Marine Ecology

Deep reefs are not universal refuges: Reseeding potential varies among coral species

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Deep coral reefs (that is, mesophotic coral ecosystems) can act as refuges against major disturbances affecting shallow reefs. It has been proposed that, through the provision of coral propagules, such deep reefs may aid in shallow reef recovery; however, this “reseeding” hypothesis remains largely untested. We conducted a genome-wide assessment of two scleractinian coral species with contrasting reproductive modes, to assess the potential for connectivity between mesophotic (40 m) and shallow (12 m) depths on an isolated reef system in the Western Atlantic (Bermuda). To overcome the pervasive issue of endosymbiont contamination associated with de novo sequencing of corals, we used a novel subtraction reference approach. We have demonstrated that strong depth-associated selection has led to genome-wide divergence in the brooding species Agaricia fragilis (with divergence by depth exceeding divergence by location). Despite introgression from shallow into deep populations, a lack of first-generation migrants indicates that effective connectivity over ecological time scales is extremely limited for this species and thus precludes reseeding of shallow reefs from deep reefs. In contrast, no genetic structuring between depths (or locations) was observed for the broadcasting species Stephanocoenia intersepta, indicating substantial potential for vertical connectivity. Our findings demonstrate that vertical connectivity within the same reef system can differ greatly between species and that the reseeding potential of deep reefs in Bermuda may apply to only a small number of scleractinian species. Overall, we argue that the “deep reef refuge hypothesis” holds for individual coral species during episodic disturbances but should not be assumed as a broader ecosystem-wide phenomenon.

INTRODUCTION

Tropical coral reefs are in global decline, with many under immediate threat because of a rapidly changing climate and the accumulation of local stressors (1, 2). Even in remote areas with strong legal protection, the increase in frequency and magnitude of large-scale bleaching and storm events has severely affected coral reefs (3). Coral reef persistence increasingly depends on local areas that can offer refuge against such disturbances (4) and provide propagules to recolonize affected areas (5). Deeper sections of coral reefs (below depths of 30 to 40 m), also referred to as “mesophotic coral ecosystems,” arguably represent potentially critical ecological refuges. Although certainly not immune to disturbance (6–9), they are commonly buffered from major bleaching and storm events (10, 11) and have a near-ubiquitous presence directly adjacent to the world’s shallow coral reefs (12). Although their ability to escape disturbances has been relatively well documented, the hypothesis that deep reefs harbor reliable reproductive sources remains largely untested (11, 13–15) and this was recently highlighted as a key knowledge gap in a report by the United Nations Environment Programme (16).

The “reseeding” potential of mesophotic coral reefs firstly depends on the extent of species overlap with their shallow-water counterparts. Although greater depths often confer greater protection from disturbances (11), the community similarity between shallow and deep reefs also decreases with depth, as a reflection of changing environmental conditions (17). Given this trade-off, the “deep reef refuge” hypothesis (DRRH) pertains primarily to upper mesophotic depths (~30 to 60 m)—deep enough to escape disturbances but shallow enough to guarantee sufficient overlap in species composition (11, 18, 19). Species overlap between shallow and upper mesophotic communities can be substantial, with an estimated 25 to 40% of Caribbean coral species and ~20 to 40% of Acropora species (on the Great Barrier Reef) representing depth-generalist species (11, 20). However, initial assessments have demonstrated that depth-generalist coral species are not necessarily composed of a single panmictic population over depth (14, 21–25). The three coral species thus far assessed for shallow-mesophotic connectivity (which includes two brooding and one broadcasting species) exhibit ambiguous patterns across geographic regions, with genetic differentiation observed in certain locations but not others (14, 22, 23, 25). This geographic variation has consequently hampered a more general assessment of the DRRH as the overall role of deep reefs in shallow reef recovery remains unclear (16, 26).

To understand the reseed potential on an ecosystem-wide scale (that is, across many species), it is essential to identify the nature of barriers to vertical connectivity and how these barriers may vary across coral species with distinct life history traits. The relative roles of depth and location have remained difficult to disentangle in past assessments of vertical connectivity, often because of geographic separation of sampled shallow and mesophotic populations (14, 22, 25, 27). Nonetheless, depth-associated selection could pose an important ecological barrier to vertical connectivity (24, 28) even in the absence of physical barriers (for instance, on those reef slopes where shallow and mesophotic communities are directly adjacent). Depth-associated selection does not necessarily restrict gene flow; however, in species with localized sperm and larval dispersal (such as in most brooding corals), localized dispersal could lead to assortative mating and therefore facilitate a divergence-with-gene-flow scenario (28, 29). Similarly, assortative mating could facilitate divergence in broadcasting species if depth-associated selection is accompanied by temporal reproductive isolation, either through pleiotropic effects or through depth-related differences in spawning cues (24, 27, 30). The role of such ecological barriers to vertical connectivity remains poorly understood, and the

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impact of depth-associated selection on vertical connectivity has remained largely obscured because genomic sampling has been restricted to mostly to microsatellite sequencing in presumably neutral parts of the genome [but see the work of Brazeau et al. (23)].

Reduced-representation approaches to genome sequencing [for example, restriction-site associated DNA sequencing (RAD-seq)] allow for sampling of both adaptive and neutral genetic variation across the genome (31, 32). In combination with a replicated sampling design (sampling the same habitats across multiple locations in a region), such reduced-representation approaches have the potential to identify the nature of barriers (that is, ecological or physical) that might hamper cross-habitat connectivity. Nonetheless, uptake of such genotyping-by-sequencing approaches in symbiotic marine invertebrates has been slow [but see related studies (33–37)] largely due to the nonspecific nature of these methods (for example, targeting universal restriction site motifs) in light of pervasive contamination by the obligate endosymbionts. In an aposymbiotic (that is, lacking symbionts) reference (38), with the hosted Symbiodinium representing a significant proportion of the overall biomass (39). Endosymbiont contamination might not necessarily affect inferred genetic patterns for the host; however, this can only be ascertained when comparing it against an aposymbiotic (that is, lacking symbionts) reference (36). Contamination is of particular concern when evaluating horizontal versus vertical connectivity, given the prevalence of geographic differentiation might not necessarily affect inferred genetic patterns for the other deep populations (PD, JD, and GD), with, again, the shallow Princess Beach population (PS) exhibiting a much lower corallite density. Sequencing of the COX1 (mitochondrial) region of associated endosymbionts showed that A. fragilis and S. intersepta associated with distinct endosymbiont types. Further characterization of a short chloroplast minicircle locus indicated that A. fragilis associated with a single endosymbiont haplotype, while S. intersepta associated with two different Symbiodinium haplotypes (Fig. 1, C and D). However, the vast majority of S. intersepta colonies (94%) associated with a single Symbiodinium haplotype (sint_chl_a); a small proportion associated either with the alternative haplotype (sint_chl_b) or with a combination of both haplotypes (nucleotide positions ambiguous for the mutations separating the two genotypes).

Contaminant filtering, missing data, and clonality

Sequencing of nextRAD libraries resulted in an average of ~1.4 million reads for each individually bar-coded sample of the two focal species (n = 213; not including failed samples). Using alignment-based clustering, we obtained 12,145 nextRAD sequence loci (hereafter referred to as “RAD loci”) for A. fragilis and 7591 for S. intersepta under initial PyRAD conditions. From the recovered RAD loci, 10% of the A. fragilis and 14% of the S. intersepta data sets were removed because they represented contaminants matching the Symbiodinium subtraction reference. Additional contaminants (~2% of RAD loci) matching non-cnidarian references (with a notable abundance of the proteobacteria Ralstonia sp.) were also removed before downstream filtering. After quality control and minimal representation filtering, we obtained 2568 biallelic single-nucleotide polymorphisms (SNPs) from 1579 RAD loci for A. fragilis and 7547 biallelic SNPs from 2187 RAD loci for S. intersepta (excluding 108 multiallelic SNPs that were included in analyses that support multiallelic data). Missing data average after filtering was 15% of SNPs for A. fragilis and 19% of SNPs for S. intersepta (Fig. 2). Although no potential clonemates were identified for S. intersepta (maximum allelic similarity, 86%), four groups of potential clones were identified for A. fragilis (allelic similarities, 94 to 98%). These represented two triplets and two pairs and always occurred within the same population (Fig. 2). In addition, a small recruit (<1.5 cm) sampled directly adjacent to an A. fragilis colony collected from a depth of 67 m was found to represent a clonemate. A single representative of each group of potential clones was kept in the data set for population-level analyses.

Genome-wide patterns of differentiation and outlier analyses

Global mean $F_{ST}$ (fixation index, a measure of genetic differentiation between populations) varied nearly two orders of magnitude between A. fragilis ($F_{ST}$ = 0.06998) and S. intersepta ($F_{ST}$ = 0.00081), with many individual SNPs exhibiting high $F_{ST}$ values for A. fragilis (Fig. 3A). From the 175 SNPs initially identified as $F_{ST}$ outliers for A. fragilis in the overall Fdist analysis, 56 were also identified as outliers by BayeScan. From those, 25 SNPs located on 12 RAD loci were identified as potentially under depth-associated selection (outliers in at least two pairwise shallow-deep comparisons but never in within-depth pairwise comparisons). These depth-associated outliers exhibited very distinct genotype frequencies between shallow and deep populations, with a minor allele private to the deep populations (but not fixed) in nearly

RESULTS

Sample collections, skeletal morphology, and endosymbiont associations

Coral colonies from the brooding species A. fragilis and the broadcasting species S. intersepta were sampled from shallow (~12 m) and mesophotic depths (~40 m) at four different locations around Bermuda (Fig. 1A and table S1). Phenotypic characterization of two skeletal properties (corallite density and diameter; Fig. 1B) showed similar skeletal values for A. fragilis specimens from all three eastern deep populations (PD, JD, and GD; Fig. 1, A and B). In contrast, the two shallow populations and the western deep population had either a much lower corallite density (PS) or a larger diameter (GS and WD). For S. intersepta, samples from the western population (WD) were similar to those from the other deep populations (PD, JD, and GD), with, again, the shallow Princess Beach population (PS) exhibiting a much lower corallite density.
half of these (Fig. 3B). Depth-outlier SNPs on the same RAD locus exhibited (near-)identical patterns, and genotype frequencies are therefore given for a single representative SNP. Two of the depth-outlier RAD loci matched sections of the Acropora digitifera genome, encoding an uncharacterized protein and a phospholipase enzyme (D1-like) (42). Visual assessment of the remaining outlier SNPs (fig. S1) suggests that an additional ~13 SNPs each had a distinct allele occurring in high frequency at the Western Blue Cut population (WD). For S. intersepta, 138 outlier SNPs were identified in the Fdist analysis, 4 of which were also identified as outliers by BayeScan, but none were identified as potentially under depth-associated selection using the criterion of outliers in multiple pairwise shallow-deep comparisons (Fig. 3, C and D).

**Genetic structuring in* A. fragilis* across depths and locations**

Genetic structure was identified for* A. fragilis* (G-statistic Monte Carlo test for all SNPs: G_{obs} = 56,315, P < 0.01) and supported by all approaches [Bayesian clustering, principal components analysis (PCA), and pairwise genetic distances between individuals; Figs. 2 and 4], indicating clear genetic differentiation between shallow and deep populations of* A. fragilis*, as well as differentiation of a deep population on the northwestern side of the island. The existence of three distinct clusters was further supported using the MedMeaK and MedMedK estimators for STRUCTURE (that is, maximum of three clusters detected; table S2). To explicitly test for divergence with depth, we categorized measures of differentiation into four groups of three pairwise population comparisons (fig. S2 and table S3): within-shallow (GS-JS, JS-PS, and PS-GS), within-deep (GD-JD, JD-PD, and PD-GD), within-deep-west (WD-GD, WD-JD, and WD-PD), and between-depths (GS-GD, JS-JD, and PS-PD). A nonparametric Kruskal-Wallis test confirmed the presence of an overall difference between groups [Kruskal-Wallis χ²(3) = 499.43, P < 2.2 × 10⁻¹⁶], and individual nonparametric Wilcoxon rank sum tests indicated that differentiation of the between-depths group was significantly greater than that of the within-shallow and within-deep population groups (W = 20,749,000 and 21,537,000, P < 2.2 × 10⁻¹⁶). However, differentiation of the between-depths group was significantly different from that between the western (WD) and the eastern (GD, JD, and PD) deep populations. No significant differences were observed between the level of differentiation of the within-shallow and that of the within-deep populations, apart from differentiation of the western deep population (W = 17,196,000, P < 2.2 × 10⁻¹⁶).
the shallow and deep cluster, respectively ($K = 2$). A total of 108 SNPs (in addition to the identified 25 depth-outlier SNPs) exhibited strong allelic differences ($\Delta p \geq 0.5$) between these “purebred” groups. Plotting the genotypes of all individuals for these SNPs across the ancestry assignment gradient reveals introgression from shallow into deep individuals, which is also reflected in the overall ancestry assignments (Fig. 5). In contrast, introgression from deep into shallow appears very limited (for example, only minor assignment of shallow individuals to the deep cluster). The genetic diversity (that is, expected heterozygosity) of deep populations is also consistently higher than that of shallow populations (table S1). Despite the shallow-to-deep introgression, no migrants were observed, indicating limited migration over ecological time scales.

**Lack of genetic structuring in *S. intersepta***

No genetic structuring across populations was identified for *S. intersepta* (G-statistic Monte Carlo test for all SNPs: $G_{\text{obs}} = 107,590, P = 0.3$). The lack of genetic structure was further confirmed by assessing pairwise genetic distances between individuals (Fig. 2), and the lack of clustering identified by PCA (Fig. 4D), DAPC, and STRUCTURE (maximum MedMeaK and MedMedK of one cluster; Fig. 4C and table S2). When running STRUCTURE for markers in the 90th to 100th percentile of overall $F_{ST}$ (that is, most divergent SNPs), clear structuring between all seven populations was observed ($K = 7$), but the same structuring could be achieved when shuffling population assignments, highlighting that this apparent recovery of population delineation is an artifact of a priori population assignment (fig. S3). Genetic diversity (that is, expected heterozygosity) was slightly higher in shallow populations (table S1). Despite the lack of genetic differentiation among locations and depths, the Bermuda population was highly differentiated from the two *S. intersepta* samples from Curaçao (which were included as an out-group; Fig. 2). The Curaçao individuals contained 52 additional SNPs that were fixed in the Bermuda population.

**Natural species densities and relative contribution to overall community structure**

The species *A. fragilis* and *S. intersepta* are depth-generalists that were consistently encountered at both shallow (~12 m) and deep (~40 m) depths around Bermuda (Fig. 6A). Two additional species were also found to be common at both sampling depths: *M. cavernosa* [focus of a previous vertical connectivity study (25)] and *Orbicella franksi*. The population densities of these four depth-generalist species were highly variable across locations (Fig. 6A), but for all species other than *O. franksi*, population densities were higher at depth (except for *A. fragilis* at Gurnet Rock). In terms of the overall species composition at the three locations (Fig. 6B), the four depth-generalist species
should not be assumed to be a broader ecosystem-wide phenomenon. Ecologically relevant to these individual species, deep-water coral reseeding restricted to only a subset of depth-generalist coral species; although species intensive gene flow and lack of differentiation characterize the broadcasting for endosymbiont contamination), we demonstrate that selection drives reduced-representation genome sequencing approach (which accounts sampling of adjacent shallow and mesophotic populations and a vary greatly between species within a single reef system. Using replicated tested (13), our study demonstrates that vertical connectivity can vary greatly between species within a single reef system. Using replicated sampling of adjacent shallow and mesophotic populations and a reduced-representation genome sequencing approach (which accounts for endosymbiont contamination), we demonstrate that selection drives depth differentiation in the brooding species Agaricia fragilis. In contrast, extensive gene flow and lack of differentiation characterize the broadcasting species S. intersepta. Overall, the reseed potential of deep reefs appears restricted to only a subset of depth-generalist coral species; although ecologically relevant to these individual species, deep-water coral reseeding should not be assumed to be a broader ecosystem-wide phenomenon.

The strong genome-wide differentiation between shallow and mesophotic populations of Agaricia fragilis was linked to depth-associated selection for a select number of RAD loci (Fig. 3, A and B). A pattern of local-scale genetic differentiation is commonly observed among brooding species and is generally associated with the more localized dispersal of both sperm and larvae (45, 47–50). In the case of Agaricia fragilis, geographic distance cannot account for the strong observed differentiation between shallow and deep eastern populations, because differentiation between depths within each of the three locations consistently exceeded that observed for the same depth between locations (average horizontal distances of ~1.5 km versus ~12 km, respectively). Outlier analyses identified strong depth-associated selection across a number of RAD loci (n = 12; 0.8% of those examined). In the presence of substantial gene flow, such divergent selection across opposing environments is expected to result in heterogeneous genomic divergence (or “genomic islands of divergence”) (51). Instead, for Agaricia fragilis, the background levels of genetic differentiation were also observed to be high (with genetic structuring remaining after removing outliers), indicating that the depth-associated selection in Agaricia fragilis has led to genome-wide “resistance” to shallow-deep gene flow.

Given the observed lack of shallow-deep migrants, immigrant inviability (that is, selection against migrants between locally adapted
Fig. 4. Genetic structuring across depths and locations. (A) STRUCTURE diagram (K = 3) for *A. fragilis* as inferred from the overall (top) and neutral (bottom) data set. (B) PCA for *A. fragilis* as inferred from the overall data set. (C) STRUCTURE diagram (K = 2) for *S. intersepta* as inferred from the overall data set (top) and discriminant analysis of principal components (DAPC) for the overall data set (bottom). (D) PCA for *S. intersepta* as inferred from the overall data set. STRUCTURE and DAPC bar graphs indicate ancestry proportions for individuals in each population. In the PCA, sampling depth of individuals is indicated by hue [shallow (light) and deep (dark)], and location is denoted by symbol shape (following Fig. 1A).

Fig. 5. Admixture between shallow and deep populations in *A. fragilis*. SNP genotypes for the 25 outlier SNPs (indicated in red hues) (left) and 108 additional highly divergent SNPs (indicated in grayscale) (middle) for all shallow and deep individuals (eastern populations only) sorted by depth and overall ancestry assignment (right). Hues indicate genotype [following Fig. 3 (B and D)] with a white color indicating missing data. In the overall ancestry assignment, light green refers to the “shallow” cluster, and dark green corresponds to the “deep” cluster, with the horizontal solid line separating individuals originating from shallow and deep reefs. Highly divergent SNPs were selected as those with an allele frequency difference (Δp) of ≥0.5 between shallow and deep individuals with >0.98 ancestry assignment (threshold indicated by the dashed lines).
populations) is likely an important contributor to this resistance to gene flow in *A. fragilis*, reducing the chances of between-ecotype mating encounters and leading to assortative mating (52). Despite this apparent gene flow barrier, we observed introgression between populations at shallow and mesophotic depths, which may be facilitated through ecologically rare events of migration and interbreeding that were detectable with our relatively small sample sizes. Alternatively, limited introgression could be facilitated by stepping-stone populations at intermediate depths (not sampled), effectively enabling "multigenerational vertical connectivity" (15). Either way, the extent of vertical connectivity does not appear to be ecologically relevant.

Introgression between populations of *A. fragilis* was asymmetrical (Fig. 5), matching observations from two other studies reporting directional gene flow from shallow to deep populations in the broadcasting species *M. cavernosa* and the octocoral *Eunicella flexuosa* (24, 25). In the case of *E. flexuosa*, higher gamete production in shallow water (due to greater abundances and average colony sizes) was offered as a likely explanation for the asymmetric introgression (24). However, for *A. fragilis*, this explanation appears unlikely given that population densities at two of three locations (Fig. 6) appear lower at shallow depths. Instead, the lower population densities and the near-complete or complete fixation of the major allele in 8 of 12 outlier RAD loci at shallow depths (Fig. 3) indicate that selection against immigrants may be stronger at shallow depths and locations. Differential selective pressures could be a contributor to the observed asymmetric introgression (Fig. 5) and associated higher genetic diversity at mesophotic depths (table S1).

The observed introgression and the lack of alternatively fixed polymorphisms in *A. fragilis* appear to reflect a population-level rather than a species-level divergence. This is further corroborated by the observation that shallow-depth differentiation (eastern locations) is comparable to that observed between the western and eastern deep populations (fig. S2). Although distance and/or seascape resistance (for example, in relation to prevailing currents) may have played a role in the differentiation of this Western Blue Cut (WD) location, several outlier loci were identified in association with this specific population, perhaps indicative of environment-based selection (fig. S1). The western population also exhibited a phenotypic skeletal signature distinct from the other deep locations (Fig. 1B).

Despite this study focusing on vertical connectivity, the identification of both neutral differentiation and outlier loci in association with this western location reiterates the broader role of environment-based selection in cross-habitat connectivity (21, 28, 53, 54).

For the gonochoric broadcasting species *S. intersepta*, no genetic structuring was observed across depths and locations (Figs. 2 and 4). Strong genetic differentiation was only observed for the two out-group individuals from Curaçao (Fig. 2), which is expected given their origin near the opposite end of the latitudinal distribution of this species. Eggs released by *S. intersepta* (as well as by *M. cavernosa*) have been observed to exhibit high rates of fertilization immediately upon release (55), which appears to be facilitated by intratentacular fertilization (56). Although this rapid fertilization may not directly affect larval dispersal, it would result in more localized cross-fertilization similar to that in species with a brooding reproductive mode. Assuming such localized spermcasting, the observation of a single panmictic population across the reef platform of Bermuda indicates considerable larval migration between both depths and locations. Similar genetic connectivity over depth has been inferred in Bermuda for the broadcasting species *M. cavernosa* (4 to 58 m depth) (25). In contrast to the brooding species *A. fragilis*, no RAD loci were identified to be under depth-specific selection [although, of course, only a reduced proportion of the genome was queried (57)]. In combination, the observed panmixia, apparent morphological plasticity, and association with a single endosymbiont highlight the depth-generalist nature of *S. intersepta* and the potential for deep populations of this species to act as a larval source for shallow reefs.

The subtraction method allowed us to successfully apply a de novo reduced-representation genome sequencing approach (in this case, nextRAD) on endosymbiont-contaminated samples. Despite our attempt to reduce the endosymbiont contamination through multiple steps of centrifugation before DNA extraction, we still found 10 to 14% of RAD loci to be of *Symbiodinium* origin, highlighting the necessity of further steps to identify and remove such contamination. Only a small fraction of these sequences could be identified solely by comparison with *Symbiodinium* reference genomes, which is not unsurprising given their potential level of divergence (particularly in the case of the *Symbiodinium minutum* "B1" genome). This indicates that comparison with unrelated reference genomes alone is not a viable approach to eliminating endosymbiont contamination. Thus, although genotyping-by-sequencing approaches have demonstrated their de novo potential in nonmodel organisms, population genomic assessments of endosymbiotic marine invertebrates still require either an aposymbiotic

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**Fig. 6. Depth-generalist species abundances and overlap in community structure.** (A) Species densities of the four depth-generalist coral species common to shallow (left) and deep (right) reefs in Bermuda. (B) Proportion of the four depth-generalist species of the overall coral community on shallow and deep reefs in Bermuda (with the proportion of other species indicated in white). Shading behind pie graphs indicates the proportion of species with vertical connectivity potential: *M. cavernosa* (25), *S. intersepta* (this study), and *O. franksi* (based on reproductive mode and depth distribution).

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host reference [by sequencing aposymbiotic sperm or larvae; for example, see related studies (35–37)] or an endosymbiont subtraction reference (as in this study). Because obtaining aposymbiotic tissue can be difficult, particularly in brooding species, our subtraction method based on fluorescence-activated cell sorting provides an easy and cost-effective solution to this pervasive issue.

Overall, we demonstrate that coral species with similar ecological distributions can exhibit very distinct potentials for vertical connectivity at a single location. Geographic congruence in patterns of vertical connectivity in M. cavernosa and P. astreoides (despite their opposing reproductive modes) highlighted the importance of location-specific extrinsic factors in determining vertical connectivity (14). Our study focused on a single geographic region to specifically evaluate the role of depth, controlling for location-specific effects (by sampling replicate pairs of proximate shallow and mesophotic populations) and sampling genetic variation across the genome. The observed pattern for A. fragilis demonstrates how depth-associated selection can hamper vertical connectivity in the absence of physical dispersal barriers and corroborates the emerging notion that divergence by depth is prevalent in species with a brooding reproductive mode (21, 22, 29, 58). In contrast, we found that the broadcasting species S. intersepta formed a single panmictic population across the same depths and locations. Nonetheless, divergence by depth has been observed in both brooding and broadcasting species (14, 21–25), and the hypothesis that vertical connectivity is more prevalent in broadcasters than in brooders (11) cannot be evaluated until more species and locations are assessed. Regardless, these results demonstrate that the reseeding potential of deep reefs only applies to a subset of depth-generalist species.

Of the ~20 zooxanthellate scleractinian coral species described for Bermuda (59), 6 species appear to be depth-generalists occurring at both shallow and mesophotic depths (44, 60). Among these species, A. fragilis, Madracis decactis, and Scolymia cubensis are brooders, while S. intersepta, M. cavernosa, and O. franksi are broadcasters (61). Although one of the brooders, A. fragilis, occurs in reasonable abundance at both shallow and mesophotic depths, our study showed little evidence for vertical connectivity between shallow and deeper-water populations. On the basis of their reproductive mode, such vertical connectivity may be similarly hampered for M. decactis and S. cubensis. Regardless, S. cubensis was rare at both depths, and M. decactis was only common at intermediate and mesophotic depths along the southern shore (and therefore not individually reported in Fig. 6A), further indicating a limited importance of these species in a reseeding context. Similarly, P. astreoides (for which vertical connectivity was observed across 4 to 26 m depth) (14) was relatively rare at mesophotic depths (~40 m). In contrast, all three broadcasting species occurred in reasonable abundance at both shallow (12 m) and upper mesophotic (40 m) depths (Fig. 6A), and mesophotic populations of these species may represent a viable source of recruitment for the shallow reefs in this isolated reef system (although O. franksi has not been assessed). Nonetheless, although these three species constitute a considerable proportion of the diversity and composition of the upper mesophotic reef, they only represent a relatively small fraction of the diversity (~15% of species) and community (~15% of coral colonies) in shallow water (Fig. 6B).

The DRRH postulates that deep reefs (i) are protected or dampened from disturbances that affect shallow reefs and (ii) can provide a reproductive source after disturbance [sensu Bongaerts et al. (11)]. Although deep reefs can escape certain disturbances, they are not entirely immune to disturbance (6–9, 26). In fact, even their ability to escape bleaching and storm events can vary spatially and temporally, and their greater depth does not confer long-term protection against disturbances associated with climate change (9). Similarly, disturbance events can vary strongly in their impact over depth and in their impact on specific species, which can vary in thermal susceptibility or structural fragility. Thus rather than the term “refugia” (as currently used in “deep reef refugia hypothesis”), which Keppel et al. (62) classify as large geographic areas offering an escape over evolutionary time scales (for example, climate change refugia), we propose that the term “refuge(s)” is probably more appropriate (that is, “deep reef refuge hypothesis”), reflecting the role of deep reefs in providing temporal and/or spatial protection from disturbances on ecological time scales. In addition, this study confirms that the active reseeding potential is specific to only a subset of species (~15% here) and does not apply to a significant proportion of the species (diversity) that make up shallow reefs. Therefore, we conclude that the “deep reef refuge hypothesis” should be considered as an ecological concept relevant to individual species, rather than being referred to as an ecosystem-wide phenomenon.

**MATERIALS AND METHODS**

**Study locations, specimen collections, and skeletal morphometrics**

We focused on the isolated Bermuda platform in the Western Atlantic, where we targeted shallow (12 ± 2 m) and deep (40 ± 2 m) habitats at four different reef locations: Gurnet Rock, John Smith’s Bay, and Princess Beach on the southeastern side and Western Blue Cut on the northwestern side of the island (table S1). Specimens (n = 30 per depth at each location) of A. fragilis (n = 208) and S. intersepta (n = 209) were collected under collection permit no. 130901b from the Department of Environmental Protection of the Bermuda Government, as part of the “XL Catlin Seaview Survey” in September 2013. Two additional samples of S. intersepta from Curaçao were included for use as a comparative out-group. Small fragments of the collected specimens were stored in salt-saturated buffer solution containing 20% dimethyl sulfoxide and 0.5 M EDTA, and for a subset of specimens, a skeletal voucher was bleached, rinsed in freshwater, and dried. For these skeletal vouchers (A. fragilis, n = 49; S. intersepta, n = 37), corallite diameter and density were assessed across five pseudoreplicates per specimen.

**Separation of host fractions and DNA extraction**

To counter the issue of pervasive endosymbiont contamination, we separated the tissue from the skeleton and performed multiple centrifugation steps separating out the supernatant to eliminate intact Symbiodinium cells. Genomic DNA (gDNA) was extracted using a modified version of the “Wayne’s method” (for details, see electronic notebook). Quality and yield of gDNA were assessed using gel electrophoresis and a Qubit fluorometer (Invitrogen) to select a subset of high-quality samples (n = 16 per species per depth at each location) for downstream sequencing (A. fragilis, n = 110; S. intersepta, n = 113; Symbiodinium, n = 15). Selected samples were normalized (gDNA quantity) and purified using AMPure XP beads to remove potential inhibitors.

**Isolation of Symbiodinium**

Given that a certain degree of endosymbiont contamination remained in the “host” samples [confirmed through polymerase chain reaction (PCR) amplification with Symbiodinium-specific primers, likely due to lysed cells], we isolated individual Symbiodinium cells from the original pellet (which was, in turn, contaminated with a certain extent of host cells) to be sequenced separately as a subtraction reference. Symbiodinium isolation was carried out for aliquots of a shallow and deep specimen...
from each of the two species using fluorescence-activated cell sorting (BD FACSAria cell sorter) at the Queensland Brain Institute, separating out ~300,000 individual cells based on their distinct fluorescence signature (for details, see electronic notebook). In addition, Symbiodinium cells from several other coral host species were isolated from samples intended for other species-specific studies. Extractions of gDNA for the isolates were carried out as for the host.

Library preparation, sequencing, and clustering
Library preparation was carried out using the nextRAD method (Nextera-fragmented, reductively amplified DNA; SNPsaurus LLC), which uses a selective primer sequence (rather than restriction enzymes) to genotype loci consistently between samples (63). gDNA was fragmented and ligated with adapter sequences using Nextera reagent (Illumina Inc). Fragmented DNA was then PCR-amplified (73°C for 26 cycles) with one of the primers matching the adapter and extending into the gDNA using a 9-base pair (bp) selective sequence (“GTGTAGAGG”). Libraries were sequenced on three HiSeq 2500 (Illumina Inc) lanes using 100-bp single-end chemistry and following the manufacturer’s recommended protocol. Quality control, clustering, and SNP variant calling were carried out using PyRAD v3.0.6 (64) to allow for indel variation (common in marine invertebrates). After initial runs at different thresholds, we separately analyzed both species using a clustering threshold of 90%, a minimum coverage of 6, a maximum of three sites with a PHRED quality below 20, and a minimum of 10 samples in a final locus. Sequence data from the isolated Symbiodinium were filtered using PyRAD, allowing for a maximum of five sites with a PHRED quality below 20 (step 1), and reads were then clustered within samples using an 85% clustering threshold (step 3). Clusters with a coverage of ≥2 (to eliminate singletons) were reduced to a single representative sequence and used to compile a BLASTN database (n = 697,146 sequences). Parsing and analyses were carried out using custom Python (v3.4.5) scripts (available through https://github.com/pimbongaerts/radseq), unless otherwise indicated, with statistical analyses and plotting performed in R (v3.3.1) (electronic notebook available through https://github.com/pimbongaerts/bermuda-rad).

Identification of coral host loci and QC
One representative sequence was extracted for each coral RAD locus, which was then BLASTN-searched against our Symbiodinium RAD database, and two culture-based Symbiodinium genomes [Symbiodinium B1 from Shoguchi et al. (65) and Symbiodinium C1 from Chan et al., draft genome], with positive matches (maximum E-value of ≤10−15) removed from the coral RAD data set. An additional BLASTN comparison was carried out against the National Center for Biotechnology Information (NCBI) nonredundant nucleotide database (with the particular aim of removing any microbial contamination, if applicable), with taxonomic IDs of positive matches (maximum E-value of ≤10−4) extracted and classified to phylum using the NCBI Taxonomy database. RAD loci matching to non-cnidarian taxa were subsequently removed. Given an overrepresentation of SNPs toward the end of reads, RAD loci were then truncated to 75 and 80 bp for A. fragilis and S. intersepta, respectively. SNPs representing singletons/doubletons were removed (rather than filtering for a minimum allele frequency), as were SNPs that were genotyped for <50% of individuals in each population (rather than using an overall representation filter). Genetic distance/similarity (Hamming-based) was then calculated between all pairs of individuals to identify clones, with clonal groups reduced to a single representative per population. The remaining SNPs were then evaluated for significant deviations from the Hardy-Weinberg equilibrium using arlecore (v3.5.2.2) (66), with SNPs that deviated (excess or deficit) in more than five populations removed, as well as RAD loci that contained an SNP exhibiting a significant excess in at least one population (to filter out potential paralogous RAD loci).

Outliers and private alleles
Initially, outliers were identified for the overall data set and for pairwise population comparisons (considering only biallelic SNPs) using the Fdist approach (67) as implemented in Lositan (v. 1.0) (68) and BayeScan (v.2.1) (69). To conservatively identify depth-outliers, we considered those SNPs that were identified in the overall outlier analyses by both Fdist [outside the 99% confidence interval (CI); 50,000 simulations] and BayeScan (using a Bayes factor cutoff of 0.01), as well as those that were identified in at least two of three independent Fdist outlier analyses between individual shallow and deep populations within a location (but never in Fdist outlier analyses within the same depth between different locations). Genotype frequencies of outliers were calculated, and sequences of depth-outlier RAD loci were searched (BLASTN) against the NCBI nonredundant nucleotide database for potential gene region identification. To identify a subset of SNPs that was more representative of neutral genomic diversity, we compiled a separate data set (neutral), eliminating SNPs identified in one or both overall outlier analyses using more lenient cutoffs (95% CI for Fdist and a Bayes factor cutoff of 0.05 for BayeScan).

Genetic structure
To assess the genetic structure for the outlier and nonoutlier data set, we wrote a multiprocessing wrapper (“structure_mp”) for STRUCTURE (v.2.3.4) (70), which takes a Variant Call Format file and converts it into distinct replicate STRUCTURE data sets, each with a single SNP per RAD locus, which are then run in parallel through STRUCTURE and permuted and summarized in CLUMPP (v.1.1.2) (71). Four estimators of the number of clusters (MedMeaK, MaxMeaK, MedMedK, and MaxMedK) were determined using a membership coefficient threshold of 0.5 [sensu Puechmaille (72)]. STRUCTURE runs were conducted using the admixture model with correlated allele frequencies and not considering priors (burn-in of 100,000 and 50,000 reps). Additional STRUCTURE runs were carried out for the eastern populations of A. fragilis (K = 2) for downstream ancestry analysis and for the most differentiated SNPs (90th to 100th percentile) of S. intersepta (based on overall FST calculated for the original data and when using random population assignments). Genetic structure was also assessed for both species through PCA and DAPC, both using the adegenet R package (v1.3-1) (73).

Genomic intergradation of shallow and deep individuals
Patterns of SNP ancestry were assessed for shallow and deep individuals of A. fragilis (eastern populations only) by classifying the data set into two putative “parental” groups (those with an ancestry assignment of >0.98) and an admixed group. We then extracted genotypes for the identified depth-outlier SNPs, as well as other SNPs that had a reference allele frequency differential (Δp) between parental groups of at least 0.5, and plotted for all the individuals sorted by ancestry assignment (similar to the visualization in the R package “introgress”) (74).

Population genetics statistics
To test for overall population structure in both species, we implemented Goudet’s G-statistic Monte Carlo test (75) using the “hierfstat” R...


SYNTHETICAL MATERIALS
Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/3/2/e1602373/DC1

fig. S1. Genotype frequencies of SNP outliers for A. fragilis.
fig. S2. Genome-wide pairwise differentiation (F ST ) of populations.
fig. S3. Genetic structuring of S. intersepta populations based on most divergent SNPs.
table S1. Details of the sampled A. fragilis and S. intersepta populations.
table S2. Estimators of the number of STRUCTURE clusters for A. fragilis and S. intersepta.
table S3. Genome-wide pairwise differentiation (F ST ) of populations.

REFERENCES AND NOTES
Limited gene flow in the mussel hill coral (Porites astreoides) from different reef environments.


the fieldwork and specimen collections. R.B. conducted the laboratory work and photographic analysis. P.B. performed the data analyses and wrote the manuscript, with contributions from C.R. All authors edited the manuscript before submission. **Competing interests:** The authors declare that they have no competing interests. **Data and materials availability:** All data and analyses are available as an electronic notebook (https://github.com/pimbongaerts/bermuda-rad), with scripts available through https://github.com/pimbongaerts/radseq. Raw Illumina sequence data can be obtained from the NCBI Sequence Read Archive (SRA) under the accession code PRJNA361144 (https://www.ncbi.nlm.nih.gov/bioproject/361144).

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