Review Article

Deep thinking: a systematic review of mesophotic coral ecosystems

Joseph A. Turner,1,2* Russell C. Babcock,1,2,3 Renae Hovey,1 and Gary A. Kendrick1

1School of Biological Sciences, University of Western Australia, 35 Stirling Highway, Crawley, WA 6009, Australia
2CSIRO Oceans and Atmosphere, Indian Ocean Marine Research Centre, University of Western Australia, M097, 35 Stirling Highway, Crawley 6009, Australia
3CSIRO Oceans & Atmosphere, Dutton Park, Brisbane, Queensland 4001, Australia

*Corresponding author: tel: +61 8 9333 6021. e-mail: joseph.turner@research.uwa.edu.au

Turner, J. A., Babcock, R. C., Hovey, R., and Kendrick, G. A. Deep thinking: a systematic review of mesophotic coral ecosystems. – ICES Journal of Marine Science, 74: 2309–2320.

Received 16 November 2016; revised 1 May 2017; accepted 2 May 2017; advance access publication 26 May 2017.

Mesophotic coral ecosystems (MCEs) occur at depths beyond those typically associated with coral reefs. Significant logistical challenges associated with data collection in deep water have resulted in a limited understanding of the ecological relevance of these deeper coral ecosystems. We review the trends in this research, covering the geographic spread of MCE research, the focus of these studies, the methods used, how MCEs differ in terms of species diversity and begin to assess connectivity of coral populations. Clear locational biases were observed, with studies concentrated in a few discrete areas mainly around the Atlantic region. The focus of MCE studies has diversified in recent years and more detailed aspects of MCE ecology are now being investigated in particular areas of research. Advances in technology are also reflected in the current range of research, with a wider variety of methods now employed. However, large information gaps are present in entire regions and particularly in relation to the threats, impacts and subsequent management of MCEs. Analysis of species diversity shows that initial definitions based on depth alone may not be appropriate globally, while further taxonomic resolution may also be required to deduce the full biodiversity of major groups in certain regions. Genetic studies to date show species-specific results, although distinct deeper populations do appear to exist, which raises questions regarding the potential of MCEs to act as refugia.

Keywords: connectivity, coral reefs, mesophotic coral ecosystems, twilight zone.

Introduction

Coral reefs are in worldwide decline, due to increased mass disturbance events brought about by climate change and anthropogenic activities (Hughes et al., 2003; Bellwood et al., 2004; Hoegh-Guldberg et al., 2007). However, a majority of the data on which these projections are based are from coral reefs shallower than 20–30 m, while the trends below this depth remain unknown (Bak et al., 2005; Bridge et al., 2013). Deeper mesophotic coral ecosystems (MCEs) are defined as tropical and sub-tropical light-dependent communities occurring from approximately 30 m to the lower limit of the photic zone, extending as deep as 150 m in some locations (Hinderstein et al., 2010). These reefs are perceived as continuations of the shallow reef communities, with a similarly diverse range of taxa (Lesser et al., 2009; Hinderstein et al., 2010). Communities are primarily structured by light (Sheppard, 1982; Lesser et al., 2009) although there are other factors at play, including topography (Bridge et al., 2010), temperature (Kahng et al., 2012), sedimentation, and water movement (Goreau and Goreau, 1973; Sheppard, 1982).

Mesophotic communities have been broadly described (Bushy, 1966; Goreau and Goreau, 1973; Bouchon, 1981; Sheppard, 1982; Fricke and Meischner, 1985; Colin et al., 1986; Fricke and Knauer, 1986; Thresher and Colin, 1986) but remain relatively unexplored compared to shallow water reefs, particularly in relation to ecological characteristics and functions. This is primarily due to their location, lying beyond recreational SCUBA diving limits, and therefore posing increased logistical challenges (Lesser et al., 2009; Kahng et al., 2010). Mesophotic reefs are starting to
gain more attention as modern technological advances make them increasingly accessible (Lesser et al., 2009; Kahng et al., 2010). Advances in habitat mapping and technologies such as Remotely Operated underwater Vehicles (ROVs) and Autonomous Underwater Vehicles (AUVs) can provide a useful platform for monitoring these systems (Singh et al., 2004; Armstrong et al., 2006; Bridge et al., 2011a). Increased interest in MCEs is evident in the exponential increase in publications following recent workshops and special journal theme sections (Loya et al., 2016). During the past decade in particular, knowledge of these systems has moved on significantly.

MCEs have only been studied in a few areas of the world resulting in little generalizable knowledge of the drivers of their structure, function, connectivity and refugia role for shallow reefs globally. There is a poor understanding of the role environmental factors have in influencing spatial patterns in community structure, and therefore how MCEs respond to anthropogenic threats and climate change (Puglise et al., 2009; Kahng et al., 2014). MCEs can harbour diverse biological assemblages of corals, fish and other invertebrates consisting of a range of “deep-specialist” and “depth-generalist” species (Bongaerts et al., 2010a; Kahng et al., 2014). Some species are endemic to these systems, highlighting the importance of MCEs in contributing to and maintaining global biodiversity (Heyward et al., 2010; Bridge et al., 2011b; Kane et al., 2014; Muir et al., 2015). As more studies are completed, the limitations to our current knowledge have become evident. For example, studies investigating mesophotic areas of the Great Barrier Reef identified that submerged reef habitat may have been underestimated by as much as 100% (Harris et al., 2012) and new species records for Australia have also been found (Muir et al., 2015).

It has been suggested that MCEs function as refugia, where communities are sheltered from perturbations in shallow waters including high temperature, sedimentation, storm damage and fishing and so may re-seed more frequently disturbed shallow reefs (Bongaerts et al., 2010a; Hinderstein et al., 2010). Larval connectivity needs to be understood, in order to assess the extent of re-seeding potential, including whether species are present in deep and shallow water and how they are connected through the movement of currents (Lesser et al., 2009; Slattery et al., 2011; Baker et al., 2016). A zone that harbours both shallow reef and mesophotic species appears to occur in a number of the locations studied, generally between 30 and 60 m (Lesser et al., 2009; Slattery et al., 2011), often termed the upper-mesophotic. However, in deeper areas, high levels of depth-endemism would suggest greater habitat specialization, and possibly limited larval exchange with shallower waters and a reduced ability to replenish shallow habitats (Slattery et al., 2011). If the community structures between deep mesophotic and shallow coral reefs are different then re-seeding will not be possible.

Information on the distribution and extent of MCEs, the factors that determine their distributions, and the organisms found in these ecosystems, are all critical to inform biodiversity management (Puglise et al., 2009; Baker et al., 2016). The spatial distributions of rare and ecologically important habitats are required to adequately design networks of Marine Protected Areas and ensure representation of all habitat types (Bridge et al., 2016a). Mesophotic reefs are likely to provide similar ecosystem services to those of shallow water reefs and can contribute to fisheries, tourism, and pharmaceutical uses (Eyal et al., 2015; Baker et al., 2016). Identifying the key ecosystem services provided by these systems is important in order to gain support for their protection (Puglise et al., 2009). Adopting a broad, ecosystem-wide approach that encompasses deep reefs is most likely to have many environmental, social and economic benefits (Bridge et al., 2013).

This review investigates the current literature regarding MCEs. The term “deep coral reef” is often used to refer to much deeper water ecosystems of aphytic species associated with colder water, e.g. Lophelia pertusa, defined as living without light (Freiwald et al., 2004). While these ecosystems may occasionally occur in mesophotic depth ranges at high latitudes, such as in Norwegian fjords, or at similar latitudes but at much greater depths (Roberts et al., 2006), they function differently from shallower coral ecosystems; with the term cold-water corals coined to differentiate them from tropical coral reefs (Freiwald and Roberts, 2005). For this review, MCEs will be defined as in Hinderstein et al. (2010) as light-dependent coral-dominated systems in tropical regions that form extensions of shallow coral reefs. The aims are to: (1) Characterize study locations and global hotspots of MCE research, (2) Identify trends in MCE research topics, (3) Identify the methods used, including how they have changed over time, (4) Describe how mesophotic biodiversity may differ between locations, and (5) Describe connectivity trends across shallow reefs to mesophotic depths. Assessing the work done so far will allow us to identify and characterize the key aspects of MCEs as well as identifying the key gaps in our understanding to inform future research direction.

**Methods**

A literature review was carried out following the systematic methods outlined in Pickering and Byrne (2014) and Pickering et al. (2015). The databases Google Scholar, Web of Science, and Scopus in May 2016 and February 2017 were searched using the search terms:

mesophotic

AND

reef OR coral OR fish OR sponge OR connectivity OR ecolog* OR community OR recruit* OR impact OR disturbance

The specialist database at mesophotic.org (http://www.mesophotic.org/publications/), maintained by field experts, was also utilized, and all papers were screened for content. As “mesophotic” is a relatively new term to be applied to reef ecosystems, defined in Puglise et al. (2009), we further checked references from recent review articles (Lesser et al., 2009; Kahng et al., 2010; Kahng et al., 2014; Baker et al., 2016; Loya et al., 2016) to ensure all relevant papers were acquired. Still, the search was conservative and some papers that did not use the search terms we utilized would not have been identified. Results were limited to those with an English title and abstract.

Studies were screened to ensure relevance in a two-step process outlined below, and results are shown in Figure 1.

(1) Titles and abstracts were required to mention or contain information on:

(a) mesophotic or deep/twilight reef,

(b) tropical habitats, and

(c) coral reef ecology.
Following screening, the full texts of relevant articles were obtained and reviewed. Studies were excluded if the main aims did not concentrate on mesophotic depths and communities (e.g., "deep" areas can relate to less than 10 m in some studies and so would not meet the aims of this review). A study had to encompass a proportion of the 30–150 m depth range defined for mesophotic areas in order to be considered relevant.

In order to gather data to meet Aims 1–3 the following information was recorded for all papers:

- Authors and Title
- Geographic location (including coordinates), split into region [based on those used by Burke et al. (2011)], country and study area
- Year
- Primary research focus (Table 1)
- Methods used for data collection
- Depth range investigated

Only a subset of papers provided information to investigate Aims 4 and 5. Mesophotic species diversity, deepest records, or the depth at which significant changes in community structure occur was included in the database when available. Any study that concentrated on genetic differences was investigated for Aim 5. This is currently the most effective method to quantify connectivity between deep and shallow areas: we extracted information on species, whether there was a genetic change with depth and if so what depth the changes occurred.

Data manipulation and analysis was conducted in R (R Core Team, 2010) and figures were constructed using the ggplot2 package (Wickham, 2009). Aims 1–3 involved summarizing the information by location (map produced in ArcGIS 10.4), research focus and method. To address Aim 4 summary statistics were calculated for species diversity and transition depths between regions. Due to the relatively few data points, a rigorous statistical analysis was not possible for Aims 4 and 5.

**Results**

A total of 349 papers were classified in this study, spanning from 1966 to 2017. A majority of the studies on mesophotic reefs have been completed since 2010 (56%) (Figure 2), with 54 studies (15%) completed in 2016 alone. Research is concentrated in specific regions and countries (Figure 3) with over half (57%) of global mesophotic studies having been carried out in the Atlantic region, particularly in the Caribbean.

Research into mesophotic reefs is globally very regionally localized. For example, while research spans a number of countries in the Atlantic (Table 2) effort is disproportionally split across them. Additionally, studies can concentrate in specific countries; with Israel (Middle East) and Hawaii, USA (Pacific) contributing to 91 and 71% of the entire studies for that region respectively. A single country, the USA has the greatest number of studies (18%, split over two regions) although they are focussed in geographically small areas with almost all studies occurring in Hawaii (Pacific) and Florida (Atlantic). Australia (13% of global studies) has observed significant modern research interest with 70% of Australian studies occurring since 2010. Again, studies are localized with 50% occurring on the Great Barrier Reef. The Indian Ocean and Southeast Asia are significantly under-represented (1 and 2% of global studies, respectively).

Descriptive studies dominate the MCE literature (Table 3). However, research focus has shifted over time (Figure 4). The proportion of descriptive studies drops from 92% before 1980 to 33% post-2010. Research focus has also diversified, with increases observed in all other research categories between 2001 and 2011 onwards. Studies investigating molecular ecology have seen the largest increase, from zero before 2000 to 7% then 16% in 2001–2010 and post-2010, respectively. These studies are mostly conducted in the Atlantic (47%, exclusively in the Caribbean), Australia (24%) and the Pacific (21%). Life history studies and research focussing on impacts are in their infancy, only occurring since 2000. This work is currently highly concentrated in the Atlantic region with 78% of life history and 59% of impact (including natural and anthropogenic) studies taking place there.

A variety of methods are used to study MCEs (Figure 5). Although lying beyond recreational diving depths, SCUBA diving is the most common method used for most year categories (Second to Submersibles in the 1980s and 1990s). Even prior to 1980, 69% of the studies were completed using SCUBA-based observations, before advances in technical diving such as closed-circuit rebreathers (CCRs), and when health and safety regulations were less conservative. Methods have diversified widely since 2001 as more techniques have become available. Recent studies appear to be utilizing a number of methods as the research focus diversifies. Experimental and genetic labwork are now used more widely, as finer ecological details of MCEs are explored, with large increases in the use of these methods observed from 2011. Technological advances are observed with the arrival of ROVs, AUVs, and Baited Remote Underwater Video (BRUVs) from 2001 onwards. SCUBA remains the most popular method, accounting for 33% of studies post-2010 although labwork, including identification, experiments, and genetics, totals 31%.
Like their shallow-water counterparts, MCEs vary with respect to their location around the globe (Baker et al., 2016). While accurate species numbers are rarely reported, results that could be extracted are summarized in Table 4. No data were available for the Indian Ocean and South-East Asia regions but MCEs harbour high benthic and fish biodiversity in all other regions. Atlantic MCEs are less species-rich in terms of scleractinian corals, but macroalgal and sponge diversity is high. Challenges in species identification are a likely cause of low numbers of sponge and algal studies, particularly as remote methods become increasingly used. Reported transition depths, defined as the boundary where significant changes in species composition are observed, thus representing the transition between upper and lower mesophotic communities, appear variable (Table 5). Fish transition depths appear shallower, however, this is probably an artefact of most fish studies including surveys at shallower depths (Mean start depth = 22.7 ± 4.4 m) with 61% of studies completed using SCUBA. This could be interpreted better as the transition between shallow and “mesophotic associated” fish communities. Most mesophotic benthic studies start at greater depths (Mean start depth = 40.5 ± 4.6 m) covering the entire mesophotic range (Mean end depth = 218.3 ± 37.6 m). Benthic communities in the Atlantic transition to more deep-specialized communities at 60 m. This does not hold globally, with this change occurring at greater depths in the Pacific and Australia.

We have reviewed studies looking at genetic differences between corals and their associated *Symbiodinium* to describe connectivity patterns between MCEs and shallow reefs, and these studies showed distinct differences with depth (58% of records), between and within genera (Table 6; Figure 6). Most genetic differences appear to occur below 30 m potentially implying shallow and deep populations. Six genera (*Acropora, Eusmilia, Helioseris, Meandrina, Montipora,* and *Mycetophyllia*) showed no genetic differences across depth, though most were only sampled in a single study (except n = 2 for *Helioseris*).

**Discussion**

Studies of Mesophotic coral ecosystems (MCEs) are currently highly location and region specific and not represented in all oceans globally. While this is also the case with shallow reefs (Fisher et al., 2011) the imbalance is not as great. A result of the strong locational bias is that there is not enough evidence to suggest an understanding of the ecological role of MCEs in a global context. Data collection in these ecosystems is still relatively expensive, as most methods require specialized equipment and training. It seems likely that this is the main reason why mesophotic studies are concentrated in areas where the initial investments have been made and equipment is available to enable specific research groups to explore these ecosystems. Huge regional gaps are apparent, showing that almost no studies have been conducted in the Indian Ocean and South-East Asia regions. This is of particular concern given the known high biodiversity of shallow coral ecosystems in these regions and the threats they face (Burke et al., 2011).

MCE research has been mainly focussed in the exploratory phase, aiming to characterize the communities in different locations. What we know from these descriptive studies is that there is a common depth/light attenuation pattern in MCE benthic community structure indicating that upper mesophotic depths have a dominance, in terms of percentage cover, of phototrophic taxa, predominantly corals, shifting to primarily heterotrophic communities, made up of sponges and octocorals, of the lower mesophotic (Lesset et al., 2009; Bongaerts et al., 2010a; Kahng et al., 2010; Baker et al., 2016). It is also well understood that light, topography, and temperature stand out as three main factors that influence the structure of MCE communities. Light is the major factor, with the deepest zooxanthellate coral records associated with areas known for clear water (Kahng et al., 2010; Baker et al., 2016). Topography is also important, with local bathymetric features, such as slope, influencing benthic community structure (Bridge et al., 2010; Locker et al., 2010; Sherman et al., 2010; Englebert et al., 2017). Temperature is influenced by local upwelling (Bridge et al., 2010) and internal waves (Kahng et al., 2016).
and Kelley, 2007; Kahng et al., 2012). This affects depth limits of organisms (Kahng et al., 2012) particularly at higher latitudes (Grigg, 2005) where corals are already residing close to their physiological limits. As the processes associated with MCEs have become better understood, fewer descriptive studies are being carried out and a greater proportion are focused on understanding ecological processes. Moving forward there is more need for studies to be targeted in locations with varying combinations of these influencing factors, as well as proving these trends hold in currently unstudied regions.

We still know little about the pressures that MCEs face, from both anthropogenic and natural sources. The “deep reef refugia” hypothesis suggests that mesophotic areas are more remote from these threats and may re-seed impacted shallow areas (Bongaerts et al., 2010a; Hinderstein et al., 2010). Locational differences occur for natural impacts, for example coral bleaching and disease are reported mainly in the Caribbean (Garcia-Sais et al., 2007; Nemeth et al., 2008; Smith et al., 2015) whereas storm impacts are common in western Pacific areas (Harmelin-Vivien and Laboute, 1986; Bongaerts et al., 2013b; White et al., 2013). Human impacts are currently poorly documented and although localized studies are occurring (Appeldoorn et al., 2015), not enough evidence is available to discuss global or regional trends and threats. Additionally, recovery rates appear to be largely unknown. This is inevitable given that current impacts on MCEs are likely unnoticed or unquantified. This kind of longitudinal information is crucial for effective management of these systems. Additional gaps lie around the direct measurements of life history characteristics and post-settlement processes of benthic organisms at mesophotic depths. Further work into life history dynamics of mesophotic organisms will give an insight into resilience and recovery when faced with disturbances. Conflicting results have been found in terms of fecundity and spawning synchrony of mesophotic coral colonies (Holstein et al., 2015; Prasetia et al., 2016) that also vary between species and locations (Eyal-Shaham et al., 2016). These variations highlight that we know little regarding this subject, which is a concern for managers.

Technological advances have made a range of techniques available for studying MCEs; however, the cost of many of these techniques impacts on the extent to which they are used for data collection. Technical SCUBA diving, despite the training and equipment required, tends to be a cheaper option hence its popularity. The advantage of diving is that it permits investigation of organisms in situ, allowing easier species identification and more precise sample collection. ROVs are commonly utilized for sample collection in the more inaccessible Cold-Water Coral (CWC)
ecosystems where the simultaneous use of video allows increased sampling precision and minimal damage (Fosseå et al., 2005). There is a lack of precise benthic species diversity data, in particular for difficult to identify macroalgae and sponges, so it is important that this taxonomic detail is not lost, and rather targeted to assess specific community structures. However, diving only allows small areas to be surveyed, which may not meet management goals, and additional health and safety concerns associated with technical diving may mean remote methods are more appropriate in some areas.

Acoustic methods cover large areas and allow geophysical variables to be measured at fine scales, where reef corals show distinct bathymetric signatures (Brown et al., 2011). Acoustic data have proven to be highly successful for identifying the extent of CWC ecosystems (Fosseå et al., 2005; Roberts et al., 2009; Buhl-Mortensen et al., 2015). Estimates of total habitat area can be deduced and the information can be used for habitat suitability modelling to identify areas of likely occurrence, which has performed well when applied to mesophotic habitats, given the knowledge of the key structuring variables (Bridge et al., 2012; Costa et al., 2015). Detailed bathymetric information allows for planning of future surveys and can assist with ROV navigation, particularly in areas of high rugosity (Fosseå et al., 2005).

ROVs are often used to provide qualitative visual information to explore new areas (Kahng and Kelley, 2007; Bongaerts et al., 2011a; Blythe-Skyrme et al., 2013; Englebert et al., 2014), commonly prior to committing divers or to survey depths >150 m. Samples collected from ROVs have allowed the detailed taxonomy of mesophotic corals (Muir et al., 2015) as well as further lab experiments (van Oppen et al., 2011). CWC ecosystems have utilized ROVs to deploy additional equipment and set up in situ experiments (Roberts et al., 2009) which are an approach that should be considered for MCEs. ROVs, while unable to collect samples, can offer a more quantitative approach to obtaining imagery. Hundreds of thousands of accurately georeferenced images may be collected, as well as accompanying environmental information (Williams et al., 2012; Pizarro et al., 2013), while also having the advantage of running independently to the deployment vessel. AUVs, while unable to collect samples, can offer a more quantitative approach to obtaining imagery. Hundreds of thousands of accurately georeferenced images may be collected, as well as accompanying environmental information (Williams et al., 2012; Pizarro et al., 2013), while also having the advantage of running independently to the deployment vessel. AUVs have the capability to accurately perform repeat monitoring surveys and relocate colonies (Pizarro et al., 2013; Ferrari et al., 2016) which enable an insight into processes such as growth rates in the future. In terms of costs per area surveyed remote methods may be cheaper, although their ability to fill data gaps surrounding life history traits may be limited; however they may be complemented by diving surveys and experiments.

High biodiversity is common across MCEs of all regions (Baker et al., 2016) although there is still further biodiversity to be discovered as mentioned above. High taxonomic resolution is required to assess connectivity, as will be discussed, where species-specific differences are observed. Depths at which communities change appear to be area-specific and evidence appears to suggest that using depth alone as a basis for universal definitions may not be appropriate. The depth of the transition zone, representing a shift in upper and lower mesophotic assemblages also varies between locations. While 60 m is commonly reported (Fricke and Meischner, 1985; Liddell and Ohlhorst, 1988; Bongaerts et al., 2010a; Bridge et al., 2010; Slattery et al., 2011) on average this only applies to the Atlantic region. In the Pacific and the Coral Sea, the transition zone depth extends past 80 m (Kahng and Kelley, 2007; Pyle et al., 2016; Englebert et al., 2017). Equally, mesophotic depths are shallower for locations with lower light regimes, such as Ningaloo, Australia (Rees et al., 2004) or

![Figure 4. Research focus of mesophotic coral ecosystem studies over time.](https://academic.oup.com/icesjms/article-abstract/74/9/2309/3855117/2314)
Pohnpei, Micronesia (Muir and Wallace, 2016) and reduced temperature, such as Bermuda (Fricke and Meischner, 1985). This raises questions about the ecological relevance (Laverick et al., 2015; Andradi-Brown et al., 2014; Lindfield et al., 2016a) so as to allow comparisons to be made. High proportions of fish species are common to both shallow and lower mesophotic areas across regions (Bejarano et al., 2014; Wagner et al., 2014; Lindfield et al., 2015), while genetic similarities are also described (Tenggardjaja et al., 2014). Ontogenic movements are also reported (Brokovich et al., 2006; Rosa et al., 2015; Andradi-Brown et al., 2016b) suggesting movement is common across depths. Given these findings, reported transition depths seem to represent the change from shallow water to mesophotic associated fish communities, in contrast to benthic communities where transition depths represent the change from upper to lower mesophotic; having already seen a shift from shallow waters. While changes in the benthic composition are likely to affect distributions of fish species (García-Sais et al., 2007; Brokovich et al., 2008; García-Sais, 2010): corals may decrease but sponges and other benthic organisms can provide structural habitat (Bell et al., 2013) at depth. Other factors may be structuring fish communities, such as food availability, given the distinct changes in functional groups observed (Bridge et al., 2016b).

Assessing connectivity between shallow and deep reefs is a primary focus in the published literature, and more studies are being undertaken in this research area. Vertical connectivity will ultimately determine whether MCEs can re-seed shallow coral reefs following chronic disturbances. Questions do remain over how genetic changes in Symbiodinium correlate with that of their hosts. However, given that host specificity is common and specific adaptation to environmental conditions are likely to have evolved (LaJeunesse et al., 2004; Frade et al., 2008b; Finney et al., 2010) differences probably indicate genetic separation of shallow and MCEs (Bongaerts et al., 2010b,c). Populations below 30 m are reported as unconnected to shallower conspecifics, with distinct shallow and deep genetic populations found (Brazeau et al., 2013). Deeper coral populations are specialized to lower light conditions, showing changes in morphology (Fricke and Meischner, 1985; Einbinder et al., 2009; Nir et al., 2011), photosynthetic efficiency (Lesser et al., 2010; Mass et al., 2010; Nir et al., 2011; Einbinder et al., 2016), and alternative nutrient sources (Muscatine et al., 1989; Einbinder et al., 2009; Crandall et al., 2016). Isolated reefs appear to have higher vertical genetic

### Table 4. Mean species richness at mesophotic depths (>30 m) for each region ± Standard Error.

| Region | Macroalgaes | Scleractinian coral | Fish | Sponge |
|--------|-------------|---------------------|------|--------|
| Atlantic | 90.3 (± 25.9, n = 4) | 16.3 (± 1.7, n = 18) | 77.3 (± 9.2, n = 16) | 79.6 (± 28.8, n = 9) |
| Australia | 62.0 (± 3.0, n = 3) | 32.0 (± 20.3, n = 3) | 139 (n = 1) | 240.4 (± 37.7, n = 5) |
| Middle East | 48.0 (± 20.3, n = 3) | 27.6 (± 6.6, n = 5) | 132.7 (± 6.6, n = 7) | |

Blanks show no data available for that region.

### Table 5. Mean transition depth between benthic and fish communities for each region ± Standard Error.

| Region | Benthic | Fish |
|--------|--------|------|
| Atlantic | 60.9 m (± 4.68, n = 12) | 63.1 m (± 4.9, n = 8) |
| Australia | 75.3 m (± 3.4, n = 15) | 49 m (n = 1) |
| Middle East | 50.0 m (± 0.0, n = 2) | |
| Pacific | 74.7 m (± 3.6, n = 7) | 61.3 m (± 16.1, n = 9) |

Blanks show no data available for that region.
**Figure 6.** Depths at which genetic changes in corals and their *Symbiodinium* occur.

**Table 6.** Numbers of records of whether genetic differences in corals of their *Symbiodinium* occur (Yes) or do not occur (No) with depth.

| Genera        | Species                | No | Yes | Depth range investigated                              | References                                                                 |
|---------------|------------------------|----|-----|-------------------------------------------------------|---------------------------------------------------------------------------|
| Acropora      | Acropora elegans       | 1  | 1   | 10–48 m                                              | (Bongaerts et al., 2011c)                                                 |
| Agaricia      | Agaricia agaricites     | 2  | 1   | 5–50 m                                               | (Bongaerts et al., 2013a, 2015a)                                          |
|               | Agaricia fragilis       | 1  | 1   | 12–40 m                                              | (Bongaerts et al., 2017)                                                  |
|               | Agaricia grahamae       | 1  | 1   | 15–90 m                                              | (Bongaerts et al., 2013a, 2015b)                                          |
|               | Agaricia lamarcki       | 3  | 1   | 10–70 m                                              | (Bongaerts et al., 2013a, 2015a; Lucas et al., 2016)                     |
|               | Agaricia undata         | 1  | 1   | 15–90 m                                              | (Bongaerts et al., 2015b)                                                |
| Echinophyllia | Echinophyllia aspera    | 1  | 1   | 10–62 m                                              | (Bongaerts et al., 2011c)                                                 |
| Eusmilia      | Eusmilia fastigiata     | 1  | 1   | 5–40 m                                               | (Bongaerts et al., 2015a)                                                 |
| Galaxea       | Galaxea astreata        | 1  | 1   | 10–55 m                                              | (Bongaerts et al., 2011c)                                                 |
| Helioseris    | Helioseris cucullata    | 2  | 1   | 25–45 m                                              | (Bongaerts et al., 2013a, 2015a)                                          |
| Leptoseris    | Leptoseris hawaiensis   | 1  | 1   | 10–70 m                                              | (Bongaerts et al., 2011c)                                                 |
|               | Leptoseris spp.         | 1  | 3   | 1–127 m                                              | (Chan et al., 2009; Luck et al., 2013; Pochon et al., 2015; Ziegler et al., 2015) |
| Madracis      | Madracis carmabi        | 1  | 1   | 5–40 m                                               | (Frade et al., 2008b)                                                    |
|               | Madracis decatis        | 2  | 1   | 5–40 m                                               | (Frade et al., 2008b; Bongaerts et al., 2015a)                           |
|               | Madracis formosa        | 2  | 1   | 5–60 m                                               | (Frade et al., 2008a,b; Bongaerts et al., 2015a)                         |
|               | Madracis mirabilis      | 1  | 1   | 5–40 m                                               | (Bongaerts et al., 2015a)                                                |
|               | Madracis pharensis      | 5  | 1   | 5–90 m                                               | (Frade et al., 2008a,b; Bongaerts et al., 2015a,b)                       |
|               | Madracis senaria        | 2  | 1   | 5–40 m                                               | (Frade et al., 2008a,b)                                                  |
| Meandrina     | Meandrina meandrites    | 1  | 1   | 5–40 m                                               | (Bongaerts et al., 2015a)                                                |
| Montastrea    | Montastrea cavernosa    | 3  | 3   | 3–91 m                                               | (Lesser et al., 2010; Brazeau et al., 2013; Bongaerts et al., 2015a)     |
| Montipora     | Montipora spp.          | 1  | 1   | 10–70 m                                              | (Bongaerts et al., 2011c)                                                 |
| Mycetophyllia | Mycetophyllia ferox     | 1  | 1   | 25–40 m                                              | (Bongaerts et al., 2015a)                                                |
| Orbicella     | Orbicella faveolata     | 1  | 1   | 5–25 m                                               | (Bongaerts et al., 2015a)                                                |
|               | Orbicella franksi       | 1  | 1   | 10–25 m                                              | (Bongaerts et al., 2015a)                                                |
| Pachyseris    | Pachyseris speciosa     | 1  | 2   | 1–62 m                                               | (Bongaerts et al., 2011c; Cooper et al., 2011; Ziegler et al., 2015)     |
| Pavona        | Pavona spp.             | 1  | 1   | 10–59 m                                              | (Bongaerts et al., 2011c)                                                |
| Porites       | Porites astrooides      | 2  | 3   | 2–30 m                                               | (Bongaerts et al., 2015a; Serrano et al., 2016; Reich et al., 2017)      |
|               | Porites spp.            | 1  | 1   | 1–70 m                                               | (Bongaerts et al., 2011c; Ziegler et al., 2015)                          |
| Seriatopora   | Seriatopora hystrix     | 3  | 4   | 2–57 m                                               | (Bongaerts et al., 2010b, 2011b,c; Cooper et al., 2011; Nir et al., 2011; van Oppen et al., 2011) |
| Siderastrea   | Siderastrea siderea     | 1  | 1   | 2–50 m                                               | (Bongaerts et al., 2015a)                                                |
| Stephanocoenia| Stephanocoenia intersepta| 1  | 1   | 10–60 m                                              | (Bongaerts et al., 2015a, 2017)                                           |
connectivity, possibly due to the importance of localized recruitment for sustaining populations (Serrano et al., 2016) or reduced competition following disturbance which may prevent localized extinctions (van Oppen et al., 2011; Sinninge et al., 2012; Muir et al., 2015). However, using Bermuda as an example, not all species show this pattern of strong vertical connectivity (Bongaerts et al., 2017). The reproductive mode may give some insight, with broadcast spawning genera generally showing reduced genetic partitioning with depth (Bongaerts et al., 2011c, 2017) although this is not exclusive (Bongaerts et al., 2015a). Local environmental conditions also play a role, and light levels will ultimately influence the upper and lower limits of coral species and their Symbiodinium types due to functional adaptations (Frade et al., 2008a,c). This again calls into question the use of only depth to define deep and shallow MCEs. Overall, the findings in this review show that differences in vertical connectivity patterns at species and genera level are common across MCEs globally. These results highlight our limited knowledge, and the need for these studies to be done at both localized scales, for a detailed analysis of local populations, and across biogeographic ranges.

Conclusions
The importance of mesophotic areas is now recognized in the scientific community. There is a clear locational bias of the existing research to the Atlantic, and specifically the Caribbean, which makes the extrapolation of findings to the rest of the world difficult. Definitions coined from data in this region alone need to be redefined as more studies are completed globally. A clear priority is to collect data for MCEs in South East Asia and the Indian Ocean. Remote methods are clearly advancing research in this field, though it is important to not lose taxonomic detail, given the apparent species and location specificity of connectivity patterns. If shallow and deep populations are separate, then management plans need to accommodate this in order to conserve the different biodiversity of both of these light-mediated ecosystems. The current lack of information about the threats and impacts on MCEs needs to be addressed immediately so that they can be identified at local, regional, and global scales so that effective management can be implemented. Further prioritization of such studies, as well as those investigating connectivity at both local and regional scales, is clearly required, to ensure adequate protection of these ecosystems and their shallow water counterparts, for which relying on MCEs as refugia may not be appropriate.

Acknowledgements
We thank our funding agency BHP Billiton-CSIRO Ningaloo Outlook Marine Research Partnership for support of this work. The views expressed herein are those of the authors and do not necessarily reflect the views of BHP Billiton or CSIRO. We would also like to thank Tom Bridge and Tim Cooper for their comments on the manuscript prior to submission.

References
Andradi-Brown, D., East, A., Shepherd, L., Stockdale, E., and Rogers, A. 2016a. Challenges and opportunities in mesophotic reef research. Reef Encounter: The News Journal of the International Society for Reef Studies, 31: 26–31.
Andradi-Brown, D., Gress, E., Wright, G., Exton, D., and Rogers, A. 2016b. Reef fish community biomass and trophic structure changes across shallow to upper-mesophotic reefs in the Mesoamerican barrier reef, Caribbean. PLoS One, 11:e0156641.
Appeldoorn, R., Ballantine, D., Bejarano, I., Carlo, M., Nemeth, M., Otero, E., Pagan, F., et al. 2015. Mesophotic coral ecosystems under anthropogenic stress: a case study at Ponce, Puerto Rico. Coral Reefs, 35: 63–75.
Armstrong, R. A., Singh, H., Torres, J., Nemeth, R. S., Can, A., Roman, C., Eustice, R., et al. 2006. Characterising the deep insular shelf coral reef habitat of the Hind Bank Marine Conservation District (US Virgin Islands) using the Seabed Autonomous Underwater Vehicle. Continental Shelf Research, 26: 194–205.
Bak, R. P. M., Nieuwland, G., and Meesters, E. H. 2005. Coral reef crisis in deep and shallow reefs: 30-years of constancy and change in reefs of Curacao and Bonaire. Coral Reefs, 24: 475–479.
Baker, E. K., Puglise, K. A., and Harris, P. T. 2016. Mesophotic Coral Ecosystems—A Lifeboat for Coral Reefs? The United Nations Environment Programme and GRID-Arendal, Nairobi and Arendal.
Bejarano, I., Appeldoorn, R. S., and Nemeth, M. 2014. Fishes associated with mesophotic coral ecosystems in La Parguera, Puerto Rico. Coral Reefs, 33: 313–328.
Bell, J. J., Davy, S. K., Jones, T., Taylor, M. W., and Webster, N. S. 2013. Could some coral reefs become sponge reefs as our climate changes? Global Change Biology, 19: 2613–2624.
Bellwood, D. R., Hughes, T. P., Folke, C., and Nystrom, M. 2004. Confronting the coral reef crisis. Nature, 429: 827–833.
Blaythe-Skyrme, V. J., Rooney, J., Parrish, F., and Boland, R. 2013. Mesophotic Coral Ecosystems—Potential Candidates as Essential Fish Habitat and Habitat Areas of Particular Concern. Pacific Island Fisheries Science Center, National Marine Fisheries Science Center Administrative Reports, H-13-02: 53 p.
Bongaerts, P., Bridge, T. C. L., Kline, D. I., Muir, P. R., Wallace, C. C., Beaman, R. L., and Hoegh-Guldberg, O. 2011a. Mesophotic coral ecosystems on the walls of Coral Sea atolls. Coral Reefs, 30: 335–335.
Bongaerts, P., Carmichael, M., Hay, K. B., Tonk, L., Frade, P. R., and Hoegh-Guldberg, O. 2015a. Prevalent endosymbiont zonation shapes the depth distributions of scleractinian coral species. Royal Society Open Science, 2: 140297.
Bongaerts, P., Frade, P. R., Hay, K. B., Englebert, N., Latijnhouwers, K. R., Bak, R. P., Vermeij, M. J., et al. 2015b. Deep down on a Caribbean reef: lower mesophotic depths harbor a specialized coral-endosymbiont community. Scientific Reports, 5: 7652.
Bongaerts, P., Frade, P. R., Ogier, J. J., Hay, K. B., van Beijswijk, J., Englebert, N., Vermeij, M. J., et al. 2013a. Sharing the slope: depth partitioning of agariciid corals and associated Symbiodinium across shallo and mesophotic habitats (2-60 m) on a Caribbean reef. BMC Evolutionary Biology, 13: 205.
Bongaerts, P., Muir, P., Englebert, N., Bridge, T. C. L., and Hoegh-Guldberg, O. 2013b. Cyclone damage at mesophotic depths on Myrmaidon Reef (GBR). Coral Reefs, 32: 935–935.
Bongaerts, P., Ridway, T., Sampayo, E. M., and Hoegh-Guldberg, O. 2010a. Assessing the ‘deep reef refugia’ hypothesis: focus on Caribbean reefs. Coral Reefs, 29: 309–327.
Bongaerts, P., Riginos, C., Brunner, R., Englebert, N., Smith, S. R., and Hoegh-Guldberg, O. 2017. Deep reefs are not universal refuges: Reseeding potential varies among coral species. Science Advances, 3: e1602373.
Bongaerts, P., Riginos, C., Hay, K. B., van Oppen, M. J., Hoegh-Guldberg, O., and Dove, S. 2011b. Adaptive divergence in a scleractinian coral: physiological adaptation of Seriatopora hystrix to shallow and deep reef habitats. BMC Evolutionary Biology, 11: 303.
Bongaerts, P., Riginos, C., Ridgway, T., Sampayo, E. M., van Oppen, M. J., Englebert, N., Vermeulen, F., et al. 2010b. Genetic divergence across habitats in the widespread coral Seriatopora hystrix and its associated Symbiodinium. PLoS One 5: e10871.
Bongaerts, P., Sampayo, E. M., Bridge, T. C. L., Ridgway, T., Vermeulen, F., Englebert, N., Webster, J. M., et al. 2011c.
Symbiodinium diversity in mesophotic coral communities on the Great Barrier Reef: a first assessment. Marine Ecology Progress Series, 439: 117–126.

Bouchon, C. 1981. Quantitative study of the Scleractinia coralline communities of a fringing reef of Reunion Island (Indian Ocean). Marine Ecology Progress Series, 4: 273–288.

Brazeau, D. A., Lesser, M. P., and Slattery, M. 2013. Genetic structure in the coral, Montastraea cavernosa: assessing genetic differentiation among and within Mesophotic reefs. PLoS One, 8: e65845.

Bridge, T., Beaman, R., Done, T., and Webster, J. 2012. Predicting the location and spatial extent of submerged coral reef habitat in the Great Barrier Reef world heritage area, Australia. PLoS One, 7: e48203.

Bridge, T. C., Grech, A. M., and Pressey, R. L. 2016a. Factors influencing incidental representation of previously unknown conservation features in marine protected areas. Conservation Biology, 30: 154–165.

Bridge, T. C., Luiz, O. J., Coleman, R. R., Kane, C. N., and Kosaki, R. K. 2016b. Ecological and morphological traits predict depth-generalistfishes on coral reefs. Proceedings of the Royal Society of London B: Biological Sciences, 283: 20152332.

Bridge, T. C. L., Done, T. J., Beaman, R. J., Friedman, A., Williams, S. B., Pizarro, O., and Webster, J. M. 2010. Topography, substratum and benthic macrofaunal relationships on a tropical mesophotic shelf margin, central Great Barrier Reef, Australia. Coral Reefs, 30: 143–153.

Bridge, T. C. L., Done, T. J., Friedman, A., Beaman, R. J., Williams, S. B., Pizarro, O., and Webster, J. M. 2011a. Variability in mesophotic coral reef communities along the Great Barrier Reef, Australia. Marine Ecology Progress Series, 428: 63–75.

Bridge, T. C. L., Fabricius, K. E., Brogaerts, P., Wallace, C. C., Muir, P. R., Done, T. J., and Webster, J. M. 2011b. Diversity of Scleractinia and Octocorallia in the mesopotic zone of the Great Barrier Reef, Australia. Coral Reefs, 31: 179–189.

Bridge, T. C. L., Hughes, T. P., Guinotte, J. M., and Bongaerts, P. 2013. Call to protect all coral reefs. Nature Climate Change, 3: 528–530.

Brokovich, E., Einbinder, S., Kark, S., Shashar, N., and Kiflawi, M. 2006. A deep nursery for juveniles of the zebra angelfish Gени cantidad caudovittatus. Environmental Biology of Fishes, 80: 1–6.

Brokovich, E., Einbinder, S., Shashar, N., Kiflawi, M., and Kark, S. 2008. Descending to the twilight-zone: changes in coral reef fish assemblages along a depth gradient down to 65 m. Marine Ecology Progress Series, 371: 253–262.

Brown, C. J., Smith, S. J., Lawton, P., and Anderson, J. T. 2011. Benthic habitat mapping: a review of progress towards improved understanding of the spatial ecology of the seafloor using acoustic techniques. Estuarine, Coastal and Shelf Science, 99: 502–520.

Buhl-Mortensen, L., Buhl-Mortensen, P., Dolan, M. J. F., and Gonzalez-Mireles, G. 2015. Habitat mapping as a tool for conservation and sustainable use of marine resources: some perspectives from the MAREANO Programme, Norway. Journal of Sea Research, 100: 46–61.

Burke, L., Reyta, K., Spalding, M., and Perry, A. 2011. Reefs at Risk Revisited. World Resources Institute, Washington, DC.

Busby, R. 1966. Ocean Bottom Reconnaissance Off the East Coast of Andros Island, Bahamas. US Navel Oceanographic Office, Technical Report 20390, Washington DC.

Chan, Y. L., Pochon, X., Fisher, M. A., Wagner, D., Concepcion, G. T., Kahng, S. E., Toonen, R. J., et al. 2009. Generalist dinoflagellate endosymbionts and host genotype diversity detected from mesophotic (67-100 m depth) coral Leptoseris. BMC Ecology, 9: 21.

Colin, P. L., Devaney, M., Hills-Colinvaux, L., Suchanek, T. H., and Harrison, J. T. 1986. Geology and biological zonation of the reef slope, 50-360 m depth, at Enewetak Atoll, Marshall Islands. Bulletin of Marine Science, 38: 111–128.

Cooper, T. F., Ulstrup, K. E., Dandan, S. S., Heyward, A. J., Kuhl, M., Muirhead, A., O’Leary, R. A., et al. 2011. Niche specialization of reef-building corals in the mesophotic zone: metabolic trade-offs between divergent Symbiodinium types. Proceedings of the Royal Society of London B: Biological Sciences, 278: 1840–1850.

Costa, B., Kendall, M. S., Parrish, F. A., Rooney, J., Boland, R. C., Chow, M., Lecky, J., et al. 2015. Identifying suitable locations for mesophotic hard corals offshore of Maui, Hawai’i. PLoS One, 10: e0130285.

Crandall, J. B., Tece, M. A., Estes, B. A., Manfrino, C., and Ciesla, J. H. 2016. Nutrient acquisition strategies in mesophotic hard corals using compound specific stable isotope analysis of sterols. Journal of Experimental Marine Biology and Ecology, 474: 133–141.

Einbinder, S., Gruber, D. F., Salomon, E., Liran, O., Keren, N., and Tchernov, D. 2016. Novel adaptive photosynthetic characteristics of mesophotic symbiotic microalgae within the reef-building coral, Stylophora pistillata. Frontiers in Marine Science, 3(Article 175): 1–9.

Einbinder, S., Mass, T., Brokovich, E., Dubinsky, Z., Erez, J., and Tchernov, D. 2009. Changes in morphology and diet of the coral Stylophora pistillata along a depth gradient. Marine Ecology Progress Series, 381: 167–174.

Englebert, N., Bongaerts, P., Muir, P., Hay, K. B., and Hoegh-Guldberg, O. 2014. Deepest zooxanthellate corals of the Great Barrier Reef and Coral Sea. Marine Biodiversity, 45: 1–2.

Englebert, N., Bongaerts, P., Muir, P. R., Hay, K. B., Pichon, M., and Hoegh-Guldberg, O. 2017. Lower mesophotic coral communities (60-125 m depth) of the northern Great Barrier Reef and Coral Sea. PLoS One, 12: e0170336.

Eyal-Shaham, L., Eyal, G., Tamir, R., and Loya, Y. 2016. Reproduction, abundance and survivorship of two Alveopora spp. in the mesophotic reefs of Eilat, Red Sea. Scientific Reports, 6: 20964.

Eyal, G., Wiedenmann, J., Grinblat, M., D’Angelo, C., Kramarsky-Winter, E., Treibitz, T., Ben-Zvi, O., et al. 2015. Spectral diversity and regulation of coral fluorescence in a mesophotic reef habitat in the Red Sea. PLoS One, 10: e0128697.

Ferrari, R., Bryson, M., Bridge, T., Hustache, J., Williams, S. B., Byrne, M., and Figueira, W. 2016. Quantifying the response of structural complexity and community composition to environmental change in marine communities. Global Change Biology, 22: 1965–1975.

Finney, J. C., Pettay, D. T., Sampayo, E. M., Warner, M. E., Oxenford, H. A., and Lajuneesse, T. C. 2010. The relative significance of host–habitat, depth, and geography on the ecology, endemism, and speciation of coral endosymbionts in the genus Symbiodinium. Microbial Ecology, 60: 250–263.

Fisher, R., Radford, B. T., Knowlton, N., Brainard, R. E., Michaelis, F. B., and Caley, M. J. 2011. Global mismatch between research effort and conservation needs of tropical coral reefs. Conservation Letters, 4: 64–72.

Fossa, J. H., Lindberg, B., Christensen, O., Lundåv, T., Svinglen, I., Mortensen, P. B., and Alsvåg, J. 2005. Mapping of Lophelia reefs in Norway: experiences and survey methods. In Cold-Water Corals and Ecosystems, pp. 359–391. Ed. by A. Freiwald, and J. M. Roberts. Springer Berlin Heidelberg, Berlin, Heidelberg.

Frade, P. R., Bongaerts, P., Winkelhagen, A. J. S., Tonk, L., and Bak, R. P. M. 2008a. In situ photobiology of corals over large depth ranges: a multivariate analysis on the roles of environment, host, and algal symbiont. Limnology and Oceanography, 53: 2711–2723.

Frade, P. R., De Jongh, F., Vermeulen, F., Van Bleijswijk, J., and Bak, R. P. M. 2008b. Variation in symbiont distribution between closely related coral species over large depth ranges. Molecular Ecology, 17: 691–703.
Kahng, S. E., Englebert, N., Faria, J., Visser, P. M., and Bak, R. P. M. 2008c. Distribution and photobiology of Symbiodinium types in different light environments for three colour morphs of the coral *Madracis pharensis*: is there more to it than total irradiance? Coral Reefs, 27: 913–925.

Freiwald, A., Fossa, J. H., Grehan, A., Koslow, T., and Roberts, J. M. 2004. Cold-Water Coral Reefs. UNEP-WCMC, Cambridge, UK.

Freiwald, A., and Roberts, J. M. 2005. Cold-Water Corals and Ecosystems: Preface. Springer, Heidelberg, VII–XII.

Fricke, H. W., and Knauer, B. 1986. Diversity and spatial pattern of coral communities in the Red Sea upper twilight zone. Oecologia, 71: 29–37.

Fricke, H. W., and Meischner, D. 1985. Depth limits of Bermudan scleractinian corals: a submersible survey. Marine Biology, 88: 175–187.

Garcia-Sais, J. R. 2010. Reef habitats and associated sessile-benthic and fish assemblages across a euphotic–mesophotic depth gradient in Isla Desecheo, Puerto Rico. Coral Reefs, 29: 277–288.

Garcia-Sais, J. R., Castro, R., Sabater-Clavel, J., Carlo, M., and Esteves, R. 2007. Characterization of benthic habitats and associated reef communities at Bajo de Sico Seamount, Mona Passage, Puerto Rico. Report to the Caribbean Fishery Management Council.

Goreau, T. F., and Goreau, N. I. 1973. The ecology of Jamaican coral reefs. II. Geomorphology, zonation, and sedimentary phases. Bulletin of Marine Science, 23: 400–464.

Grigg, R. W. 2005. Depth limit for reef building corals in the Au’au Channel, S.E. Hawaii. Coral Reefs, 25: 77–84.

Harmelin-Vivien, M. L., and Laboute, P. 1986. Catastrophic impact of hurricanes on atoll outer reef slopes in the Tuamotu (French Polynesia). Coral Reefs, 5: 55–62.

Harris, P. T., Bridge, T. C. L., Beaman, R. J., Webster, J. M., Nichol, S. L., and Brooke, B. P. 2012. Submerged banks in the Great Barrier Reef, Australia, greatly increase available coral reef habitat. ICES Journal of Marine Science, 70: 284–293.

Heyward, A., Fromont, J., Schönberg, C. H. L., Colquhoun, J., Radford, B., and Gomez, O. 2010. The sponge gardens of Ningaloo Reef, Western Australia. The Open Marine Biology Journal, 4: 3–11.

Hinderstein, L. M., Marr, J. C. A., Martinez, F. A., Dowgiallo, M. J., Puglise, K. A., Pyle, R. L., Zawada, D. G., et al. 2010. Theme section on 'Mesophotic Coral Ecosystems: Characterization, Ecology, and Management'. Coral Reefs, 29: 247–251.

Hoech-Guldberg, O., Mumby, P. J., Hooten, A. J., Steneck, R. S., Greenfield, P., Gomez, E., Harvell, C. D., et al. 2007. Coral reefs under rapid climate change and ocean acidification. Science, 318: 1737–1742.

Holstein, D. M., Smith, T. B., Gyory, J., and Paris, C. B. 2015. Fertile fathoms: deep reproductive refugia for threatened shallow corals. Scientific Reports, 5: 12407.

Hughes, T. P., Baird, A. H., Bellwood, D. R., Card, M., Connolly, S. R., Folke, C., Grosberg, R., et al. 2003. Climate change, human impacts, and the resilience of coral reefs. Science, 301: 929–933.

Kahng, S. E., Copus, J. M., and Wagner, D. 2014. Recent advances in the ecology of mesophotic coral ecosystems (MCEs). Current Opinion in Environmental Sustainability, 7: 72–81.

Kahng, S. E., Garcia-Sais, J. R., Spalding, H. L., Brokovich, E., Wagner, D., Weil, E., Hinderstein, L., et al. 2010. Community ecology of mesophotic coral reef ecosystems. Coral Reefs, 29: 255–273.

Kahng, S. E., and Kelley, C. D. 2007. Vertical zonation of megabenthic taxa on a deep photosynthetic reef (50–140 m) in the Au’au Channel, Hawaii. Coral Reefs, 26: 679–687.

Kahng, S. E., Wagner, D., Lantz, C., Vetere, O., Gove, J., and Merrifield, M. 2012. Temperature related depth limits of warm-water corals. Proceedings of the 12th International Coral Reef Symposium, Cairns, Australia.

Kane, C., Kosaki, R. K., and Wagner, D. 2014. High levels of mesophotic reef fish endemism in the Northwestern Hawaiian Islands. Bulletin of Marine Science, 90: 693–703.

Lajeunesse, T. C., Thornhill, D. J., Cox, E. F., Stanton, F. G., Fitt, W. K., and Schmidt, G. W. 2004. High diversity and host specificity observed among symbiotic dinoflagellates in reef coral communities from Hawaii. Coral Reefs, 23: 596–603.

Laverick, J. H., Andradi-Brown, D. A., Exton, D. A., Bongaerts, P., Bridge, T. C. L., Lesser, M. P., Pyle, R. L., et al. 2016. To what extent do mesophotic coral ecosystems and shallow reefs share species of conservation interest? Environmental Evidence, 5: 16.

Lesser, M. P., Slattery, M., and Leichter, J. J. 2009. Ecology of mesophotic coral reefs. Journal of Experimental Marine Biology and Ecology, 375: 1–8.

Lesser, M. P., Slattery, M., Stat, M., Ojimi, M., Gates, R. D., and Grottoli, A. 2010. Photoacclimatization by the coral *Montastraea cavernosa* in the mesophotic zone: light, food, and genetics. Ecology, 91: 990–1003.

Liddell, D., and Ohllhorst, S. L. 1988. Hard Substrata Community Patterns, 1-120 M, North Jamaica. Palaos, 3: 413–423.

Lindfield, S. J., Harvey, E. S., Halford, A. R., and McIlwain, J. L. 2015. Mesophotic depths as refuge areas for fishery-targeted species on coral reefs. Coral Reefs, 35: 125–137.

Locke, S. D., Armstrong, R. A., Battista, T. A., Rooney, J. J., Sherman, C., and Zawada, D. G. 2010. Geomorphology of mesophotic coral ecosystems: current perspectives on morphology, distribution, and mapping strategies. Coral Reefs, 29: 329–343.

Lombardi, M., and Godfrey, J. 2011. In-Water Strategies for Scientific Diver-Based Examinations of the Vertical Mesophotic Coral Ecosystem (vMCE) from 50 to 150 meters. In Diving for Science 2011. Proceedings of the American Academy of Underwater Sciences 30th Symposium. Ed by. Pollock, N.W. AAUS, Dauphin Island, AL.

Loya, Y., Eyal, G., Treibitz, T., Lesser, M. P., and Appeldoorn, R. 2016. Theme section on mesophotic coral ecosystems: advances in knowledge and future perspectives. Coral Reefs, 35: 1–9.

Lucas, M. O., Stat, M., Smith, M. C., Weil, E., and Schizas, N. V. 2016. Symbiodinium (internal transcribed spacer 2) diversity in the coral host Agaricia lamarcki (Cnidaria: Scleractinia) between shallow and mesophotic reefs in the Northern Caribbean (20–70 m). Marine Ecology, 37: 1079–1087.

Luck, D. G., Forsman, Z. H., Toonen, R. J., Leicht, S. J., and Kahng, S. E. 2013. Polyphyly and hidden species among Hawai’i’s dominant mesophotic coral genera, Leptoseris and Pavona (Scleractinia: Agaricidae). Peer Journal, 1: e132.

Mass, T., Kline, D. I., Roopin, M., Veal, C. J., Cohen, S., Iluz, D., and Levy, O. 2010. The spectral quality of light is a key driver of photosynthesis and photoacclimation in Stylphora pistillata colonies from different depths in the Red Sea. The Journal of Experimental Biology, 213: 4084–4091.

Muir, P., Wallace, C., Bridge, T. C., and Bongaerts, P. 2015. Diverse staghorn coral fauna on the mesophotic reefs of north-east Australia. PLoS One, 10: e0117933.

Muir, P. R., and Wallace, C. C. 2016. A rare ‘deep-water’ coral assemblage in a shallow lagoon in Micronesia. Marine Biodiversity, 46: 543–544.

Muscatine, L., Porter, J. W., and Kaplan, I. R. 1989. Resource partitioning by reef corals as determined from stable isotope composition. Marine Biology, 100: 185–193.

Nemeth, M., Smith, T. B., Blondeau, J., Kadison, E., Calnan, J. M., and Gass, J. 2008. Characterization of deep water reef communities within the marine conservation district, St. Thomas, US Virgin Islands. Report to the Caribbean Fisheries Management Council.

Nir, O., Gruber, D. F., Einbinder, S., Kark, S., and Tchernov, D. 2011. Changes in scleractinian coral Seriatopora hystrix morphology and its endocellular Symbiodinium characteristics along a...
bathymetric gradient from shallow to mesophotic reef. Coral Reefs, 30: 1089–1100.

Pickering, C., and Byrne, J. 2014. The benefits of publishing systematic quantitative literature reviews for PhD candidates and other early-career researchers. Higher Education Research & Development, 33: 534–548.

Pickering, C., Grignon, J., Steven, R., Guitart, D., and Byrne, J. 2015. Publishing not perishing: how research students transition from novice to knowledgeable using systematic quantitative literature reviews. Studies in Higher Education, 40: 1756–1769.

Pizarro, O., Williams, S. B., Jakuba, M. V., Johnson-Roberson, M., Mahon, I., Beyson, M., Steinberg, D., et al. 2013. Benthic monitoring with robotic platforms—the experience of Australia. Underwater Technology Symposium (UT), 2013 IEEE International: 1–10.

Pochon, X., Forsman, Z. H., Padilla-Gamino, J. L., Smith, C. M., and Gates, R. D. 2015. Depth specialization in mesophotic corals (Leptoseris spp.) and associated algal symbionts in Hawai`i. Royal Society Open Science, 2: 140351.

Prasetia, R., Sinniger, F., and Harii, S. 2016. Gametogenesis and fecundity of Acropora tenua (Brook 1892) in a mesophotic coral ecosystem in Okinawa, Japan. Coral Reefs, 35: 53–66.

Puglise, K. A., Hinderstein, L. M., Marr, J. C. A., Dowgiliallo, M. J., and Martinez, F. A. 2009. Mesophotic coral ecosystems research strategy: international workshop to prioritize research and management needs for mesophotic coral ecosystems, Jupiter, Florida, 12–15 July 2008. NOAA Technical Memorandum NOS NCCOS 98 and OAR OER 2.

Pyle, R. L., Boland, R., Bolick, B., Bowen, B. W., Bradley, C. J., Kane, C., Kosaki, R. K., et al. 2016. A comprehensive investigation of mesophotic coral ecosystems in the Hawaiian Archipelago. Peer Journal, 4: e2475.

R Core Team. 2010. R: A Language and Environment for Statistical Computing. R Core Team, Vienna, Austria.

Rees, M., Heyward, A., Cappo, M., Speare, P., and Smith, L. 2004. Ningaloo Marine Park—Initial Survey of Seabed Biodiversity in Intermediate and Deeper Waters. Australian Institute of Marine Science, Crawley, Australia.

Reich, H. G., Robertson, D. L., and Goodbody-Gringley, G. 2017. Do the shuffle: changes in Symbiodinium consortia throughout juvenile coral development. PLoS One, 12: e0171768.

Roberts, J. M., Wheeler, A. J., and Freiwald, A. 2006. Reefs of the deep: the biology and geology of cold-water coral ecosystems. Science, 312: 543–547.

Roberts, J. M., Wheeler, A. J., Freiwald, A., and Cairns, S. D. 2009. Cold-water Corals: The Biology and Geology of Deep-sea Coral Habitats. Cambridge University Press, Cambridge, 334 p.

Rosa, M. R., Alves, A. C., Medeiros, D. V., Coni, E. O. C., Ferreira, C. M., Ferreira, B. P., de Souza Rosa, R., et al. 2015. Mesophotic reef fish assemblages of the remote St. Peter and St. Paul's Archipelago, Mid-Atlantic Ridge, Brazil. Coral Reefs, 35: 113–123.

Serrano, X. M., Baums, I. B., Smith, T. B., Jones, R. J., Shearer, T. L., and Baker, A. C. 2016. Long distance dispersal and vertical gene flow in the Caribbean brooding coral Porites astreoides. Scientific Reports, 6: 21619.

Sheppard, C. 1982. Coral populations on reef slopes and their major controls. Marine Ecology Progress Series, 7: 83–115.

Sherman, C., Nemeth, M., Ruiz, H., Bejarano, I., Appeldorn, R., Pagán, F., Schärer, M., et al. 2010. Geomorphology and benthic cover of mesophotic coral ecosystems of the upper insular slope of southwest Puerto Rico. Coral Reefs, 29: 347–360.

Singh, H., Armstrong, R. A., Gilbes, F., Eustice, R., Roman, C., Pizarro, O., and Torres, J. 2004. Imaging coral I: imaging coral habitats with the SeaBED AUV. Subsurface Sensing Technologies and Applications, 5: 25–42.

Sinniger, F., Morita, M., and Harii, S. 2012. "Locally extinct" coral species Seriatopora hystrix found at upper mesophotic depths in Okinawa. Coral Reefs, 32: 153–153.

Slattery, M., Lesser, M. P., Brazeau, D., Stokes, M. D., and Leichter, J. J. 2011. Connectivity and stability of mesophotic coral reefs. Journal of Experimental Marine Biology and Ecology, 408: 32–41.

Smith, T. B., Gyory, J., Brandt, M. E., Miller, W. J., Jossart, J., and Nemeth, R. S. 2015. Caribbean mesophotic coral ecosystems are unlikely climate change refugia. Global Change Biology, 22: 2756–2765.

Tenggardjaja, K. A., Bernardi, G., and Bowen, B. W. 2014. Vertical and Horizontal Genetic Connectivity in Chromis verater, an Endemic Damselfish Found on Shallow and Mesophotic Reefs in the Hawaiian Archipelago and Adjacent Johnston Atoll. Figshare. PLoS One, 9: e1154963.

Thresher, R. E., and Colin, P. L. 1986. Trophic structure, diversity and abundance of fishes of the deep reef (30-300 m) at Enewetak, Marshall Islands. Bulletin of Marine Science, 38: 253–272.

van Oppen, M. J., Bongaerts, P., Underwood, I. N., Peplow, L. M., and Cooper, T. F. 2011. The role of deep reefs in shallow reef recovery: an assessment of vertical connectivity in a brooding coral from west and east Australia. Molecular Ecology, 20: 1647–1660.

Wagner, D., Kosaki, R. K., Spalding, H. L., Whitton, R. K., Pyle, R. L., Sherwood, A. R., Tsuda, R. T., et al. 2014. Mesophotic surveys of the flora and fauna at Johnston Atoll, Central Pacific Ocean. Marine Biodiversity Records, 7: e68.

White, K. N., Ohara, T., Fujii, T., Kawamura, I., Mizuyama, M., Montenegro, J., Shikiba, H., et al. 2013. Typhoon damage on a shallow mesophotic coral in Okinawa, Japan. Peer Journal, 1: e151.

Wickham, H. 2009. ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag, New York.

Williams, S. B., Pizarro, O., Jakuba, M. V., Johnson, C. R., Barrett, N., Babcock, R., Kendrick, G. A., et al. 2012. Monitoring of benthic reference sites using an autonomous underwater vehicle. Robotics & Automation Magazine, 19: 73–84.

Ziegler, M., Roder, C. M., Buchel, C., and Voolstra, C. R. 2015. Mesophotic coral depth acclimatization is a function of host-specific symbiont physiology. Frontiers in Marine Science, 2: 4.