et al. 2004, Goreau & Hayes 2005) is having a severe impact on important species in tropical coral reef ecosystems, concurrent with an increase in coral and sponge diseases (Goreau et al. 1998, Wulff 2000, Porter et al. 2001, Webster et al. 2002). Because the precise mechanisms of these mortalities have mostly not been identified (Green & Bruckner 2000), many of these diseases affecting hermatypic corals are currently referred to as syndromes (Goreau et al. 1998). Some of these syndromes have been suggested as being triggered by unusual stresses from natural or man-made causes such as temperature fluctuations, sedimentation, and pollution (Harvell et al. 1999, Cervino et al. 2004). *I. basta* is one such marine organism that this research found to be affected by a disease.

In 1996 we noticed scattered and sparse lesions on *Ianthella basta* during a 1996 coral health monitoring survey in west New Britain, the largest island in PNG (Fig. 1). Subsequent monitoring trips in the Kimbe Bay study locations from 1998 to 2000 showed a decline in the health and abundance of *I. basta*. The appearance of this pathology in *I. basta* is similar to widespread disease in Caribbean Sea fans that is caused by the soil fungus *Aspergillus* (Smith et al. 1996). This paper documents the bacteria associated with diseased sponges in PNG. The sponge decline corresponds to a decrease in reef fish and coral cover abundance reported in a recently published 8 yr study of Kimbe Bay (Jones et al. 2004).

**MATERIALS AND METHODS**

**Fieldwork.** In August of 1996, 1998, and 2000, 7 in-shore and 10 remote reef sites at latitude 5.5° S and longitude 150.1° E were visited in Kimbe Bay to track the extent of necrosis among the *Ianthella basta* population and to sample the sponges in order to identify microbes that may be associated with the necrosis. To conduct the surveys, we used the belt transect method while diving with SCUBA. Belt transects allow for an

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**Fig. 1.** Two examples of lesions from *Ianthella basta* taken within Kimbe Bay (Restoff Island) over time
instantaneous survey of a predefined area within which the number of organisms of interest are recorded. For this research, multiple belt transects of 10 m in length by 1 m in width were performed at each reef site wherever sponges were seen over the course of a 45 to 50 min dive. This width is narrow enough for the observer to be able to easily estimate the distances being measured, while allowing for a large enough sample area. Transects were conducted at depths of approximately 3 and 16 m.

To perform the belt transects, a waterproof, tape-measured line was stretched horizontally at each location where sponges were witnessed. It should be noted that *Ianthella basta* sponges grow in sparse quantities and in scattered locales compared to many corals and other reef-based organisms. As 1 diver held onto the roll-up tape measure, the other swam forward with the tape at a constant rate for a distance of 10 m, counting the number of healthy and necrotic sponges visible in a 1 m width and recording the data on an underwater slate. Sponge colonies in Kimbe Bay were photographed in August of 1996, 1998, and 2000 using an Underwater Nikon RS (SLR System). The average lesion size on each affected sponge was measured.

Sponge tissue samples were collected in 2000 using SCUBA. Several types of samples measuring 5 cm² were cut from the unhealthy sponges using stainless steel scissors. First, samples of both degenerated and healthy sponge tissue were taken from within the circumference of the lesion on affected sponges. Zones of degenerated tissue often contain some healthy tissue within their boundaries. Normal tissue was also cut from the degenerated sponges, from a zone near to, but outside of, the decayed tissue area. Samples were also taken from healthy *Ianthella basta* sponges with no visible sign of necrosis. At each survey site, 25 samples of normal sponge tissue and 25 samples of affected sponge tissue were collected, all from different sponges. The samples were first placed in plastic Ziploc bags while underwater, then transferred to 100 ml polyethylene bottles containing fresh filtered (0.5 µm) seawater and kept in ice coolers. The tissue samples were stored at a temperature of ~2°C for future analysis.

**Lab work.** Researchers placed a 10 ml needle-less syringe next to the sponge’s healthy and diseased surfaces and applied gentle suction to collect the surface mucus and possibly particles of necrotic tissue (Fig. 2a,b). Upon surfacing, syringe samples were placed on ice in coolers (2°C) on the boat. The samples were transferred to media plates with Glycerol Artificial Sea Water (GASW). All samples were transported by airplane while being kept in a cooler. Upon return to the laboratory, samples were re-plated, streaked and separated to obtain pure colonies and incubated on Glycerol Artificial Sea Water (GASW), in duplicate. Plates were then incubated for 72 h at 28°C. Bacteria were re-isolated from the plates, chosen to represent the distinct morphological and growth characteristics such as color and colony shapes. Those isolates were then sub-cultured to obtain pure cultures. In total, 99 pure cultures were isolated, 35 originating from control sponge samples (the healthy samples) and 64 originating from diseased sponge samples. A 48 h agar culture of each isolate was then subjected to Carbon Source Utilization Pattern (CSUP) analysis (Ritchie et al. 2001) in order to cluster bacteria based on their ability to utilize a panel of sole carbon sources. This was accomplished using BIOLOG GN1 96-well MicroPlates.
(Biolog 1989). Resulting data from the tests with the isolates from *Ianthella basta* were organized into a dendrogram using the ML3 BIOLOG software clustering analysis program, MLClust, with default settings. The 16S of the rRNA gene sequencing was used to identify the microbes from groups of interest (Godfrey et al. 2002). DNA isolation from the bacterial isolates was followed by PCR amplification of the 16S rRNA gene using universal bacteria 16S rRNA gene primers: 27F (5'-AGA GTT TGA TCM TGG CTC AG-3') and 1492R (5'-TAC GGY TAC CTT GTT ACG ACT T-3'), and sequenced using primers 27F, 536F (5'-GTG CCA GCC GCG GTA ATW C-3'), 928F (5'-TAA AAC TYA AAK GAA TTG ACG GG-3'), 907R (5'-GCC CCC GTC AAT TCM TTT RAG TTT-3') (Weidner 1996), and/or 1492R. All 16S rDNA sequences were then subjected to BLAST searches of the National Center for Biotechnology Information nucleotide database (Altschul et al. 1997) and Sequence Match analysis using the Ribosomal Database Project's (RDP) database (Cole et al. 2003). The dendrograms displaying phylogenetic positions of bacteria isolated in this study were based on alignment of 1200 bp (*Bacillus* spp.) and 953 bp (*Pseudomonas* sp.) of the 16S rRNA gene. Sequence alignments were performed with DNA Star's Lasergene MEGALIGN software (v5.06) using the CLUSTALW method. The tree was generated by the neighbor-joining method (Saitou et al. 1987) with sequence dissimilarity determined by the method of Jukes & Cantor (1969) using MEGA3 software (v3.1). Bootstrap values are based on 500 replicates. Reference strain sequences were obtained from the RDP (Cole et al. 2003). The 16S rRNA sequences generated during this project were deposited in GenBank and assigned accession numbers as follows: SDC2A (DQ323743, DQ323744), SDG1A (DQ323745), SDA3A (DQ323746), SDA21A (DQ323747), and SDB21A (DQ323748).

**Infection experiments.** Six healthy sponge samples were collected at 3 m depth in Kimbe Bay and immediately brought back to Mahonia Na Dari research laboratory and placed in 750 l tanks for 2 d to monitor the health of the sponges. Each aquarium was filled with fresh seawater and temperatures were kept constant at 27°C. Sponges were held at the bottom of the aquarium with clothespins and mounted to stay weighted down at the bottom of the aquarium tank. Water quality was monitored and kept within the following ranges: temperature of 27 ± 2°C, pH of 8.1 to 8.3, salinity of 35 ± 2 g l⁻¹, and dissolved oxygen of 8.0 ± 0.1 mg l⁻¹. Filtration was conducted using biological/mechanical ‘box’ type filters. Inoculation experiments were commenced on the third day. Single-isolate inoculation tests were not successful in showing the disease signs; therefore, all bacterial inoculations included the 5 bacterial species during the second trial. Bacterial isolates were cultured on GASW media and then transferred to small, 2 cm cut-triangles of fresh media. Each triangle contained all 5 species. No bacteria were swabbed onto the control media. Each media triangle was held on the sponge surface for 72 h, attached with a needle and thread. Each of the 6 sponges had 4 triangles on them, 2 had no bacteria (representing controls), and the other 2 had the 5 bacteria swabbed onto the media triangle (Fig. 3).

**RESULTS**

Field observational data from 1996, 1998, and 2000 show that the sponge infections spread increasingly further within and between sponges as time passed. Seven study sites in Kimbe Bay were visited in each of the 3 yr for which data were collected. They are as follows with their respective distances from shore: Christine’s Reef, 15 km; Restoff Island, 16 km; Vanessa’s Reef, 17 km; Jeffrey’s Reef, 20 km; Inglis Shoal, 27 km; Joelle’s Reef, 32 km; and Kimbe Island, 41 km (Fig. 4). An additional 10 dive sites located more than 41 km from shore (beyond Kimbe Island) were visited in 1998, and *Ianthella basta* sponges there showed no signs of necrosis. Of the 7 sites visited within Kimbe Bay during 1996, only 4 of 28 sponges exhibited lesions (14%). The average lesion size was 0.11 ± 0.28 cm² (Fig. 5). During 1998, many more holes or lesions were observed on sponges at 3 study sites close to shore. The average lesion size was much larger than in 1996, increasing to 6.48 ± 3.69 cm² on average (Fig. 5). The lesions develop within a biofilm and tangled mat of filamentous cyanobacteria (Fig. 6). Upper edges and large central zones on degenerated sponges appeared rotted, with thinned lattice, brown discoloration, and in some cases with purplish blue cyanobacteria filaments. The most
severely affected sites observed in 1998 were: Restoff Island (78% of the sponges had lesions, or 32 of 41 sponges counted in 5 transects), Jeffrey’s Reef (27% with lesions, or 20 of 93); and Vanessa’s Reef (15% with lesions, or 13 of 88) (Fig. 7). No lesions were evident on *Ianthella basta* at reef study sites located more than 20 km from shore.

By August 2000, tissue decay was evident on a larger quantity of *Ianthella basta* sponges at reefs within 20 km of shore. Lesions became more prevalent at Restoff Island, Vanessa’s Reef and Jeffrey’s Reef. At Restoff, 94% of *I. basta* had lesions (48 of 51); at Vanessa’s, 33% of *I. basta* had lesions (31 of 94); and Jeffrey’s, 30% of *I. basta* had lesions (31 of 105). As of 2000, the average lesion size had increased to 36.39 ± 16.54 cm² (Fig. 7). No lesions appeared on *I. basta* at the 4 other reef sites, 3 of which were located further from shore than the impacted reefs (Inglis Shoal, 27 km; Joelle’s Reef, 32 km; and Kimbe Island, 41 km). Unexpectedly, no lesions were found on *I. basta* at Christine’s Reef, which is closer to shore (15 km) than the other reef study sites. Reports from Mahonia Na Dari Research Center, a scientific research facility located on the Kimbe Bay shoreline, indicate that *I. basta* sponges located at these 4 unaffected reefs remained healthy throughout 2000, with no lesions appearing on their surfaces.

### Bacterial isolates

Carbon Source Utilization Pattern (CSUP) analyses of the bacterial isolates revealed 3 groups of bacteria present in diseased samples of *Ianthella basta*, that were not represented in healthy sponge samples (Fig. 8). These 3 metabolic groupings were termed B, I, and J based on their positions on the CSUP dendrogram. Groups I and J were found in 4 isolates each and, despite being clustered next to each other on the...
dendrogram, share only ~20% metabolic similarity. Group B is remote from both groups I and J, sharing only ~2% similarity with those groups, and was present in 11 isolates (Table 1).

Sequencing of the 16S rRNA genes from 5 bacterial isolates belonging to Groups B, I, and J showed that they correspond with the genera *Bacillus*, which is ubiquitous in the environment, and *Pseudomonas*, which is common in many habitats including soil and water. The closest bacterial strain matches, when compared to the Ribosomal Database Project II at Michigan State University (Cole et al. 2003) using Sequence Match Analysis, are as follows:

1. Isolate SDG1A: *Pseudomonas* sp. SMCC DO715. SDG1A hails from the J group as identified in the CSUP analyses (Fig. 9).
2. Isolate SDC2A: *Pseudomonas pseudoalcaligenes*. SDC2A is a member of the B group (Fig. 9).
3. Isolate SDA21A: *Pseudomonas* sp. MBIC2027. SDA21A is a member of the B group (Fig. 9).
4. Isolate SDB21A: *Bacillus cereus, thurengiensis/anthracis*. SDB21A is a member of the B group (Fig. 10).
5. Isolate SDA3A: *Bacillus pumilus*. SDA3A is a member of the I group (Fig. 10).

Infection experiments from the 2 sponges revealed paling of the sponge tissue as seen during thermal bleaching in 2 out of the 6 sponges. The remaining sponges showed necrotic decay similar to field specimens. However, complete holes or openings were not established during the course of this experiment. Ostial openings were smothered with brown mucus surrounded by blue-green algae. The control triangles showed no signs of disease. Trying different combinations of the bacteria was not an option due to time con-

<table>
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<th>Group</th>
<th>Total isolates</th>
<th>Healthy (%)</th>
<th>Diseased (%)</th>
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<td>0</td>
</tr>
<tr>
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<tr>
<td>Total</td>
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Fig. 6. *Ianthella basta*. Detail of lesions around the ostial pores

![Image of sponge with lesions](image-url)

<table>
<thead>
<tr>
<th>Location</th>
<th>Disease Progression (%)</th>
</tr>
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<tbody>
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<td>Restoff 1998</td>
<td>78</td>
</tr>
<tr>
<td>Restoff 2000</td>
<td>94</td>
</tr>
<tr>
<td>Vanessa's 1998</td>
<td>15</td>
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<td>Vanessa's 2000</td>
<td>33</td>
</tr>
<tr>
<td>Jeffrey's 1998</td>
<td>27</td>
</tr>
<tr>
<td>Jeffrey's 2000</td>
<td>30</td>
</tr>
</tbody>
</table>

Fig. 7. *Ianthella basta*. Time series of disease progression between 1996 and 2000
Fig. 8. Dendrogram constructed from carbon source utilization patterns. Each item in the dendrogram represents a bacterial isolate. Isolates shaded light gray were from diseased samples. All other samples are from healthy sponges.

Fig. 9. NCBI accession numbers of non-type strains are parenthetically noted. Strains isolated in this study (I); Type strains (T). Dendrogram showing the results of 16S rRNA gene sequencing of bacterial isolates from *Bacillus* spp. Phylogenetic positions of these putative pathogens isolated in this study are based on 16S rRNA gene sequence.

Fig. 10. NCBI accession numbers of non-type strains are parenthetically noted. Strains isolated in this study (I); Type strains (T). Dendrogram showing the results of 16S rRNA gene sequencing of bacterial isolates from *Pseudomonas* spp. Phylogenetic positions of these putative pathogens isolated in this study are based on 16S rRNA gene sequence.
strains and weather conditions. Therefore, since we were not able to obtain results using single isolates within 48 h, we immediately applied all 5 bacterial groups together. There was no opportunity to re-isolate the bacteria that caused the lesions in aquaria, and DNA comparison of bacterial communities were not sampled and re-tested. Therefore, these results show only infectivity and are not complete confirmation of Koch’s postulates. Samples were brought back to the University of South Carolina Coral Pathology laboratory for further testing; however, all specimens were in advanced stages of stress, preventing further re-testing of this experiment for accurate analysis.

**DISCUSSION**

**Environmental patterns and disease**

The inner fringing reefs of southwestern Kimbe Bay are being subjected to various stress factors that cause coral reef decline, including sedimentation, algal overgrowth, and rising sea surface temperatures. The inner reefs are particularly impacted by river-transported sediments and fertilizer, which enter the waterways from land that is being increasingly deforested and developed for palm oil plantations (S. Seeto pers. comm.). The inshore fringing reef area is quite turbid and muddy as a result, and a high sediment load covers barrel sponges and corals on the reefs closest to land. The sediment and algal abundance are much lower on the land-facing side of the offshore bank reefs, located only 50 to 100 m offshore; and are nearly absent on the seaward side of those banks.

Between 1996 and 2000, lesions on *Ianthella basta* and die-offs of those sponges increased by 90% at Restoff Island. The bases of the sponges, or ‘holdfasts’, appeared wilted and unable to support their structures. Although the focus of this research was *I. basta*, degenerated tissue witnessed on other sponge species is relevant. Degenerated tissue and sponge bleaching was also seen on the undersides and bases of *Jaspis* sp., *Xestospongia* sp. and *X. testudinaria* (*Haplosclerida*) at all locations. Upper portions of these sponges appeared normal but their bases were rotted away. Sponge bleaching was also seen during September 1998, coincident with coral bleaching (J. M. Cervino & T. Goreau pers. obs.).

It is worth noting that 2 of the affected sites closest to shore (Restoff and Vanessa’s) experienced the largest increase in degenerated sponges between 1998 and 2000. Observations of a less dramatic increase in degenerated sponges at some of the other study sites (Jeffrey’s Reef), and a lack of increase at the remaining sites, support a possible link between sponge disease prevalence and proximity to shore. The one exception to that theory during this research was the absence of diseased sponges at Christine’s Reef, the study site closest to shore. A confluence of 3 rivers that flow into Kimbe Bay may be a source of land-derived bacteria and pollutants for near-shore habitats (Fig. 4). We hypothesize that the ocean current patterns are taking the effluent north of Christine’s reef, which is the furthest south of all the study sites. This is one explanation for why no diseased sponges were found there. However, ocean current patterns for this specific area have not been directly determined.

**Bacterial isolates**

We studied the 3 distinct groups of bacteria that we found to be specific to the diseased samples of *Ianthella basta*, to investigate their possible role as pathogenic agents. Each bacterial isolate from the diseased sponge samples belonged to 1 of 3 CSUP Groups: 11 isolates belonging to CSUP Group B, 4 belonging to CSUP Group I, and 4 belonging to CSUP Group J. CSUP Group B included the genera *Bacillus* and *Pseudomonas*. CSUP Group I included the genus *Bacillus*, and CSUP Group J included the genus *Pseudomonas*. Interestingly none of the diseased samples had more than 1 bacterial pathogen group present: diseased samples that tested positive for Group B bacteria, for example, did not show the presence of bacteria from either Group I or J (Fig. 8). Still, the possibility that multiple pathogens may be acting in tandem to cause symptoms in *I. basta* should not be ruled out. Black Band disease of scleractinian corals has been shown to be caused by a consortium of microorganisms (Richardson 1996, Richie et al. 2001). A consortium of 4 unidentified *Vibrio* sp. was also found to induce the signs of yellow blotch/band disease in corals in the Caribbean Sea (Cervino et al. 2004).

**Bacillus**

Members of 2 bacterial genera were isolated only from diseased sponges, *Bacillus* and *Pseudomonas*. *Bacillus* is a genus of Gram-positive bacteria that are ubiquitous in soil, water, and airborne dust. They are also common in marine habitats (Johnson et al. 1994, Ivanova et al. 1999). Two *Bacilli* were isolated as possible pathogens from degenerated tissue of *Ianthella basta* infected tissue (Fig. 9). The first *Bacillus* isolated from diseased sponges was most closely related to the *B. cereus* group, which includes several species of bacteria which are closely related and difficult to differentiate, including *B. cereus*, *B. thuringiensis*, *B. anthracis*, and *B. mycoides*. This motile
bacterium is also found in marine samples (Kobayashi et al. 2004). Strains of *B. cereus* are used as pesticide to kill mosquitoes (Finlay et al. 1999). Another member of the *B. cereus* group, *B. thuringiensis* (Bt), is widely used in a broad-spectrum insecticide spray, and genetically engineered into plants to control pests (Hothe & Whiteley 1989, Lereclus et al. 1993, Huang et al. 1999). Bt is widely claimed to be a ‘natural’ and ‘safe’ insecticide and is commonly sprayed on a wide variety of crops to control insect pests (Baum et al. 1999). It is specifically recommended to control the major insect pest in oil palm plantations (Hoong et al. 1992), which occupy much of the lowland areas of New Britain. Bt bacterium is claimed to be safe because it only attacks specific insect pests (Syed & Shah 1976, Bledsoe et al. 2004) by producing a crystal protein toxin that kills the cells lining the gut of susceptible insects. However, recent evidence shows that Bt affects a wide variety of invertebrates, including crustaceans and nematodes (Ivanova et al. 1999, Helgason et al. 2000), and that Bt persists in soils and in waters (Jensen et al. 1994). It is therefore not surprising that if terrestrial insecticide sprays containing Bt are leached into Kimbe Bay, *I. basta* and other marine invertebrates might be affected.

The second strain of *Bacillus* sp. isolated from infected *Ianthella basta* showed closest homology to *B. pumilus*, which is also found commonly in soils. Strains of *B. pumilus* are used as a fungicide on many crops including soybeans (Ouoba et al. 2003). It is very interesting that both *Bacillus* strains associated with our diseased sponges are often used for agricultural pest control. This is especially pertinent to this study because the diseased and dying sponges were located in such close proximity to agricultural activity sites and to river mouths that drain farmed land. While further studies are recommended to track the sources of the bacteria, anthropogenic introduction via agricultural activities seems a convincing potential source.

**Pseudomonas**

Members of the genus *Pseudomonas* form part of a large, heterogeneous and ubiquitous group of highly versatile, metabolically bioactive colonizers of surfaces. Pseudomonads are Gram-negative chemoheterotrophs. Many strains are bioactive, fast-growing and are able to suppress or out-compete pathogenic and other deleterious microorganisms. *Pseudomonas* species are also opportunistic invaders of plants, and cause blight disease in bean plants as well as lethal blight in palms, inducing slow soft rot in plant tissue upon inoculation (Willis et al. 2001).

Three types of *Pseudomonas* were isolated only from diseased tissue (Fig. 7b). The first isolate is very close to *P. alcaliphila*, found in cold marine waters (Yumoto et al. 2003). It is also closely related to *P. mendocina* and *P. pseudoalcaligenes*, which are found in soils and waters and used in the degradation of aromatic hydrocarbons as well as being a bivalve and human pathogen (Lodeiros et al. 1992, Wang et al. 2001). The next closest relatives are the second *Pseudomonas* isolated from *Ianthella basta* diseased tissue and strain K49 (Wei et al. 1995). The consortium or interactions between bacterial species are similar to bacterial *hrp* genes, which encode a large set of regulatory proteins broadly conserved among plant and animal pathogens and constituting a type III secretion pathway known as the ‘*Hrp* pathway’ in phytopathogenic bacteria (Kim et al. 1997). It is also important to note that at least 2 proteins secreted via the *Hrp* pathway *hrp* genes are present in all gram-negative necrogenic plant pathogens and are created by transposon mutagenesis of *Pseudomonas syringae* (Bogdanove 2002). This leads to rapid tissue necrosis at the site of the pathogen, as appears to be occurring with *I. basta*. The kin group to the above strain is comprised of *P. alcaligenes*, a very common marine bacterium (Lorenz & Sikorski 2000) including forms that are toxic to fish (Kobayashi et al. 2004), shrimp, and scallops (Alavandi et al. 2004); *P. fulva*, found in leaf litter on soils (Luz et al. 2003) and used in fungicide sprays on rice and other crops (Xie et al. 2003); and *P. putida* and other *Pseudomonas* spp., which are also used as anti-fungal sprays and have forms that cause tomato disease (Pedley et al. 2004) and are used for biodegradation of DDT (Kamanavalli et al. 2004). The third *Pseudomonas* isolate belonged to a more distantly related group. It was related only to an uncultured bacterium, S17 sBac13m, for which no further information was found.

*Pseudomonas* species thrive under moist conditions in soil, particularly in association with plants, as well as in sewage sediments and aquatic environments (Maidigan et al. 1997). Surface water runoff and wind currents are among the environmental conditions that may affect the dissemination of *Pseudomonas* species. For example, conditions of high humidity and temperature (80 to 90% humidity, 27°C) favored the colonization of this species on lettuce and bean plants (OECD 1997). While the marine environment is hostile to many terrestrial microbial pathogens and will cause most to rapidly die off, as seen in the Great Barrier Reef (Webster et al. 2002), some of these pathogens may accumulate in filter-feeding organisms such as sponges (Webster et al. 2002, J. Lopez pers. comm.). In marine waters, species of *Pseudomonas* and other bacterial families have been detected in elevated numbers during fish kills associated with red tides (www.ukmarinesac.org.uk/index.htm).
A temperature increase has occurred in PNG waters as shown in satellite-derived sea surface temperature records of Kimbe Bay, studied since 1982 (Goreau & Hayes 2005) (Fig. 11). Recent overall coral mortality of offshore reefs at Kimbe Bay was estimated to be in the range of 10 to 20% in 2001, based on filmed transects of numerous sites during this study, and much of this may be due to bleaching events. A coral bleaching-like syndrome that correlates with temperature stress may also be caused by, or associated with, pathogens (Ben-Haim et al. 2003, Cervino et al. 2004). These coral diseases are more virulent and faster spreading at high temperatures (Cervino et al. 2004). Some of the sponge die-offs may be similarly linked to local temperature stress, or may be due to the dissemination of novel or newly adapted pathogenic agents in the aquatic environment (Vacelet 1994, Goreau et al. 1998, Harvell et al. 1999) or a combination of both pathogens and elevated temperature and nutrient enrichment (LaPointe et al. 1992, Ben-Haim et al. 2003, Cervino et al. 2004). Jones et al. (2004) recently conducted an 8 yr study which indicated that over 75% of reef fish species declined in abundance in Kimbe Bay in what the authors term a catastrophic decline, and showing that 50% of reef fish species declined to less than half of their original numbers. Field research also showed that there has been a decline in scleractinian coral cover from 66% in 1996 to a low of 7% in 2002. This corresponds to the time of our study (1996–2000). The authors also point out that this decline is attributed to anthropogenic habitat degrada-
tion as a result of pollution and thermal bleaching events. However, we note that Jones et al. (2004), studied only inshore fringing reefs, and did not examine the offshore reefs sites investigated in this Ianthella basta study. We note that the live coral cover in offshore reefs was an order of magnitude higher than that of the inshore reefs reported by Jones et al. (2004), and that our estimates of cover on inshore fringing reefs is similar to theirs. We therefore concur with Jones et al. (2004) that marine reserves are not by themselves an effective management strategy for protecting coral reefs and marine biodiversity from large-scale pollution or global warming.

**Future work**

The findings of this paper suggest that the bacteria causing Ianthella basta disease may be closely related to pathogenic terrestrial bacteria, some of which are widely applied in land-based agricultural activities. Direct tests of the possible links to land-based sources of pollution as well as to climate change need to be conducted. Future research will test pathogenicity of various isolates or combinations of isolates from the groups of putative pathogens that we have identified in infected tissues, followed by re-culture and sequencing of these isolates. The putative pathogens will be tested on healthy sponges in attempt to fulfill the requirement of Koch’s postulates (Koch 1882) in its completion; to date, we have only satisfied infectivity of the bacteria isolated from diseased sponges. The
potential inability to culture putative pathogens to satisfy Koch’s postulates should not prevent the pathology of *I. basta* from being classified as a disease, if a disease is defined as any impairment, interruption, cessation, proliferation or other disorder of vital cellular functions, systems or organs (Willis et al. 2001).

CONCLUSION

Microbiological analyses have shown that 5 distinct bacteria in the *Bacillus* and *Pseudomonas* families are strongly correlated with a disease that spread through the *Ianthella basta* sponge population in west New Britain, PNG from 1996 through 2000. Die-offs of *I. basta* may have increased during periods of elevated sea-surface temperatures. The die-offs were most abundant on reefs close to shore. Therefore, we speculate that land deforestation activities, bacterial pesticide pollution from agriculture, and thermal bleaching events may have introduced new pathogenic bacterial strains that affected marine sponges, allowing invasive cyanobacterial communities to flourish on the sponges’ ostial surfaces. A common pattern was seen in the 2 *Bacillus* species and 3 *Pseudomonas* species isolated only from disease sponge tissue. They were closely related to species that are known to be human pathogens, or that are widely used as pesticides and sprayed on many agricultural crops, including oil palms. These species were also closely related to known marine pathogens affecting fish and invertebrates. It is possible that horizontal gene flow from terrestrial species, some pathogenic to many organisms (Helgason et al. 2000, Lorenz & Sikorski 2000) and used as wide-spectrum pesticides, could occur into related marine species that could then infect marine invertebrates. If so, the allegedly ‘safe natural’ insecticides widely used in Integrated Pest Management could be having serious and unanticipated effects in marine ecosystems. These possible connections need to be thoroughly examined in future research.

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LITERATURE CITED


Biolog (1989) GN MicroPlate: instructions for use. Biolog, Hayward, CA


**Bacillus cereus** and **Bacillus thuringiensis**—one species on the basis of genetic evidence. Appl Environ Microbiol 66:2627–2630


Vacelet J (1994) Control of the severe sponge epidemic—Near East and Europe: Algeria, Cyprus, Egypt, Lebanon, Malta, Morocco, Syria, Tunisia, Turkey, Yugoslavia, technical report: the struggle against the epidemic which is decimating Mediterranean sponges, FT/TCP/RAB/8853, FAO, Rome, p 1–39


Wei ZM, Beer SVJ (1996) hrP activates *Erwinia amylovora* hrp gene transcription and is a member of the ECF subfamily of sigma factors. Bacteriol 177:6201–6210


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