Lesion bacterial communities in American lobsters with diet-induced shell disease

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ABSTRACT: In southern New England, USA, shell disease affects the profitability of the American lobster Homarus americanus fishery. In laboratory trials using juvenile lobsters, exclusive feeding of herring Clupea harengus induces shell disease typified initially by small melanized spots that progress into distinct lesions. Amongst a cohabitated, but segregated, cohort of 11 juvenile lobsters fed exclusively herring, bacterial communities colonizing spots and lesions were investigated by denaturing gradient gel electrophoresis of 16S rDNA amplified using 1 group-specific and 2 universal primer sets. The Bacteroidetes and Proteobacteria predominated in both spots and lesions and included members of the orders Flavobacteriales (Bacteroidetes), Rhodobacterales, Rhodospirillales and Rhizobiales (Alphaproteobacteria), Xanthomonadales (Gammaproteobacteria) and unclassified Gammaproteobacteria. Bacterial communities in spot lesions displayed more diversity than communities with larger (older) lesions, indicating that the lesion communities stabilize over time. At least 8 bacterial types persisted as lesions developed from spots. Aquimarina ‘homaria’, a species commonly cultured from lesions present on wild lobsters with epizootic shell disease, was found ubiquitously in spots and lesions, as was the ‘Candidatus Kopriimonas aquarianus’, implicating putative roles of these species in diet-induced shell disease of captive lobsters.

KEY WORDS: Shell disease · Aquimarina · Denaturing gradient gel electrophoresis · DGGE · Flavobacteria · Herring diet · Homarus americanus

INTRODUCTION

Shell diseases causing cuticle damage and melanization occur ubiquitously in crustaceans. In the late 1990s, epizootic shell disease (ESD) emerged in the American lobster Homarus americanus H. Milne Edwards, 1837 fishery off the southern coast of New England, USA (Castro & Angell 2000), where it caused significant financial losses (Hsu & Smolowitz 2003). The average prevalence of ESD from south of Cape Cod to central Long Island Sound was approximately 20 to 30% from 1998 to 2005 (Cobb & Castro 2006). However, prevalence >70% has been recorded in ovigerous females in eastern Long Island Sound in 2002 (Cobb & Castro 2006). Lobsters that molt out of their shells affected by ESD usually survive (Smolowitz et al. 2005). However, lobsters affected even mildly by ESD are generally unmarketable due to the grotesque appearance of erosions in the dorsal cephalothorax and abdomen that extend laterally in an irregular pattern uncharacteristic of other known forms of shell disease (Smolowitz et al. 2005). Histological analyses of ESD lesions are typified by shallow erosions in the epicuticle and exocuticle, but in severe cases these erosions can penetrate into the uncalcified endocuticle or further into the underlying epithelium causing ulceration (Smolowitz et al. 2005).
Bacteria, fungi, protistans and nematodes have been observed in ESD lesions (Chistoserdov et al. 2005, Quinn et al. 2009). However, high numbers of bacteria generally occur at lesion edges and within pits in the lesions that develop in the cuticle (Hsu & Smolowitz 2003, Smolowitz et al. 2005). The culturable bacterial community existing within ESD lesions has been defined using various media (Chistoserdov et al. 2002, 2005). Of the bacteria identified, Pseudoalteromonas gracilis and members of the Flavobacteriaceae were particularly prevalent, with Shewanella spp., Vibrio spp., Pseudoalteromonas spp. and Alteromonas arctica less prevalent (Chistoserdov et al. 2005). Using DNA-based community profiling techniques, Aquimarina ‘homaria’ (Flavobacteriaceae) and a specific alphaproteobacterium belonging either to the genus Thalassobius or a sister genus (‘Marinosulfuromonas’) have been found to occur ubiquitously in ESD lesions (Chistoserdov et al. 2009). Culturing of bacteria from lesions of lobsters with impoundment shell disease has predominantly identified Vibrio, Aeromonas, Beneckea and Pseudomonas spp. (Malloy 1978, Stewart 1980, Getchell 1989, Smolowitz et al. 1992). Many of the aforementioned bacteria are known to be chitinoclastic (Chistoserdov et al. 2005). Koch’s postulates have not yet been fulfilled for any single bacterial species as the primary cause of any shell lesion type. Thus, the possibility exists that a complex microbial community might become established in and induce erosions as a result of specific environmental and host factors that compromise lobster health, carapace composition or immune competence (Smolowitz et al. 2005, Castro et al. 2006, Tlusty et al. 2007). One such factor that can induce shell disease and mortality in juvenile Homarus americanus is inadequate diet (Tlusty et al. 2008). A diet of herring Clupea harengus has been shown to induce high levels of shell disease and mortality and a mixed diet of crabs, mussels and commercial shrimp feed and diets containing 50% herring also induce shell disease, although at slightly reduced prevalence (Tlusty et al. 2008).

Here we use the diet-induced shell disease (DISD) method developed by Tlusty et al. (2008) to track the constituents of bacterial communities that become established in spots and lesions induced in juvenile Homarus americanus. To do this, PCR with universal- and group-specific 16S rDNA primer sets was used in conjunction with denaturing gradient gel electrophoresis (DGGE) to compare band profiles and to identify bacteria inhabiting spots and lesions of lobsters affected by DISD.

MATERIALS AND METHODS

Lobster rearing

Lobsters were reared from eggs at the New England Aquarium Lobster Research and Rearing Facility (Boston, MA). Larvae were fed Spirulina-enriched frozen adult brine shrimp (Artemia spp., San Francisco Bay Brand) and Cyclopeeze (Argent Chemical Laboratories). Upon metamorphosis to Stage IV, individual lobsters were placed into 4.5 cm diameter containers. From ~1 yr of age onwards, lobsters were fed a gelatin-based diet containing Progression 3 shrimp aquaculture feed (Salt Creek), krill meal, soy lecithin, various vitamins and minerals, bone meal, kelp meal, Spirulina algae, and frozen Spirulina-enriched adult Artemia.

Induction of shell lesions

At 3.5 to 4 yr of age (23.5 to 33.2 mm carapace length, 7.8 to 25 g body weight), 11 lobsters were housed individually in 13.5 × 7.2 × 9 cm lidded containers that contained side slots for water flow and top circular holes large enough to add feed but prevent escape. The 11 containers were placed in a shallow tank (193 × 53 × 10 cm) employing a semi-closed re-circulating seawater system (15% water replacement daily) and maintained for 180 to 245 d; therefore, although physically separated, the animals shared a common water supply. Our experimental design induced a single outbreak of shell disease, so that the sample size is 1, with a subsample size of 11 lobsters. Lobsters serving as controls were fed a balanced diet as per Tlusty et al. (2008) and never developed DISD even though they were maintained in the same system. The water source and treatment is as described in Tlusty et al. (2008). Over this period, water salinity varied minimally (30.33 to 31.73‰) and temperatures ranged between 10.6 and 17.5°C. The diet was made by homogenizing 150 g frozen herring (Walcan Seafood) and mixing the slurry with 90 ml of hot dissolved Knox gelatin (7.1 g of gelatin in distilled water) to create water soluble blocks of food, which were stored at −20°C. Each lobster received a piece of frozen herring homogenate approximately half the size of its last abdominal segment each day, and any uneaten food was removed after 4 h.

Lobsters were observed for shell disease on Day 0 and Day 50 and bi-weekly thereafter until sampled. Because Tlusty et al. (2008) had shown that lobsters succumb to shell disease as severity increases, a
severity score system was used along with time since onset of disease to ensure lobsters had a sufficient degree of shell disease without a possibility of death. The system was based on the number of carapace locations exhibiting disease spots and/or lesions (lobster carapace was divided into 27 different sections). Using this system 3 lobsters (1–3) were sampled after 180 d, 4 lobsters (4–6 and 9) after 201 d, and 4 lobsters (7, 8, 10 and 11) after 245 d. Lobsters were rinsed in sterile seawater before either regions of normal carapace (6 lobsters) or regions displaying spots or lesions (all 11 lobsters) were scraped using a sterile razor blade. Typically material from all spots (~5.5 per animal) or all lesions (~5.7 per animal) were pooled for each animal. The entire carapace surface was scraped to sample unaffected sections of the carapaces. As less matter was visibly collected from normal carapace and spots compared to lesions, material was suspended in either 50 or 200 µl buffer (50 mM Tris-HCl pH 8.8, 1 mM EDTA), respectively.

**DNA extraction**

Suspended shell matter was made to 1 mg ml⁻¹ egg white lysozyme (Amresco®) and incubated at 37°C for 30 min before addition of sodium dodecyl sulphate to a 2% final concentration and 1.25 mg ml⁻¹ Proteinase K (Fisher Bioreagents®). Following digestion at 50°C for 15 min, the lysate was freeze-thawed (−80°C and 50°C) 3 times before being mechanically disrupted using 3 × 0.1 mm Zirconia/Silica beads in a Mini-Beadbeater™-8 (BioSpec Products). The pulverized lysate was then extracted with an equal volume of phenol buffered with Tris-HCl, pH 7 (Sigma-Aldrich) followed by an equal volume of chloroform (Fisher-Biotech), and DNA was precipitated at −80°C following addition of 1/10 volume 3 M sodium acetate pH 8.5 and 2.2 volume cold ethanol. Following centrifugation, DNA pellets were washed in 70% ethanol, dried and resuspended in 200 µl sterile ddH₂O and stored at −20°C.

**PCR**

For DGGE analysis, 16S rDNA variable regions V3, V4 and V5 were amplified by PCR using the universal primers 341FM (5'-CCT ACC GGD GGC WGC AG-3') and 907RM (5'-CCG YCW ATT CMT TTG AGT TT-3') modified slightly from sequences described previously (Muyzer 1999). The 5' GC-clamp, designed by Muyzer et al. (1993), was attached to the 5' terminus of the 341FM primer. The resulting primer is designated 341FM-GC. Each PCR (50 µl) contained 25 µl GoTaq® Green Master Mix (Promega), 1.5 µM 341FM-GC and 0.5 µM 907RM, an additional 1.0 mM MgCl₂ and 1.5 µl DNA estimated to contain either ~1 ng or ~10 ng DNA from the extractions of melanized spots or lesions, respectively. The touchdown thermal cycling conditions were 95°C for 5 min, 20 cycles of 95°C for 1 min, 65°C for 1 min touchdown −0.5°C per cycle to 55°C for 1 min, 72°C for 3 min, 15 cycles of 95°C for 1 min, 55°C for 1 min, and 72°C for 3 min followed by 72°C for 7 min. To amplify 16S rDNA regions V1 and V2 of bacteria, the universal primers 27F (5'-AGA GTT TGA TCM TGG CTC AG-3') and 355R (5'-GCT GCC TCG TCC CGT AGG AGT-3') were used. The 5' GC-clamp, designed by Muyzer et al. (1993), was attached to the 5' terminus of the 27F primer. The resulting primer is designated 27F-GC. The PCR was performed as above except using 0.5 mM of each primer and the thermal cycling conditions 95°C for 5 min, 30 cycles of 95°C for 30 s, 59°C for 30 s, and 72°C for 30 s followed by 72°C for 5 min. Bacteroidetes communities were amplified using primer 341FM-GC in conjunction with the Bacteroidetes-specific primer CFB721R (5'-CTG CCT TCG CAA TCG G-3') modified slightly from a sequence described elsewhere (Weller et al. 2000). Thermal cycling conditions were 95°C for 3 min followed by 30 cycles of 95°C for 30 s, 57°C for 45 s, and 72°C for 45 s followed by 72°C for 3 min. PCRs to amplify Aquimarina 'homaria' and an unidentified Bacteroidetes used primers designed to 16S rDNA sequences determined for these bacteria, and the primer sequences were unique compared to those of other bacteria found associated with ESD lesions or to any sequences deposited in GenBank or the Ribosomal Database Project (RDP) database. The A. 'homaria' PCR primers were AhomF (5'-CAG TAT TAC GTG TAA TAC T-3') and AhomR (5'-TCA ATG GCA ATT TTC CG-3') and the unidentified Bacteroidetes PCR primers were AcompF (5'-CCC ACC ACG TGT GGG-3') and AcompR (5'-AAT ATA ACT CAA GAC AAC AAC-3'). Thermal cycling conditions were 95°C for 3 min, 30 cycles of 95°C for 30 s, 48.5°C for 30 s for A. 'homaria' or 57°C for 30 s for Bacteroidetes, and 72°C for 30 s followed by 72°C for 3 min.

**DGGE and band processing**

Gel electrophoresis of PCR products employed a polyacrylamide-based DGGE system (CBS Scientific)
and 1× Tris-acetate EDTA buffer, pH 7.8 (Fisher-Biotech) at 60°C. All DGGE gels (20×17.6 cm, 1.5 mm thick) were prepared using 6% polyacrylamide and concentration gradients of 100% denaturing solution (7 M urea and 40% formamide). Gradients of 20 to 80% were employed in gels to resolve 341F+907R PCR products, 20 to 65% to resolve 27F+355R PCR products and 35 to 60% to resolve 341F+C35721R PCR products. Gels were subjected to electrophoresis at 80 V at 60°C for 14 h, stained with 0.1 mg ml⁻¹ ethidium bromide for 20 min and banding patterns were visualized using a UV transilluminator. Bands of interest were excised using a scalpel blade, placed into microcentrifuge tubes containing 0.2 g sterile 2 mm glass beads (Biospec Products) and 500 µl ddH₂O and pulverized at high speed for 3 min using a Mini Beadbeater™ (Biospec Products) before being stored at 4°C overnight for DNA extraction.

DNA sequencing

Aliquots (1 µl) of excised bands pulverized in ddH₂O were reamplified by PCR using the same reverse primer in conjunction with a forward primer of the same sequence except for the absence of the 5‘GC-clamp. These PCRs (50 µl) contained 25 µl GoTaq® Green Master Mix, 0.5 µM each primer and 1.5 µl DNA and used the thermal cycling conditions described earlier. PCR products were purified using the Wizard® SV PCR Clean-Up System (Promega) and sequenced using one of the aforementioned forward primers and a BigDye® terminator V3.1 cycle sequencing kit. Nucleotide sequences were generated using a 3130xl Genetic Analyzer (Applied Biosystems). BLASTn (Altschul et al. 1997) was used for nucleotide sequence searches of GenBank and the RDP database (http://rdp.cme.msu.edu) to assign phylogenies. Bacterial 16S rDNA sequences were deposited in GenBank (Accession nos. JF297188–JF297220 and JF304026), analyzed using Lasergene 5.3 (DNA*Star) and examined for possible chimeras using the RDP release 8.1 CHIMERA_CHECK program (rdp.cme.msu.edu).

Statistical analysis of community profiles

Jaccard’s similarity index was used to quantify similarities of DGGE profiles of the microbial community identified among the melanized spots and lesions examined (Jaccard 1908). The index measures similarity based on the number of DGGE bands shared between 2 samples divided by the number of bands identified as unique in the 2 samples (Real & Vargas 1996). Gel images were digitized and TotalLab TL100 version 2008.01 1D gel analysis (TotalLab) was used to identify shared and unique bands. Because band similarities of 1 sample were determined against several other samples, a randomization approach was used to evaluate their statistical significance compared to averaged index values determined for either spot samples or lesion samples (Manly 1991).

Individual bands from the lesion and spot samples were pooled into a single sample (this is comparable to assuming there is no difference between the samples). Individuals were then randomly reassigned to 2 samples, of the same sizes as the original spot and lesion samples. The average similarity index was then calculated for these 2 new sets of randomized samples, and the difference between the average similarity was obtained. Procedures in Statistical Analysis System (SAS 2008) for randomizing data sets were used to generate these differences in average index values for 10 000 data permutations. These 10 000 differences between similarity indices represent the variation in differences between index values that would be expected if there was no difference in the community similarities of the lesson and spot samples. If the difference between average similarity index values observed from our original samples was >95% of values obtained through randomization, we can conclude that the observed differences in community similarities were statistically significant (Manly 1991).

Procedures in SAS (2008) for randomizing data sets were used to generate index values for difference distributions for 10 000 data permutations.

Histology

Tissue pieces surrounding individual spots were sampled from 2 lobsters and processed for histology. Individual lesions were sampled from 6 lobsters for histology. These individuals were separate from those sampled for DNA extraction to ensure their morphology was not compromised. The carapace was removed by inserting a scalpel and making a circular cut at distance of ~3 mm away from a lesion or a spot. The scalpel penetrated through the shell into the hemocoel to cut out approximately 2 to 3 mm thick slices perpendicular to the surface of the lesion. The resulting shell slices also contained the attached underlying epithelium and connective tissues. Tis-
sues fixed in 10% formalin in seawater were decalci-
ified using an EDTA solution pH 8.0, processed into
paraffin, 5 µm sections were stained using hema-
toxylin and eosin (Howard et al. 2004) and lesion
pathology observed was described using terminol-
ogy defined by Smolowitz et al. (2002).

RESULTS

Shell disease development

Among the 11 lobsters fed exclusively on herring,
the times at which shell disease developed varied
from 48 to 201 d (mean ± SD = 90.1 ± 51.9 d). Lobsters
initially presented melanized spots <1 mm diameter
with no conspicuous signs of shell erosions, and these
generally developed over time into larger, more
advanced melanized lesions in which the shell had
eroded (Fig. 1). Lesions most often developed on the
chelipeds, uropods, and carapace. While most spots
appeared as uniform circles, lesions on the cutting
and crushing regions of the cheliped were often elon-
gated along the length of the claw. Over the duration
of the trial the general health of lobsters declined as
evidenced by them losing chelipeds and becoming
moribund.

Histopathology

Histology on single lesions from 6 lobsters showed
severe erosions that mostly extended through the
uncalciﬁed endocuticle but did not penetrate
through the epithelium (Fig. 2). They also displayed a
thick surface pseudo-membrane and an underlying
new cuticle layer formed by the cuticular epithelium.
All pseudo-membrane surfaces and adjacent degener-
ating carapace regions were melanized. Mats of
bacteria occurred on some lesion surfaces (Fig. 2A)
and small unidentified protozoans often occurred in
conjunction with bacteria on surfaces of deep ero-
sions where the cuticle had degenerated. The single
spots examined from 2 lobsters were melanized to
varying degrees, one at the epicuticle and superficial
exocuticle surfaces and the other at deeper areas of
the degenerated exocuticle (Fig. 2B). Bacterial mats,
individual bacteria and some small protozoans were
noted in affected areas of the spots.
**Primer bias**

Due to known amplification biases of the universal primers (Hong et al. 2009), 2 universal primer sets were tested. When applied to lesions, the 341FM+907RM primer set amplified both Bacteroidetes and Proteobacteria, whereas the 27F+355R primer set appeared to favor Proteobacteria (Figs. 3 & 4). For example, with the lesion from Lobster 5, the 341FM+907RM primer set clearly amplified at least 4 different species of Bacteroidetes, whereas the 27F+355R primer set amplified only 1. In contrast, with the lesion from Lobster 2 where the 341FM+907RM primer set amplified 7 Proteobacteria species, the 27F+355R primer set amplified a much higher diversity (up to 19 bands) of Proteobacteria. Both primer sets were thus used across all lesion and spot samples to ensure more complete coverage of all major bacterial lineages, and a second Bacteroidetes-specific primer set was used to ensure good resolution of this group due to the bias of the 27F+355R primer set.

**DGGE analysis of healthy carapace and melanized disease spots**

No PCR products were amplified from DNA extracted from healthy carapace scrapings using either the Bacteroidetes group-specific or the 2 universal primer sets, indicating the presence of few if any bacteria. In disease spots from juvenile lobsters with DISD, 16S rDNA PCR in conjunction with DGGE and band sequence analysis identified bacteria from the Bacteroidetes and Proteobacteria phyla and 1 species of Actinobacteria. The 341FM+907RM primer set amplified both Bacteroidetes and Proteobacteria, whereas the 27F+355R primer set appeared to favor Proteobacteria (Figs. 3 & 4). For example, with the lesion from Lobster 5, the 341FM+907RM primer set clearly amplified at least 4 different species of Bacteroidetes, whereas the 27F+355R primer set amplified only 1. In contrast, with the lesion from Lobster 2 where the 341FM+907RM primer set amplified 7 Proteobacteria species, the 27F+355R primer set amplified a much higher diversity (up to 19 bands) of Proteobacteria. Both primer sets were thus used across all lesion and spot samples to ensure more complete coverage of all major bacterial lineages, and a second Bacteroidetes-specific primer set was used to ensure good resolution of this group due to the bias of the 27F+355R primer set.

**Fig. 3.** Denaturing gradient gel electrophoresis showing 16S rDNA bands amplified from lesion bacterial communities using the universal PCR primer set 341FM+907RM. Lane numbers 1 to 6 identify individual lobsters *Homarus americanus*. ‘C’ bands were amplified from both spots and lesions and ‘L’ bands were amplified only from lesions as described in Tables 1 & 2, respectively.

**Fig. 4.** Denaturing gradient gel electrophoresis showing 16S rDNA bands amplified from (A) lesion and (B) spot bacterial communities using the universal primer set 27F+355R. Lane numbers 1 to 11 identify individual lobsters *Homarus americanus*. ‘C’ and ‘L’ bands are labeled as in Fig. 3, and ‘S’ bands were amplified exclusively from spot bacterial communities as described in Table 3. Lane Ah includes reference bands of (A) *Aquimarina homaria* I32.4, and (B) *A. homaria* I32.4 and *Thalassobius* sp. I31.1.
Table 1. List and bacterial phylogenies inferred from 16S rDNA sequences of denaturing gradient gel electrophoresis (DGGE) bands amplified by PCR from DNA extracted from scrapings of single spots and single lesions of individual diet-induced shell disease-affected juvenile lobsters *Homarus americanus*. Sample sizes show number of individual lobsters from which samples were obtained. Band designations correspond to Figs. 3, 4 & 5, although not all samples from primer set 341FM-GC+907RM are shown in Fig. 3. BLAST: basic local alignment search tool; RDP: Ribosomal Database Project; na: not available.

<table>
<thead>
<tr>
<th>Band</th>
<th>Closest BLASTn match (Accession no.)</th>
<th>Presence in DGGE of PCRs using various primer sets</th>
<th>RDP-based phylogeny</th>
<th>Accession nos.</th>
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<td></td>
<td></td>
<td>(n=10)</td>
<td>(n=11)</td>
<td>(n=11)</td>
</tr>
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<td>C1</td>
<td><em>Aquimarina macrophalphi</em> (ABS17144)</td>
<td>9</td>
<td>8</td>
<td>11</td>
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<td>C2</td>
<td>Uncultured bacterium clone EPR9368-08a-bc46 (EU497113)</td>
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<td>0</td>
<td>10</td>
</tr>
<tr>
<td>C3</td>
<td><em>Flavobacteriaceae</em> bacterium ACEMC 10-9 (FM162878)</td>
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<td>0</td>
<td>6</td>
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<td>C4</td>
<td><em>Aquimarina</em> sp. MOLA 71 (EU345609)</td>
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<td>2</td>
<td>3</td>
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<td>C5</td>
<td>Uncultured bacterium clone AR87 (F3656480)</td>
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<td>na</td>
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<tr>
<td>C6</td>
<td><em>Koprimonas byunusaeensis</em> (DQ162724S)</td>
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<td>na</td>
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<tr>
<td>C7</td>
<td>Uncultured <em>Rhodospirillaceae</em> bacterium BME47 (DQ917824)</td>
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<td>C8</td>
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<td>C9</td>
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<td>7</td>
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confirmed following re-amplification. An unclassified Flavobacteriaceae sp. (Band C2) was amplified from 10 of 11 spots (Fig. 5). Other Bacteroidetes identified relatively frequently included the Krokinobacter sp., an unclassified Aquimarina sp., and various other Flavobacteriaceae (Table 2).

**DGGE and PCR analyses of bacterial communities from disease lesions**

In lesions of juvenile lobsters with DISD, 16S rDNA PCR in conjunction with DGGE and band sequence analysis identified bacteria from the Bacteroidetes

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**Table 2. List and bacterial phylogenies inferred from 16S rDNA sequences of denaturing gradient gel electrophoresis (DGGE) bands amplified by PCR from DNA extracted from scrapings of single lesions of individual diet-induced shell disease-affected lobsters Homarus americanus.** Sample sizes show number of individual lobsters from which samples were obtained. Band designations correspond to Figs. 3, 4 & 5. BLAST: basic local alignment search tool; RDP: Ribosomal Database Project

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<th>Presence in DGGE of PCRs using various primer sets</th>
<th>RDP-based phylogeny</th>
<th>Accession no.</th>
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<td>L1</td>
<td>Marine bacterium MSC5 (EU753120)</td>
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<td>Flexibacter aurantiacus subsp. Copepodarum (AB078044)</td>
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<tr>
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<td>JF297194</td>
</tr>
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</table>

**Fig. 5. Denaturing gradient gel electrophoresis showing 16S rDNA bands amplified from Bacteroidetes sp. present in (A) lesion and (B) spot bacterial communities.** Lane numbers 1 to 11 identify individual lobsters Homarus americanus and Lane Ah is Aquimarina homaria I32.4 reference DNA. Bands are labeled ‘C’, ‘L’ and ‘S’ as in Figs. 3 & 4 and as described in Tables 1, 2 & 3.
and Proteobacteria phyla. The Bacteroidetes were exclusively from the Flavobacteraeae, whereas Proteobacteria encompassed several families of Alpha- and Gammaproteobacteria.

As the 11 lobsters were held communally within a shallow tank but segregated as individuals within interconnected cubicles, bacterial communities that established in lesions of different lobsters were expected to be similar. Therefore, particular attention was paid to identifying the nature of bacteria that established commonly across the lobsters. Both universal primer sets amplified the Band C1 in a high number of lesions (Fig. 3, Table 1). Analysis of the 16S rDNA sequence of Band C1 from all 3 primer sets showed it to be identical to that of Aquimarina ‘homaria’ cultured from ESD lesions of wild lobsters (Chistoserdov et al. 2005). Both universal PCR primer sets also amplified and indicated a common presence of a Pelagibius sp., a Kiloniella sp. (Table 1) and a Hoflea sp. (Table 2). The 27F+355R primer set also identified the common presence of an unclassified Rhodobacteraeae sp. (sp. 1), 2 different unclassified Hyphomicrobiaceae sp., a Kiloniella sp., an Octadecabacter sp., a Hoflea sp., a Pelagibius sp., other Rhodobacteraeae and ‘Candidatus Homarophilus dermatus’ (Tables 1 & 2). In addition, the 341FM+907RM primer set identified the common presence of a Thalassobius sp. that the 27F+355R primers did not reveal (Table 1). Other bacteria types were detected but only in lesions of 1 of the 11 lobsters, and as such have not been described as they are unlikely candidates for being involved in shell disease.

During the experiment, DNA from 1 of the 11 lesions degraded (Lobster 8, Fig. 5) and was therefore not amplifiable with the Bacteroidetes primer set. However, of the remaining 10 lesions, Aquimarina ‘homaria’ DNA was amplified from all lobsters (Fig. 5). Band C2, belonging to the same unclassified and unclassified Flavobacteraeae sp., was also amplified from all 10 lesions (Fig. 5). Interestingly, this sequence was not amplified using either of the 2 universal primer sets. The Bacteroidetes-specific PCR primer set also amplified species detected by the other tests including the Krokinobacter sp., 2 unidentified Maribacter spp., 2 Tenacibaculum spp. and another Aquimarina sp. (Tables 1 & 2).

Comparison of bacterial communities from spots and lesions

Using the 27F+355R and 341FM+CFB721R PCR primer sets, direct comparisons between bacterial 16S rDNA sequences amplified from spots with those amplified from lesions were possible. The Flavobacteraeae and Alpha- and Gammaproteobacteria community profiles of spots and lesions were similar overall even though different species were often identified. This similarity may reflect their cohabitation in the same tank water. In both spots and lesions, 8 species were found consistently (Band C2 is a chimera), 9 species were unique to spots and 9 species were unique to lesions (Tables 1, 2 & 3). Of the 8 common species, only 2, Aquimarina ‘homaria’ and the Kiloniella sp., were identified in spots or lesions from 9 of the lobsters (Table 1).

Comparisons of the bacteria existing in each spot and each lesion of each individual lobster revealed that community profiles were similar for some, but dissimilar for others. For example, 2 DNA products with identical sequences were amplified in abundance from the spot and the lesions sampled from Lobster 10. In contrast, with Lobster 9, a diverse range of DNA products were amplified from the lesion (Fig. 4). Similar phenomena occurred in Bacteroidetes community profiles. For example, DNA band profiles between spots and lesions were similar from Lobsters 1 and 7 but vastly different from Lobsters 2 and 9 (Fig. 5).

Differences in Jaccard similarity indices determined for the 11 spots (mean = 0.385) and the 11 lesions (mean = 0.459) sampled from the 11 lobsters indicated that spots shared fewer common 16S rDNA products than did lesions and, thus, possessed higher bacterial diversity. Based on analyses of a 10 000 randomized data sets, the magnitude of the difference in Jaccard similarity indices (0.074) between spots and lesions was significant (p = 0.0022).

Detection of specific bacteria in lesions

As each of the 2 universal primer sets amplified Aquimarina ‘homaria’ and as the 341FM+CFB721R primer set amplified the unclassified Flavobacteraeae sp. in all spot and lesion samples, specific PCR primer sets were designed to confirm the high prevalence of these bacteria. Additional data on A. ‘homaria’ was also obtained. Using the PCR specific for A. ‘homaria’, a 16S rDNA product was amplified from all 11 spots and all 11 lesions sampled from the 11 lobsters as well as from healthy carapace scrapings from 2 of 6 control lobsters. In contrast, using the PCR specific for the unclassified Flavobacteraeae sp. sequence (Band C2), no product for the
16S rDNA was amplified from any spots or lesions sampled from the 11 lobsters, including those from which a product was amplified by the 341FM+CFB721R primer set. As this bacterium has not been cultured, a gel-purified DGGE band amplified from a lesion was used as a positive-control DNA and was effectively amplified in abundance by the specific PCR primer set. Based on these data, the unclassified Flavobacteriaceae (Band C2) DNA product amplified by the universal PCR primer set was designated as a DNA chimera and was indicated as such in silico.

**DISCUSSION**

Bacterial infestation of compromised lobsters has been suggested as the cause of lobster shell disease (Tlusty et al. 2007, Vogan et al. 2008). Evidence for this was shown by Tlusty et al. (2008) where lobsters fed exclusively on herring invariably became moribund and developed shell lesions (Tlusty et al. 2008). The same diet-based approach was used here to induce shell disease in *Homarus americanus* lobsters to investigate the makeup of bacterial communities that established in early-stage melanized spots and subsequent lesions.

Several bacterial types were particularly associated with DISD, with 2 species, *Aqumarina homaria* and *Kiloniella* sp., found ubiquitously in both spots and lesions. To some extent, homogeneity in bacterial types that established in spots and lesions was expected due to the experimental design. For practical reasons, the cohort of 11 lobsters studied was maintained in a single tank where individuals were housed in cubicles to avoid direct contact. Based on this design, all lobsters were exposed to the same water and environmental conditions at a single location. Thus, this study represented one outbreak of DISD with a subset of 11 individuals sampled for study of their microbial communities. The same bacteria present in spots or lesions of multiple lobsters

### Table 3. List and bacterial phylogenies inferred from 16S rDNA sequences of denaturing gradient gel electrophoresis (DGGE) bands amplified by PCR from DNA extracted from scrapings of single melanized spots from individual diet-induced shell disease-affected lobsters *Homarus americanus*. Sample sizes show no. of individual lobsters from which samples were obtained.

<table>
<thead>
<tr>
<th>Band</th>
<th>Closest BLASTn match (Accession no.)</th>
<th>Presence in DGGE of PCRs using various primer sets</th>
<th>RDP-based phylogeny</th>
<th>Accession no.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>27F-GC+ (n = 10) 355R 341FM-GC+ (n = 11) 907R CFB721R (n = 11)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S1</td>
<td>Uncultured <em>Rhodobacter</em> bacterium (EF215773)</td>
<td>9 0 na</td>
<td>Proteobacteria; Alphaproteobacteria; <em>Rhodobacter</em> sp.</td>
<td>JF297205</td>
</tr>
<tr>
<td>S2</td>
<td>Uncultured alpha proteobacterium (AM412521)</td>
<td>8 0 na</td>
<td>Proteobacteria; Alphaproteobacteria; <em>Rhodobacter</em> sp.</td>
<td>JF297206</td>
</tr>
<tr>
<td>S3</td>
<td><em>Rhodobacteraceae</em> bacterium strain 197 (AJ810843)</td>
<td>8 0 na</td>
<td>Proteobacteria; Alphaproteobacteria; <em>Rhodobacter</em> sp.</td>
<td>JF297207</td>
</tr>
<tr>
<td>S4</td>
<td>Uncultured bacterium clone HH1_a4 (FN401203)</td>
<td>7 0 na</td>
<td>Proteobacteria; Alphaproteobacteria; <em>Rhizobiales</em>; <em>Hyphomicrobiaceae</em>; <em>Hyphomicrobiium</em> sp.</td>
<td>JF297208</td>
</tr>
<tr>
<td>S5</td>
<td>Uncultured alphaproteobacterium G3-37 (EU005299)</td>
<td>5 0 na</td>
<td>Proteobacteria; Alphaproteobacteria; <em>Rhodobacter</em> sp.</td>
<td>JF297209</td>
</tr>
<tr>
<td>S6</td>
<td>Uncultured bacterium Crozet_s_404 (FM213885)</td>
<td>3 0 na</td>
<td>Proteobacteria; <em>unclassified</em>; <em>Alphaproteobacteria</em> sp.</td>
<td>JF297210</td>
</tr>
<tr>
<td>S7</td>
<td>Uncultured bacterium clone A32 (GQ215673)</td>
<td>6 0 na</td>
<td>Proteobacteria; <em>Gammaproteobacteria</em>; Xanthomonadaceae; Xanthomonadaceae; <em>Lysobacter</em> sp.</td>
<td>JF304026</td>
</tr>
<tr>
<td>S8</td>
<td>Uncultured <em>Xanthomonadaceae</em> bacterium (GQ926873)</td>
<td>5 0 na</td>
<td>Proteobacteria; <em>Gammaproteobacteria</em>; Xanthomonadaceae; <em>Lysobacter</em> sp.</td>
<td>JF297211</td>
</tr>
<tr>
<td>S9</td>
<td><em>Propionibacterium acnes</em> ATCC6919 (AB042288)</td>
<td>10 0 na</td>
<td>Actinobacteria; <em>Actinobacteria</em>; <em>Propionibacteriaceae</em>; <em>Propionibacterium</em> acnes</td>
<td>JF297212</td>
</tr>
</tbody>
</table>
were due to their presence in the shared tank water, whereas differences between bacterial communities may be due to individual host factors, such as the variation in time since shell disease onset and, subsequently, sampling.

Bacteria detected in spots and lesions of lobsters with DISD were almost invariably from the Bacteroidetes or Proteobacteria phyla. Bacteria in these phyla have also been identified as common inhabitants of lesions in lobsters with epizootic shell disease collected at multiple sites in New England and Atlantic Canada over a 9 yr period (Chistoserdov et al. 2005, 2009). Propionibacterium acnes was amplified at low levels from all of the spot samples. It can potentially be a pathogen initiating the infection since it was found in 10 out of 10 samples. However, an alternative explanation is that it is a contaminant, because it was only detected in spot communities that yielded low DNA quantities. Thus, small quantities of Propionibacterium acnes cells introduced by us during sampling could have contributed DNA and become detectable by PCR. This bacterium is a normal resident of human skin (Grice & Segre 2011), and possibly entered the system through regular water maintenance of the system. Propionibacterium acnes are slow growing, fermenting strict anaerobes, making them unlikely inhabitants of lobster surfaces.

The ubiquitous detection of Aquimarina ‘homaria’ in lesions of different shell disease types in wild and experimental lobsters (Chistoserdov et al. 2005, 2009), including the early-stage diet-induced spots examined here, suggests its potential role in initiating shell disease. A 16S rDNA sequence of a hypothetical unclassified Bacteroidetes bacterium belonging to Flavobacteriaceae was also detected in all samples using the Bacteroidetes-specific primer set. The universal bacterial primer set did not amplify DNA with a similar sequence, nor did specific primers for the sequence, thus indicating that it was most likely a DNA chimera (PCR artifact), which can be generated when group-specific primers are used to amplify large pools of 16S rDNA with similar sequences (Acinas et al. 2005). Indeed, the CHIMERA_CHECK program identified that its sequence was chimeric, although computer-based chimera analyses are mostly suggestive. The 16S rDNA sequences of several additional Flavobacteriaceae genera (Aquimarina sp., Krokinobacter sp., Maribacter spp., Tenacibaculum spp.) were amplified from spots and lesions, mostly with the Bacteroidetes-specific PCR primers. However, their greater prevalence in lesions suggested that these Flavobacteria may be secondary colonizers that establish as spots become lesions.

Various Proteobacteria, an unclassified Hyphomicrobiaceae sp., a Pelagibius sp., a Roseovarius sp. and the ‘Candidatus Homophilus dermatus’ were also amplified from both spots and lesions of multiple lobsters but were not truly ubiquitous, suggesting their opportunistic colonization of these lobsters. In contrast, the Kiloniella sp. was a dominant constituent of almost every spot or lesion community and an unclassified Rhodobacteraceae sp. 1 was a minor constituent of all lesion communities. The 16S rDNA sequence determined for the Kiloniella sp. was 96% identical to that of Kiloniella laminariae LD815, the only cultured Kiloniellaceae member and an aerobic chemoheterotrophic spirillum isolated from the marine brown alga Laminaria saccharina in the Baltic sea (Wiese et al. 2009). It was also 99% identical to ‘Kopriimonas byanusanensis’, a novel species not described formally, and from this high sequence similarity we propose the name ‘Candidatus Kopriimonas aquaticus’ for this newly identified Kiloniella relative. Unlike Aquimarina ‘homaria’, the ‘Candidatus Kopriimonas aquaticus’ identified in diet-induced lesions has not been detected in lesions from free-living lobsters with ESD (Chistoserdov et al. 2009). Additional culturing of bacterial communities associated with lobster shell disease should allow enrichment for Kiloniella spp. to determine whether they infest lesions of wild lobsters with ESD and to assess their relevance to ESD more directly. The 16S rDNA sequence of the Rhodobacteraceae sp. 1 matched distantly (90% identity) to the cultured Roseovarius pelophilus, but as it has also not been found in wild lobsters with ESD, its importance to DISD is not obvious.

DNA extracted from healthy carapace matter scraped from lobsters unaffected by DISD did not amplify by PCR using universal bacterial 16S rDNA primers. Thus few bacteria colonized the healthy cuticle of juvenile lobsters in our system, as the exact same method was used on healthy carapaces of wild adult lobsters and bacterial communities were readily amplifiable (Chistoserdov et al. 2009). However, Aquimarina ‘homaria’ was amplified in low abundance from 2 of 6 lobsters using specific PCR primers, indicating that its abundance might have been below the detection threshold for the universal PCR primers. Whether this is normal and due possibly to juvenile lobsters molting more frequently than adults or to their maintenance in an artificial aquarium environment is not known. However, consistent with this, DNA yields from melanized spot scrapings were much lower than those from lesion scrapings, indicating that bacterial loads increase in lesions.
The analysis of bacterial communities between spots and lesions from the 11 lobsters with DISD showed a more diverse assemblage in spots than in lesions, suggesting that the community constituents stabilize over time or that some constituents overgrow less abundant members. Only 2 bacteria, *Aquimarina ‘homaria’* and *Candidatus Kopriimonas aquarianus*, were found to persist universally as spots progressed to lesions (Table 1). Other than these bacteria, the bacterial communities are apparently dynamic in composition, with some bacteria having colonized spots being lost and others establishing as lesions develop and progress in severity.

While these observations on the bacterial community give clues to future research directions to identify community dynamics over shorter time scales, which bacteria escalate in communities as spots transition to lesions and, ultimately, which bacteria contribute to disease, the caveat remains that the data were gathered from a small cohort (n = 11) of lobsters cohabitated and exposed to exactly the same feed and water conditions. Thus, in our single experimental system, the source of the bacteria identified in association with DISD-induced shell pathology is likely either the herring used to prepare the diet or the bacterial community in the aquarium water taken in from Boston harbor; as such, extrapolations to bacterial communities of shell disease in wild lobsters are not easily made. It is notable however, that some of the same bacteria detected in DISD lesions here are also commonly detected in wild outbreaks of shell disease (Chistoserdov et al. 2005, 2009) and wild lobsters are known to consume high levels of fish from baited traps (Bethoney et al. 2011).

It is apparent from this and others studies that lobster shell disease lesions comprise complex communities of microbes possibly acting in concert to induce pathology (Chistoserdov et al. 2002, 2005, Smolowitz et al. 2005, Quinn et al. 2009). The experimental system in which the lobsters were maintained used seawater that was circulated through a biofilter at a rate of ~15% water exchange per day and therefore does not contain bacterial diversity likely to exist in natural environments. While the data on DISD cannot be used directly to infer what causes ESD in wild lobsters, the ubiquitous detection of *Aquimarina ‘homaria’* in DISD lesions adds further weight to it being involved in shell lesion development in lobsters irrespective of how it is induced (Chistoserdov et al. 2002, 2005, Smolowitz et al. 2005). The common detection of other bacteria in lesions, including *Candidatus Kopriimonas aquarianus* and various members of the *Rhodobacteraceae*, *Rhodospirillaceae*, and *Hyphomicrobiaceae*, also implicated these as possible agents contributing to DISD, and it will be interesting to determine whether these bacteria also occur commonly in lesions of wild lobsters affected by ESD. It will be interesting to compare the results from this study to those with more controlled microbial communities, such as sterile sea water or tanks inoculated with different bacterial strains to monitor the complex dynamics of lesion communities as shell disease progresses.

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**LITERATURE CITED**

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