Self-replacement and community modification by the southern bull kelp Durvillaea antarctica

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ABSTRACT: Stands of the southern bull kelp Durvillaea antarctica (Chamisso) Hariot provide considerable biomass and a major habitat in the lower intertidal zone of exposed shores on the austral land masses. Whiplash effects of adult fronds (up to 10 m long) can affect recruitment, growth and survival of understorey species and potentially large brown algal competitors, thereby affecting community development. In southern New Zealand, D. antarctica is one of several species of large brown algae inhabiting the low intertidal zone. Effects of its canopy and its associated understorey coralline algae on community development were tested at 2 sites (Moeraki and Kaikoura) at different times of year between February 1999 and October 2001. Removal of D. antarctica canopies had surprising results compared to most studies where canopies of large brown algae were removed. The greatest initial recruitment of bull kelp occurred beneath intact canopies, usually in areas where corallines were removed. Recruitment was highly variable through time, with peaks occurring in June and October (austral winter–spring), depending mostly on when canopies were removed. There was an order of magnitude difference in recruitment between sites. The cover of turfing coralline algae, however, increased in all canopy removal treatments. A major source of mortality of young recruits was grazing by the herbivorous fish Odax pullus. Its distinct grazing marks were seen on recruits, almost exclusively outside the canopy of bull kelp where 80% of recruits were grazed. We show that D. antarctica has the ability to recruit beneath adult canopies, but that survival and growth ultimately depend on the extent of canopies, underlaying benthic algae and escapes from grazing by herbivorous fish.

KEY WORDS: Durvillaea antarctica · Recruitment · Intertidal algae · Substratum · Canopy · Fish grazing

INTRODUCTION

Large brown algae dominate many marine rocky shores world-wide, providing biomass, food and bio-genic habitat that support most inshore species. Although many species of algae, particularly those of the order Fucales, are desiccation-resistant and occur throughout the intertidal zone, the large and dense stands usually begin along the intertidal–subtidal fringe (Stephenson & Stephenson 1972). Here, aerial exposure occurs only during the lowest of tides, but on high-energy shores, wave splash ameliorates this effect. As few of the species that occur at this fringe can survive either higher in the intertidal zone (cf. Schonbeck & Norton 1978) or lower in the subtidal zone (cf. Choat & Schiel 1982, Chapman 1995), this habitat is likely to be unique in the combination of processes that structure and maintain it. If this is the case, general models accounting for community structure (Menge et al. 1997) will require modification. Furthermore, a general understanding of structuring processes must include regional or global differences in the important taxa (Menge & Branch 2001).

One of the largest species of algae in the southern hemisphere occurs at the intertidal–subtidal fringe of exposed shores. The southern bull kelp Durvillaea antarctica (Chamisso) Hariot is abundant on most southern land masses at latitudes between 45 and

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while their biomass reaches up to 80 kg m–2 (Santelices 1990a). This species is the largest member of the order Fucales (Hurd 2000) and is surpassed in size only by a few of the largest laminarian algae. Fronds of this species can reach 10 m in length, passed in size only by a few of the largest laminarian species of the order Fucales (Hurd 2000) and is surpassed in size only by a few of the largest laminarian species of the order Fucales (Hurd 2000). 1980, Santelices 1990a). This species is the largest sub-antarctic islands (Hay & South 1979, Santelices et al. 1980, Santelices 1990a). For example, Edgar & Burton (2000) found 23 macro-invertebrate taxa associated with *D. antarctica* holdfasts on the subantarctic Heard Island. However, *D. antarctica* can also reduce or eliminate other species through the whiplash effects of its fronds (Santelices et al. 1980), leaving only a few species of tough foliose and coralline algae intact near canopies of *D. antarctica*.

Unlike Chile, however, where most of the potentially competing species are laminarian algae, particularly the tough *Lessonia nigrescens* (Santelices et al. 1980, Santelices & Ojeda 1984, Santelices 1990a), southern New Zealand has a rich flora of fucalean species as well as a wide variety of red algae in the lowest intertidal zone (Naylor 1953, Hay 1977, Schiel 1990, Nelson 1994, Schiel et al. 1995). *Durvillaea antarctica* occurs almost exclusively at exposed, outer coast sites, occasionally intruding into semi-exposed areas (such as behind headlands), but never into sheltered areas (Santelices 1990a, Taylor & Schiel 2003). There are usually few other foliose algae present, except hardy turfing coralline and other red algae, in the vicinity of adult bull kelp. Recruitment can be extensive. In southern New Zealand, Hay & South (1981) found that dense recruitment occurred in patches where plants were removed. This tended to be seasonal because in southern New Zealand, *D. antarctica* reproduces during austral autumn and winter (April to September) (Hay 1977) and greatest recruitment occurs during late winter–early spring.

Algal canopies can affect spore dispersal, light and nutrient supply to areas below, and the whiplash of fronds and the extensive areas occupied by holdfasts can pre-empt successful recruitment (Kennelly 1983, Dayton et al. 1984, Santelices & Ojeda 1984, Schiel & Foster 1986, Foster & Schiel 1987, Schiel 1988, 1990, Santelices 1990b, Connell 2003a). Consequently, large brown algae often recruit poorly beneath adult canopies compared to gaps outside them. For example, Santelices & Ojeda (1984) found that canopy effects and grazing combined to inhibit recruitment of *Lessonia nigrescens* Bory in central Chile. For multi-species assemblages, however, much of our understanding of the effects of large brown algae canopies comes from subtidal studies. For example, Dayton et al. (1984) followed demographic patterns of populations in southern California over a 10 yr period, removed canopies and seeded areas with sporogenic material of several algal species. They found a dominance hierarchy for light competition determined by adult canopy height, but a trade-off in the ability of higher canopies to withstand wave stresses. Of overriding importance, however, were the life-history constraints such as dispersal abilities and growth rates that determined the ability of each species to invade and persist under canopies of other algal species.

The relationship between recruitment of large brown algae and the corallines that occur beneath them is unclear. Coralline algae, a common feature of the understorey in lower intertidal and subtidal habitats, have been found both to inhibit the recruitment of some species and facilitate others (Connell 2003a,b). For example, Camus (1994) suggested that encrusting coralline algae reduced recruitment of *Lessonia nigrescens* in northern Chile by shedding epithallial cells. In other studies, turfing corallines facilitated recruitment of fucoid algae by providing suitable micro-habitat conditions (Brawley & Johnson 1991, 1993, Benedetti-Cecchi & Cinelli 1992).

Here, we seek to elucidate the structuring processes in one of the most dominant and extensive assemblages in southern New Zealand (Nelson 1994). We test the effects of *Durvillaea antarctica* canopies, understorey corallines and the influence of different timing of disturbances on algal community development. These are then discussed with reference to the prevailing understanding of these processes in other related assemblages.

**MATERIALS AND METHODS**

**Study sites.** Two algal-covered platforms on the central eastern coast of South Island of New Zealand were used. First Bay, located on the NE end of the Kaikoura site (42°25′S, 173°44′E), is a mudstone platform that extends 70 m from the land. At its eroded edge are small fragments of harder reef covered in *Durvillaea antarctica* (Hay 1979). Waves are rarely less than 1 m in height and oceanic swells hit the reef largely unimpeded. The other study site was a platform of volcanic rock on the northern end of Moeraki (45°11′S, 170°98′E, 400 km south of Kaikoura). It is around 200 m long and 60 m wide. It has large beds of *D. antarctica* on flat but broken reef along its seaward margin, which is exposed to oceanic swells. At both sites, *D. antarctica* occurs at the 0.1 to 0.25 m tidal level. Both sites are further described in Schiel & Taylor (1999).
The understorey at both sites is dominated by a complex of turfing and encrusting coralline algae, especially Haloptilum roseum (Lamarck) Garbary et Johansen, Jania micrarthrodia JV Lamouroux, Corallina officinalis Linnaeus and Lithothamnion spp., which together covered over 95% of the primary substrata at both sites (Table 1). Bare substratum was scarce (<5%), usually occurring in any abundance only where adult holdfasts had detached.

**Experimental design.** The intertidal–subtidal fringe of exposed shores is notoriously difficult to work in. It is only fully exposed on the lowest tides, but swell conditions can limit access and the ability to do some types of manipulative experiments. For example, we trialled several types of cages and fences to test grazing effects of gastropods, but all devices were pummelled and destroyed, usually within a month, by waves and the fronds of *Durvillaea antarctica*. Our experiments were, therefore, limited to canopy and substratum manipulations.

To test the effects of canopies and coralline algae on recruitment, 3 fully crossed factors were used: sites (n = 2, treated as random), canopies (+/−; fixed) and coralline algae (+/−; fixed). Plots (30 × 30 cm) were marked under a canopy or in areas where canopies had been cleared, and randomly assigned as plus or minus coralline algae. Plots were marked with a plastic anchor plug. The coralline algae removal treatments were scraped back to bare rock using a tile hammer. To ensure that there were no ‘whiplash’ effects of the *Durvillaea antarctica* canopy in the clearance plots, canopies were removed for several metres away from adult plants so that none of the adult fronds were able to reach the clearance treatment plots. There were 4 replicates of each treatment.

We tested the effect of disturbance at different times, by initiating the full design 5 times: in summer (January 1999), autumn (May 1999), winter (August 1999), spring (October 1999) and the following summer (January 2000). As the design was pre-planned, we were able to randomise all treatments through time and as these experiments were placed into the lowest tidal zone on exposed shores, monitoring had to be performed opportunistically when swell and tide conditions permitted. This was done prior to the experiment and then 5 times between May 1999 and October 2001. The first 4 sampling times were 5 to 7 mo apart. The last sample (October 2001) was to determine the result of the matured experimental treatments.

Treatments were monitored using a 30 × 30 cm quadrat divided into 100 equally sized squares. Each treatment was visually assessed for the percentage cover of large brown algae, other foliose algae, encrusting coralline algae, turfing coralline algae, and bare rock. Invertebrate herbivores and large brown algal recruits were counted. In March and October 2001, the total length (base of holdfast to tip of blade) of each surviving *Durvillaea antarctica* recruit was measured. The number of recruits with lunate grazing marks was also recorded.

**Data analysis.** Analysis of variance models were used to test for treatment effects on the number and percentage cover of new *Durvillaea antarctica* recruits at 4 sampling periods. Analysis of variance models were also used to test for treatment effects on the percentage cover of turfing coralline algae, encrusting coralline algae and all other foliose algae at the end of the experiment in October 2001. The main factors were Site (random), Season cleared (summer 1998/99, autumn 1999, winter 1999, spring 1999, summer 1999/2000, fixed), Canopy (+/−, fixed) and Substratum (+/− coralline algae, fixed). Data were tested for homogeneity of variances using Cochran’s test prior to analysis. If significant, data were square-root-transformed, arcsine-transformed or log-transformed (log [n + 1]) prior to analysis. Where appropriate, Tukey HSD tests were used to locate differences among treatments following univariate analyses. Data were analysed in Statistica 5® (StatSoft).

### RESULTS

**Recruitment of *Durvillaea antarctica***

Recruitment was highly variable across sites, treatments and sampling times (Table 2, Fig. 1). The summer 1998/99 treatments had far greater recruitment in Moeraki than in Kaikoura through the first 3 samples (Fig. 1A,F), and for most sampling times, the magnitude of treatment effects differed between sites (Table 2). On the last sampling date (October 2001), the most recruits appeared in Kaikoura in the treatment where the canopy and substratum were removed. The treat-

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**Table 1. Percentage cover (mean, SE) and abundance data (mean per 0.09 m²) for dominant species and functional groups in all treatments at Moeraki and Kaikoura before treatment initiation. Data are averages from all clearance times.**

<table>
<thead>
<tr>
<th>Unit</th>
<th>At Day 0 Moeraki</th>
<th></th>
<th>Kaikoura</th>
</tr>
</thead>
<tbody>
<tr>
<td>% All turfing corallines</td>
<td>40.8 (6.10)</td>
<td>10.2 (1.73)</td>
<td></td>
</tr>
<tr>
<td>% Jania spp.</td>
<td>32.6 (4.88)</td>
<td>1.3 (0.24)</td>
<td></td>
</tr>
<tr>
<td>% Haloptilum roseum</td>
<td>6.5 (1.46)</td>
<td>6.9 (1.23)</td>
<td></td>
</tr>
<tr>
<td>% All encrusting corallines</td>
<td>57.0 (6.09)</td>
<td>85.1 (1.61)</td>
<td></td>
</tr>
<tr>
<td>% Other foliose algae</td>
<td>7.9 (1.80)</td>
<td>5.8 (1.34)</td>
<td></td>
</tr>
<tr>
<td>% Bare substratum</td>
<td>1.6 (0.34)</td>
<td>4.2 (0.72)</td>
<td></td>
</tr>
<tr>
<td>% <em>Durvillaea antarctica</em> can</td>
<td>100.0 (0.05)</td>
<td>99.8 (0.09)</td>
<td></td>
</tr>
<tr>
<td>% <em>Durvillaea antarctica</em> rec</td>
<td>2.2 (0.79)</td>
<td>1.5 (0.47)</td>
<td></td>
</tr>
<tr>
<td># <em>Durvillaea antarctica</em></td>
<td>2.1 (0.83)</td>
<td>1.1 (0.66)</td>
<td></td>
</tr>
<tr>
<td># Molluscan grazers</td>
<td>1.5 (0.42)</td>
<td>1.5 (0.29)</td>
<td></td>
</tr>
</tbody>
</table>
ments initiated in autumn 1999 also recruited greatly by the following spring (October 1999) and mostly at Moeraki. There were significant Site × Season of clearance and Site × Canopy interactions for most sampling periods, but these often depended on the season of clearance. Most recruits appeared in the +Canopy –Substratum treatments at Moeraki but in the –Can –Sub treatment in Kaikoura. As for the Summer 1998/99 treatments, about 6× as many recruits appeared at Moeraki than at Kaikoura. The treatments initiated in winter 1999 did not recruit well until the following winter (June 2000) (Fig. 1C,H). Again, most recruits appeared in Moeraki, in treatments where the canopy was intact but the underlying substratum algae were removed. Treatments initiated in spring 1999 had low-level recruitment the following winter at Moeraki and did not recruit until the following year in Kaikoura (Fig. 1D,I). The treatments initiated in summer 1999/2000 had recruitment during the following 2 yr but it was highly variable among all factors and there were no significant effects.

Overall, recruitment of *Durvillaea* occurred sooner at Moeraki than at Kaikoura. Almost invariably, the poorest recruitment occurred in Controls (+Can +Sub) and, overall, the greatest number of recruits appeared beneath an intact canopy where the substratum had been cleared.

**Total recruitment**

Over the entire experiment, the cumulative recruitment of *Durvillaea* gave an overall indication of treatment effects and showed significant Site × Canopy ($F_{1,120} = 21.47$, $p < 0.001$) and Site × Clearance time ($F_{4,120} = 2.51$, $p = 0.046$) interactions (Fig. 2). The greatest total recruitment occurred at Moeraki in the Summer 1998/99, autumn and winter clearances, mostly where the adult canopy remained intact. At Kaikoura, the greatest total recruitment was in the winter clearances where the canopy had been removed. Substratum alone also had a significant effect on recruitment ($F_{1,1} = 577.60$, $p = 0.03$); over all other treatments, there was an average of 73 (±12) *Durvillaea* recruits where substratum coralline algae were removed but only 17 (±2) where the substratum remained intact.

**Percent cover of *Durvillaea* recruits**

The percent cover of recruits within treatments was highly variable (Fig. 3) and was affected by the variation in plant length (Fig. 4B). It generally mirrored recruitment patterns at Moeraki but not at Kaikoura where grazing had a greater effect on plant cover. The cover was greater in treatments where the substratum algae were removed at Moeraki, regardless of canopy presence or absence (Fig. 3A–E). The cover of recruits at Kaikoura was more variable and was often initially greater in canopy removal treatments (Fig. 3F–J). In several cases, the cover of recruits in the –Can treatments peaked and then declined or did not increase (Fig. 3B,D,E,G). This was because recruits were often grazed, leaving only the holdfast and a short stipe. At the end of the experiment, the cover of *Durvillaea*

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**Table 2. *Durvillaea antarctica*. Summary of ANOVA of the total number of new recruits with Site, Season cleared (Sn), Canopy (Can) and Substratum (Sub) as factors. Significance levels: *p = 0.05; **p = 0.01; ***p = 0.001. Data were (a) log (n+1) or (b) square-root-transformed. Bold indicates significant effects. Cochran’s tests were not significant.**

<table>
<thead>
<tr>
<th>Source</th>
<th>October 1999a</th>
<th>June 2000a</th>
<th>March/April 2001a</th>
<th>October 2001b</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>df MS</td>
<td>F</td>
<td>df MS</td>
<td>F</td>
</tr>
<tr>
<td>Site</td>
<td>1 22.62</td>
<td>19.64***</td>
<td>1 43.91</td>
<td>85.57***</td>
</tr>
<tr>
<td>Season cleared</td>
<td>3 13.70</td>
<td>3.12</td>
<td>4 6.68</td>
<td>1.73</td>
</tr>
<tr>
<td>Canary</td>
<td>1 0.08</td>
<td>0.04</td>
<td>1 0.32</td>
<td>0.04</td>
</tr>
<tr>
<td>Substratum</td>
<td>1 22.11</td>
<td>7.81</td>
<td>1 8.25</td>
<td>0.83</td>
</tr>
<tr>
<td>Site × Sn</td>
<td>3 4.39</td>
<td>3.81*</td>
<td>4 3.85</td>
<td>7.51***</td>
</tr>
<tr>
<td>Site × Can</td>
<td>1 1.96</td>
<td>1.70</td>
<td>1 7.94</td>
<td>15.47***</td>
</tr>
<tr>
<td>Sn × Can</td>
<td>3 4.56</td>
<td>0.82</td>
<td>4 2.00</td>
<td>2.06</td>
</tr>
<tr>
<td>Site × Sub</td>
<td>1 2.83</td>
<td>2.46</td>
<td>1 9.88</td>
<td>19.25***</td>
</tr>
<tr>
<td>Sn × Sub</td>
<td>3 13.20</td>
<td>2.51</td>
<td>4 5.28</td>
<td>0.93</td>
</tr>
<tr>
<td>Can × Sub</td>
<td>1 6.37</td>
<td>0.81</td>
<td>1 14.23</td>
<td>1.93</td>
</tr>
<tr>
<td>Site × Sn × Can</td>
<td>3 5.59</td>
<td>4.85**</td>
<td>4 0.97</td>
<td>1.90</td>
</tr>
<tr>
<td>Site × Sn × Sub</td>
<td>3 5.26</td>
<td>4.56**</td>
<td>4 5.67</td>
<td>11.04***</td>
</tr>
<tr>
<td>Site × Can × Sub</td>
<td>1 7.88</td>
<td>6.84*</td>
<td>1 7.37</td>
<td>14.37***</td>
</tr>
<tr>
<td>Sn × Can × Sub</td>
<td>3 3.15</td>
<td>2.81</td>
<td>4 2.82</td>
<td>1.06</td>
</tr>
<tr>
<td>Site × Sn × Can × Sub</td>
<td>3 1.12</td>
<td>0.98</td>
<td>4 2.67</td>
<td>5.20***</td>
</tr>
<tr>
<td>Can × Sub</td>
<td>Error</td>
<td>96 1.15</td>
<td>120 0.51</td>
<td>120 1.03</td>
</tr>
</tbody>
</table>
Fig. 1. *Durvillaea antarctica*. Mean number of newly appearing recruits per 0.09 m$^2$ (± 1 SE) at each sampling time in the 5 initiation times in summer 1998 (A,F), autumn 1999 (B,G), winter 1999 (C,H), spring 1999 (D,I) and summer 1999 (E,J) at Moeraki and Kaikoura. Treatments are canopy intact/removed (+Can/–Can) and understorey coralline algae intact/removed (+Sub/–Sub).
recruits was minimal in the spring 1999 and summer 1999/2000 treatments, and resulted in a significant effect of season of clearance (Table 3). There was also a significant Site × Canopy interaction effect. There did not appear to be much difference within similar treatments across clearance dates at Moeraki, with the greatest cover in the +Can –Sub treatments. However, at Kaikoura the single treatment with high cover was –Can +Sub. A significant Site × Substratum interaction effect was due to greater cover in the –Sub treatments at Moeraki. Overall, the quickest recovery occurred in the autumn and spring clearances at Moeraki. Summer clearances either did not recover (Fig. 3E,F,J) or took around 2 yr to reach 60 to 90% cover.

When results at the end of the experiment were pooled across all clearance times, there were 2 significant treatment effects (Fig. 4A). The greatest % cover overall occurred in the clearances performed during winter, when *Durvillaea* was reproductive (*F*4,4 = 12.8, *p* = 0.015). These winter recruits covered up to 50% of the substratum in October 2001, while none of the other treatments had more than 30% cover of recruits. There was also a Site × Canopy interaction (F1,119 = 23.99, *p* < 0.001). Recruitment plants had a greater cover outside of adult canopies at Kaikoura but beneath canopies at Moeraki (Fig. 4A). The percent cover of recruits mirrored the patterns for plant length (Fig. 4B).

**Durvillaea lengths**

At the end of the experiment, there were 2 significant treatment interactions involving the lengths of *Durvillaea* recruits. The highest order effect was the interaction of clearance times, substratum and canopies (*F*4,4 = 6.68, *p* = 0.046). Over these factors, plants tended to be largest under adult canopies where substratum coralline algae had been initially removed (Fig. 5). Pooled data across sites showed that plants in the –Sub +Can treatments averaged from 300 to 380 mm in length in the summer 1998/99 and autumn clearances, while plants in the +Sub +Can treatments were longest in the spring clearances (Fig. 5). In canopy removal treatments, plants in the Autumn and winter treatments averaged between 100 and 320 mm. Overall, plants in the winter –Sub –Can treatments were significantly longer than the same treatments started in summer 1998/99, and summer 2000 treatments (Fig. 5; Tukey HSD, *p* < 0.05). There was also a significant Site × Canopy interaction (*F*1,119 = 27.98, *p* < 0.001). Plants at Kaikoura were longer outside of canopies, while those at Moeraki were longer beneath canopies (Fig. 4B; Tukey HSD, *p* < 0.05).

One of the greatest sources of mortality of developing *Durvillaea* recruits was grazing by what was later identified as the butterfish *Odax pullus*. Grazed plants had distinct lunate bites, which are characteristic of this herbivorous fish. Usually, most of the lamina was grazed, leaving only the stipe. Grazing was particularly evident where adult canopies had been removed (Fig. 4C). At Kaikoura, 60% of –Can recruits were grazed.
Fig. 3. *Durvillaea antarctica*. Mean percent cover (±1 SE) of recruits at each sampling time at the 5 initiation times in summer 1998 (A,F), autumn 1999 (B,G), winter 1999 (C,H), spring 1999 (D,I) and summer 1999 (E,J) at Moeraki and Kaikoura.
grazed but only 28% at Moeraki. However, the grazing effect was relatively stronger at Moeraki. Up to 28× more plants were grazed outside than under canopies at Moeraki, but only 6× as many in Kaikoura.

**Understorey foliose algae**

Only a few recruits of other fucalean algae (*Carposphyllum maschalocarpum*, *Cystophora torulosa*, *Hormosira banksii*, *Xiphophora chondrophylla*) recruited during the experiment, and none of them survived for more than a few months. However, there was considerable recruitment and great variation in the cover of foliose red, green and brown algae across clearance times, treatments and sites (Fig. 6). In most cases, these small, non-coraline species recruited within a few months, particularly where substratum clearances were performed and regardless of whether or not the *Durvillaea* canopy was intact. This was particularly pronounced at Moeraki after the first 3 initiation times (Fig. 6A–C). An exception was the autumn 1999 treatments at Kaikoura, which had poor recruitment of foliose algae (Fig. 6G). By the end of the experiment (October 2001), however, the substratum effects were no longer evident and the treatments outside of the intact *Durvillaea* canopy had the greatest cover of algae for 3 of the 5 treatment times (Table 4). Generally, few algae recruited into controls, where the canopy and substratum were left intact.

There were differences in the mixtures of species at the 2 sites. At Kaikoura, the most frequent species were brown algae *Halopteris congesta*, *Carpophyllum maschalocarpum* and the ephemeral *Scytosiphon lomentaria*, and the red alga *Gelidium caulacanthemum*. At Moeraki, there were occasionally recruits of *Macroystis pyrifera*, *Cystophora torulosa*, *Hormosira banksii* and *Halopteris congetsea*. In treatments where canopies were removed and the coraline algae were left intact on the substratum, the most common species were the red algae *Lophothamnion hirtum* and *Ballia callitricha*, and the brown *Glossophora kunthii*.

![Fig. 4. *Durvillaea antarctica*. (A) Mean percent cover, (B) total length and (C) percent grazed by the butterfish *Odax pullus* for recruits (±1 SE) in canopy intact treatments and canopy removal treatments at Moeraki and Kaikoura in October 2001](image-url1)

![Fig. 5. *Durvillaea antarctica*. Mean total length of recruits across +/- substratum coraline algae treatments in (A) canopy-intact treatments and (B) canopy-removal treatments within the 5 clearance times in October 2001](image-url2)
The cover of turfing coralline algae (Corallina officinalis, Jania spp., Halimeda roseum) generally recovered slowly after initial substratum clearances, although this tended to vary between sites (Fig. 7, Table 4). Moeraki had a greater initial cover of turfing corallines ranging from 30 to 60%, compared to <20% at Kaikoura. Where the canopy was cleared (–Can –Sub), the turfing corallines grew back within several months. In all cases at Moeraki, the greatest cover at the end of the experiment occurred in the treatments where Durvillaea canopies were removed but the coralline substratum was left intact. At Kaikoura, however, the results were far more variable. Generally, the recovery of turfing corallines did not depend on the presence or absence of an intact D. antarctica canopy, but several canopy removal treatments had increased cover of turfing corallines by the end of the experiment. By October 2001, the substratum treatments were still readily discerned at Moeraki but not at Kaikoura, where the overall cover was less.

**Encrusting coralline algae**

Encrusting coralline Lithothamnion spp. recovered slowly after the initial substratum clearances although, again, this varied between sites and clearance times (Fig. 8, Table 4). Generally, Kaikoura had a greater initial cover of encrusting coralline algae ranging from 70 to 90%, compared to 45 to 80% at Moeraki. In all clearances, the removal of the Durvillaea canopy resulted in a decrease in encrusting coralline algae (Fig. 7A–J). Encrusting corallines generally recovered best in the treatment where the Durvillaea canopy was left intact. Where the adult canopy was removed, there was an initial burn-off of the exposed corallines. The later decrease seen in many treatments was due to the expansion of turfing corallines. There was poor recovery in the treatments initiated in summer 1999/2000.

**DISCUSSION**

The ability of Durvillaea antarctica to recruit beneath adult canopies has important implications for its self-replacement capabilities and the structure of understorey communities. Few other large brown algae and no filter-feeding assemblages recruited into any of the treatments, despite spanning the reproductive and settlement periods of all species. The effects of underlying coralline algae were complex but over time, across treatments, there was a strong negative effect of these species on recruitment. Due to its relatively discrete reproductive period, the time of clearance affected the ability of D. antarctica to successfully re-capture space. Initial recruitment was greatest in winter treatments when D. antarctica is reproductive. These results have important implications for understanding the recovery of populations of this habitat-forming alga following natural disturbances and harvesting.
Fig. 6. Mean cover of all noncoralline algae per 0.09 m$^2$ (±1 SE) (includes all other foliose brown, red and green algae) at each sampling time at the 5 initiation times in summer 1998 (A,F), autumn 1999 (B,G), winter 1999 (C,H), spring 1999 (D,I) and summer 1999 (E,J) at Moeraki and Kaikoura.
Fig. 7. Mean percent cover of all turfing coralline algae (±1 SE) at each sampling time at the 5 initiation times in summer 1998 (A,F), autumn 1999 (B,G), winter 1999 (C,H), spring 1999 (D,I) and summer 1999 (E,J) at Moeraki and Kaikoura.
Fig. 8. Mean percent cover of encrusting coralline algae (±1 SE) at each sampling time at the 5 initiation times in summer 1998 (A,F), autumn 1999 (B,G), winter 1999 (C,H), spring 1999 (D,I) and summer 1999 (E,J) at Moeraki and Kaikoura.
Intra-specific interactions

The process of self-replacement varies between species of large brown algae (Chapman 1986, Schiel 1988, Santelices 1990b). A common feature of this ecologically important process is a tendency for greatest recruitment to occur immediately outside adult canopies. Surprisingly, we found that *Durvillaea antarctica* recruitment can occur directly under adult canopies, particularly if coralline algae were removed. However, there were differences between sites in the canopy and substratum effects. Between-site differences in substratum heterogeneity and rock type may have been responsible for these differences. The broken mudstone reef at Kaikoura may have provided more opportunities for recruits to survive outside adult canopies despite fish grazing effects.

That *Durvillaea antarctica* recruitment occurred under adult canopies is contrary to many studies of large brown algae but, in this high-energy environment, it may be an effective strategy for self-replacement outside usual recruitment times. In central Chile, recruitment has been stated to occur all year round (Santelices et al. 1980, Santelices 1990a,b). It is possible that these observations resulted from small fugitive recruits from beneath adult canopies remaining when canopies were removed by storms. As adult plants are frequently removed by storm events (Hay 1977, Santelices et al. 1980), the presence of small fugitive recruits under canopies could ensure immediate recovery of populations despite reproduction only occurring in discrete periods.

The overall poor growth and survival under canopies, however, suggests the habitat is suboptimal for recruitment. In the subtidal zone, factors such as shading, and possibly nutrient depletion, have been shown to inhibit intra- and interspecific recruitment under adult canopies (Kennelly 1983, 1987 Dayton et al. 1984, Reed & Foster 1984, Schiel 1988, Clark et al. 2004). However, the canopy effects on algal recruitment vary with species and some large brown algae are capable of recruiting into low-light, sub-canopy habitats (Clark et al. 2004). Schiel (1981), for example, found that the subtidal fucoid *Landsburgia quercifolia* (Hooker et Harvey) Harvey was able to recruit under adult canopies of *Ecklonia radiata* (C. Agardh) in NE New Zealand.

Rather than any particular facility to withstand low-light situations, the ability of *Durvillaea antarctica* to recruit under canopies is probably related to inter-holdfast distances and escapes from whiplash effects of adult canopies (cf. Santelices & Ojeda 1984). Sweeping and whiplash effects of adult canopies can greatly affect recruitment of large brown algae in the turbulent intertidal environment (Santelices & Ojeda 1984). In central Chile, Santelices & Ojeda (1984) showed that recruitment of the large brown alga *Lessonia nigrescens* was most likely to occur in an optimal canopy gap size large enough to reduce the sweeping and whiplash effects of adults, but small enough to prevent grazing by sea urchins. In the case of *Durvillaea antarctica*, the erect stipe of adults may elevate the blades enough to allow an area close to the holdfast where whiplash effects may not be so great.

Interspecific interactions

In general, recruitment of *Durvillaea antarctica* was greater in treatments where understorey coralline algae had been removed, regardless of canopy treatment. The relationship between recruitment of large perennial algae and coralline algae can be complex and variable. Encrusting coralline algae appear to inhibit recruitment of other algae through sloughing of epithallial cells (Johnson & Mann 1986, Vadas et al. 1992, Camus 1994). For example, Camus (1994) suggested that, through the shedding of epithallial cells, encrusting corallines reduced the recruitment of *Lessonia nigrescens* in central Chile. In another example, Worm & Chapman (1996) found that the recruitment of *Fucus evanescens* was inhibited by the crustose alga *Chondrus crispus* Stackhouse that prevented successful attachment of young plants. However, turfing coralline algae have been shown to facilitate the recruitment of several species of fucoid algae (Brawley & Johnson 1991, 1993, Benedetti-Cecchi & Cinelli 1992). For example, using agarose beads, Brawley & Johnson (1991) found that the microhabitat provided by turfing corallines provided protection from desiccation for zygotes of *Pelvetia fastigiata*.

Turfing coralline algae are often among the first species to colonise disturbed areas in lower intertidal zones (Hay 1981, Menge et al. 1993). At both Moeraki and Kaikoura, cover of turfing corallines increased once *Durvillaea antarctica* canopies were removed, probably due to a reduction in whiplash effects. Other studies have shown that macroalgal canopies can maintain and facilitate encrusting coralline habitat, while inhibiting turfing coralline algae (Connell 2003a,b). For example, in a recent study in south Australia, Melville & Connell (2001), using reciprocal transplants of turf and encrusting coralline boulders, found that *Ecklonia radiata* canopies, rather than simply co-occurring with encrusting coralline algae, actually facilitated its growth and survival by inhibiting turfing coralline algae.

Few other species of large brown algae recruited into canopy removal treatments and those that did, disappeared soon after recruitment of *Durvillaea antarc-
Recruitment of algae into disturbed intertidal areas is largely determined by clearance time, size of clearance, grazing, the dispersal and reproductive characteristics of the algal species and growth (Paine 1984, Sousa 1984, Schiel 1988, Farrell 1989, Menge et al. 1993, Kim & DeWreede 1996, McCook & Chapman 1997). We found that other species of foliose algae were more likely to recruit into treatments in which *D. antarctica* canopies had been removed and were initiated outside the winter reproductive period of *D. antarctica*.

The recruitment of other algae into canopy clearance areas was closely related to the proximity of other algal habitats and may be related to the dispersal abilities of surrounding species. The fucoid algae that surround *Durvillaea* habitats are relatively poor dispersers due to their relatively large egg sizes (Clayton 1990, 1992, Chapman 1995). Consequently, zygotes rarely settle more than a few meters from adult plants (D.I.T. unpubl. data). Moreover, the ability of fucoid species to stick quickly and remain attached in wave-exposed situations can vary between species. For example, Taylor & Schiel (2003) found that the fucoid algae *Cystophora torulosa* and *Hormosira banksii* took longer to attach and had poorer survival across wave exposures than *D. antarctica*.

At Moeraki, large beds of *Cystophora torulosa*, *Hormosira banksii* and *Xiphophora chondrophylla* on the periphery of the *Durvillaea antarctica* habitat resulted in the recruitment of small plants of these species into turf-covered canopy clearance treatments, which disappeared quickly as *D. antarctica* recruited. At Kaikoura, other species such as *Carposiphon maschalocarpum* and *Halopteris congesta* were more abundant on the edge of *D. antarctica* canopies, and a small number of plants recruited into the same treatments but eventually disappeared following recruitment of *D. antarctica*.

Ephemeral algae and diatomaceous films were the first species to colonise coralline removal treatments. In this study, no immediate effects of ephemeral algae on the recruitment of *Durvillaea antarctica* were found. Hay (1979) documented similar results in a limpet removal experiment in Kaikoura. He found *D. antarctica* quickly recruited into areas dominated by ephemeral algae if limpets were removed during the *D. antarctica* reproductive season, even though limpet removals were above the usual distribution of *D. antarctica*. These results are in contrast to those of several studies of algal succession in the higher intertidal zone where recruitment of fucoid algae is often suppressed by ephemeral algae, slowing succession (Lubchenco & Menge 1978, Underwood & Jernakoff 1981, Hawkins & Hartnoll 1983, Cubit 1984, Kim 1997).

For example, Kim (1997) found recruitment of *Fucus gardneri* Silva, in the upper intertidal zone in British Columbia, was slowed when limpet grazers were excluded and a cover of ephemeral algae dominated.

The effects of invertebrate grazers that inhabit *Durvillaea antarctica* holdfasts and geniculate coralline assemblages were not examined in this study. Molluscan grazers are abundant under *D. antarctica* holdfasts. For example, Edgar & Burton (2000) found 23 macro-invertebrate taxa, many of which were herbivorous limpets and chitons, under holdfasts on the subantarctic Heard Island off Australia. An alternative explanation for the lack of recruitment of other species of large brown algae into treatments could be that such grazers selectively removed these species. Furthermore, it is also possible that a release from grazing by the suites of micro-grazers that inhabit turffing coralline assemblages (Duffy & Hay 2000), might also explain the greater recruitment of *D. antarctica* into coralline removal treatments and possibly some of the between-site differences.

We observed that post-recruitment mortality of *Durvillaea antarctica* was greatly affected by the grazing effects of the herbivorous fish *Odax pullus*. In some cases, *D. antarctica* canopies appeared to afford some protection from fish grazing and possibly desiccation, but in most treatments, the great number of recruits ensured that 1 or 2 plants reached what appeared to be a size refuge from grazing.

Many *Durvillaea antarctica* recruits from outside adult canopies had some evidence of fish grazing (lunate bite marks along the margin of the lamina) and plants were regularly grazed back to stipes. This resulted, in most cases, in the eventual death of the recruits (D.I.T. pers. obs.). In some cases, plants regrew from the remaining blade on the margin of the fish bite marks. Hay (1977) also documented similar growth patterns of *Durvillaea* recruits from Otago and Kaikoura in southern New Zealand following butterfish grazing. In their example of harvesting effects, Hay & South (1981) showed that if fronds were pruned above the holdfast, plants recovered to only 32% of pre-harvest biomass after 15 mo. However, in central Chile, Santelices et al. (1980) found that *D. antarctica* lacked the ability to regenerate from storm-damaged stipes.

While there are several examples of invertebrate grazers controlling algal abundance in the lower tidal zones (Sousa et al. 1981, Choat & Schiel 1982, Andrew & Underwood 1989, Jones & Andrew 1990, Andrew 1993), there are very few examples of herbivorous fish controlling the abundance of intertidal algae in temperate climates. The majority of examples are from subtidal studies. For example, in the Mediterranean sublittoral, Sala & Boudouresque (1997) found that experimental reduction of fish grazing led to dramatic
changes in algal community structure, with significant increases in the abundance of fleshy erect algae. In *Ecklonia radiata* forests in New South Wales, Andrew & Jones (1990) describe the formation of patches by *Odax cyanomelas*. They found that, during spring, adult female *O. cyanomelas* cleared patches of adult *Ecklonia* near territorial males creating single age cohorts in patches. This is in contrast to the effects of *Odax pululus* in our study, where grazing occurred all year round during calm periods and mainly on recruit stages.

In general, examples of fishes controlling intertidal algal abundance are not common in the literature. However, Ojeda & Munoz (1999) provided an experimental example of fish grazing determining algal abundance in a temperate intertidal environment on the coast of central Chile. Their results show that grazing by the herbivorous fish *Scartichthys viridis* had significant effects on the algal community structure in the mid-intertidal zone by reducing the cover of the green alga *Ulva rigida* and the red alga *Gelidium chilense*. No examples of fish controlling intertidal brown algal abundance could be found in the literature.

**Summary**

Overall, *Durvillaea antarctica* appears to dominate these highly disturbed environments by arriving and recruiting in great numbers near adult canopies. We found that this is most likely to occur when free space occurs during winter and, because of winter storms, this is the time when free space is most readily available. The whiplash effects of *D. antarctica* canopies appears to modify the understorey community by suppressing understorey turfing coralline algae and excluding all other species of large brown algae.

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