Kelp Forest Restoration in Australia

Cayne Layton1*, Melinda A. Coleman2,3, Ezequiel M. Marzinelli4,5,6, Peter D. Steinberg5,6,7, Stephen E. Swearer8, Adriana Vergés5,7, Thomas Wernberg9 and Craig R. Johnson1

1 Institute for Marine and Antarctic Studies, University of Tasmania, Hobart, TAS, Australia, 2 Department Primary Industries, NSW Fisheries, Coffs Harbour, NSW, Australia, 3 National Marine Science Centre, Southern Cross University, Coffs Harbour, NSW, Australia, 4 The University of Sydney, School of Life and Environmental Sciences, Coastal and Marine Ecosystems, Sydney, NSW, Australia, 5 Sydney Institute of Marine Science, Mosman, NSW, Australia, 6 Singapore Centre for Environmental Life Sciences Engineering, Nanyang Technological University, Singapore, Singapore, 7 Centre for Marine Science and Innovation, School of Biological Earth and Environmental Sciences, The University of New South Wales, Sydney, NSW, Australia, 8 National Centre for Coasts and Climate, School of Biosciences, The University of Melbourne, Melbourne, VIC, Australia, 9 UWA Oceans Institute and School of Biological Sciences, The University of Western Australia, Crawley, WA, Australia

Kelp forests dominate the rocky coasts of temperate Australia and are the foundation of the Great Southern Reef. Much like terrestrial forests, these marine forests create complex habitat for diverse communities of flora and fauna. Kelp forests also support coastal food-webs and valuable fisheries and provide a suite of additional ecosystem services. In many regions of Australia and around the world, kelp forests are in decline due to ocean warming, overgrazing, and pollution. One potential tool in the conservation and management of these important ecosystems is habitat restoration, the science and practice of which is currently undergoing substantial expansion. We summarize the present state of Australian kelp forests and emphasize that consideration of the initial drivers of kelp decline is a critical first step in restoration. With a focus on Australian examples, we review methods, implementation and outcomes of kelp forest restoration, and discuss suitable measures of success and the estimated costs of restoration activities. We propose a workflow and decision system for kelp forest restoration that identifies alternative pathways for implementation and acknowledges that under some circumstances restoration at scale is not possible or feasible. As a case study, we then apply the Society for Ecological Restoration’s 5-star evaluation to Operation Crayweed, Australia’s primary example of kelp forest restoration. Overall, no single method of kelp forest restoration is suitable for all situations, but outcomes can be optimized by ameliorating the driver(s) of kelp decline and achieving ongoing natural recruitment of kelp. Whilst scalability of kelp forest restoration to the seascape-scale remains a considerable challenge, the present review should provide a platform for future restoration efforts. However, it is also crucial to emphasize that the challenges of restoration place a high value on preventative conservation and protection of existing kelp forest ecosystems – prevention is invariably better than cure.

Keywords: canopy, Great Southern Reef, ecosystem, habitat-forming, macroalga, rehabilitation
THE ROLE OF KELP FORESTS

Kelp\(^1\) dominate rocky coastal environments in temperate and subpolar latitudes around the world (Smale et al., 2013; Krumhansl et al., 2016; Wernberg et al., 2019). These habitat-forming macroalgae occur in intertidal and subtidal habitats and range in size from less than a meter to over 40 m in length. Much like terrestrial forests, kelp forests are complex three-dimensional habitats with modified sub-canopy conditions (Gaylord et al., 2007; Layton et al., 2019) that support diverse communities of associated flora and fauna (Steneck and Johnson, 2014; Teagle et al., 2017; Miller et al., 2018). Kelp also act as the trophic foundation of coastal food-webs by providing food for a suite of grazers, detritivores, and microbes (Schiel and Foster, 2015; Wernberg et al., 2019) – the effects of which can reach to adjacent reef, seagrass, and sediment communities (Bustamente et al., 1995; Bishop et al., 2010), as well as to deep waters and beyond the continental shelf (Harrold et al., 1998; Thompson et al., 2011; Filbee-Dexter et al., 2018).

Most shallow (<30–50 m) rocky reefs in temperate Australia are dominated by kelp (Table 1; Marzinelli et al., 2015; Bennett et al., 2016; Coleman and Wernberg, 2017; Wernberg et al., 2019). Altogether, these kelp-dominated rocky reefs form an ~8,000 km long interconnected system known as the Great Southern Reef (GSR, Bennett et al., 2016), and sustain high levels of biodiversity and productivity (Ling, 2008; Bennett et al., 2016; Wernberg et al., 2019). One remarkable feature of biodiversity on the GSR is the high levels of endemism, and this is particularly true for macroalgae, with the GSR a global hotspot of macroalgal biodiversity and endemism (Womersley, 1987, 1994; Phillips, 2001; Kerswell, 2006). The GSR is also a global biodiversity hotspot for bryozoans, chordates, crustaceans, echinoderms, molluscs and sponges, with rates of endemism across these taxa ranging from ~20–60% (Bennett et al., 2016).

Australian kelp forests also have high economic value and support many fisheries, including the rock lobster and abalone fisheries that contribute >US$600 million p.a. to the national economy (ABARES, 2019). Beyond direct economic outputs, a lack of data makes it difficult to quantify the full value of ecosystem services provided by kelp forests in Australia. Notably, indirect commercial and social benefits arising from kelp forests are likely to be substantial, especially in coastal communities. These include indirect effects on fisheries (e.g., influence on coastal food-webs and prey species), recreational fishing, ecotourism, and other forms of marine recreation (e.g., scuba-diving) (Bennett et al., 2016). Despite their significant value, Australia temperate marine ecosystems are conspicuously underfunded and understudied relative to their tropical counterparts (Bennett et al., 2016).

STATUS OF AUSTRALIAN KELP FORESTS AND DRIVERS OF DECLINE

A strong rationale for considering restoration of Australia kelp forests is that they are in decline in many regions globally. In Australia, significant declines of kelp have occurred in Western Australia (Smale and Wernberg, 2013; Wernberg et al., 2016), South Australia (Connell et al., 2008), Tasmania (Ling, 2008; Johnson et al., 2011), Victoria (Kriegisch et al., 2016; Carnell and Keough, 2019), New South Wales (Andrew and O’Neill, 2000; Coleman et al., 2008; Vergés et al., 2016), and Southern Queensland (Phillips and Blackshaw, 2011). In these areas, and in many locations globally, drivers of kelp forest decline include both physical and biological factors (also see Krumhansl et al., 2016), and these must be understood for restoration efforts to be effective.

In Western Australia, an extreme marine heatwave over the 2010/2011 summer, in combination with southward range extension of subtropical herbivorous fishes associated with ocean warming, resulted in the loss of Ecklonia radiata (the dominant kelp across the GSR) and Scytothalia dorycarpa forests from ~100 km of coastline between Kalbarri and Geraldton (Smale and Wernberg, 2013; Wernberg et al., 2016). In South Australia, kelp forest losses have been mostly attributed to urbanization and increased runoff of sediments and nutrients (Connell et al., 2008; Gorman and Connell, 2009). Consequently, kelp forests within ~25 km of Adelaide, consisting mostly of E. radiata, have been largely replaced by less complex and less productive turf algae habitats (Figure 1). There has also been widespread loss of E. radiata in Port Phillip Bay, Victoria, particularly along the western and northern coastlines near the metropolitan areas of Geelong and Melbourne (Kriegisch et al., 2016; Carnell and Keough, 2019). Here, overgrazing by Heliocidaris erythrogramma urchins is the primary cause of kelp destruction, but subsequent proliferation of turf algae, in part due to high nutrient levels, also acts to inhibit kelp recruitment and recovery (Kriegisch et al., 2016; Reeves et al., 2018).

Destructive grazing by urchins is also a significant driver of kelp forest loss across Tasmania and the Bass Strait (Johnson et al., 2005, 2011; Ling, 2008), where urchin barrens formed by Centrostephanus rodgersii are now extensive and have replaced formerly lush kelp forests (Ling and Keane, 2018). This urchin, previously only abundant along the New South Wales coast, has undergone southern range extension over the last several decades due to increasing poleward penetration of the East Australia Current (EAC, Johnson et al., 2005, 2011; Ling, 2008). While warming waters and a strengthened EAC are responsible for the incursion of the urchin into southern waters, their local proliferation is linked to ecological overfishing of large southern rock lobster (Jasus edwardsii), which are the primary predator of C. rodgersii urchins in Tasmania (Ling et al., 2009; Johnson et al., 2011). While overgrazing by urchins has mostly affected E. radiata kelp forests, Tasmania has also suffered extensive losses of giant kelp (Macrocystis pyrifera) forests (Johnson et al., 2011). The loss of these iconic underwater forests (Figure 2) is mostly attributed to the increasing influence of the warm, nutrient-poor
TABLE 1 | The primary genera/species of habitat-forming kelp in Australia. Unlike the northern hemisphere where kelp canopies primarily consist of “true” laminarian keps, canopy-forming species in Australia constitute both laminarian and fucalian algae, with a larger diversity of fucoids than laminarians.

| Species (order) | Common name | Australian distribution | Description | References |
|----------------|-------------|-------------------------|-------------|------------|
| Cystophora spp. (Fucales) | cystophora | Sheltered to exposed reefs, 0–48 m. | Grows to 4 m. A widespread and diverse genus found only in Australasia. Can be locally abundant and dominant. May rise vertically (due to air-filled floats) or lay across the substrata | Huisman (2000), Edgar (2008), and Coleman and Wernberg (2017) |
| Durvillaea potatorum and D. anathea (Fucales) | bull kelp | Exposed reef, 0–30 m. Cape Jaffa, SA, to Tathra, NSW, and around TAS | Grows to 8 m. Massive, thick, and leathery kelp that lie prostrate across the substrata. The dominant species around low-tide level on exposed coastlines. Recently revised into two distinct species | Huisman (2000), Edgar (2008), Coleman and Wernberg (2017), and Weber et al. (2017) |
| Ecklonia radiata (Laminariales) | common kelp | Moderately exposed reef, 0–60 m. | Grows to 1.5 m. Most widespread and abundant kelp in Australia, with a distribution that mirrors the extent of the GSR. Very often the dominant kelp on the reef. Has a long rigid “stipe” (i.e., stem) that holds the fronds above the substrata | Huisman (2000), Edgar (2008), and Wernberg et al. (2019) |
| Lessonia corrugata (Laminariales) | strapweed | Exposed reef, 0–20 m. Phillip Island, VIC, and around TAS | Grows to 1.5 m. Occasionally locally abundant and dominant, typically in shallower and more exposed locations than *E. radiata*. Typically lies across the substrata | Huisman (2000) and Edgar (2008) |
| Macrocystis pyrifera (Laminariales) | giant kelp, string kelp | Moderate to exposed reef, 0–28 m. Cape Jaffa, SA, to Walkerville, VIC, and around TAS | Grows taller than 35 m. Has air-filled floats and can form immense underwater forests, often with a floating surface-canopy. Can be locally abundant and dominant. Has a shorter ecotype (∼10 m, form angustifolia) that typically grows in shallower locations | Huisman (2000), Edgar (2008), and Schiel and Foster (2015) |
| Phyllospora comosa (Fucales) | crayweed | Moderate to exposed reef, 0–20 m. | Grows to 3 m. Among the most common and dominant kelp on shallow and exposed sections of coastline. Has air-filled floats and typically floats just above the substrata. Often forms a dense band shallower than the zone dominated by *E. radiata*. | Huisman (2000), Edgar (2008), and Coleman and Wernberg (2017) |
| Sargassum spp. (Fucales) | sargassum | Sheltered to exposed reefs, 0–48 m. | Grows to 1.5 m. A diverse genus with global distribution that occur throughout tropical and temperate Australia. Can be locally abundant and dominant. May rise vertically (due to air-filled floats) or lay across the substrata | Huisman (2000), Edgar (2008), and Coleman and Wernberg (2017) |
| Scytosphaera dorycarpa (Fucales) | western crayweed | Moderate to exposed reef, 0–44 m. | Grows to 2 m. Fulfills a similar role to *P. comosa*, especially in Western Australia. | Huisman (2000), Edgar (2008), and Coleman and Wernberg (2017) |
| Undaria pinnatifida (Laminariales) | Japanese kelp | Moderately exposed reef, 0–10 m. Port Phillip and Apollo Bays, VIC, and D’Entrecasteaux Channel to Coles Bay, TAS | Grows to 1 m. An introduced and invasive species. Occasionally locally common and dominant but highly seasonal, almost disappearing throughout summer and autumn. Has a short rigid “stipe” (i.e., stem) that holds the fronds above the substrata | Edgar (2008) and South et al. (2017) |

waters of the EAC in Tasmania, although urchin overgrazing is likely to have exacerbated the problem (also see Ling, 2008; Ling and Keane, 2018) and be impeding recovery in some areas. Overall, more than 95% of Tasmania’s surface canopy-forming giant kelp forests (which also occur to a lesser extent in parts of Victoria and South Australia) have been lost over recent decades, to be replaced by *E. radiata* forests or urchin barrens (Johnson et al., 2011; Ling and Keane, 2018). Consequently, in 2012 the giant kelp forests of southeast Australia became the first marine community listed as endangered under the Australian Federal Government Environment Protection and Biodiversity Conservation Act (Evans et al., 2017). There is still no Federal Recovery Plan prepared for this community.

The *C. rodgersii* urchin has also contributed to extensive barren formation across its historical range in New South Wales (Andrew, 1993; Andrew and O’Neill, 2000). Urchin barrens are estimated to extend across >50% of shallow rocky reef habitats along the central and southern coastlines of the state (Andrew and O’Neill, 2000), suggesting widespread losses of the two dominant kelp in the region, *E. radiata* and *Phyllospora comosa*. These urchin barrens likely formed over many decades – possibly due to overfishing of urchin predators such as eastern rock lobster (*Sagmariasus*...
losses of E. radiata (see below). Losses of now being restored under the aegis of Operation Crayweed wastewater infrastructure and water quality, these forests are 1980’s (Coleman et al., 2008). Following improvements in P. comosa in the local extinction of Mabin et al., 2013). Following improvements in warming (Andrew and O’Neill, 2000; Coleman et al., 2008; Mabin et al., 2013).

Urban development on the coasts of metropolitan Sydney, and in particular untreated sewage outfalls, were implicated in the local extinction of P. comosa forests throughout the 1980’s (Coleman et al., 2008). Following improvements in wastewater infrastructure and water quality, these forests are now being restored under the aegis of Operation Crayweed (see below). Losses of E. radiata forests have also occurred on islands off northern New South Wales, toward the northern limit of the species’ distribution, and have been attributed to warming waters and overgrazing by herbivorous subtropical fishes (Vergés et al., 2016). This poleward shift of sub/tropical species into temperate waters is referred to as tropicalization and is expected to increase in the future as oceans continue to warm (Vergés et al., 2016; Zarco-Perello et al., 2017). Indeed, it seems that increasing ocean temperatures – especially in southeast and southwest Australia, which represent global hotspots of ocean warming (Hobday and Pecl, 2014) – are likely to cause continued poleward range contractions of Australian kelp, to be replaced by smaller subtropical macroalgae (Wernberg et al., 2016; Coleman et al., 2017; Martínez et al., 2018) and substantially altering ecosystem functioning (Vergés et al., 2019).

**FIGURE 1** | Example of Ecklonia radiata kelp forests (left) replaced by turf algae (right). Putative drivers include increased nutrients, and inhibition of kelp recruitment by the turf algae and entrained sediments. Photos by TW.

**FIGURE 2** | Examples of healthy (left) and degraded (right) giant kelp (Macrocystis pyrifera) forests in Tasmania. Photos reproduced with permission of Matthew Ramaley (left) and Matthew Doggett (right).

**BENEFITS AND VALUES OF KELP FOREST RESTORATION**

Since kelp forests are the foundation of Australia’s rocky reef ecosystems, it follows that maintaining or increasing the abundance and health of kelp forests via restoration could lead to concomitant benefits for production, biodiversity, and fisheries. Work has demonstrated that following urchin removal, small areas of recovered E. radiata kelp forests can support similar communities to natural E. radiata forests (Ling, 2008). However, this is not always the case and “recovered” kelp-dominated communities can be dissimilar to the pre-loss state (Valentine and Johnson, 2003). On artificial reefs in Tasmania, transplanting of adult E. radiata encouraged recruitment of economically and ecologically valuable invertebrates and fishes (including Ostrea angasi oysters and J. edwardsii rock lobsters), and facilitated development of diverse assemblages of flora and fauna (Layton et al., 2019a; Shelamoff et al., 2019). Whilst restoration of P. comosa forests by Operation Crayweed has had cascading benefits on epifaunal community composition, lags in system dynamics mean that restored communities can require time (i.e., years) to match natural P. comosa forests (Marzinelli et al., 2016).

Coastal macroalgal beds – of which kelp are the largest component by biomass – have also been identified as potentially important sinks of marine carbon (so called blue carbon) (Krause-Jensen and Duarte, 2016; Macreadie et al., 2019; Queirós et al., 2019). Recent work suggests that a significant portion of kelp forest biomass can be transported to coastal sediments and the deep ocean (Filbee-Dexter et al., 2018; Queirós et al., 2019). Furthermore, since macroalgal beds comprise such massive biomass, even conservative estimates of carbon transport/storage suggest that sequestration by kelp-dominated macroalgae beds can be a considerable contributor to blue carbon sinks (Macreadie et al., 2019; Queirós et al., 2019).

High biomass and fast growth rates ensure kelp forests also have critical roles in coastal nutrient-cycling (Smale et al., 2013; Bennett et al., 2016) and have great potential to absorb nutrients for bioremediation. Integrated multi-trophic aquaculture is a rapidly emerging field that might be able to utilize kelp to absorb excess nutrients associated with shellfish or finfish aquaculture (Buschmann et al., 2017; Hadley et al., 2018). Other emerging technologies and investments are also positioning kelp and macroalgae as a cornerstone of future blue economy applications, including as food for human consumption, livestock feed, biofuel, nutraceuticals, and pharmaceuticals (Buschmann et al., 2017; Froehlich et al., 2019). These high-value products represent market opportunities and economic incentives to help fund restoration.
efforts and thus contribute to the “restoration economy” (BenDor et al., 2015).

Kelp forests can also modify local hydrography and bolster coastal defenses by dampening ocean swell and decreasing erosion (Lovás and Torum, 2001; Gaylord et al., 2007). This service should be given special consideration with regards to forecast increases in sea level and storm activity due to climate change (IPCC, 2014).

Kelp and associated macroalgae also play an important role in Indigenous Australian culture and tradition (reviewed by Thurstan et al., 2018). Uses include ceremonial activities, medicinal practices, clothing, food, shelter, and as domestic devices. Archival records of the use of bull kelp (i.e., *Durvillaea* spp.) are particularly numerous, and there is considerable contemporary use of this kelp by Indigenous practitioners in artistic and knowledge-sharing activities. Additionally, the culturing, outplanting and monitoring that large-scale kelp forest restoration efforts require, provides ideal opportunities for Indigenous employment, management and custodianship (also see McLeod et al., 2018), and to establish skills and knowledge that underpin macroalgae farming.

**KELP FOREST RESTORATION ATTEMPTS: SUCCESSES AND FAILURES**

The restoration of kelp forests globally has typically followed two broad strategies: assisted recovery and active restoration. Assisted recovery – where natural kelp recovery is facilitated by either the removal of the agent of decline (e.g., culling of sea urchins, Ling, 2008; House et al., 2018) or the installation of substrata for kelp colonization (e.g., artificial reefs, Carter et al., 1985; Ambrose, 1994; Terawaki et al., 2001) – has been successful at increasing kelp recruitment and development of a kelp canopy over the short-term. Results are nonetheless highly variable and site-dependent, and projects involving removal of the agents of decline have seemingly had greater success than those that only provide novel substratum. This may be due to the unsuitability of some artificial substrata for kelp colonization, and/or effects of other colonizing organisms (e.g., filamentous turf algae, mussel, and barnacles) on kelp recruitment. Critically, assisted recovery approaches are often hindered by resource constraints and by hysteresis and feedbacks in the ecological dynamic (see Scheffer et al., 2001; Marzloff et al., 2011), which impair kelp recruitment and reestablishment even after the initial driver of decline has been ameliorated (Gorman and Connell, 2009; Johnson et al., 2017). We are aware of only one example where assisted recovery in isolation has resulted in the long-term restoration of kelp forests (giant kelp in California, see Reed et al., 2006, 2017).

Active restoration efforts have typically had greater success, and typically involve transplanting adult or juvenile kelp from a donor site, or outplanting lab-cultured kelp (North, 1976; Hernández-Carmona et al., 2000; Perkol-Finkel et al., 2012; Zarcó-Pérello et al., 2017; Verdura et al., 2018). The long-term success of this approach is reliant on either continued transplantation of kelp – which can be cost-prohibitive and dependent on a healthy donor population (North, 1976; Devinny and Leventhal, 1979) – or adequate natural recruitment of juvenile kelp. In the latter case, recruitment could occur from nearby populations of kelp and/or from the transplanted kelp itself (see Operation Crayweed below). Notably, the planting of juvenile kelp (whether lab-cultured or otherwise) has had little success (but see Perkol-Finkel et al., 2012) unless it is combined with the outplanting of adult kelp (North, 1976; Devinny and Leventhal, 1979; Layton et al., 2019a). This may be due to increased herbivory, competition, or abiotic stressors that cause mortality of juvenile kelp in the absence of adults (Hernández-Carmona et al., 2000; Konar and Estes, 2003; Vergès et al., 2016; Layton et al., 2019a).

Within Australia there have been few attempts to restore kelp forests. The earliest reported work comes from the Seacare community group (Sanderson, 2003) who attempted to restore areas of giant kelp (*M. pyrifera*) in Tasmania. Multiple techniques were used, including transplanting juvenile kelp from donor populations; transplanting artificial substrata on which juvenile giant kelp were growing following natural recruitment; transplanting sporophylls (i.e., the reproductive fronds of giant kelp); and outplanting ropes seeded with small (~5 mm) lab-cultivated juvenile kelp. *Centrostephanus rodgersii* urchins were also removed at some restoration sites and improved the likelihood of positive outcomes. Nonetheless, the project realized only marginal success and outcomes varied markedly across the 10 + sites. A single patch of giant kelp was established at one site but subsequently disappeared, in keeping with the ongoing decline of giant kelp in southeast Australia. The methods employed at the site of success did not differ from those at other sites (i.e., transplanting ~100 juvenile kelp and 3 fertile sporophylls), but it did have the most similar community composition to the donor site and was the most exposed and southerly location of the restoration sites.

Operation Crayweed is the only other reported example of targeted kelp forest restoration in Australia of which we are aware (although there are several projects currently underway, as discussed below). This ongoing project began in 2011 and aims to restore crayweed (*P. comosa*) forests to metropolitan Sydney where the species was once abundant (Coleman et al., 2008; Campbell et al., 2014; Marzinelli et al., 2014, 2016). Adult crayweed are transplanted from donor populations outside of metropolitan Sydney to restoration sites, with the primary aim to establish sufficient adult individuals to promote recruitment of juvenile crayweed. Despite high variability among sites, survival of transplanted crayweed is typically comparable to natural mortality (Campbell et al., 2014) and, as of 2019, transplanted crayweed has reproduced in six locations such that multiple generations are now identifiable, often hundreds of meters from the original restored patches. These restored crayweed forests have become self-sustaining without the need for additional cost or maintenance, which is a rare result in marine restoration. This relatively small-scale intervention has translated into a large-scale impact/benefit, with crayweed populations continuing to expand and colonize substantial areas, and beginning to function as natural forests (Marzinelli et al., 2016).

Additional work has employed aspects of active restoration and assisted recovery to improve understanding of kelp forests
and ecological restoration. Work by Valentine and Johnson (2005) found that even after the removal of urchins, heavy inoculation with *E. radiata* kelp spores was insufficient to achieve kelp reestablishment – presumably due to recruitment inhibition by the turf algae and sediments that had proliferated in the absence of the kelp (also see Layton et al., 2019b). However, Gorman and Connell (2009) showed that recovery of *E. radiata* kelp can naturally occur where turf algae are removed. Others have illustrated that urchin removal can facilitate natural recovery of kelp and other macroalgae on Australian temperate reefs when healthy kelp forests are nearby the denuded areas (Fletcher, 1987; Ling, 2008). Layton et al. (2019a) demonstrated successful transplanting of >500 adult *E. radiata* on artificial reefs in Tasmania. Survivorship of transplants was comparable to natural reefs, and abundant recruitment of juveniles (>750) ensured that many patches became self-sustaining. Crucially however, it was only patches above a certain size and density of adult kelp that facilitated adequate recruitment to maintain the kelp canopy; illustrating the importance of minimum patch sizes and densities when transplanting *E. radiata*, and likely other kelp species.

Given the rate of environmental change that is influencing coastlines worldwide, and that some drivers of kelp decline cannot be easily ameliorated (e.g., ocean warming), there is growing recognition of the need to plan adaptively and to “future-proof” marine restoration efforts, and potentially even consider the restoration of novel, more-suitable, species (van Oppen et al., 2015; Coleman and Goold, 2019; Wood et al., 2019). While we are aware of no published work implementing these strategies for kelp restoration, research in Australia is currently pioneering the identification of warm water-tolerant seaweed genotypes as the basis of future restoration efforts (IMAS, 2019; Gurgel et al., 2020).

**ESTIMATION OF THE COSTS OF RESTORATION**

Estimating the costs of implementing effective kelp forest restoration in Australia is difficult given the few examples to date. For Operation Crayweed, workers initially transplanted six 2 m$^2$ patches of *P. comosa* at each restoration site, with adult kelp densities of 15 m$^{-2}$. Initial transplanting efforts required ~5 days at each site and included site marking/preparation, securing of mesh mats for crayweed attachment, collection of adult crayweed from the donor population, and the transplanting itself. Costs of these efforts are estimated at ~US$6,850 per restoration site (i.e., ~US$570 m$^{-2}$), which cover a 4-person team, boat and tow-vehicle, SCUBA tank fills, basic equipment and consumables. Project management and monitoring of the multiple Operation Crayweed sites is estimated at an additional ~US$18,500 p.a. Note that these costs do not include the science necessary to underpin decisions such as choice of donor site, size of patch, etc. Active restoration efforts typically occur at small to medium scales, and not the seascapes at which kelp forest loss can occur. And so, while Operation Crayweed has demonstrated the translation of small-scale efforts into large-scale outcomes, efficient up-scaling of active restoration remains as a key ongoing consideration.

Assisted recovery techniques such as urchin culling are typically suited to tactical interventions at local spatial scales, such as reducing kelp loss to maintain/bolster the resilience of existing forests (Ling and Johnson, 2012; Layton et al., 2019a), remove incipient barrens (Ling, 2008; Tracey et al., 2015), or support active restoration efforts (Sanderson, 2003). Economic projections indicate that culling of *C. rodgersii* urchins from densities of 1.5 urchins/m$^2$ to 0.1 urchins/m$^2$ (i.e., the maximum density estimated to allow kelp recovery) across a 1 km$^2$ area of reef and from depths of 0–20 m would take two divers 685 days and cost ~US$980,478 or US$1,431 day$^{-1}$ (Tracey et al., 2014). These projections are nonetheless conservative given that urchin densities on barrens are typically closer to 2 urchins/m$^2$ (Ling, 2008; Ling and Johnson, 2012). Novel technology is promising to improve the scalability and cost-effectiveness of urchin culling, and trials of autonomous underwater vehicles designed to locate and kill urchins are in planning. Other alternatives to enhance the value or reduce the costs of urchin-culling operations may involve working in partnership with fisheries/aquaculture industries (Pert et al., 2018) or enlisting the help of citizen scientists, as has been done in Japan (Watanuki et al., 2010) and the United States (House et al., 2018).

Overall, the impetus to consider kelp forest restoration would benefit greatly from environmental accounting to ascertain the (currently unknown) value of kelp forests to human society, and which could underpin rigorous benefit-cost analysis (e.g., Rogers et al., 2018). This is especially since the costs of restoration operations can likely be reduced by minimizing diver labor and increasing automation/efficacy of mass seeding techniques (e.g., mass dispersal of lab-cultured kelp propagules from boat-mounted pumps; North, 1976).

**KELP FOREST RESTORATION: A WORKFLOW**

The loss of kelp forests in Australia, and indeed the world, is complex due to high levels of geographic variation and the multitude of different stressors present in any given location. Accordingly, it is useful to apply a workflow and decision framework when approaching restoration (Figure 3), especially where this can also provide an indicator of potential local outcomes. Development and preliminary-testing of this workflow utilized examples of kelp loss from Tasmania, where local restoration interventions must consider two species of dominant kelp (i.e., *Ecklonia radiata* and *Macrocystis pyrifera*) and multiple drivers of kelp forest decline (e.g., ocean-warming, overgrazing by urchins).

Our novel workflow and decision support system illustrates multiple alternative pathways and endpoints of restoration, and critically, helps to identify circumstances where restoration is not possible or advisable (also see Johnson et al., 2017). This diversity of pathways and endpoints exists because of environmental factors that are beyond the control of practitioners, such as whether hysteresis is present in the system (e.g., Gorman and Connell, 2009; Johnson et al., 2017). The multiple pathways of kelp forest restoration are also reflective of the diversity of drivers of decline, variability in the resources available for restoration
efforts, and scalability of the intervention. For example, the efficacy of assisted recovery techniques is often hindered by resource constraints (e.g., Tracey et al., 2015), while active restoration efforts can be reliant on ongoing transplantation of kelp, which has limited scalability.

The workflow highlights that at each node, research is needed to inform decision making and progression to the next stage, and thus may also help identify knowledge gaps in baseline ecosystem knowledge. Using this workflow to plan restoration efforts should also help practitioners and managers to ensure that interventions are considered within resource constraints and that the driver(s) of kelp forest decline has been addressed.

Progressing through the workflow toward the point of successful kelp restoration incorporates several key questions as decision points. Firstly, is it possible to return the environment to its pre-loss state, such as the improvements in water quality that preceded Operation Crayweed? If not, it will be essential to select and facilitate kelp to survive in the new environmental state, for example, the selection for thermally tolerant kelp from remaining healthy individuals (see IMAS 2019). However, if it is not possible to ameliorate or adapt to the novel ecosystem state, restoration efforts will be, at best, limited to the local scale (Figure 3). Secondly, is there hysteresis present in the dynamics of the system? Knowledge of the capacity for hysteresis is critical since it can prove one of the biggest challenges to kelp forest restoration (see Gorman and Connell, 2009; Marzloff et al., 2011; Johnson et al., 2017). Lastly, for successful restoration to occur at scale, efforts to overcome hysteresis and/or provide a novel source of propagules must also be scalable, and ideally commensurate with the scale of the initial degradation (Steinberg et al., 2016; Johnson et al., 2017; Wood et al., 2019).

Overall, ensuring natural recruitment of juvenile kelp and the continuation of self-sustaining generations is critical to long-term restoration success. Thus, it is important that kelp forest restoration focus on restoring the positive feedbacks that initiate recruitment and facilitation cascades.
and which promote ecosystem stability (Halpern et al., 2007; Layton et al., 2019a). To this end, restoration actions should also be considered as tools to bolster resilience in existing and/or partially degraded kelp forests where smaller interventions may be adequate to conserve/restore these feedbacks.

### MEASURING THE SUCCESS OF KELP FOREST RESTORATION

Since kelp are foundation species that support diverse ecological communities, comparison of community structure between restored and natural “reference ecosystems” can provide comparative indicators of restoration success at the community-level. However, in some instances reference ecosystems may not be identifiable or available, for example, when there are shifting baselines and poor understanding of ecosystem dynamics (Dayton et al., 1998; Johnson et al., 2017), or where ecosystem loss occurs prior to collection of adequate data (e.g., many Australian shellfish reefs, McLeod et al., 2018). Likewise, reference ecosystems may not be sensible targets for restoration when current/predicted environmental parameters (e.g., water temperature) are different to the historical state or community (Perring et al., 2015; Wood et al., 2019).

These diverse circumstances are acknowledged by the Society for Ecological Restoration (SER) who recognize that a reference ecosystem may instead be a conceptual model synthesized from numerous locations, indicators, and historical and predictive records (McDonald et al., 2016). SER have developed International Standards for the practice of ecological restoration, central to which is the “5-star recovery system” (see McDonald et al., 2016). This tool provides a consistent set of criteria against which key ecosystem attributes can be assessed, and acts as a conceptual framework for restoration practitioners, managers, and regulators. The 5-star system also provides a framework for habitat-specific indicators and metrics to be developed. Such indicators for kelp forest restoration might include transplant survival, growth rates, and condition (e.g., fouling, bleaching, and photosynthetic efficiency), genetic diversity, and recruitment. Certainly, recruitment of juvenile kelp is one of the greatest indicators of ongoing success and kelp forest resilience. In most cases, the ideal goal, as demonstrated by Operation Crayweed, is kelp forest recovery and reestablishment beyond the restoration footprint due to spill-over of natural recruitment.

### OPERATION CRAYWEED – A CASE STUDY OF THE SER 5-STAR RECOVERY SYSTEM

Operation Crayweed has been the most successful kelp restoration program in Australia to date. It has evolved from pilot restoration attempts in 2011 (2 sites) through to restoration at the scale of loss in 2018 (11 sites across ~70 km) with further expansion ongoing. It provides an ideal opportunity to apply the 5-star recovery system to a kelp restoration project, especially since science was used to rigorously design and test restoration approaches, thus allowing objective assessment of some recovery attributes. While the recovery wheel is site, scale, and temporally specific, here we apply it to the overall Operation Crayweed initiative (~9 years following initial restoration began) (Figure 4).

First, we know that some key aspects of ecosystem function (i.e., recruitment) in restored patches have reached levels found in natural areas (Campbell et al., 2014), warranting a 5-star rating for these attributes. Moreover, species composition of some associated organisms in restored crayweed forests is approaching that of natural forests (Marzinelli et al., 2016) warranting a rating of 4 for “habitat and interactions.” Additionally, ongoing monitoring illustrates that the structural diversity of restored crayweed populations are approaching natural levels (Marzinelli et al., 2016; Steinberg et al., 2016), warranting ratings of 3–4. Whilst the physical conditions that likely precipitated the initial decline (i.e., poor water quality, Campbell et al., 2014) have dramatically improved (following improvements to wastewater infrastructure), given the highly urbanized nature of the restoration locations in metropolitan Sydney there are likely to be ongoing threats from contamination; and therefore...
warrant a rating of 4 for “absence of threats” (e.g., invasive species and contamination) and “physical conditions” (e.g., water chemo-physical). Moreover, these external attributes cannot be controlled within the restoration framework, but are managed and regulated by government.

Targets for other ecosystem attributes are less well defined, have not been measured, or cannot be measured within the temporal scale defined here (due to prolonged response times). Thus, the recovery rating for some attributes is subjective and based on expert opinion of the Operation Crayweed team (MAC, EMM, PDS, and AV). For example, it is premature to assess “external exchanges” (e.g., gene flow) between restored and pre-existing crayweed populations because insufficient time has passed to restore those connections. The collective scientific knowledge of these systems suggests however that restoration of external connections is occurring, since crayweed cover and extent is expanding at some sites without additional intervention (hence the 2–3 star rating). While recovery across multiple trophic levels is being monitored, it is possible that recovery of higher trophic levels may take decades (e.g., Babcock et al., 2010; Reed et al., 2017), especially for species with limited dispersal (e.g., abalone; Marzinelli et al., 2014). The long timescales over which some attributes respond to restoration may therefore be reflected as a low rating, but can still be indicative of an appropriate trajectory of recovery.

The Society for Ecological Restoration considers the mean attribute score as an appropriate measure of a project's progress along a trajectory of recovery. However, depending on project and ecosystem-specific requirements, certain criteria could be weighted to provide a more subtle assessment. For Operation Crayweed, the mean attribute score was 3.7, indicating that restoration of crayweed forests at the local of the initial loss is well under way, with high levels of recruitment and good progress toward development of associated communities and ecosystem functions that are on a self-sustaining trajectory. While the mean score and individual recovery attributes could be used as a tool to identify areas requiring ongoing active restoration to accelerate recovery, we believe that it is sufficient to restore populations to a point whereby they can continue on a recovery trajectory naturally, without additional intervention. This is not only cost effective but ensures that limited resources can be strategically directed to maximize restoration efforts across species and ecosystems in need. Regardless, recovery of restored crayweed populations will continue to be measured and assessment of the need for additional interventions will be re-examined on an ongoing basis. The SER recovery system provides assessment of the trajectory of ecosystem recovery but also identifies research gaps. For Operation Crayweed, areas that would benefit from future research include assessing whether crayweed restoration has resulted in recovery of broader ecosystem functions and services (e.g., productivity and recreational fishing) and whether there is increased connectivity among adjacent non-kelp forest habitats (e.g., seagrass).

CONCLUSION

While kelp forest restoration is not achievable or feasible in all situations, use of a decision framework and consideration of the initial drivers of decline should increase the likelihood of success and the appropriate use of resources. And while past examples of kelp forest restoration are as notable for the failures as the successes, it seems that under many circumstances, small to medium scale restoration is achievable. For example, Operation Crayweed has demonstrated positive ecological outcomes at a scale beyond their initial restoration intervention. However, increasing the scalability of kelp forest restoration to the seascape-scale remains a considerable challenge, as does restoration in response to climate change where drivers of decline cannot be readily ameliorated (e.g., ocean warming). Optimal results will be achieved via thorough planning of restoration interventions and where positive feedbacks in the dynamics of kelp forests can be harnessed to promote habitat resilience and recruitment of juvenile kelp.

It follows however that the challenges and costs inherent in restoring kelp forests ecosystems places a great emphasis on the importance of maintaining and conserving kelp habitats. The difficulty of restoring heavily degraded environments – such as expansive urchin barrens and turf algae habitats – reinforces the notion that “an ounce of prevention is worth more than a ton of cure.” Ultimately, the management and conservation of Australian kelp forests should adopt policies that facilitate early warning and intervention for kelp environments under threat, with the aim to maintain and restore resilience in these critically important habitats.

AUTHOR CONTRIBUTIONS

CL and CJ conceived the initial idea for the review. CL prepared the manuscript with input from all other authors. All authors contributed to manuscript revision.

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Author/s:
Layton, C; Coleman, MA; Marzinelli, EM; Steinberg, PD; Swearer, SE; Verges, A; Wernberg, T; Johnson, CR

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