



REPORT

Studying functions on coral reefs: past perspectives, current conundrums, and future potential

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Abstract Function-based studies have opened a new chapter in our understanding of coral reefs. Unfortunately, we are opening this chapter as the world’s reefs rapidly transform. In this context, one of the most important roles of function-based studies is to inform coral reef conservation. At this critical juncture, we have a chance to reflect on where we have come from, and where we are going, in coral reef functional ecology, with specific consideration of what this means for our approaches to conserving reefs. As focal examples, we examine the role of corals on reefs, and the practice of culling crown-of-thorns starfish, from a functional perspective. We also consider how the papers in this special issue build on our current understanding. Ultimately, we highlight how robust scientific investigation, based on an understanding of ecosystem functions, will be key in helping us navigate reefs through the current coral reef crisis.

Keywords Active interventions and reef restoration · Coral reef management · Crown-of-thorns starfish · Ecosystem function · Ecosystem process · Reef growth

Introduction

Compared to other disciplines in ecology, the study of ecosystem functions harbours a unique suite of challenges. In addition to species-specific abundances, studies of function require detailed information on organismal traits (e.g. size, diet, mobility), as well as the role of species in ecosystem-scale processes (e.g. herbivory, predation). While this need for additional information can impede the pace at which research on function progresses, the rewards for integrating this information are clear. This is especially true for high-diversity coral reefs, where strong biotic interactions can lead to a handful of species disproportionately contributing to ecosystem processes. For example, studies of coral reef functions have revealed that *Bolbometopon* parrotfish can erode the reef at a rate equal to total calcification, with each individual producing over 5 tonnes of sand per year (Bellwood et al. 2003), and that 60% of all fish flesh consumed on reefs may come from ‘unseen’ cryptobenthic fishes (Brandl et al. 2019b). Such information is critical in our quest to decipher how coral reefs operate and, increasingly, to inform coral reef conservation efforts.

We have now entered a critical period for coral reef conservation, where the actions we take today will dictate the nature of coral reefs tomorrow and for many decades to come. Successful coral reef conservation requires that these actions are grounded in robust science, in which functional studies can play an important role (Kuffner and Toth 2016; Bellwood et al. 2019a). Given that we have reached this critical juncture, it is prudent to take stock and reflect on where we have come from, and where we are going, in terms of function-based studies on coral reefs, with specific consideration of what this means for the conservation of coral reefs in the Anthropocene. We also consider how the papers included in this special issue *Functional studies on*

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coral reefs: insights from a changing world have changed our understanding of coral reef ecosystem functioning.

The evolution of functional studies on coral reefs

There is a long history of research studying functions on coral reefs, and observations with functional relevance began centuries ago, including observations by Darwin (1842) who's quantitative reasoning on links between coral growth and reef production are still relevant today. Nevertheless, although studies of coral reef functions are scattered throughout most of the history of coral reef ecology, systematic and cohesive efforts to understand functions have only recently begun to emerge. From the 1700s until the 1970s, most reef research was centred on taxonomy and basic ecological descriptions of which organisms occurred where (e.g. Randall 1955; Goreau 1959; Talbot 1965). From the 1970s onwards, the increased prevalence of SCUBA diving in science underpinned a burgeoning interest in the quantification of marine organisms, with increasingly sophisticated analytical approaches facilitating a growing interest in documenting abundance patterns, be they at biogeographic or population levels (e.g. Sheppard 1980; Anderson et al. 1981; Williams and Sale 1981; Done 1982; Alevizon et al. 1985). Documenting spatiotemporal variation was in vogue; indeed, reef scientists were primarily interested in understanding patterns (e.g. community composition), rather than examining processes (i.e. functions). The term 'function' was largely left to functional morphologists who, at this time, were exploring the potential for combining morphology and ecology in the rapidly expanding field of eco-morphology (Motta 1988; Wainwright 1988), or in the delineation of 'functional groups' of coral reef organisms (Steneck 1983, 1988). At this time, the term functional was most commonly used when examining how organisms worked, not ecosystems.

As our understanding of coral reef organisms grew, our approaches to quantifying where they occur on reefs, and why, became increasingly structured. In the 1980s, coral work increasingly focused on distribution and abundance censuses (quadrats, line intercept methods etc.) (Bak and Luckhurst 1980; Done 1982). Moreover, on the Australian Great Barrier Reef (GBR), the Australian Institute of Marine Science was implementing new cross-shelf fish survey approaches that captured the different environmental contexts in which reefs, and their associated communities, existed—from turbid inshore reefs to clear-water high energy offshore reefs (Fig. 1); each with their own characteristic faunas and associated regimes of ecosystem processes such as calcification and erosion (Done 1983; Kleypas et al. 1999). At this time, the best available fish data were based on either mass of fishes per family from explosive samples

(Williams and Hatcher 1983) or early underwater visual census (UVC) methods, e.g. using point censuses (Bohnsack and Bannerot 1986) or a Log₃ abundance scale (e.g. Russ 1984a, 1984b). Fish numbers (i.e. abundance) and identities were the focus; ecologically informative traits, like body size, were often not recorded. Thus, identification and counting were still relatively crude and, indeed, the first full fish identification book for the GBR was not published until 1990 (Randall et al. 1990).

Our understanding of what reef organisms actually did was even less refined and there was still no united field of 'coral reef functional ecology'. Nevertheless, the process-focussed foundations for the subdisciplines within this field, that would ultimately become major avenues of research, had already been laid. This included threads of ecological understanding emerging from earlier feeding observations and dietary analyses (e.g. Hiatt and Strasburg 1960; Randall 1967; Jones 1968; Choat 1969; Robertson et al. 1979). Moreover, a basic understanding of system-wide coral reef productivity and trophic energy flows was established (Odum and Odum 1955), as well as the roles of reef fishes in controlling (Randall 1961) and utilizing (Bardach 1959) this productivity. There was an early appreciation of the central role of corals in reef growth (Goreau 1963; Stoddart 1969; Chave et al. 1972; Stearn et al. 1977; Chappell 1980). Ecological aspects of reef growth and carbonate budgets (Smith and Kinsey 1976; Stearn et al. 1977), as well as the role of fishes in these processes (Bardach 1961), was also already underway.

Building on this pioneering work, the study of processes operating on coral reefs rapidly expanded in the late 1980s and into the 1990s. The field of reef fish research shifted from documenting communities to understanding where they came from, through population biology and recruitment-oriented work. A focus on the 'black-box' of the pelagic realm (e.g. Doherty and Williams 1988; Sale 2004) developed into work on population connectivity, especially pertaining to marine protected areas (Jones et al. 2009; Harrison et al. 2020), while benthic studies showed an increased interest in settlement, growth, and demography (e.g. Hughes et al. 1999; Pratchett et al. 2015). Other major advances included the direct measurement of reef processes, like herbivory and benthic productivity (e.g. Hatcher and Larkum 1983; Steneck 1983; Hay 1984; Carpenter 1986; Russ 1987; Klumpp and McKinnon 1989). These early findings were so fundamental to our understanding of herbivory that today we are still drawing on the lessons learnt during this period (see Tebbett et al. 2023b in this special issue for an overview). Discoveries also expanded beyond the reef benthos, with increased attention to off-reef habitats such as seagrasses and mangroves and, perhaps most importantly, the pelagic realm. Pioneering work by Hamner et al. (1988) shaped our understanding of planktivory on

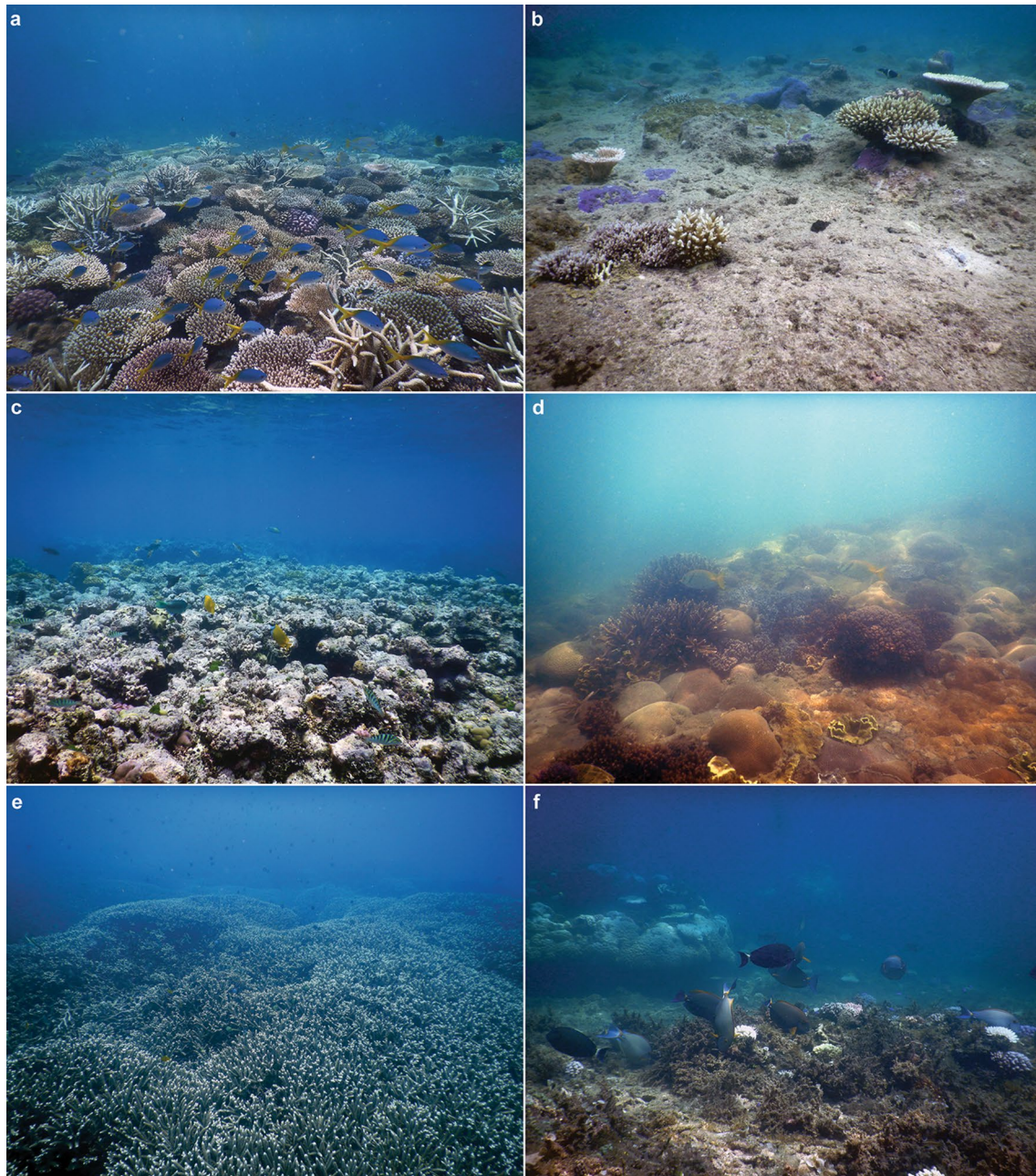


Fig. 1 ‘Coral reefs’ come in a variety of forms, all of which are embedded within their own environmental context. **a** Tropical, high coral cover reef system (Linnett Reef, Great Barrier Reef [GBR], 2021), **b** low coral cover, high turf cover reef system (Lizard Island, GBR, 2021 post disturbance by cyclones and bleaching events), **c** high-energy, high crustose coralline algae cover reef system (Yonge

Reef, GBR, 2021), **d** turbid, nearshore reef system (Bowen, GBR, 2019), **e** low diversity, subtropical reef system (Flinders Reef, South East Queensland, Australia, 2020), and **f** high macroalgal cover, low coral cover reef system (Turtle Group, GBR, 2021) (Photographs: SB Tebbett)

reefs, with recent studies (e.g. Leray et al. 2019 and Gahan et al. 2023 in this special issue) expanding our insights into the nutritional value of key planktonic groups for fishes. This period also saw a major expansion in our understanding of reef growth, primarily through coring, in both the Indo-Pacific and Western Atlantic (Adey 1978; Scoffin and Dixon

1983), as well as the role of key bioeroding organisms in these processes (Hutchings 1986; Scoffin 1992; Bak 1994; Bellwood 1995). Indeed, researchers were already focusing on understanding the interaction between reef growth and global climate change/sea level rise (Buddemeier and Smith 1988), with this field expanding in the twenty-first century as

the effects of climate change intensified (Perry et al. 2018; Kench et al. 2022; Toth et al. 2023).

Consequently, by the late 1980s, for both fishes and corals, many of the methods that would lay the foundations for studies of coral reef functions, as we know them today, were already in place. However, a unified and cohesive understanding of reef functions was still lacking, and it was not until the mid-1990s that the broader concept of reef ‘ecosystem function’, was more formally expressed (see Done et al. 1996). Unfortunately, in many areas, this pioneering research on coral reef ecosystem functions was not immediately followed up. This may be, in part, due to the changes that began rapidly unfolding on the world’s coral reefs. The halcyon days of interest-based science on coral reefs ended in 1998 with the first global coral bleaching event (Fig. 2). The 1998 event triggered a great deal of concern, monitoring, and funding (Goreau et al. 2000; Obura et al. 2019), but not necessarily new ideas of what to measure or how. Indeed, such was the parlous state of reef function studies that it was not until 2019 that the term ‘function’ was defined (Bellwood et al. 2019b) and clarified (Brandl et al. 2019a), and only in 2022 was the related concept of ‘functional traits’ similarly clarified (Streit and Bellwood 2023). In the meantime, we continued to quantify coral reef declines at increasingly large spatial scales, especially in terms of ‘coral cover’ (e.g. Gardner et al. 2003; De’ath et al. 2012; Hughes et al. 2017b; Tebbett et al. 2023c) (Fig. 2). To this day, monitoring is largely based on fish counts (often of a select group of fishes in a select set of reef habitats; Bellwood et al. 2020) and coral cover data (in groups separated by growth morphology); all on relatively short transects (e.g. 30–50 m).

The current special issue explores the growing diversity of methods for quantifying reef functions, including video-based methods (Collins et al. 2023; Magneville et al. 2023), or the integration of remote-sensing (Lutzenkirchen et al. 2024); two areas in which information on reef functions has benefited greatly from technological advances (e.g. DiFiore et al. 2019; Madin et al. 2019; Schiettekatte et al. 2022b). Nevertheless, with ongoing global bleaching events, it has become increasingly clear that reefs are in serious danger (Hughes et al. 2017a; Sully et al. 2019) and that, despite many advances, we still have little idea of: (i) how reefs function and (ii) what the implications of coral bleaching are for critical reef processes. What is missing is a clear, quantitative, understanding of reef processes, i.e. ecosystem functions, and their respective importance across reef types and environmental conditions. Currently, we are left with rapidly transforming ecosystems, largely non-transformed scientific methods, and little idea of which ecosystem functions (processes) are still operating, needed, or manageable (Bellwood et al. 2019a). To focus research and to address these shortcomings, it may pay to re-examine the three key

questions: (i) what do we want to protect on coral reefs? (ii) why? and (iii) how can this be achieved?

What do we want to protect on reefs and why?

There is no simple, single answer to the questions: what do we want to protect on coral reefs and why? Answers are invariably context dependent (cf. Bellwood et al. 2019b), with people valuing coral reefs for a plethora of different services in different parts of the world (Lau et al. 2019; Woodhead et al. 2019). However, arguably one of the most ubiquitous features of coral reefs that we want to protect is their physical structure (Hoegh-Guldberg et al. 2019; Perry and Alvarez-Filip 2019), which is tied to nearly every key service that we derive from reefs, including their capacity to (a) protect coastal communities from hydrodynamic activity (Ferrario et al. 2014), (b) enhance fisheries productivity (Rogers et al. 2018), and (c) support tourist/recreational activities (Santavy et al. 2021), including the organismal biodiversity for which reefs are renowned (Siqueira et al. 2023). Indeed, the importance of focusing on this goal was recently expressed by Kuffner and Toth (2016) when they stated: “*A better understanding of the processes that control the long-term resilience of reefs as geomorphic structures, not just as ecological communities, may help optimize management activities aimed at increasing both reef longevity and the delivery of critical ecosystem services*”.

Importantly, to protect the physical structure of reefs, we know that reef growth via carbonate deposition/production needs to match or exceed erosion. This may not always require sustained vertical accretion (e.g. Toth et al. 2018, 2022), but in the wake of climate change and sea-level rise, sufficient accretion will be needed to accommodate future sea-level changes (Kennedy et al. 2013; Kench et al. 2022). We thus have a near-universal key function of interest—reef growth (accretion). With this clear focus at hand, functional studies offer a rare opportunity to critically evaluate whether our conservation priorities match natural ecological processes and our increasingly prevalent response to climate change: reef restoration.

The vast majority of reef restoration approaches have a single target: to increase coral cover (Hughes et al. 2023). This appears to be a logical goal, the presumed rationale being that: (a) corals have suffered the greatest impact from global warming, and (b) corals provide critical ecosystem functions. Most importantly, the latter includes both the promotion of reef growth/accretion and ancillary functions, such as maintaining fish populations, increasing fisheries productivity, increasing reef complexity for the thousands of associated reef creatures, and, finally, looking attractive to humans (Pratchett et al. 2014b; Rogers et al. 2018). Despite its instinctive appeal, it is important to evaluate whether

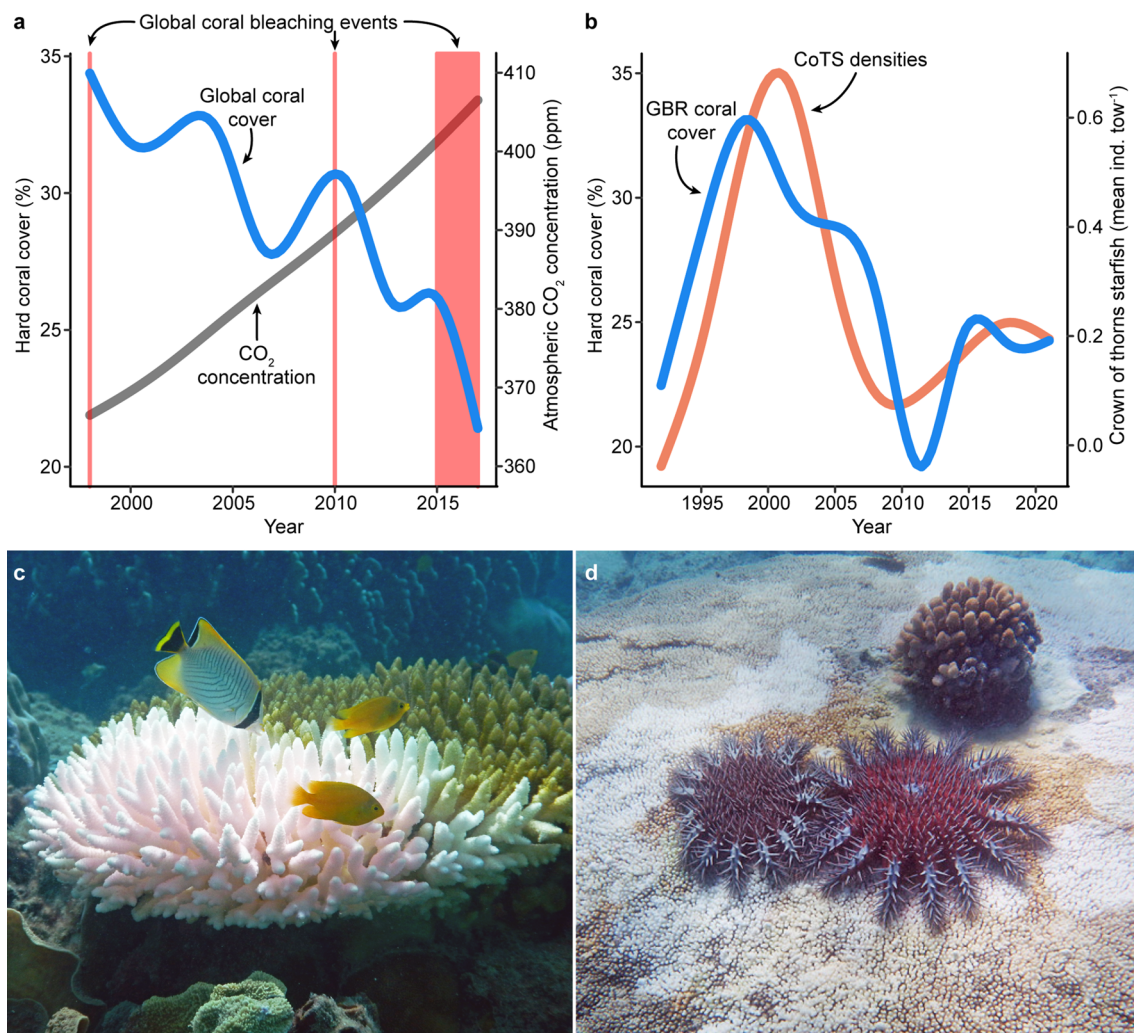


Fig. 2 Coral cover dynamics on Anthropocene coral reefs. **a** Temporal variation in coral cover on the world's coral reefs (blue line) and atmospheric carbon dioxide concentration (grey line) from the onset of the first global bleaching event (1998) to the end of the most recent global bleaching event (2015–2017). Global coral bleaching events during this period are shown as red bars (Skirving et al. 2019). **b** Temporal variation in coral cover (blue line) on Australia's Great Barrier Reef (GBR) and crown-of-thorns starfish (CoTS) densities (orange line) on the GBR. **c** A corallivorous butterflyfish and two yellow damselfishes, interacting with a bleached *Acropora* coral at Lizard Island during the 2016 bleaching event (photograph: RP Streit), and **d** two corallivorous CoTS feeding on an *Acropora* coral (white areas are feeding scars) (photograph: A Hoggett). These two corallivorous organisms have fundamentally different reputations; one will make it onto postcards (the butterflyfish in **c**), while the other is

the subject of a multi-million dollar culling program (the CoTS in **d**). Coral cover data in **a** were sourced from a dataset compiled in Tebbett et al. (2023c) (see this reference for original data sources and relevant citations) and are based on 15,668 mean site level coral cover datapoints (collected using point-intercept or photo-quadrat transect methods) from reef crest and slope habitats at depths < 15 m. The atmospheric CO₂ concentration data in **a** were sourced from Keeling et al. (2001, 2023) and were based on in situ air measurements from the Mauna Loa Observatory, Hawaii. Coral cover data in **b** were based on photo transects and were sourced from AIMS (2015a), while CoTS density data were based on manta tows and sourced from AIMS (2015b). All lines in **a** and **b** were fitted using gam smoothers for visualisation purposes in *ggplot2* (Wickham 2016) in the software R (version 4.2.2; R Core Team 2022)

this focus on corals is based on ecological evidence or our intuition. If it transpires that reef functions (e.g. reef growth or the creation of 3D habitat) are not directly dependent on the corals that are being targeted for restoration, then can current practices of restoring corals be justified in a functional context (at least from the perspective of reef growth)?

It is undoubtedly true that a large proportion of the carbonate in reef cores comes from corals (Montaggioni 2005; Gischler 2014; Webster et al. 2018; Toth et al. 2019) and that corals are one of the dominant calcifiers on coral reefs (Perry et al. 2018; Perry and Alvarez-Filip 2019; Cornwall et al. 2021). It thus appears logical that the loss of corals would result in diminished reef growth and eventually

the loss of reef structures. And, indeed, this may well be the case in many regions, including the East Pacific, where, for example, the loss of corals, reduced coral growth, and exceptionally high urchin-based bioerosion have been associated with the disappearance of reef structures (Cortés 1997; Manzello et al. 2017; Glynn et al. 2018; Enochs et al. 2021). However, the relationship between coral abundance, diversity, or growth rates and reef accretion may not be linear and in extreme cases, may not be present at all.

Indeed, recent research suggests that coral growth rates and coral diversity may be largely unrelated to reef growth, at least across geological timescales. For example, reefs in the Atlantic have just 7% of the coral species richness found in the Indo-Pacific (and have just two fast-growing *Acropora* species) (Roff 2021). They have also had 6000 years less time with high sea-levels in which to grow (Gischler 2010). Thus, in theory, Indo-Pacific reefs should be 500–1000 m wider than their Caribbean counterparts; yet the widths of Caribbean and Indo-Pacific reefs are almost identical (Lutzenkirchen et al. 2023). This suggests that horizontal shallow water reef area and coral diversity are not inherently correlated; instead, reef size appears to be constrained by hydrodynamics, not coral species richness nor sea-level stability (Lutzenkirchen et al. 2023; also see Adey 1978; Montaggioni 2005).

A similar pattern has been documented when looking at reef growth through vertical accretion. In the Caribbean, vertical reef accretion was not found to be affected by significant coral extinction events (i.e. a 50% loss of species; Johnson et al. 2008). Similarly, on Indo-Pacific reefs, vertical accretion appears to be independent of the growth rates of the local dominant framework-building coral species (Roff 2020). Indeed, it is widely accepted in geological contexts that coral growth rates and diversity are not directly correlated with the onset or extent of reef development or growth (Pandolfi and Kiessling 2014; Pomar et al. 2017). The lack of a clear relationship between coral diversity and reef growth also means that, if the goal of restoration activities is to enhance reef growth, then selecting the key coral species is likely to be extraordinarily difficult on high-diversity Indo-Pacific reefs (cf. Madin et al. 2023).

If coral diversity and coral growth rates are poorly correlated with reef accretion, perhaps coral abundances are a better indicator? Today, and for the last few thousand years, Caribbean reef accretion has depended largely on just two *Acropora* species; with recent losses raising concerns over the functional persistence of these structures (Toth et al. 2018, 2022). However, the disconnects between coral growth and reef growth may also manifest over shorter timescales and incorporate local coral abundances. Today, global mean coral cover is only approximately 25% (Tebbett et al. 2023c), yet even then some Indo-Pacific corals, especially branching *Acropora*, may contribute little to the consolidated

reef matrix and net accretion processes in critical habitats, such as on the high-energy reef crest. For example, Morais et al. (2022) found that the vast majority of *Acropora* colonies disappeared completely within five years, with no detectable contribution to net in-situ accretion. Presumably, these corals contribute primarily to reef growth in lower energy habitats via production of sediment and rubble that contribute to infilling (with corals invariably constituting the dominant component of cores, and being the major source of carbonate, in such habitats). Indeed, detailed quantification of reef growth over ecological timescales has revealed that accretion can continue despite coral loss, with the contribution of crustose coralline algae to this process being particularly notable (Kench et al. 2022).

Unfortunately, to date, virtually all contemporary reef growth/reef budget studies have focused on the generally high coral cover reef slope (especially ecological studies) or the down-current reef flat habitat (especially coring studies) where dead coral skeletons often accumulate, with far less attention given to reef growth in the reef crest habitat; the primary place where the reef breaks the waves (see Kench et al. 2022 for an overview). Indeed, comparisons between reef habitats raise the possibility that there may be two functionally distinct types of reef growth: ‘reef growth’ version 1 sensu lato, i.e. growth of the total reef structure, especially in terms of backreef progradation—as detected by most cores and which is indeed coral dominated versus ‘reef growth’ version 2 sensu stricto, i.e. in terms of the formation of the structure that breaks waves and allows the backreef to exist. The latter type of growth may depend to a substantial extent on crustose coralline algae and encrusting corals, and results in the actual presence of a reef based on the definition provided by Hopley and Smithers (2008): “*Coral reefs are biogenic limestone structures built by corals and other carbonate-producing organisms in shallow tropical and subtropical marine settings, where they grow upwards or towards sea level as landforms able to resist wave action*” (for other definitions see Kleypas et al. 2001). Beyond the reef structure that breaks the waves, reef growth sensu lato as defined above (version 1), is largely infilling and widening (notably the substantial amount of carbonate produced by corals appears to be key in these reef-building processes). The study by Kench et al. (2022) appears to be one of the first ecological studies to comprehensively quantify growth on the critical wave-breaking reef crest, though this study is restricted to one location at small spatial and temporal scales (1 m² over 2 years). Additional studies of reef growth in the most exposed reef habitats are clearly needed.

This evidence inevitably raises the question: are corals always necessary for reef growth (especially vertical growth at the reef crest)? The answer is: probably not. However, reefs built by non-coral organisms are likely to be functionally different, even though they can deliver the key function

of reef construction. In this respect, there is growing recognition within coral reef geology and ecology that crustose coralline algae may be critical for reef growth and carbonate production (e.g. Rasser and Riegl 2002; Nash et al. 2013; Vargas-Ángel et al. 2015; Weiss and Martindale 2017; Teichert et al. 2020; Cornwall et al. 2023; Raja et al. 2023), following recognition of this important association decades earlier (e.g. Goreau 1963; Womersley and Bailey 1969; Adey and Macintyre 1973; Littler and Littler 1984). Indeed, some physically complex biogenic tropical reefs function with virtually no corals, such as the Atol das Rocas in Brazil (Gherardi and Bosence 2001). Similarly, coralline algae also dominate framework construction in various other scenarios, including in the formation of the critical algal ridge/rim habitat, which breaks the waves in high-energy settings (Adey 1975; Littler and Doty 1975; Bosence 1983; Rees et al. 2005). These reef structures constructed by coralline algae may offer a glimpse into potential benthic configurations in the Anthropocene, if non-coral calcifiers are not as sensitive as corals to climate change (note the response of CCA to future climate change conditions have been varied [see Cornwall et al. 2019, 2022 for reviews on this topic]). Therefore, it could be that, under some circumstances, some of the functions of corals on coral reefs can be sustained independent of coral diversity (e.g. long-term reef crest development) and perhaps also independent of coral composition or abundance (e.g. short-term carbonate production). From a functional perspective, this opens the possibility of a conceptual shift from ‘coral reefs’ to ‘reefs with corals’ (also see Womersley and Bailey 1969; Kleypas et al. 1999). As continued anthropogenic impacts decimate coral populations on reefs this conceptual shift may become more apparent.

Ultimately, this also means, that if the goal of coral restoration is to maintain reef growth, then our method of doing this (increasing the cover of ‘weedy’ corals in the Indo-Pacific; i.e. fast growing, highly fecund, rapidly recruiting, and highly competitive *Acropora*) may not be the most appropriate strategy (cf. Madin et al. 2023); especially if we take this action before we stop the main driver of coral declines (i.e. climate change [Hughes et al. 2023]). Even in the Caribbean/Western Atlantic, where benthic systems operate in fundamentally different ways to the Indo-Pacific (Roff and Mumby 2012; Tebbett et al. 2023c), and where two *Acropora* species can be easily identified as the main contributors to the carbonate framework, care is needed. Even if *A. palmata* and *A. cervicornis* can be reared, transplanted, and kept alive through successive bleaching events and disease outbreaks, they may not adequately perform the processes required for consolidation and reef accretion in monocultures. Indeed, an inherently coral-centric perspective (perhaps guided by their high vulnerability) may have

left us with only a partial understanding of the impacts of climate change and its implications for coral reefs.

The key problem is that a reef with corals is likely to be functional, but adding corals may not always guarantee reef health or growth. Accepting that the coral contributions to reef growth may be negligible is challenging, as is the fact that many reefs are not dominated by corals (Vroom 2011; Tebbett et al. 2023d). But this changing mindset may be essential if we are to accurately and responsibly evaluate our actions to preserve reefs into the future; simply adding corals, while an attractive and intuitive solution, may not be enough or the best approach. Viewing reefs as the outcome of collective calcification of a range of organisms, including algae, with scleractinian corals offering a boost under the right circumstances, offers a new perspective on what we may need to do to preserve and sustain reefs (cf. Adey 1998). While there may be strong socioeconomic incentives for coral restoration (Hein et al. 2019), if we are primarily interested in maintaining coral reef accretion and structure, restoration efforts that raise and outplant colonies of corals that are highly susceptible to marine heatwaves, and have limited contributions to in-situ accretion, are likely to be an extraordinarily expensive and ultimately futile undertaking (Hughes et al. 2023).

It should be acknowledged that restoration practitioners may focus on restoring corals as an indirect approach to sustain or restore other reef functions, such as the productivity of fish communities. However, such indirect approaches invariably rely on the existence of strong links between fishes and coral cover, which can be context-dependent and weaker than often assumed (e.g. Wismer et al. 2019a, 2019b; Muruga et al. 2024). Indeed, reef fish productivity appears to be more closely related to structural complexity, rather than coral cover (e.g. Rogers et al. 2018; Morais and Bellwood 2019; Hamilton et al. 2022). Given that the structure provided by many fast-growing corals is likely to be relatively short-lived (due to the sensitivity of such corals to disturbance [Loya et al. 2001; Hughes et al. 2017b] and the fact that their skeletons rapidly erode after death [Morais et al. 2022]), then restoring these fast-growing corals may prove to be a relatively inefficient way of restoring reef functions such as fish productivity. Clearly there is a need to better understand the roles that corals and structural complexity play in supporting fish populations, especially in respect to separating co-occurrence from more causal, functional, interdependence.

How do we currently manage reef functions?

To date, a coral-centric view of coral reefs has largely shaped our approaches to managing functions, both directly and indirectly. Historically, we have aimed to secure

‘healthy’ high-coral cover reefs by managing the behaviour of local human populations (e.g. fishing activities). Certain functional groups of fishes were identified as critical to maintaining resilient reefs with high coral cover, such as herbivorous parrotfishes, that are thought to control the growth of competitive algae (Bellwood et al. 2004; Hughes et al. 2007; Graham et al. 2013). To maintain these fishes on reefs, we implemented actions that managed human behaviour (such as establishing no-take marine protected areas [MPAs]), in the belief that this would conserve these fishes and ultimately the ecosystem benefits they delivered (Hughes et al. 2007; Mumby and Steneck 2008). However, the links from management action to the ultimate goal of ecosystem resilience and functioning appear to have been too indirect and multifaceted to deliver easily discernible, consistent, ecosystem-scale benefits (reviewed in Bruno et al. 2019). This became especially evident in the wake of climate change and associated marine heatwaves, which do not respond to MPA boundaries delineated on paper (Jones et al. 2004; Hughes et al. 2017b; Bates et al. 2019; Graham et al. 2020).

The overarching goal of preserving coral cover on reefs has also spawned some of the largest scale and most well-funded attempts to directly manage a coral reef ecosystem function: reducing coral consumption by the corallivorous crown-of-thorns starfish (CoTS) in the Indo-Pacific (Fig. 2d) (Great Barrier Reef Marine Park Authority 2020). By 2018 over 1 million CoTS had been killed on the GBR (Pratchett et al. 2019), with the Australian Government investing ~\$70 million US in direct funding to the culling program between 2012 and 2022 (Pratchett and Cumming 2019), with an additional ~\$109 million US recently awarded (David and Bathgate 2022). The GBR program is, however, dwarfed by the most extensive culling effort undertaken to date in Japan, where ~13 million CoTS were removed from reefs in southern Japan in the 1970s and 1980s (Yamaguchi 1986). The cost of this activity was over 600 million yen in 1970; which is equivalent to more than \$2.5 billion US today [Pratchett and Cumming 2019]. Despite the popularity of these interventions, an important conundrum has remained unresolved for decades (e.g. Kenchington and Kelleher 1992): are these active management interventions ecologically justifiable?

The conundrum of CoTS control revolves around the fact that these animals are a native species, in their natural habitat, expressing natural behaviour, and they are part of the biodiversity for which coral reefs are renowned. On the GBR, CoTS have been present for 1000 s of years (Fabricius and Fabricius 1992), with regular outbreaks occurring since at least the early-mid 1900s (Moran et al. 1992; Pratchett et al. 2014a). Indeed, CoTS have probably been interacting with coral reefs, and delivering the function of coral consumption, for 10 million years (Hall et al. 2017; Yuasa

et al. 2021). Yet, because of their capacity to eat corals, CoTS have been labelled a pest (Hall et al. 2017) (with the Merriam-Webster dictionary defining a pest as: “*a plant or animal detrimental to humans or human concerns [such as agriculture or livestock production]*”). This pejorative view of CoTS is reinforced by the belief that outbreaks are caused by, or exacerbated by, anthropogenic activities. In this respect, in 1992 the first chairman and chief executive officer of the Great Barrier Reef Marine Park Authority (GBRMPA), Graeme Kelleher, noted: “*The Authority’s policy on controlling COTS is not to interfere on a large scale unless it can be shown that outbreaks are caused or exacerbated by human activity*” (Kelleher 1992). Although it was clearly considered fundamental for science to establish if anthropogenic activities caused or exacerbated CoTS outbreaks before decisions were made to actively regulate population densities, we are still far from a conclusive answer (Pratchett et al. 2017, 2021). Indeed, after reviewing 30 years of research and > 1200 research articles, Pratchett et al. (2017) concluded that the question “*Why [CoTS] outbreaks occur and whether they are natural or unnatural phenomena?*” was largely unresolved. Nevertheless, despite a lack of clear scientific evidence that humans have modified CoTS outbreaks, the decision to systematically cull CoTS populations has already been made (discussed in Pratchett et al. 2017).

But, what does the mass killing of CoTS mean from a functional perspective? At face value the outcomes appear obvious: a decrease in coral consuming starfishes will yield higher coral cover (Plagányi et al. 2020; Westcott et al. 2020; Castro-Sanguino et al. 2023). However, if we consider that CoTS are a natural component of many coral reefs, we can dive deeper and ask: could coral consumption by CoTS be an important function on reefs? In answering this question, the argument could be made that CoTS preferentially target fast growing, competitively dominant, ‘weedy’ corals (albeit often moving onto other coral species once they have exhausted the supply of their preferred prey) (De’ath and Moran 1998; Pratchett et al. 2009; Pratchett 2010; Kayal et al. 2012; Keesing et al. 2019). It has even been suggested (by Porter 1972) that such consumption of highly competitive, fast-growing corals may release slower-growing species from overgrowth and shading and increase long-term benthic diversity (note Pratchett 2010 found overall coral diversity declined during a moderate outbreak of CoTS). In this respect, it is interesting to note that over evolutionary timescales, the fast-growing Acroporidae has been associated with the greatest rates of extinction in non-acroporid corals (Siqueira et al. 2022). Moreover, by consuming fast-growing corals, CoTS: (a) release benthic space for increased benthic productivity, which is critical in coral reef food webs (Russ and St. John 1988; Klumpp and McKinnon 1989; Hatcher 1990), (b) provide the preferred

habitat for many reef crustacea, dead coral (Kramer et al. 2014; Fraser et al. 2021), and (c) free up space for other critical benthic coverings, such as crustose coralline algae (Adey 1998; Cornwall et al. 2023).

We are currently entering a period in which fast growing corals, especially *Acropora*, are likely to be subjected to unstable boom-and-bust dynamics (Wilson et al. 2019; Pratchett et al. 2020; Morais et al. 2021). Indeed, while many fast-growing corals are highly vulnerable to heat stress, their comparatively high growth rates may ultimately (and counterintuitively) bestow an ecological advantage. Specifically, by heating the planet, we may artificially increase the competitive advantage of fast-growing corals (e.g. *Acropora*) in some contexts, with such corals being some of the few that can ‘recover’ during the much shorter ‘recovery windows’ between the large disturbance events that coral reefs now experience (Hughes et al. 2018; Lough et al. 2018; Morais et al. 2023). Could it be that CoTS moderate the instability of these boom-and-bust periods via coral consumption akin to the important role fires play in some Australian native forests (as noted by Kenchington and Kelleher 1992)? Are we in effect, actively preventing the establishment of diverse, resilient reefs by culling *Acropora* consuming CoTS? Alternatively, could large fluctuations in CoTS densities, combined with their variable effects on coral assemblages, add to instability in *Acropora* abundance and overall coral cover? Answering these critical ecological questions is essential if we wish to understand how unstable, disturbed, coral reefs function, and the roles of CoTS therein.

In a nutshell, CoTS may perform a key function on coral reefs. Ecological evidence (or rather the lack thereof) warrants, at the very least, further investigation into their true functional role, despite the current established view that CoTS are detrimental to reef functioning because they decrease live coral cover. Could it be that across the Indo-Pacific we have spent the equivalent of billions of dollars trying to prevent a natural and functionally important process? In this context, the prescience of Graeme Kelleher’s call for caution (in the same article as above) is striking: “*there is the possibility that our massive interference in what might be a natural element of the Great Barrier Reef system could have major unforeseen ecological effects*” and that the “*conclusion is that the risks from adopting a policy of massive destruction of crown-of-thorns starfish are very significant and that the only potential benefits may turn out to be costs in the long run*” (Kelleher 1992). Changing priorities of the management agency, GBRMPA, have since overturned these initial considerations (cf. Morrison 2017).

Currently it is unclear if CoTS do, or do not, perform a key function on reefs. Nevertheless, we have already labelled these organisms as pests, a defamatory characterization that is facilitated by their toxic spines, hostile appearance, and the religious overtones of their common name. Such

value-laden terms can cloud scientific judgement (Johns and DellaSala 2017; Morton 2017) and may mean that we do not approach the study of CoTS with an open mind, nor entertain the possibility of them performing a key function on reefs. These subconscious biases can be critical in shaping how we study coral reefs and, as scientists, we must be aware of their potential to shape our decisions of what to study and why (Bellwood et al. 2020). Our perceptions of the parrotfish *Bolbometopon muricatum*, relative to CoTS may be indicative of such potential biases.

After CoTS, *B. muricatum* are arguably the single largest predator of corals on Indo-Pacific reefs (when present in their original, natural, densities; Bellwood et al. 2003, 2012; McCauley et al. 2014). In contrast to CoTS, they actively reduce the coral reef framework and net storage of calcium carbonate. *B. muricatum* can erode the reef at a rate of 5.69 tonnes per individual per year, with intact parrotfish populations capable of consuming the reef at a rate that matches local accretion; thereby maintaining a dynamic carbonate balance (Bellwood et al. 2003) (Fig. 3). Yet, despite consuming a vast quantity of coral and reef structure each year, *B. muricatum* are widely considered to perform important, beneficial, ecosystem functions on reefs via the removal of carbonate and the production of sand (Bellwood et al. 2004; Perry et al. 2015; Morgan and Kench 2016; Thomson et al. 2021).

However, during the mid-1900s *B. muricatum* were so heavily fished on most reefs across the Indo-Pacific that there are now few reefs left that support natural densities of these parrotfish (Bellwood et al. 2012). Thus, across an entire ocean basin, we have directly modified the critical process of bioerosion—few reefs across the Indo-Pacific experience the high levels of bioerosion that they evolved under (Fig. 3). In response, the overharvesting of *B. muricatum* is widely viewed as a key conservation issue, with a growing push for better management strategies to conserve the individuals that remain (Donaldson and Dulvy 2004; Comeros-Raynal et al. 2012; Hamilton et al. 2016, 2019; Friedlander et al. 2023). The contrast between a ‘conservation priority’ (*B. muricatum*) and a ‘pest’ (CoTS), that both deliver, fundamentally, a similar ecosystem function (in terms of coral mortality), is remarkable.

Ultimately, there is no doubt that coral loss is a concerning trend that is reshaping the world’s reefs (Fig. 2) and that human activity is the fundamental driver of these trends. Scientists and managers alike recognize the critical need to halt and reverse the decline of coral reefs (De’ath et al. 2012; Bellwood et al. 2019a; Bruno et al. 2019). The only problem is that we may be making questionable decisions and for the wrong reasons, especially if our decision-making process is grounded not on robust scientific evidence but is instead shaped by aesthetic, sentimental, economic, or political values. Indeed, it seems startling to think that during an era

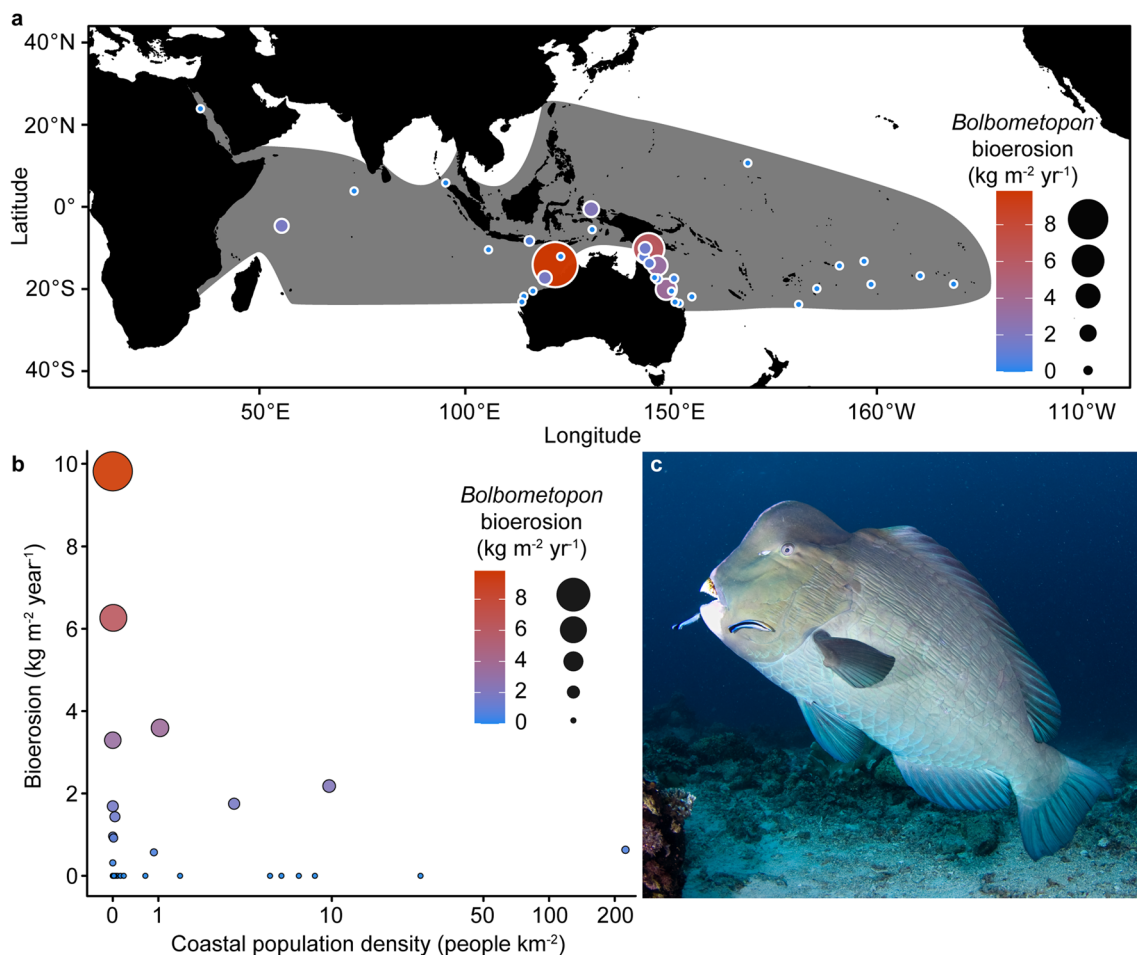


Fig. 3 Bioerosion by the excavating parrotfish *Bolbometopon muricatum*. **A** The spatial distribution of mean bioerosion by *B. muricatum* at 35 coral reef locations within their natural range (denoted by the grey area) based on the capacity of each individual *B. muricatum* to erode 5.69 tonnes of carbonate per year (Bellwood et al. 2003). **B** *B. muricatum* bioerosion rates compared to coastal human population densities from within a 100 km radius buffer. **C** *B. muricatum* (pho-

tograph: JP Krajewski). *B. muricatum* abundance data were sourced from the publicly available Reef Life Survey dataset (Edgar and Stuart-Smith 2014; Edgar et al. 2020). Only locations with 20 or more transects at shallow depths (<15 m) were included. Coastal population data were sourced from Center for International Earth Science Information Network–CIESIN–Columbia University (2018)

of the twenty-first century, when humanity’s effort to put a person on the moon is becoming a distant memory, our greatest achievement in directly managing an ecosystem function on coral reefs is the mass killing of a naturally occurring echinoderm. Will such approaches hold up to scrutiny when future generations ask us why we managed coral reefs the way we did?

Future outlook and conclusions

Considering reefs from a functional perspective may change our perceptions, including the way we perceive both corals and their predators. Going forward, studies that have an objective focus on ecosystem functions will be critical in developing and appraising our decision-making processes

when it comes to reef conservation. In this respect, a first step may be to carefully consider if our coral-centric view of the world’s coral reefs is justified and if corals will be the best indicators for coral reef conservation in the future. It has been repeatedly noted that many reefs are not typically dominated by corals (based on 2D planar extent) (e.g. Vroom 2011; Bruno et al. 2014; Tebbett et al. 2023d), but instead are, in the words of Adey (1998), “*algal structured and mediated ecosystems*”. Such algae, including crustose coralline algae and turfs, are typically the most abundant component of reefs now, and are likely to be even more so into the future (Agostini et al. 2021; Cornwall et al. 2023; Tebbett et al. 2023c, 2023d). Some algae play pivotal functional roles on reefs, including a key role in facilitating reef growth (Kench et al. 2022; Cornwall et al. 2023) and in shaping the settlement of reef organisms (Evans et al. 2020;

Ricardo et al. 2021; Doll et al. 2023). However, despite their prevalence, and the fact that they are central to how coral reefs operate, such algae have been largely overlooked in coral reef science, conservation, and restoration endeavours (especially relative to corals). Ultimately, in our efforts to date, we may have focused on studying a vulnerable and conspicuous subset of the reef's inhabitants, in a subset of places, with the high likelihood that this has shaped our perceptions of how coral reefs operate (Bellwood et al. 2020). Realigning our efforts to understand other functionally important, yet largely overlooked, components of coral reefs may hold the key to helping us navigate reefs through a changing world.

Beyond realigning scientific efforts, the second major step consists of fully appreciating the markedly different contexts in which coral reefs exist (Fig. 1) and ensuring the different biases we may have with regards to reefs in different contexts do not cloud our judgement (cf. Bellwood et al. 2019b). Indeed, it is becoming increasingly clear that lessons learnt about how reefs function in one context or location, may not translate to reefs in different contexts (Roff and Mumby 2012; Brandl et al. 2019a; Tebbett et al. 2023c). This means there is no 'one size fits all' approach to reef management, with the goals of any management strategy dependent on how the specific ecosystem in question functions given the social and environmental context within which it exists (Bellwood et al. 2019a; Williams et al. 2019). Moreover, examining coral reefs in extreme and marginal contexts can yield key insights into how reefs operate outside typical domains, and may yield functionally informative insights that question widely held perceptions (although this is invariably dependent on how analogous marginal/extreme reefs are to disturbed coral reefs) (Burt et al. 2020; Schoepf et al. 2023; Tebbett et al. 2023a). For example, the Galápagos Islands are renowned for their unique, endemic diversity. In this special issue, however, Tebbett et al. (2024) showed that the largely overlooked surgeonfish *Prionurus laticlavus* in the Eastern Galápagos Islands disproportionately contributed to the productivity of herbivorous fish assemblages, underpinning productivity that was more than threefold higher compared to typical, high-diversity reef systems. Importantly, such intriguing results are not out of the ordinary, with functional studies on coral reefs having a long history of revealing fascinating and unexpected insights into how these ecosystems operate across a variety of contexts (e.g. Bellwood et al. 2006; McWilliam et al. 2018; Brandl et al. 2019b; Tebbett et al. 2020; Schiettekatte et al. 2022a).

Overall, this special issue has built on the work of pioneering reef researchers (e.g. Odum and Odum 1955; Bardach 1961; Randall 1961; Smith and Kinsey 1976; Hamner et al. 1988) and helped showcase the potential for functional studies to contribute to our understanding

of coral reefs. Studies of reef functions, grounded in empirical data, and at appropriate scales, will be pivotal in critically appraising our perceptions and approaches to reef science, management, conservation, and governance in the Anthropocene. Ultimately, robust, independent, scientific investigation will be key in helping us navigate reefs through the current coral reef crisis.

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Declarations

Conflict of interests On behalf of all authors, the corresponding author states that there is no conflict of interest.

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