

Non-structural carbohydrate reserves of eelgrass *Zostera marina*

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ABSTRACT: The high minimum light requirement of eelgrass *Zostera marina* L. suggests that this species has difficulty in maintaining a positive carbon balance except under high light conditions. The carbon balance of *Z. marina* can be studied by following seasonal changes in non-structural carbohydrate (NSC) reserves, however, little is known about the seasonal variation in NSC reserves in seagrasses or the influence of shading on NSC reserve content and distribution. Seasonal changes in eelgrass NSC reserves were measured in a shallow coastal lagoon, Chincoteague Bay, Maryland/Virginia, USA, near the southern edge of this species' distributional range. Concentrations of sugar varied seasonally in leaves, rhizomes and roots, with maximum concentrations occurring in the rhizomes. In contrast, starch concentrations did not vary with the season, but were highest in the roots. Seasonal peaks in rhizome NSC reserves parallel the spring and fall bimodal growth patterns observed for *Z. marina* in the region. Total NSC reserves change from an estimated 52 to 73 g m⁻² in June to 4 to 18 g m⁻² in January, or a decrease of 75 to 92%. Experimental shading for 3 wk in the spring reduced ($p < 0.001$) sugar but not starch concentrations in leaves (48%), rhizomes (40%) and roots (51%). In addition, shading reduced ($p < 0.05$) leaf biomass (34%), root and rhizome biomass (23%) and density (27%). Potential NSC reserve storage during shading was reduced by an estimated 66%. Spring appears to be an important time for both growth and storage of NSC reserves in *Z. marina*, and the NSC reserves are generally depleted throughout the remainder of the year. Turbidity during this springtime 'window of opportunity' may jeopardize subsequent survival as a result of inadequate NSC reserves to maintain a positive carbon balance during the rest of the year.

KEY WORDS: Eelgrass · *Zostera* · Seagrass · Starch · Sugar · Shading · Carbon balance · Biomass

INTRODUCTION

The southern distributional range of eelgrass *Zostera marina* L. appears to be limited by high summertime water temperatures in some areas (Dennison 1987). At the southern part of its range on the east coast of the United States (e.g. Chincoteague Bay, Maryland/Virginia), water temperatures reach 30°C in the summer. Because respiration exceeds photosynthesis at temper-

atures above 25°C (Marsh et al. 1986, Dennison 1987, Zimmerman et al. 1989), summer may be a time of carbon balance depletion. Results from a number of studies show that eelgrass has a high minimum light requirement relative to other plants (Backman & Barilotti 1976, Bulthuis 1983, Goldsborough & Kemp 1988) and the maximal depth limit for survival of eelgrass is strongly linked to light availability (Burkholder & Doheny 1968, Backman & Barilotti 1976, Dennison & Alberte 1982, 1985, 1986, Wetzel & Penhale 1983, Dennison 1987). Results from these and other studies (Felger et al. 1980, Kenworthy et al. 1982, Robertson & Mann 1984) suggest that high light levels are necessary for maintenance of a positive carbon (C) balance from year to year. We hypothesize that eelgrass populations on the Maryland/Virginia border are just main-

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taining a positive annual C balance and, therefore, that any additional reduction in carbon reserves may result in increased mortality among these populations.

To maintain a positive C balance from year to year, plants must allocate photosynthate appropriately among competing sinks including roots, rhizomes, leaves, reproductive material, and storage reserves. However, the survival of eelgrass populations at the southern edge of their distributional range may depend on whether their C allocation patterns are flexible enough to withstand any additional environmental stresses, such as shading events.

Eelgrass at this latitude capitalizes on optimum environmental conditions during spring for growth (Orth & Moore 1986). It is the high light availability and cool water during that season which allow high biomass production and, we hypothesize, large accumulations of non-structural carbohydrates (NSC). What is not known is whether these plants can store enough NSC reserves to maintain a positive annual C balance throughout the year if spring shading occurs. Certainly, these reserves are critical for maintenance during the subsequent summer, autumn and winter when temperature and light conditions limit photosynthesis and/or increase respiration rates. Declines in *Zostera marina* populations, attributed to increasing turbidity in some areas (Orth & Moore 1983), led to the hypothesis that *Z. marina* is not flexible enough to change its C allocation pattern to help it survive turbidity events affecting some areas in the southern part of its range. If this is the case, allocation of photosynthate to biomass production instead of storage could eventually result in a depleted C balance.

A potentially important indicator of the C balance in eelgrass is the concentration and distribution of NSC reserves. Seasonal NSC levels in eelgrass have seldom been measured (but see Harrison & Mann 1975, Zimmerman et al. 1989, 1995) However, the seasonal variation of NSC reserves has been examined for several other seagrasses: the NSC reserves in *Thalassia testudinum*, *Syringodium filiforme*, and *Halodule wrightii* were highest in the autumn (Dawes & Lawrence 1980) suggesting that those reserves were used to sustain plants through the winter. The influence of experimental shading on NSC reserves in naturally growing populations of *Zostera marina* has not been determined, although shading of *T. testudinum* was shown to reduce NSC reserves in all plant tissues (Tomasko & Dawes 1989).

We hypothesized that NSC levels in healthy eelgrass beds in Chincoteague Bay would peak in the spring and decline during the remainder of the year as the plants used those reserves to sustain themselves through the summer, autumn and winter. Further, we hypothesized that shading during the spring would lead to a reduction in NSC reserves compared to unshaded plants. We tested these hypotheses by measuring the seasonal variation in NSC concentrations and growth in a naturally growing seagrass bed and conducted a shading experiment in an adjacent part of the bed. Finally, we used *Zostera marina* biomass estimates to produce seasonal NSC budgets and to estimate the influence of spring shading on the annual carbon balance.

METHODS

The study was conducted during 1990 to 1992 in Chincoteague Bay (37°N, 75°W), a shallow coastal lagoon located between the mouths of the Delaware and Chesapeake Bays (Fig. 1). Chincoteague Bay has a

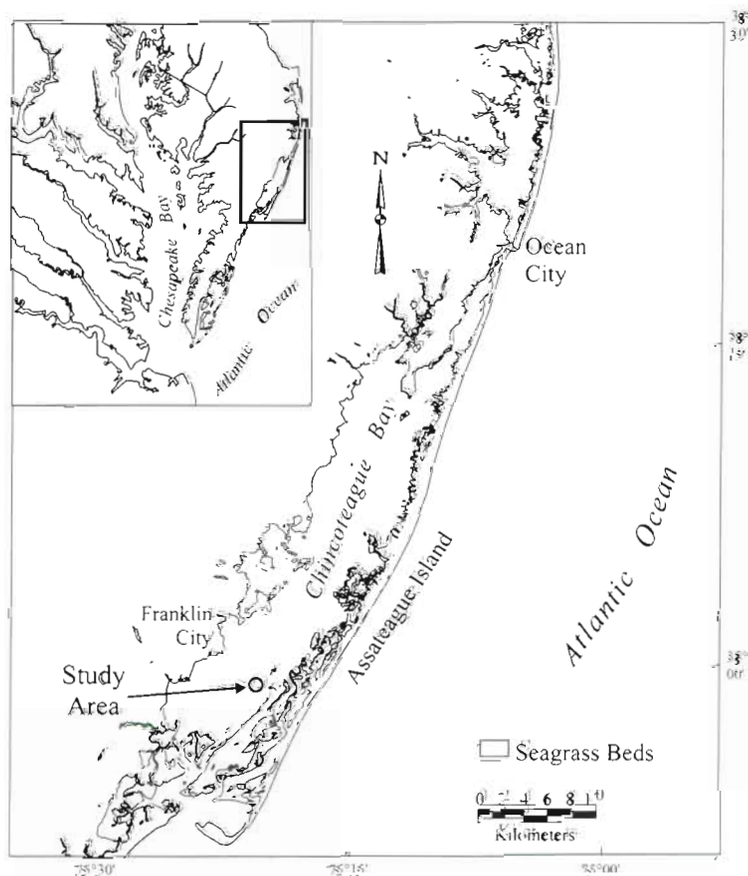


Fig. 1. Location of Chincoteague Bay on the eastern coast of the Delmarva Peninsula, USA (insert), and an enlargement of the bay, showing the distribution of eelgrass beds (shaded) and the study area

mean depth of 1.1 m and inlets are situated at the north and south ends of the bay. A large bed of eelgrass exists along the barrier island (eastern) portion of the Bay, covering 2300 ha or 7% of the total bottom. Historically, the Chincoteague Bay area supported dense stands of eelgrass and now contains some of the only eelgrass populations between New Jersey and the Chesapeake Bay.

Samples of eelgrass were collected from the study site on October 30, 1990, April 23, 1991, June 4, 1991, June 25, 1991, January 8, 1992, July 6, 1992, and September 30, 1992. On each date, 3 to 12 cores (15 cm diameter \times 20 cm) were collected using a hand-held auger. Each sample was placed in a zip-lock bag and was transported to the laboratory packed in an ice-filled cooler. Samples were stored on ice until they were processed, within 12 h of collection.

A field experiment was set up on June 3–5, 1991. At each of 5 locations paired groups of 4 cores were removed, placed in a PVC pipe, and then replaced in the original holes. Each core was 15 cm in diameter by 20 cm deep. Of each group, 2 were left unshaded, and 2 had shades erected directly above. Each shade was 1.8 m in diameter and was constructed of fine mesh fiberglass screening supported by a plastic pipe frame. The screens were buoyed at the water surface by styrofoam floats, and reduced the photosynthetically active radiation to 20% of ambient, measured using a submersible hand-held quantum sensor (Biospherical QS I-140). Fouling was kept to a minimum by cleaning screens daily.

The experimental cores were harvested on June 24–26, 1991. One of each pair was used for biomass measurements; the second was used for analysis of NSC. Two additional cores were taken from the vicinity of each replicate to compare biomass and carbon allocation patterns in the natural population and the unshaded cores. Cores were cleaned of sediment on a 1 cm mesh wire sieve by gently washing with seawater. The samples were stored on ice and processed within 12 h of collection. Samples were separated into 3 tissues: leaves, roots and rhizomes. Leaves consisted of all shoot material above the meristematic region (Sand-Jensen 1975) and roots and rhizomes were from the first 4 internodes only. Samples for NSC analysis were cleaned of sediment and organic material by briefly washing with tap water and gently scraping to remove epiphytes and sediment. The plant tissue used for the NSC analysis was then frozen and later freeze-dried to a constant weight. The freeze-dried samples were ground to a fine powder using a Crescent Wig-L-Bug dental amalgamator and stored in a freezer (-20°C).

The leaves, rhizomes and root material were analyzed for soluble carbohydrates according to Burke et al. (1992). Soluble sugars (sucrose, glucose, and fruc-

tose) were extracted from the ground tissue using hot 80% ethanol (Association of Official Analytical Chemists 1980) and starch using cold (0°C) 30% perchloric acid in a modified version of the Hansen & Möller (1975) method. Sugar and starch concentrations in the extractants were then measured using the phenol-sulfuric acid colorimetric method (Hodge & Hofneiter 1962) with an absorbance of 490 nm. NSC content was estimated by multiplying sugar and starch concentrations by biomass values obtained from this study as well as those from 3 nearby *Zostera marina* populations (Orth & Moore 1986) in a previous study.

Samples for biomass analyses were cleaned using tap water, counted and separated into leaves, roots and rhizomes. Because of the small mass of roots and difficulty in sampling them completely, roots and rhizomes were combined for biomass estimates. Epiphytes were gently removed from the leaves by scraping. Dry weight was determined after drying at 60°C for 2 d and ash free dry weight by subtraction after ashing at 550°C for 5 h.

Calculations of means, standard errors, Bartlett's test for homogeneity of variance and analysis of variance (ANOVA) were performed using Systat (Wilkinson 1988) or Stastica/Mac (StatSoft Inc. 1992). When variance was heteroscedastic, data were transformed and homogeneity of variance was insured before ANOVA were performed on the transformed data. For each significant ($p < 0.05$) ANOVA, factor level means were compared *post hoc* using Tukey's HSD. Significance levels reported are for the Tukey's HSD tests.

RESULTS

The concentration of NSC reserves (Table 1) was lowest in the leaves, highest in the rhizomes, and intermediate in the roots. Sugar was the dominant NSC form in the leaves (77%) and rhizomes (84%), while starch dominated over sugar (31%) in the roots. Because roots comprised a small (approximately 1%) proportion of the total biomass, they made a small contribution to the NSC reserves. The total NSC concentrations in leaves and rhizomes peak in the autumn and in late spring/early summer, and decrease to minimum levels in late summer.

Severing the rhizomes for the experiment did not influence the NSC concentrations: NSC concentrations in control and unshaded treatments were similar in each tissue type (Table 2). However, there was a strong experimental effect. Three weeks of shading resulted in a significant ($p < 0.001$) reduction in sugar concentrations in leaves (48%), rhizomes (40%), and roots (51%), but there were no differences in starch concentrations.

Table 1 *Zostera marina*. Mean estimates of the NSC (non-structural carbohydrate) concentrations in leaves, rhizomes and roots of field grown eelgrass on 7 dates. SE = standard error of the mean

Date	Sample size	Leaves		Rhizomes		Roots	
		Sugar mg g ⁻¹ (SE)	Starch mg g ⁻¹ (SE)	Sugar mg g ⁻¹ (SE)	Starch mg g ⁻¹ (SE)	Sugar mg g ⁻¹ (SE)	Starch mg g ⁻¹ (SE)
Oct 30, 1990	(4–8)	38.4 (4.0)	61.6 (4.2)	219.0 (16.5)	59.7 (3.3)	71.3 (6.4)	140.1 (3.7)
Apr 23, 1991	(10)	86.8 (1.2)	43.5 (0.8)	157.6 (17.0)	40.1 (1.9)	54.0 (3.4)	119.4 (2.9)
Jun 4, 1991	(10)	105.3 (4.2)	31.0 (0.7)	229.0 (5.6)	41.9 (3.3)	54.0 (0.9)	122.6 (0.1)
Jun 25, 1991	(10)	76.2 (5.6)	25.9 (2.0)	213.4 (7.4)	37.1 (2.5)	52.3 (2.4)	121.9 (1.5)
Jan 8, 1992	(3)	144.4 (69.3)	21.6 (11.4)	153.5 (12.5)	45.4 (1.0)	35.2 (6.7)	91.2 (1.0)
Jul 6, 1992	(12)	66.8 (13.5)	26.3 (1.5)	187.9 (13.7)	42.8 (2.5)	36.4 (4.6)	113.8 (3.9)
Sep 30, 1992	(10)	31.8 (3.2)	18.3 (0.7)	116.4 (10.2)	48.2 (1.2)	13.4 (2.5)	95.1 (1.5)

Table 2. *Zostera marina*. Mean estimates of NSC concentration in leaves, rhizomes and roots in eelgrass subjected to experimental shading. Control = field grown unshaded plants; unshaded = unshaded experimental plants; shaded = shaded (to 20% ambient light for 3 wk) experimental plants. n = 5. Identical superscripts indicate non-significantly different means (p > 0.05)

Date	Leaves		Rhizomes		Roots	
	Sugar mg g ⁻¹ (SE)	Starch mg g ⁻¹ (SE)	Sugar mg g ⁻¹ (SE)	Starch mg g ⁻¹ (SE)	Sugar mg g ⁻¹ (SE)	Starch mg g ⁻¹ (SE)
Control	76.2 (5.6) ^a	25.9 (2.0) ^a	213.4 (7.4) ^a	37.1 (2.5) ^a	52.3 (5.3) ^a	121.9 (3.4) ^a
Unshaded	86.1 (4.1) ^a	23.8 (0.7) ^a	214.0 (4.5) ^a	38.6 (2.6) ^a	46.2 (6.3) ^a	125.2 (5.0) ^a
Shaded	44.8 (1.6) ^b	25.6 (1.0) ^a	128.3 (9.4) ^b	44.9 (4.2) ^a	22.8 (0.9) ^b	133.6 (3.4) ^a

Table 3. *Zostera marina*. Mean biomass, density and ash free dry weight proportion estimates in eelgrass subjected to experimental shading. Control = field grown unshaded plants; unshaded = unshaded experimental plants; shaded = shaded (to 20% ambient light for 3 wk) experimental plants. n = 5. afw/dw = ratio of ash free dry weight to dry weight. Identical superscripts indicate non-significantly different means (p > 0.05)

	Leaves	Rhizomes and roots	Density	Leaves*	Rhizomes and roots*
	Dry wt g shoot ⁻¹ (SE)	Dry wt g shoot ⁻¹ (SE)	Shoots m ⁻² (SE)	afw/dw (SE)	afw/dw (SE)
Control	0.147 (0.028) ^a	0.0521 (0.0069) ^a	1518 (317) ^a	0.149 (0.008) ^a	0.249 (0.011) ^a
Unshaded	0.159 (0.039) ^a	0.0532 (0.0075) ^a	1441 (263) ^a	0.155 (0.010) ^a	0.244 (0.009) ^a
Shaded	0.105 (0.024) ^b	0.0412 (0.0085) ^b	1045 (112) ^b	0.134 (0.009) ^b	0.226 (0.010) ^b

* Arcsin square root transformed for ANOVA. Untransformed data are given

As with tissue NSC concentrations there was no difference in plant biomass between the unshaded treatments and the controls, therefore coring had no apparent effect on plant growth (Table 3). However, shading had a marked effect on the plants as leaf biomass per shoot decreased 34%, root-rhizome biomass per shoot decreased 23% and density decreased 27% with treatment (p < 0.05). This resulted in a 50% reduction in biomass on an area basis. The 3 wk of shading resulted in an estimated 66% reduction in NSC reserves: 15 g m⁻² NSC compared to 44 g m⁻² in the unshaded and 42 g m⁻² NSC in the control plants. In addition, leaf ash

free dry weight decreased from approximately 15% of leaf dry weight to 13%, and root-rhizome ash free dry mass from 25 to 23% of root-rhizome dry weight.

Based on the low and high biomass values provided by Orth & Moore (1986), NSC contents could have changed in Chincoteague Bay from 52 to 73 g m⁻² in late June to 4 to 18 g m⁻² in early January (rhizome biomass estimates were greater in the Chesapeake Bay study than in the Chincoteague Bay study). Thus, only 7 to 25% of the NSC stored in the naturally growing, unshaded population during the summer may have remained in January.

DISCUSSION

Seasonal changes in NSC reserves showed an asynchrony between carbohydrate supply and the demand for use. Carbohydrates were stored in the spring and autumn, indicating that photosynthate supply was greater than demand for growth and respiration at those times. Although the tissue specific levels of NSC were similar during the spring and autumn, shoot specific levels were much lower during the autumn as rhizome biomass is greatly reduced at this time (Orth & Moore 1986). Hence, spring is the major period for NSC accumulation and levels declined throughout the summer because respiration rates are higher than photosynthesis at this time (Evans et al. 1986).

These seasonal peaks in rhizome NSC reserves parallel the spring and fall bimodal growth pattern observed for *Zostera marina* populations in this region (Orth & Moore 1986). This pattern of growth is in contrast to eelgrass growth in other areas where lower summertime water temperatures are associated with a single peak growth period in the summer (e.g. Jacobs 1979). Zimmerman et al. (1995) concurrently measured seasonal patterns of growth, biomass and NSC reserves in transplanted eelgrass in one such area and demonstrated that in San Francisco Bay populations NSC reserves were greatest during the summer and declined to seasonal lows during the winter as photosynthetic performance decreased. Thus, periods with optimum growth conditions, whether in the spring, summer or autumn, also are important for the accumulation of NSC reserves in *Z. marina*. These reserves can be important for support of metabolic activity and growth during other seasons when environmental conditions are limiting.

Seasonal fluctuations in NSC reserves also occurred in *Syringodium filiforme* and *Halodule wrightii*, but not for the seagrass *Thalassia testudinum* (Dawes & Lawrence 1980). This suggests that some but not all seagrass species have well-defined seasons of NSC reserve storage and depletion. These fluctuations reflect periods when photosynthate availability exceeds the need for growth and/or respiration (NSC storage) versus periods when growth and/or respiration exceeds photosynthate availability (NSC depletion). The storage of reserves during one season for use during other seasons is characteristic of plants subjected to extreme seasonal conditions (Osmond et al. 1987, Chapin et al. 1990).

Consistent with previous findings for Zosteraceae (Drew 1979, Zimmerman et al. 1989), a large proportion of the NSC reserves was stored as sugar. The reason that eelgrass has large sugar reserves relative to reserves of other plants is unclear, but sugar is an immediately available substrate for respiration, and

reserves of sugar rather than starch may improve the chance of survival when unpredictable environmental stresses may occur. In addition, energy is saved by not converting sugar to starch and then back to sugar.

It is not immediately apparent why our estimates of peak NSC reserves were so much lower than the 400 and 600 mg g⁻¹ reported for leaves and rhizomes for the California populations of *Zostera marina* (Zimmerman et al. 1989, 1995). A possible explanation is that the Chincoteague populations are subject to greater extremes in temperature, up to 30°C, and the California populations experience a much smaller temperature range of 12 to 17°C. Because respiration exceeds photosynthesis at temperatures greater than 25°C (Zimmerman et al. 1989), Q₁₀ values of 2.4 and 1.6 respectively (Marsh et al. 1986), the Chincoteague *Z. marina* population may spend extended periods during the summer respiring away stored reserves. However, Zimmerman et al. (1995) reported that reduced plant photosynthetic performance combined with low light availability in the winter resulted in decreases in NSC from summertime highs of 450 to 550 mg g⁻¹ to annual minimums in NSC of 50 to 150 mg g⁻¹ in San Francisco Bay populations. These are comparable to our minimum values (Table 1), and suggest a similar annual balance between NSC reserve accrual and loss, in spite of different annual patterns of allocation.

Clear water in the spring is critical to the survival of populations of *Zostera marina* because spring is an important time for growth and accumulation of energy reserves in eelgrass. It is likely that the NSC reserves accumulated during this time of year sustain the plants through other periods when respiration may exceed photosynthesis and NSC reserves can drop by 75 to 92%. Springtime turbidity has potentially serious ramifications. Three weeks of turbidity could reduce the amount of potentially accumulated NSC reserves by 66% during that time period, increase the risk for exhaustion during subsequent periods of stress, and thus jeopardize overwintering and perhaps even summer survival. In addition, as in other angiosperms, a low carbon balance in eelgrass may enhance susceptibility to opportunistic pathogens (Short et al. 1988, Burkholder et al. 1992).

Apparently, eelgrass has limited flexibility in the way in which it allocates C among competing sinks. Under shaded conditions in the spring, allocation to new biomass still superseded allocation to storage. Subsequently, the greater reduction in NSC storage compared to growth resulted in a reduced tissue specific level of NSC reserves. Thus, there was more biomass to be maintained with less substrate, a situation analogous to the decline in carbon documented in *Zostera marina* grown in nitrate-fertilized water (Burkholder et al. 1994). Apparently, the abiotic factors of

reduced light, elevated water column nitrate, and warm temperatures act alone or in concert to depress the carbon balance in *Z. marina*. Probably, it is this compromised condition that causes eelgrass populations to decline during the warm summer months in the southern part of its distributional range.

The small amount of NSC reserves in these plants in January, 8 to 25% of spring reserve levels, illustrate the small margin of safety afforded eelgrass at this southern limit to its range. Even under optimum water quality conditions the carbon balance may approach zero in the summer as a result of net carbon loss from respiration. It is no surprise that eelgrass populations in this region decline when turbidity events coincide with the spring growing season, which represents the major 'window of opportunity' for storage of NSC reserves by eelgrass.

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