PRESENCE OF SPOROPHYLLS IN FLOATING KELP RAFTS OF *MACROCYSTIS* SPP. (PHAEOPHYCEAE) ALONG THE CHILEAN PACIFIC COAST¹

Erasmo C. Macaya, Sebastián Boltaña, Iván A. Hinojosa, Juan E. Macchiavello, Nelson A. Valdivia, Nelson R. Vásquez

Facultad Ciencias del Mar, Universidad Católica del Norte, Larrondo 1281, Coquimbo, Chile

Alejandro H. Buschmann

Centro de Investigación y Desarrollo en Ambientes y Recursos Costeros (i~mar), Universidad de Los Lagos, Puerto Montt, Chile

Julio A. Vásquez, J. M. Vega Alonso, and Martin Thiel²

Facultad Ciencias del Mar, Universidad Católica del Norte, Larrondo 1281 Coquimbo, Chile, and Centro de Estudios Avanzados en Zonas Áridas (CEAZA), Coquimbo, Chile

Some species of macroalgae continue to live for extended periods of time after detachment and may even maintain reproductive structures, yet very little is known about this process. Here, we describe the presence of sporophylls (with sporogenous tissues) on floating kelp rafts of *Macrocystis* spp. along the coast of Chile. Surveys were conducted at nine sites (18-50° S) during austral summer 2002, and floating kelp rafts were seen and collected at seven of these nine sites (between 22 and 50 $^{\circ}$ S). Fifteen (26.8%) of the 56 samples had sporophylls, indicating maintenance of sporophylls after detachment. Some of the kelp sporophytes with reproductive blades showed signs of having been afloat for long periods (indicated by the large size of attached stalked barnacles). Additionally, experiments showed that floating kelps released viable zoospores. To understand the reproductive dynamics of floating kelps, we compared these results with information from attached populations of Macrocystis spp. at nearby coastal sites. In general, attached kelp had higher proportions of sporophylls than floating rafts, suggesting that detachment may negatively affect reproductive status. Nevertheless, floating kelps remained functionally reproductive, suggesting that zoospores may be dispersed via floating rafts. Published reports on other macroalgae indicate that some species (Lessoniaceae, Fucaceae, and Sargassaceae) are fertile and probably release zoospores or zygotes while floating or drifting in ocean currents. Because dispersal distances achieved by spores of most macroalgae are relatively short, release of spores from floating algae may be an alternative mechanism of long-distance dispersal.

Key index words: Chile; detachment, dispersal; floating; kelp rafts; *Macrocystis*; Pacific Coast; sporophylls

Benthic macroalgae may become detached from primary substrata as a consequence of different biotic and abiotic factors, for example, grazing by herbivores and storms (Tegner and Dayton 1991, Dayton et al. 1992). Some of these macroalgae, for example, the sporophytes of the giant kelp *Macrocystis* spp., possess special gas-filled structures or pneumatocysts that allow them to float freely on the sea surface after detachment (Hobday 2000a). These algae can continue growing and maintain positive buoyancy for weeks or months after detachment (Norton and Mathieson 1983, van den Hoek 1987). For example, Hobday (2000b) estimated that detached individuals of Macrocystis pyrifera (L.) C. Agardh remained afloat for 63 to 109 days. Detached macroalgae may even reproduce and release spores (Hoffman 1987, Santelices 1990), yet very little is known about the biology of benthic algae after detachment.

Medium-range dispersal in *Postelsia palmaeformis* Ruprecht may be increased by transportation of fertile reproductive tissue via currents (Dayton 1973). In *Sargassum muticum* (Yendo) Fensholt, branches may continue producing gametes after becoming detached from the parent alga, rendering these floating fertile parts an important means of dispersal for this invasive species (Norton 1977, Critchley et al. 1983). Similarly, the appearance of *Macrocystis* recruits and juveniles sporophytes in previously uncolonized areas has been suggested to result from fertile floating sporophytes inoculating an area (Dayton 1985, Edgar 1987). Moreover, floating kelp rafts of *Macrocystis* may facilitate the reestablishment of kelp assemblages after disturbances (Vega et al. 2005).

Anderson and North (1966), who studied dispersal of *Macrocystis pyrifera* by examining the distribution of small sporophytes in the vicinity of adult individuals, found maximum dispersal distances of 4 m. However, kelp dispersal may also occur over larger distances as suggested by Reed et al. (2004), who reported recruitment of *Macrocystis pyrifera* 3.5 km away from the nearest supply source. This long-distance dispersal may

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²Author for correspondence: e-mail thiel@ucn.cl.

occur during storms (Reed et al. 1988), when dislodged individuals and reproductive fragments may disperse, continuously releasing spores (Gaylord et al. 2004).

In the eastern Pacific Ocean, depending on the magnitude and intensity, "El Niño" can cause localized extinctions of kelps correlated with latitude (Camus 1994, Edwards 2004), and recovery after disturbances is often explained by several mechanisms such as spore dispersal (Reed et al. 1988, Santelices 1990) or seed banks of microscopical dormant stages (Ladah et al. 1999). Floating fertile fragments might also release spores, possibly representing an alternative dispersal mechanism for some benthic macroalgae.

Floating macroalgae have been reported from many areas of the world's oceans (Kingsford 1992, Helmuth et al. 1994, Ingólfsson 1998, Hobday 2000a, Smith 2002, Thiel and Gutow 2005a), but the presence of reproductive structures usually is not mentioned. Hooker (1847), in describing the circumantarctic distribution of *Macrocystis*, already marveled at the floating habits of this alga: "What then is the function of the floating mass of the plant?" he queried. However, even though he observed abundant kelp floating in the southern ocean, he did not mention whether they harbored reproductive blades. Here, we describe the reproductive status of floating individuals of giant kelp *Macrocystis* spp. along the coast of Chile, surveying the presence of sporophylls. We also conducted germination assays to test whether these floating algae indeed produce viable zoospores. To understand the reproductive dynamics of kelp after detachment, we compared our data on floating kelp with reproductive data from attached kelp populations at nearby coastal sites.

MATERIALS AND METHODS

Collection of floating kelp rafts and analysis of samples. During austral summer (January–March) 2002, floating kelp rafts were surveyed at nine locations (Fig. 1) on the coast of Chile along a latitudinal gradient from 18° S to 50° S (Table 1). Surveys were conducted from dawn to dusk, navigating from the coast to the west up to a distance of ~50 km from the coast, turning south for ~15 km before returning directly to the coast (Thiel et al. 2003). At sites where floating kelp were seen, we took samples with a special dip-net (ring diameter 60 cm, mesh size 3 mm in the upper part and 0.5 mm in the cod end) or by a diver who carefully separated small subsamples at random from large algal patches. All sporophytes were placed in plastic bags with 5% formalin and stored for later analysis (fertile sporophylls used in germination experiments were kept without formalin, see below).

In a molecular study, Coyer et al. (2001) suggested that *Macrocystis* is a monospecific genus, yet along the Chilean coast two species of *Macrocystis* are readily distinguished based on morphological characteristics, namely *M. integrifolia* Bory de



FIG. 1. Location of sampling sites of floating (gray arrows) and attached (black arrows) *Macrocystis* spp. from surveys conducted along the Chilean coast.

Site	Coordinates	Date	Number of samples (samples with holdfast)	Samples with sporophylls (samples with holdfast)	Percent reproductive samples	Total wet weight (kg)
Arica	18° 27′ S–70° 20′ W	February 2002	0	_	_	_
Iquique	20° 12′ S–70° 09′ W	February 2002	0	_	-	-
Mejillones	23° 04′ S-70° 30′ W	February 2002	6 (3)	0	0.0	4.58
Caldera	27° 04′ S–70° 64′ W	February 2002	7 (3)	0	0.0	1.91
Coquimbo	29° 57′ S–71° 22′ W	February 2002	10(3)	4 (3)	40.0	4.86
San [^] Antonio	33° 35′ S–71° 42′ W	March 2002	6(4)	2(1)	33.3	5.31
Concepción	36° 44′ S-73° 13′ W	March 2002	11 (5)	6 (5)	54.5	11.86
Ancud	41° 46′ S–73° 58′ W	March 2002	7 (3)	3 (2)	42.9	6.21
I. Madre de Dios	50° 26′ S–75° 22′ W	March 2002	9 (1)	Ò	0.0	3.74
Total			56 (22)	15 (11)	26.8	38.47

TABLE 1. Sampling sites of floating algae, number and percent of reproductive samples, and total wet weight of samples at each sampling site (coordinates represent starting points of surveys).

Saint-Vincent, with a flat holdfasts and narrow blades, and *M. pyrifera* with a high holdfast and wide blades. Sporophytes collected in the present study were morphologically representative of both species (holdfast and blade observations), but because the taxonomic situation of the genus *Macrocystis* has not yet been satisfactorily resolved, we herein treat all samples as *Macrocystis* spp. With the exception of the northern-most site where we collected floating kelps (Mejillones), samples were monospecific at all sampling sites, with samples from Caldera and Coquimbo corresponding morphologically to *M. integrifolia* and those from San Antonio and south to *M. pyrifera*.

In the laboratory, blades or individual sporophytes were carefully washed with fresh water. For each sample we counted the total number of vegetative blades and sporophylls (Reed 1987, North 1994). The presence of sporangia was confirmed by transverse sections of the blades and microscopical examination. For each sample the total wet weight (0.1 g) of vegetative blades and sporophylls was taken separately. Additionally, when samples contained stalked barnacles these were removed and counted, and the length of the scutum was measured for the 20 largest individuals from each sample. Stalked barnacles (Lepas spp.) were used, because 1) these animals are common on floating substrata but are not found on attached kelp individuals (Chen and Lewin 1976, Bernstein and Jung 1979, Helmuth et al. 1994), and 2) the body sizes of these barnacles serve as good proxies of minimum floating time (Thiel and Gutow 2005b).

Sampling of kelp from benthic populations and analysis of samples. The number of sporophylls and abundance of kelp sporophytes in benthic populations was evaluated using data collected during the summer periods of 2000–2004 (i.e. when oceanographic conditions were similar as in the summer 2002). Information on attached kelps was obtained from Iquique (January 2000), Mejillones (March 2003), Caldera (January 2002), Coquimbo (February 2002), Los Vilos (January 2004), Concepción (January 2002), and Pto. Montt (February 2002) (Fig. 1). Data on the proportion of sporophylls from benthic populations at the southern-most site, Bahía Porvenir (53° S), were obtained from the literature (Mansilla et al. 2003).

The abundance of *Macrocystis* spp. was estimated using several replicate counts of 1-m^2 quadrates along a transect perpendicular to the coast between the shoreline and 15 m depth covering the entire depth range of kelp beds. Sporophyte abundance was expressed as the number of sporophytes per square meter. Adult sporophytes were sampled at each site, and the total numbers of vegetative blades and sporophylls per individual were counted.

Viability of zoospores from floating kelps. During austral summer and fall (January and May 2004, January and April 2005), floating kelp rafts of *Macrocystis* spp. were collected in offshore waters off the Coquimbo coast (Fig. 1). Sporophylls were separated from these algae and transported to the laboratory in a cooler. Zoospores were obtained using the method of Reed et al. (1991). To evaluate the viability of zoospores (measured as number of zoospores), three replicate Petri dishees (5 cm, diameter and grid bottom) filled with 10 mL filtered seawater were inoculated with 2 mL spore solution and maintained for 3 days under the following conditions: $15 \pm 1^{\circ}$ C, $30 \pm 5 \,\mu$ mol photons · m⁻² · s⁻¹, 16:8-h light:dark cycle. Following this incubation period, zoospore viability was measured from the entire field of the microscope (n = 10 counts per Petri dish). The measurements were conducted with an inverted microscope ($400 \times$).

RESULTS

At the two northern-most sampling sites (Arica and Iquique), no floating kelp rafts were seen, but abundant floating kelp rafts were recorded at the other sampling sites. At each of those latter sites, between 6 and 11 samples were taken (Table 1). The size of the kelp rafts varied from small pieces (only stipes with some blades) to large patches of algae (several individuals including holdfasts) of more than 100 kg wet weight. The wet weight of individual samples varied from 0.034 to 5.64 kg in Coquimbo and Concepción, respectively (Table 1). Most samples with sporophylls also contained holdfasts (Table 1). Floating kelp rafts with sporophylls were only found at the four sampling sites between 30° S and 41° S. No sporophylls were found in floating algae from the northern sites (Mejillones and Caldera) or southern-most site (Isla Madre de Dios). The highest abundance of sporophylls was found in San Antonio (33° S), with one sample containing almost 70% reproductive blades (Fig. 2a). Similarly high values were observed for some samples in Concepción (36° S). In general, sporophylls were in good conditions, with the sporangial tissue covering a large area of the blade. The sporophylls often comprised a large proportion of the entire blade biomass in a sample, and the highest biomass of sporophylls was found in Concepción where in one sample almost 80% of the total blade biomass consisted of sporophylls (Fig. 2b).



FIG. 2. Proportion of sporophylls in individual samples from each site and size of stalked barnacles associated with these samples. (a) Number of sporophylls as a percentage of the total number of blades (vegetative and sporophylls) for each sample at each site; numbers on top of horizontal bars represent the total number of blades and number below bars represent number of samples collected at each site. (b) Sporophyll biomass as a percentage of the total wet weight of blades for each sample. (c) Mean scutum length (+1SD) of the 20 largest individuals of *Lepas pectinata* in samples with and without sporophylls. (d) Mean scutum length (+1SD) of the 20 largest individuals of *Lepas anatifera* (gray bars with star) in samples with and without sporophylls. NA, no algae were found at these sites.

Four species of stalked barnacles were found on the floating algae, namely *Lepas anatifera* Linnaeus, *L. pectinata* Spengler, *L. australis* Darwin, and *Dosima fascicularis* Ellis & Solander (the latter species only occurred with few individuals in two samples from Caldera and one from San Antonio). *Lepas anatifera* occurred exclusively in samples without sporophylls principally on kelp rafts from the northern sites



FIG. 3. Comparison of data from floating versus attached algae and abundance of algae along the Chilean Coast. Percent of sporophylls of total samples from each site in floating algae and percent of sporophylls per sporophyte in attached algae. Black lines in abundance data represent SE. NA, no algae were found at these sites.

(Fig. 2d), and the maximum sizes of these individuals was 5.5 ± 0.5 mm scutum length. Only two species occurred in samples with sporophylls: L. pectinata and L. australis (Fig. 2, c and d). Large individuals were found in the samples from three sites (San Antonio, Concepción, and Ancud), with maximum sizes in Concepción and Ancud for L. australis (8.1 \pm 1.4 mm and 12.4 ± 0.5 mm scutum length, respectively). The sizes of these stalked barnacles, which only colonize floating items, suggested that kelp had been floating for relatively long time periods. Interestingly, kelp rafts from San Antonio with high proportions of sporophylls contained no or only very small barnacles (Fig. 2). Similarly, in Ancud the sample with the highest proportion of sporophylls only contained relatively small barnacles (Fig. 2).

The abundance of attached *Macrocystis* spp. along the Chilean coast showed low values in Iquique (2.0 sporophytes \cdot m⁻²). Between 23° S and 33° S, kelp abundance reached values of 8.0–11.0 sporophytes \cdot m⁻², and the highest values (37 sporophytes \cdot m⁻²) were reached in Concepción (Fig. 3). In general, the percentage of sporophylls on attached sporophytes was higher than on floating kelps (Fig. 3). The highest values with 96.5% were reached in Concepción, where the mean number of sporophylls per sporophyte was 3.2 with only four blades (vegetative plus reproductive) in total (adult individuals had lost most of their vegetative biomass due to storm-induced loss of stipes). Only in San Antonio was the percentage of sporophylls on floating kelp higher than on attached kelps (64% versus 18%, respectively). In general, the percent of sporophylls on attached kelps of *Macrocystis* was at least 50%, and we never found algae without reproductive blades. In contrast, the floating algae from several sites had no sporophylls.

All examined sporophylls from floating kelp rafts collected in offshore waters off the coast of Coquimbo (30° S) released zoospores (Table 2). At the first three sampling dates, more than 60% of all zoospores germinated, whereas in April 2005 germination rates where slightly below 50%. The mean diameter of

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TABLE 2. Germination (%) of floating samples of *Macrocystis* collected in offshore waters off the Coquimbo coast.

Date	Percent germinated	Total of zoospores counted
January 2004	80.8%	6814
May 2004	66.4%	4445
January 2005	78.1%	4487
April 2005	46.1%	4265

zoospores was 11.4 μm (\pm 6.7 μm SD), and these were mobile and capable of settlement.

DISCUSSION

Our results suggest that kelp sporophytes maintained sporophylls while floating. Some of the floating kelps that we collected were very reproductive, as judged by the high number of sporophylls, even though they had been afloat for at least 2 weeks, as indicated by the fact that they were colonized by large individuals of stalked barnacles. These observations have important implications for the dispersal and recruitment capabilities of kelp species. Below we discuss the dispersal potential of *Macrocystis* spp. and provide literature examples to show that continuing reproductive activity may be common in detached macroalgae. Finally, we discuss the importance of this process as an alternative dispersal mechanism for some marine macroalgae.

Reproductive activity of floating and attached Macrocystis spp. No kelps were found floating at the northern-most study sites (Arica and Iquique), even though Macrocystis beds extend along the entire Chilean and southern Peruvian coast (Vásquez and Buschmann 1997). One possible reason for the absence of floating algae could be the low density of individuals in kelp forests from the northern-most sites. The interannual changes in abundance of kelp species at low latitudes (23° S to 20° S) in the southeast Pacific is regulated by "El Niño" (Camus 1994, Edwards 2004) with extreme cycles of distribution and kelp abundance that can last years or decades (Martinez et al. 2003, Vega et al. 2005). During 2002, Macrocystis populations in northern Chile were still recovering from the 1997-1998 El Niño disturbance (Vega et al. 2005). Recolonization processes after El Niño along the South American coast are slow (Martinez et al. 2003) and could explain the low sporophyte abundance at the northern sites compared with southern sites. However, even though density of Macrocystis spp. was low at the northern sites, the geographic extension of kelp beds between 23° S and 20° S was high in 2002 (unpublished results). Thus, lack of supply from benthic populations cannot alone be responsible for the total absence of floating kelp in Arica (18° S) and Iquique (20° S). Floating kelp rafts at the northern sites may disintegrate rapidly after detachment due to unfavorable conditions.

During the study period in summer 2002, surface water temperatures in offshore waters of Arica and Iquique were $>20^{\circ}$ C (unpublished data). Temperatures exceeding 20° C for several weeks are probably harmful to Macrocystis stands in nature (North 1994, Vega et al. 2005). In experiments with gametophytes, Peters and Breeman (1993) also found that these high temperatures are close to the survival limits of this kelp. Deysher and Dean (1986) suggested that a lack of sporophyte production may be caused by a combination of high temperatures and low nutrient concentrations. North et al. (1986) indicated that summer nutrient depletion, together with high temperatures, is responsible for canopy die-off in Macrocystis. Additionally, light conditions have been demonstrated to be important for recruitment of Macrocystis pyrifera (Graham 1996). Similarly, survival of floating algae at the sea surface depends on several factors, including temperature (Hobday 2000a), damage caused by sunlight (Jokiel 1980), nutrient levels (Edgar 1987), excessive epibiont growth (Parr 1939), and destruction by animals (Thiel and Gutow 2005a). Thus, unfavorable conditions at the sea surface and low densities in kelp beds after the 1997-1998 El Niño may be the main reasons that no floating kelps were found in Arica and Iquique.

Floating kelp rafts were present in Mejillones (23° S) and Caldera (27° S) , but these had no sporophylls (i.e. were not reproductive). Buschmann et al. (2004) showed that benthic populations of Macrocystis from the northern coast of Chile have their main reproductive activity during spring and winter but also had sporophylls during the rest of the year. Thus, reproductive seasonality cannot explain the complete absence of reproductive activity in floating kelp from these two northern sites. Reproduction in kelps is determined by environmental factors, such as light (irradiance, light quality, and photoperiod) and temperature (Lobban and Harrison 2000). Possibly, conditions at the sea surface in Mejillones and Caldera allowed a limited survival of floating kelp rafts but may have caused a rapid loss and absence of new production of sporophylls in floating algae due to light and nutrient stress. Reed (1987) found that experimental removal of M. pyrifera canopy biomass led to a significant decrease in sporophyll biomass. Observing that attached individuals completely stopped reproducing due to an episodic, sublethal, amphipod-grazing event that stripped blades from all sporophytes, Graham (2002) suggested a trade-off between sporophyte growth and reproduction in M. pyrifera. Amphipod grazers are common on floating kelp (Kingsford 1992) and might cause decreasing reproductive activity of floating rafts.

In contrast to what was found for the northern sites, floating kelp rafts from central Chile (33° S to 42° S) contained a comparatively high percentage of sporophylls, possibly because environmental stress at the sea surface is less severe than in northern Chile. Nevertheless, in central and southern Chile, reproductive activity of floating kelps was lower than that of attached

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Species	Region	Status	Method	Distance from source	Reproductive state	Abundance	References
Ascophyllum nodosum (L.) Le Jolis	Central E	Floating	ni	$5500\mathrm{km}$	Healthy mature	Large	John 1974
Codium fragile (Suringar) Hariot	NW Atlantic	Floating	Surveyed	Nearshore	conceptactues Gametangial	Occasional	Hubbard and
Ectocarpus fasciculatus Harvey	NE Atlantic	Drifting	ni ni	Nearshore	Individuals with	ni	Garbary 2002 Russell 1967
Ectocarpus siliculosus (Dillwyn)	NE Atlantic	Drifting	ni	Nearshore	sporangia Individuals with	ni	Russell 1967
Lyngoye Gelidium corneum (Hudson) J.V.	NE Atlantic	Drifting	Samples	Nearshore	sporangia Fertile thallus	ni	Seoane-Camba 1966
Lamouroux Himanthalia elongata (L.) S E C	NE Atlantic	Drifting	ni ni	Nearshore	Sexual stage	ni	Boetius 1952–53
o.r. Gray Macrocystis spp.	SE Pacific	Floating	Samples	Offshore	Fertile algae with	Locally	This study
Myagropsis myagroides	NW Pacific	Floating	Surveyed	Nearshore	sporopuyus Floating fertile algae	Common	Segawa et al. 1959
(Mertens ex Jurner) Fensholt Pilayella littoralis (L.) Kjellman	NE Atlantic	Drifting	from boats ni	Nearshore	Individuals with	ni	Russell 1967
Postelsia palmaeformis Ruprecht Sargassum fulvellum (Turner)	NE Pacific NW Pacific	Drifting Floating	ni Surveyed	Nearshore Nearshore	sporangia Fertile algae Floating fertile algae	Common Occasional	Dayton 1973 Segawa et al. 1959
C. Agardh S. fusiforme (Harvey)	NW Pacific	Floating	trom boats Surveyed	Nearshore	Floating fertile algae	Occasional	Ohno 1984
Setchell S. hystrix J. Agardh	SE Atlantic	Floating	Samples	Offshore	Floating fertile algae	Common	Oliveira et al. 1979
S. hemiphyllum (Turner)	NW Pacific	Floating	Surveyed	Nearshore	Floating fertile algae	Common	Ohno 1984
S. horneri (Turner) C. Agardh	NW Pacific	Floating	Surveyed	Nearshore	Floating fertile algae	Common	Ohno 1984
S. micracanthum (K.) Endlicher	NW Pacific	Floating	Surveyed	Nearshore	Floating fertile algae	Occasional	Ohno 1984
S. muticum (Yendo) Fensholt	ni	Floating	Artificially	ni	Floating fertile algae	Common	Norton 1977, Critchley
S. nipponicum Yendo	NW Pacific	Floating	Surveyed	Nearshore	Floating fertile algae	Common	Ohno 1984
S. patens C. Agardh	NW Pacific	Floating	Surveyed	Nearshore	Floating fertile algae	Common	Segawa et al. 1959,
S. piluliferum (Turner)	NW Pacific	Floating	Surveyed	Nearshore	Floating fertile algae	Common	Segawa et al. 1959, Obno 1084
S. <i>platycarpum</i> Montagne	SE Atlantic	Floating	Samples	Offshore	Floating fertile algae	Occasional	Oliveira et al. 1979
S. ringoldianum Harvey	NW Pacific	Floating	Surveyed	Nearshore	Floating fertile algae	Occasional	Segawa et al. 1959,
S. serratifolium (C. Agardh)	NW Pacific	Floating	Surveyed	Nearshore	Floating fertile algae	Common	Segawa et al. 1959,
S. siliquastrum (Turner)	NW Pacific	Floating	Surveyed	Nearshore	Floating fertile algae	Common	Segawa et al. 1959, Obage 1064
C. Agatun S. thunbergii Yendo	NW Pacific	Floating	Surveyed from boats	Nearshore	Floating fertile algae	Occasional	Ohno 1984

SPOROPHYLL PRESENCE IN FLOATING KELP

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algae, indicating that conditions at the sea surface are not optimal for reproductive activity of sporophytes. An alternative explanation for the low presence of sporophylls in floating kelp rafts could be due to breakage of the stipe shortly above the holdfast, resulting in floating stipes with primarily vegetative blades and attached holdfasts with remnants of sporophylls. The fact that some of the kelp holdfasts from Concepción (36° S) had suffered breakage supports this suggestion. On the other hand, we also found entire floating individuals (including holdfast) without any sporophylls, particularly in Mejillones and Caldera. In general, survival and continuing reproduction of floating kelp may only be possible in optimal conditions, that is, in the center of their geographic range.

Dispersal potential of Macrocystis spp. and other floating macroalgae. Dispersal processes depend on the characteristics of the macroalgae species as well as on the environment (Santelices 1990). The distance over which propagules are dispersed should be roughly dependent on the time spent in the water column (Shanks et al. 2003). In Macrocystis spp., dispersal via zoospores appears to occur over distances of only tens to hundreds of meters (Reed et al. 1988, Shanks et al. 2003). However, based on hydrodynamic modeling, Gaylord et al. (2002) suggested that dispersal of M. pyrifera may routinely extend over farther distances. We found several sporophytes of M. pyrifera with sporophylls floating at considerable distances off the Chilean coast. Many of these kelp rafts had been afloat for at least 2 weeks as indicated by the sizes of attached barnacles. The largest individuals of Lepas australis and L. pectinata we found to have a scutum length of >8 mm. Typical growth rates for these two species range between 0.22 and $0.32 \text{ mm} \cdot \text{d}^{-1}$ (Thiel and Gutow 2005b). Assuming similar growth rates for the stalked barnacles collected off the Chilean coast, it appears safe to assume that some of the samples found with sporophylls had been afloat for at least 21 days. During this time period and in the Humboldt Current system, these floating algae could have traveled distances of several 100 km (Thiel 2003).

Our study showed that sporophytes of *Macrocystis* spp. floating along the coast of Chile contained sporophylls, indicating continuous reproductive effort after detachment. Germination tests with floating kelps from Coquimbo (30° S) confirmed that sporophylls from detached sporophytes indeed released a high proportion of viable zoospores during the summer months, that is, when all samples of floating kelp were taken. The germination of zoospores from floating sporophytes observed was similar to results from Candia et al. (1979) and Palacios and Mansilla (2003), who showed normal development of zoospores from benthic populations of *M. pyrifera* under laboratory conditions.

In general, the presence of reproductive structures on detached algae appears to be a common phenomenon in species from the families Lessoniaceae, Fucaceae, and Sargassaceae (Table 3). These considerations in combination with the results of our study support previous suggestions, namely that long-distance dispersal of macroalgae may occur via floating individuals or fertile algal fragments, which are transported by prevailing currents (Deysher and Norton 1982, van den Hoek 1987). High abundances of floating macroalgae have been observed in many regions of the worlds oceans (Thiel and Gutow 2005a), often at great distances from the nearest shores (van den Hoek 1987). Because dispersal distances of most algal spores or zygotes are limited (Fredriksen et al. 1995, Forrest et al. 2000, Dudgeon et al. 2001), transport of sporophylls or reproductive conceptacles in floating kelp rafts may be an important dispersal mechanism for some macroalgal species that float after detachment. In the case of species from the families Fucaceae and Sargassaceae, zygotes may be released from monoecious individuals, facilitating long-distance dispersal. However, our results also suggest that this mechanism may be restricted to particular conditions that are close to the optimal growth environment of these algae. Future studies should focus on the factors affecting survival and continuing reproductive activity in floating algae after detachment.

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