Ocean warming and acidification have complex interactive effects on the dynamics of a marine fungal disease

Gareth J. Williams1,†, Nichole N. Price1,†, Blake Ushijima2,4, Greta S. Aeby1, Sean Callahan2, Simon K. Davy5, Jamison M. Gove6,3, Maggie D. Johnson1, Ingrid S. Knapp4,5, Amanda Shore-Maggio2,4, Jennifer E. Smith1, Patrick Videau2 and Thierry M. Work7

1Scripps Institution of Oceanography, Center for Marine Biodiversity and Conservation, University of California San Diego, La Jolla, CA 92093, USA
2Department of Microbiology, and 3Joint Institute for Marine and Atmospheric Research, University of Hawaii at Manoa, Honolulu, HI, USA
4Hawaii Institute of Marine Biology, Kaneohe, HI 96744, USA
5School of Biological Sciences, Victoria University of Wellington, PO Box 600, Wellington, New Zealand
6Coral Reef Ecosystem Division (CRED), Pacific Islands Fisheries Science Center (PIFSC), NOAA, 1610 Kapiolani Boulevard, Suite 1110, Honolulu, HI 96814, USA
7US Geological Survey, National Wildlife Health Center, Honolulu Field Station, PO Box 50167, Honolulu, HI 96850, USA

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Diseases threaten the structure and function of marine ecosystems and are contributing to the global decline of coral reefs. We currently lack an understanding of how climate change stressors, such as ocean acidification (OA) and warming, may simultaneously affect coral reef disease dynamics, particularly diseases threatening key reef-building organisms, for example crustose coralline algae (CCA). Here, we use coralline fungal disease (CFD), a previously described CCA disease from the Pacific, to examine these simultaneous effects using both field observations and experimental manipulations. We identify the associated fungus as belonging to the subphylum Ustilaginomycetes and show linear lesion expansion rates on individual hosts can reach 6.5 mm per day. Further, we demonstrate for the first time, to our knowledge, that ocean-warming events could increase the frequency of CFD outbreaks on coral reefs, but that OA-induced lowering of pH may ameliorate outbreaks by slowing lesion expansion rates on individual hosts. Lowered pH may still reduce overall host survivorship, however, by reducing calcification and facilitating fungal bio-erosion. Such complex, interactive effects between simultaneous extrinsic environmental stressors on disease dynamics are important to consider if we are to accurately predict the response of coral reef communities to future climate change.

1. Introduction

Diseases alter ecosystems [1] and threaten marine community function and resilience [2]. On coral reefs, disease outbreaks are considered a key contributor to the recent global decline of reef health and resilience [3]. Both global impacts, for example sea-surface temperature anomalies, and local human impacts, for example pollution, drive disease dynamics and outbreaks in scleractinian corals on reefs [4,5]. These stressors probably increase pathogen virulence and reduce host resistance, enhancing disease establishment and progression [2,6]. Our understanding of diseases that threaten other key calcifying (reef-building) organisms, however, is rudimentary. Crustose coralline algae (CCA) serve essential functional roles in coral reef ecosystems, including facilitating reef accretion and consolidation [7], providing a settlement substrate for coral larvae [8] and forming a key successional state promoting reef recovery following acute disturbance [9]. While CCA can occupy up to 50% of the living reef
benthos [7,10], relatively little is known about their biology and ecology [11], particularly their susceptibility to and subsequent impacts from disease [12].

Diseases can cause drastic reductions in CCA populations on coral reefs, with knock-on effects that promote regime shifts to fleshy macroalgal dominance and loss of functional resilience [13–15]. Several CCA diseases have been documented, although almost nothing is known about their aetiology, spatio-temporal dynamics and relationships with extrinsic environmental drivers [12]. This information is essential if we are to actively manage CCA disease occurrence and mitigate outbreaks at a local scale on coral reefs. While CCA are able to photoacclimatize [16], they are still vulnerable to bleaching as a result of increased temperature, and their calcification, photo-physiology and survival are threatened by a lowering of pH and carbonate saturation state (Ω) as a result of ocean acidification (OA) [17–19]. However, the influence of these global-scale stressors on local-scale CCA disease dynamics and occurrence on coral reefs is virtually unknown. Here, using a CCA disease previously described from the South Pacific, we shed light on this urgent research priority for, to our knowledge, the first time.

Coralline fungal disease (CFD) was first observed in 1997 on shallow (less than 20 m) reef habitats in American Samoa. Based on gross morphology, the aetiology of the disease was identified as an undescribed fungal pathogen [20]. Since 1997, CFD has been documented throughout other parts of the Pacific, with high prevalence at remote islands in the Central Pacific, in particular at Kingman Reef and Palmyra Atoll in the northern Line Islands [12]. While the distribution of CFD throughout the Pacific appears to be highly variable, evidence suggests that variations in sea-surface temperature (SST) may, in part, be driving spatial variation in disease occurrence, with higher mean SST at islands correlating with higher CFD in part, be driving spatial variation in disease occurrence, suggests that variations in sea-surface temperature (SST) may, through changes in disease occurrence appears independent of host abundance.

We show that CFD can exhibit very rapid progression across hosts, rivalling known rates for coral diseases and that increased temperature accelerates CFD lesion progression. We show that these effects are counteracted by a reduction in pH; however, under simulated OA conditions, diseased hosts still experience greater rates of net dissolution than healthy individuals. These complex interactions highlight the challenges associated with predicting disease outbreaks and their dynamics in a changing climate.

2. Material and methods

(a) Histopathology

Fragments (n = 7, approx. 3 cm²) of CCA displaying gross signs of CFD were collected at Palmyra Atoll (05°52’ N, 162°06’ W), at 10 m depth on the forereef (Figure 1). Each sample was fixed in zinc–formaldehyde solution (Z-fix, Anatech) diluted 1:5 with ambient non-filtered seawater, decalcified using a formic acid/zinc–formaldehyde solution (Z-fix, Anatech) diluted 1:5 with ambient non-filtered seawater, decalcified using a formic acid/formaldehyde solution (Cal-Ex II, Fisher Scientific), embedded in paraffin, sectioned at 5 μm and stained with haematoxylin and eosin. Grocott’s methamine silver was used to confirm the presence of fungal hyphae.

(b) Culture conditions and phylogenetic determination of coralline fungal disease fungus

(i) Culture conditions

To culture the fungus associated with CFD, Medium B, Wickerham’s, Czapek-Dox agar and cellulose agar [24] prepared with 25, 50, 75 or 100% filtered seawater as a base were used. Media were prepared with and without the inclusion of the antibiotics ampicillin and spectinomycin at a concentration of 100 μg ml⁻¹. Media with antibiotics were used to suppress bacterial growth, and the same media were used without antibiotics to preclude any potential negative effects on fungal growth. Solid media included 0.7, 1.0, 1.2 or 1.5% agar (Fisher Scientific). Cultures were incubated at 25, 27 or 30°C with aeration (solid cultures) or without aeration (liquid cultures) for 21 days.

(ii) DNA extraction and gene amplification

Frozen, ground CFD lesions were thawed on ice, 5 μl were spread on PALM PEN membrane slides (Carl Zeiss) and individual fungal filaments were isolated with a Zeiss PALM laser microdissection system. DNA was extracted from filaments as in [25] with the following modifications: two samples of 8–10 fungal filaments in 50 μl of sterile water were lysed with a mini-beadbeater (Biospec) and extracted with phenol–chloroform. DNA was
precipitated, dried and re-suspended in 20 μl of sterile 10 mM Tris buffer, pH 8.5. A 719 bp fragment of the 18S rRNA gene was PCR amplified with fungus-specific primers designed from mixed environmental samples: nu-SSU-0817–5’ and nu-SSU-1536–3’ [26] and Phusion high-fidelity polymerase (New England Biolabs). The same primers were used for sequencing.

(iii) Phylogenetic analysis
The fungal 18S rRNA gene sequence was initially assessed using The National Center for Biotechnology Information (NCBI) and the BLAST algorithm. To further assess its relatedness to other phylum Basidiomycota members, especially the subphylum Ustilaginomycetes, 653 bp 18S RNA gene sequences were aligned in MEGA5 [27]. The fungal isolate’s sequence was aligned with 22 other NCBI sequences, chosen based on a previously published analysis of the phylum Basidiomycota [28], using ClustalW. A phylogenetic tree was constructed using the neighbour-joining method with 1000 bootstrap replicates [29]. All positions containing gaps and missing data were eliminated, leaving a total of 647 positions. Evolutionary distances (number of base substitutions per site) were computed using the maximum composite likelihood method. Rate variation among sites was modelled with a gamma distribution (shape parameter = 8). Evolutionary analyses were conducted in MEGA5.

(c) Coralline fungal disease occurrence, host cover and associated changes in seawater temperature
To quantify CFD occurrence, 59 × 200 m² transects were surveyed in July–August and October–November 2008 (total of 11 800 m² of reef). Backreef (n = 4 transects, 1–5 m depth), reef terrace (n = 25, 4–5 m depth) and forereef (n = 30, 10 m depth) habitats were surveyed within 12 permanent sites (figure 1). Along each transect, per cent CCA cover was estimated using the photoquadrat method. Each photoquadrat (n = 20 per 50 m) was 0.63 m². Per cent cover was calculated post hoc by identifying 100 points in a stratified random design for each photograph and averaging within each transect. Difficulties with delineating individual CCA crusts in situ meant a true CFD prevalence (proportion of individuals displaying signs of the disease) could not be calculated. Instead, the numbers of CFD cases were normalized to host cover (per m² of CCA) along transect. Difficulties with delineating individual CCA crusts random design for each photograph and averaging within each cover was calculated post hoc by identifying 100 points in a stratified method. Each photoquadrat (transect, per cent CCA cover was estimated using the photoquadrat were surveyed within 12 permanent sites (figure 1). Along each transect (see the electronic supplementary material, table S1) were computed using the maximum composite likelihood method. Rate variation among sites was modelled with a gamma distribution (shape parameter = 8). Evolutionary analyses were conducted in MEGA5.

(d) Coralline fungal disease vital rates during the El Niño
On the central south forereef, 13 CFD cases were photographed weekly for four weeks in October–November 2009. Individual CFD cases were initially marked with a stainless steel pin as a reference point. Photographs were taken perpendicular to the substrate to minimize angle variations among images. CFD vital rates (lesion surface area and linear progression rate) were calculated post hoc using ImageJ (http://rsweb.nih.gov/ij/). Active lesion surface area was the area displaying fungal cover (blue–black discoloration), not including dead CCA left in the lesion’s path. Calculations were averaged across three images of each lesion to account for slight variations in angle among photos.

(e) Temperature and acidification experiments
We used a factorial CO₂ bubbling and heating experiment (3–13 June 2012) to examine independent and interactive effects of OA and warming on CFD disease dynamics (lesion surface area and linear progression rate) and CCA growth (net calcification). Samples were collected, as for histopathology, from independent diseased and healthy CCA crusts. Epiphytes were removed, and each CCA genus was confirmed using a dissecting microscope. Paired CCA fragments (approx. 2 cm² Neogoniolithon sp.; diseased and healthy) were placed in 11 glass aquaria holding fresh seawater equilibrated to treatment conditions (approx. 24–48 h before) (sensu [31]). Experimental OA conditions were created by bubbling pre-mixed air enriched with excess pCO₂ (AirGas Pro) to 1124 ± 88 μatm (mean ± s.e. hereafter) to reflect atmospheric CO₂ concentrations projected in 2100 (scenario IV; [32]). In control aquaria, present-day CO₂ conditions were created using a Pabracke 12 V HP625 air compressor delivering ambient air. Aquaria were immersed in flow-through water baths at 28.06 ± 0.01 °C (seasonal average for Palmyra; ambient conditions) or 29.49 ± 0.02 °C (mean SST during the El Niño warming event at Palmyra; warming conditions). Four independent water baths (two ambient and two warmed) held experimental aquaria that were randomly assigned to elevated or present-day CO₂ conditions to create every combination of warming and OA treatment level (n = 12 per level divided evenly among the two water baths).

Aquaria were covered to prevent evaporation and rainwater from affecting salinity and placed under a shade cloth to mimic the natural light environment at 10 m on the forereef. To control for algal metabolism, two empty aquaria per water bath were subjected to the same four treatments described above. In each aquarium, temperature and light conditions were recorded every 15 min using Onset HOBO Pendant UA-002-64 light and temperature loggers (see the electronic supplementary material, figure S1). Light intensities (lux) were converted to the availability of photosynthetically active radiation (PAR) using the equation: 1 μmol quanta (400–700 nm) m⁻²s⁻¹ = 51.2 lux (sensu [33]). These conversions were validated by midday PAR measurements with an Li-Cor LI 192 4r quantum sensor (492 ± 25 μmol photons m⁻²s⁻¹). Once daily, pH seawater (resolution ±0.01), temperature (±0.1 °C) and dissolved oxygen (±0.2 mg l⁻¹; electronic supplementary material, table S2) were measured using an HACH HQ 40d handheld meter. These measurements were also taken at 6:00, 12:00, 18:00 and 24:00 h to quantify diurnal fluctuations in each aquarium (see the electronic supplementary material, table S2). Twelve water samples for total alkalinity (Aₜ), total dissolved inorganic carbon (Cₜ) and salinity were collected at days 2, 6 and 10 of the experiment in 500 ml Corning brand Pyrex bottles and fixed with 200 μl saturated HgCl₂ solution (1% headspace). Samples were collected (in duplicate) from experimental aquaria and control (empty) aquaria at each of the four treatment levels.

(f) Crustose coralline algae calcification and disease progression
Net calcification rates of all fragments of Neogoniolithon sp. were quantified with the buoyant weight method [34] (to the nearest mg) using a weigh-below basket on a balance (Denver SI-403). Changes in buoyant weight over the 10 day experiment, normalized to initial fragment mass, approximated net calcification rate. For each diseased fragment, CFD disease vital rates (lesion surface area and linear progression rate) were calculated using ImageJ as described above.

(g) Experimental water chemistry analysis
Carbonate chemistry and salinity were analysed in the Dickson Laboratory at Scripps Institution of Oceanography. Cₚ was determined using a single-operator multiparameter metabolic
analyzer (SOMMA) and a UIC Model 5011 CO₂ coulometer. A_T was determined by open-cell acid titration using a Metrohm Dosimat Model 665 and Metrohm potentiometric pH probe and meter. Salinity was determined using a Mettler Toledo Model DE45 density meter. Seawater dissolved inorganic carbon parameters (HCO₃⁻, CO₃²⁻, CO₂, pCO₂) and the saturation state of carbonate minerals (Ω-calcite and Ω-Mg calcite) were calculated based on measured C_T and A_T using the computer program SEACARB [35] and stoichiometric dissociation constants [36] (see the electronic supplementary material, table S3).

(h) Data analyses
To test for differences in CCA cover and CFD occurrence across fore reef sites in 2008, we ran a permutation-based analysis of variance and subsequent pairwise comparisons using PERMANOVA+ [37]. To test for any relation between CCA abundance and CFD occurrence in 2008, we used a permutational linear model using Distlm-forward [38]. Two-way nested analyses of variance (ANOVAs) tested whether OA (fixed) and warming (fixed) treatments independently or jointly affected net calcification; each fixed factor had two levels. Replicates from a water bath were nested within temperature treatments to test for location bias. Analyses were run independently for diseased and healthy samples. Normality and homoscedasticity were verified using the Shapiro–Wilk test. Growth responses were compared between diseased and healthy CCA within each treatment using t-tests. Proportional changes in lesion area were analysed using the Dunn’s method for joint ranking, a non-parametric approach that compares means of treatments against a control (ambient SST and air); we confirmed that variances across treatments were equal with a Brown–Forsythe test. Unless otherwise stated, all analyses were completed using R 2.15.2 (R Development Core Team, http://www.r-project.org).

3. Results
(a) Coralline fungal disease gross morphology and histopathology; phylogeny of associated fungus
CFD lesions were characterized by a diffuse area of mottled white discoloration separated from a pink CCA thallus by a blue–black band (approx. 1–3 cm wide) with irregular distinct undulating borders (figure 2a). Septated hyphae with branches originating near septal junctions were commonly observed (figure 2b). Many of the fungal filaments had highly branched structures resembling conidiophores, which in some cases appeared to have attached spherical conidia, the asexual spores of fungi. These were the dominant fungal structures in diseased CCA but were absent in healthy CCA. The CCA cuticle was overlaid and disrupted by mats of brown to colourless filamentous branching septate structures (figure 2c). These irregularly walled structures infiltrated into the CCA thallus at right angles to the cuticle to a depth of approximately 100 μm. These were morphologically compatible with fungi, and Grocott’s methenamine silver confirmed a fungal infection (figure 2c).

Isolation by microdissection followed by genetic sequencing of the fungal hyphae revealed that 653 bp of the 18S rRNA gene sequence (accession no. KF255580) shared 97–98% sequence identity with members of the phylum Basidiomycota and uncultured marine isolates. The sequence was most similar (98% sequence identity) to members of the subphylum Ustilaginomycetes, in particular Malassezia restricta strain CBS-7877 and the marine isolates KM10-BASS and CK2-BASS (see the electronic supplementary material, figure S2). Repeated attempts to culture the fungus were unsuccessful.

(b) Spatio-temporal patterns of coralline fungal disease and associated changes in seawater temperature
Atoll-wide mean (± 1 s.e.) CFD occurrence on Palmyra’s fore reef habitat equalled 0.1 ± 0.06 cases m⁻² of CCA in 2008.
Although cases of CFD were seen outside of our surveyed transects on the deeper (approx. 10–15 m) reef terrace habitat, cases were rare in comparison; no CFD cases were documented within our surveyed transects on the shallow (less than 5 m) terrace or backreef habitats (total reef area surveyed in these two habitats equalled 5800 m²). Within the forereef habitat, CFD occurrence displayed spatial heterogeneity (Pseudo-$F_{5,29} = 13.456$, $p < 0.0001$), with the central south forereef having the highest mean number of cases m$^{-2}$ of CCA (0.39 ± 0.10) in 2008 (see the electronic supplementary material, table S1). Per cent cover of CCA did not differ across forereef sites in 2008 (Pseudo-$F_{5,29} = 2.67$, $p = 0.06$, mean cover 23.5%), and there was no relationship between CFD occurrence and CCA cover (Pseudo-$F_{1,29} = 0.311$, $p = 0.751$; electronic supplementary material, table S1).

Within the permanent backreef and reef terrace transects, mean CFD occurrence remained at 0 cases m$^{-2}$ of CCA in 2009 and 2010. However, within the permanent forereef transects, CFD mean occurrence increased approximately 14-fold from 0.10 cases m$^{-2}$ of CCA in 2008 to 1.37 cases m$^{-2}$ of CCA in late 2009 in association with the El Niño ocean-warming event (figure 3; electronic supplementary material, table S1). Seawater temperature had steadily increased over the latter part of 2009 and eventually peaked in November 2009, reaching 1.25 and 1.51 °C (satellite and forereef temperature observations, respectively) above the maximum long-term monthly climatological SST for Palmyra (figure 3). CFD occurrence increased in all permanent forereef transects during the El Niño event, with the central south forereef maintaining the highest levels (3.74 cases m$^{-2}$ of CCA; electronic supplementary material, table S1). By March 2010, with a decrease in seawater temperatures, mean forereef CFD occurrence had returned to pre-El Niño levels (figure 3).

(c) Coralline fungal disease vital rates during the El Niño event

CFD lesions typically progressed in a radial manner across the surface of the CCA thallus, often crossing between individual CCA thalli but never spreading onto hard coral tissue (see the electronic supplementary material, figure S3). At time point zero, mean lesion surface area of the 13 CFD cases in situ was 108 mm$^2$ (± 25) (see the electronic supplementary material, table S4). After one week, mean lesion surface area was 136 mm$^2$ (± 23), with a mean surface area progression rate of 3.5 mm$^2$ d$^{-1}$ (± 2.3) and a mean linear progression rate of 2.4 mm d$^{-1}$ (± 0.5). Across the entire four-week time period, the maximum surface area progression rate and linear progression rate of any single CFD lesion was 12.9 mm$^2$ d$^{-1}$ and 6.5 mm d$^{-1}$, respectively (see the electronic supplementary material, table S4).

(d) Experimental effects of warming and acidification on crustose coralline algae calcification and disease progression rates

Exposure to elevated temperature and atmospheric pCO$_2$, designed to simulate OA, reduced CCA net calcification rates, but this effect was dependent on fungal infection (table 1). All CCA samples gained CaCO$_3$ mass in the ambient air treatments, while all samples lost mass in the elevated pCO$_2$ (lower pH) treatments (figure 4a). However, when exposed to both elevated pCO$_2$ and temperature, diseased CCA lost nearly
twice as much mass as when exposed to simulated OA alone (significant interaction term, table 1). Mass loss was not intensified for healthy CCA (figure 4a); net calcification rates in healthy CCA were significantly depressed only by elevated $pCO_2$ and not by elevated temperature (table 1). Accordingly, calcification rates for diseased and healthy samples were statistically similar in all treatments, except for the simultaneous acidified and warmed conditions, in which diseased CCA lost 40% more mass than healthy CCA ($t$-test, d.f. = 15, $p = 0.0343$). Visible lateral progression of the CFD lesion occurred only in the elevated temperature treatment in ambient $CO_2$ conditions where lesion size and lethality increased by 60% over one week (figure 4b).

4. Discussion

Using a previously described CCA fungal disease (CFD) [20], we demonstrate that ocean warming and acidification can have complex interactive effects on marine disease dynamics. These
relationships are to be expected, as they reflect intricate relationships among the putative pathogen, host and environment [39].

(a) Identification of coralline fungal disease-associated fungus

Fungal pathogens are prevalent throughout the marine environment [40,41], are commonly associated with the coral holobiont [42] and are known to infect tropical sea-fans [43,44] and marine algae [45]. An inability to culture fungal isolates and a reliance on morphology for identification, however, have caused fungal isolates to be misclassified and their distribution underestimated [46]. Using histopathology of CFD samples from Palmyra Atoll and genetic sequencing of the associated fungus, we confirm a fungal infection of the CCA. While species-level identification was not possible, our phylogenetic analysis strongly suggests that the CFD fungus belongs to the subphylum Ustilaginomycetes, which consists of a large number of plant parasites, including strains of smut fungi [47]. Our methods, which allowed isolation of the fungus without an axenic culture, could be used to compare the fungus present in the Palmyra CFD lesions with fungi associated with suspected CFD lesions found on other reefs. These genetic approaches allow us to better interpret spatio-temporal dynamics of this disease on coral reefs and postulate their underlying mechanisms.

(b) Disease dynamics and sea-surface temperature

Many fungal pathogens in animals and plants respond positively to elevated temperatures [48–51], and ocean warming is predicted to favour pathogens for many marine diseases [2]. At our study location, CFD displayed a dramatic (14-fold) increase in occurrence on the fore reef during an El Niño in association with sustained seawater temperatures well above the long-term climatological mean. Furthermore, we found experimentally that elevated temperature increases lateral expansion rates of CFD lesions. The positive relationship between temperature and CFD occurrence was probably the result of elevated temperatures increasing the virulence of the pathogen, and hence speeding disease progression and causing physiological stress to the CCA host that ultimately reduced resistance to infection [52]. The prevalence of many coral diseases, such as white syndromes [5], atramentous necrosis [53] and black band disease [54] are also positively related to temperature. Interestingly, Vargas-Ángel [12] documented higher overall CCA disease occurrence at islands experiencing higher mean annual SST in a Pacific-wide survey of US-affiliated coral reefs, further highlighting the importance of temperature in governing CCA disease dynamics.

Though temperature variation provides a strong explanation for temporal variation in overall CFD occurrence at our study location, it does not adequately explain the spatial variation we documented at any one point in time. If higher temperatures cause both CFD occurrence and rates of lesion progression to increase, why was the disease almost exclusively limited to the fore reef, where temperatures are lower than the shallow reef terrace [21]? There are several possible explanations. Host density is often a crucial factor determining the spatio-temporal distribution patterns of disease [39], with the prevalence of many coral and CCA diseases positively related to host abundance [5,55,56]. For example, Vargas-Ángel [12] found a positive relationship between island mean CCA cover and overall CCA disease occurrence at an archipelago scale across coral reefs of the US Pacific, including the Pacific Remote Island Areas, the geopolitical region within which Palmyra resides. However, within Palmyra’s fore reef habitat, CFD occurrence appeared to be independent of host abundance. Moreover, CCA cover peaks on Palmyra’s shallow (less than 5 m) western reef terrace [10], where CFD was virtually absent; previous surveys at Palmyra have documented CFD on the terrace habitat, but again at deeper (approx. 15 m) depths (B. Vargas-Ángel 2013, personal communication) where host cover is lower [10]. These findings suggest that host abundance alone does not explain the observed spatial variation in CFD occurrence at Palmyra. However, CCA species assemblages are known to vary spatially on reefs [57], and the peak of CFD occurrence on the fore reef may simply reflect an increase in the abundance of a preferred host species. While in situ species-specific assessments of CCA abundance would potentially resolve these issues, CCA taxonomy is difficult and requires microscopic examination, making it impossible in the field.

While potentially explaining between-habitat differences in disease occurrence, variation in host species abundance alone does not adequately explain the dramatic peak in CFD abundance on Palmyra’s central south fore reef. This CFD hotspot at Palmyra appears to be temporarily stable, corroborating previous surveys conducted in 2006 [12]. The existence of this CFD hotspot at one site suggests that disease occurrence may be governed by external inputs of the pathogen, rather than by within-population transmission [44], or by an unmeasured extrinsic forcing. Palmyra’s central south fore reef is exposed to a particularly high level of lagoon outflow during the change in tidal state (G. J. Williams 2007–2013, personal observation), perhaps acting as a pathogen source and/or supplying more nutrient-rich waters that may enhance CFD establishment and progression, as has been shown for other fungal diseases on coral reefs [4]. Additionally, the south fore reef of Palmyra has, on average, measurably higher seawater pH with less frequent or severe excursions than the north fore reef or reef terrace [58]; our results indicate that the less acidic but warmer conditions characteristic of the southern fore reef are most favourable for CFD occurrence. Regardless of the mechanisms behind the fine-scale variations in CFD occurrence, it is clear that the disease is more abundant and virulent under elevated temperatures, suggesting that predicted increases in the frequency of temperature anomalies on coral reefs may result in more frequent CFD outbreaks.

(c) Eco-physiological response of host to disease and climate change

What will be the ecological consequences of increased CFD outbreaks on coral reefs? While elevated temperature increased overall CFD occurrence in situ and lateral rates of lesion expansion under experimental conditions, under the same experimental conditions elevated pCO₂ mediated these effects of temperature and slowed lesion expansion rates. While this suggests that future increases in the frequency of temperature anomalies will result in more frequent CFD outbreaks on coral reefs, the lowering of pH as a result of OA may actually slow down overall spread of the disease across the reef landscape during such outbreaks. Importantly, however, while the lateral spread of CFD was not affected by reduced seawater pH and carbonate saturation state, all CCA thalli lost mass under OA conditions, suggesting that net dissolution was occurring. For diseased thalli, these effects were exacerbated by warming.
Synergistic effects of ocean warming and acidification that together cause greater reduction in calcification of CCA than either stressor alone have been reported elsewhere [17–19,59], but synergistic global climate change effects were only observed in this study when the CCA were also infected with the CFD fungus. Microboring organisms, or euendoliths, such as fungi or cyanobacteria, burrow and erode carbonate at rates that can exceed biogenic CaCO$_3$ precipitation, leading to the net dissolution of reef-building organisms [60,61]. OA is expected to reduce resistance to eudendolith penetration in both hermatypic corals and CCA by weakening structural integrity of the CaCO$_3$ crystals [62], reducing skeletal density [63] and facilitating chemical dissolution [64,65]. Further, the colony formation is stimulated by natural reductions in pH, so OA has the potential to radically elevate the abundance of marine fungi [66]. Not only can acidification weaken host resistance to bio-erosion, but also reduced saturation states and enhanced disease infestation of the CCA thallus could further accelerate corrosion. Thus, the synergistic interaction of pathogen infection, warming and OA may exacerbate reef degradation under projected global climate change scenarios.

(d) Conclusion

Our study represents, to our knowledge, the first attempt to understand the interactive effects of two major global stressors, ocean warming and acidification, on disease dynamics on coral reefs. Using a fungal disease affecting crustose coralline algae (CFD), we show that while outbreaks of CFD should become more common on coral reefs as temperature anomalies become more frequent, OA may ameliorate lesion progression rates but still decrease overall survivorship of diseased hosts. The ecological consequences of such interactions are difficult to predict; however, it is clear that CFD possesses a tremendous capacity for lateral spread across the reef landscape during ocean-warming events. Our results highlight the intricate nature of disease-host-environment interactions and the importance of adopting a multi-factor approach to modelling disease dynamics on coral reefs in order to accurately predict dynamics in a changing climate.

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