

# Predator distribution and habitat patch area determine predation rates on Age-0 juvenile cod *Gadus* spp.

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**ABSTRACT:** Eelgrass *Zostera marina* provides refuge to numerous fish species but is vulnerable to fragmentation through natural and anthropogenic disturbance. In Bonavista Bay, Newfoundland, eelgrass patch size was altered to measure changes in predation risk in Age-0 juvenile cod, *Gadus morhua*. Artificial eelgrass mats of 5 sizes (0.32, 1.1, 5.5, 11 and 22 m<sup>2</sup>) were deployed in duplicate at each of 2 sites in Newman Sound in Terra Nova National Park during summer/autumn in 1999 and 2000. Predator distribution was determined using a combination of weekly underwater transect surveys and biweekly seining. Relative predation rates were measured by tethering Age-0 cod at the center of each patch and recording the incidence of predation (n = 1116 tether sets). Predation rates were negatively correlated with patch size during both years, suggesting that larger patches reduce predator foraging ability. However, high predator densities in the largest eelgrass patch resulted in higher than expected rates of predation. Therefore, habitat dimension affected predation risk in juvenile cod via 2 opposing mechanisms. Our results stress the importance of considering both habitat areal extent and predator distribution when estimating the effects of habitat fragmentation on predation rates.

**KEY WORDS:** Predator-prey interactions · Edge effects · Eelgrass · Juvenile cod · Predator distribution · Habitat fragmentation

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## INTRODUCTION

Seagrass beds provide shelter to a variety of juvenile fish species in coastal environments across a range of latitudes (e.g. Orth et al. 1984, Sogard & Able 1991, Laurel et al. unpubl. data). Fragmentation of these beds occurs both naturally (e.g. seasonal die-off, wave-action, ice scour) and anthropogenically (e.g. inshore boating, nitrogen loading; Fonseca 1992). Consequently, seagrass habitats are often patchy, ranging from single plants to expansive meadows greater than 1000 m<sup>2</sup> (Robbins & Bell 1994).

Juvenile fish often use complex habitats to offset the threat of predation (e.g. Werner et al. 1983a,b, Mittelbach 1986). However, increased habitat fragmentation of seagrass beds may have negative consequences for young refuge-seeking fish. Predation risk is highest during the early life stages of marine fish (Sogard

1997) and complex habitat reduces risk by interfering with the visual and swimming capabilities of piscivores (Gotceitas & Colgan 1989). However, predators may compensate for reduced foraging efficiency by feeding along the edges of complex habitat (Peterson et al. 2001). Therefore, predation risk may be higher in fragmented habitats with increased edge to area ratios (e.g. small patches). Small, fragmented patches may also be riskier because their interior is closer to the more dangerous edges. These explanations have been invoked to explain increasing predation risk with smaller habitat patches in both terrestrial (Levenson 1981) and marine systems (Irlandi 1997).

There has been considerable variation in the results of edge effects/patch size on predation rate (reviewed by Paton 1994). Equivocal results have prompted more studies to examine the local abundance of predators and prey (Donovan et al. 1997). For example, rates of

piscivory in eelgrass habitats can be equal to or higher than unstructured sand habitats, despite the reduced foraging ability of predators in complex habitats (Linehan et al. 2001, Gregory et al. unpubl. data). Differences in predation rate were attributed to higher numbers of predators in eelgrass habitats compared to unvegetated sand. Therefore, the interaction of habitat characteristics (e.g. complexity, perimeter to edge ratios) on predator distribution must be considered when quantifying predation risk across habitats.

Atlantic cod *Gadus morhua* is a commercially important demersal species that occupies nearshore areas in the northwest Atlantic Ocean during the first year of life (Taggart et al. 1994). As juveniles, cod associate with complex habitats, such as boulders/large rock (Gregory & Anderson 1997), cobble (Tupper & Boutlier 1995), macroalgae (Keats 1990) and eelgrass (Gotceitas et al. 1997), for protection from predators, such as larger conspecifics and other piscivorous fish as well as seabirds. Laboratory experiments have shown that complex habitats can reduce predation risk for juvenile cod when exposed to actively foraging predators (Gotceitas & Brown 1993, Gotceitas et al. 1995, Lindholm et al. 1999). However, recent studies have shown that predators of juvenile cod also prefer complex habitat (Linehan et al. 2001, R. Gregory unpubl. data) and may

elevate risk in these habitats. Therefore, the role of habitat characteristics on predation risk in juvenile cod remains unclear.

No studies have been conducted in marine systems examining the effects of habitat patch size and local predator abundance on predation rates. Earlier investigations considering both habitat characteristics and predator distribution have been restricted to a few large-scale avian studies (reviewed by Donovan et al. 1997). In this study, we measured the distribution of piscivorous fish and the rate of predation on tethered Age-0 cod in varying sized seagrass patches. We addressed the following questions: (1) Does risk change with increasing eelgrass patch size? and (2) How do habitat patch area and localized predator distribution interact to affect risk in Age-0 juvenile cod?

## MATERIALS AND METHODS

**Study area.** Newman Sound is located within Bonavista Bay off the northeast coast of Newfoundland in the vicinity of Terra Nova National Park (TNNP; Fig. 1). The nearshore habitat (0 to 15 m depth) varies between eelgrass *Zostera marina*, macroalgae (e.g. *Laminaria digitata*, *Chondrus crispus*) and fine- to coarse-grain mineral habitats. We conducted experiments in 2 coves within the sound: Canning's Cove (CC) and Mt. Stamford Cove (MS). These sites were chosen mainly because they lack any complex habitats such as boulders and/or eelgrass, were of similar depth in the experimental area (1.5 to 2.5 m) and were also conducive to hauling a net along the bottom.

**Artificial eelgrass.** Artificial eelgrass was manufactured using galvanized screen (2.5 cm pore size) and green curling ribbon (0.8 cm width). Ribbon was tied onto the screen to create blades 75 cm long at a density of 400 blades  $m^{-2}$ , falling within the range of length and density for naturally occurring eelgrass (Orth et al. 1984). Eelgrass mats were cut into varying length strips to form approximately square patches 0.32, 1.5, 5.5, 11 and 22  $m^2$  in size. SCUBA divers secured replicates of each of these patch sizes to the bottom at each site using reinforcing bar (i.e. re-bar) spikes (Fig. 2). Eelgrass mats were attached flush against the substrate by staggering spikes at 0.5 m distances along their entirety. This reduced possible damage from wave action and also minimized the effect of added complexity from the wire mesh frames. Spacing between patches edges was kept constant

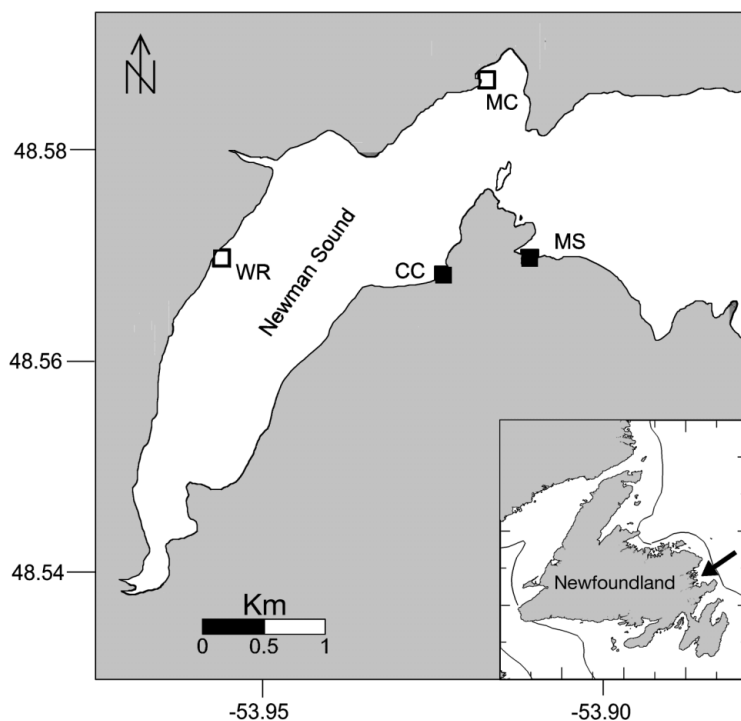


Fig. 1. Locations of eelgrass patches in Newman Sound, Bonavista Bay, Newfoundland. Artificial eelgrass locations: Canning's Cove (CC) and Mt. Stamford (MS) (■); natural eelgrass locations: White Rock (WR) and Mistaken Cove (MC) (□)

at 3 m for both sites. Patches in 1999 were deployed in July, approximately 1 mo prior to Age-0 juvenile cod settlement in coastal Newfoundland (Methven & Bajdik 1994, Grant & Brown 1998a). Artificial blades of eelgrass became fouled with epiphytes throughout the season similar to natural eelgrass. In November 1999, patches were removed, cleaned of epiphytes and stored dry before being redeployed again in July 2000. No significant degradation in the artificial eelgrass (i.e. mesh rust, blade loss, etc.) was observed in either of the 2 years of the experiment.

**Predation rates.** Predation within patches was determined using a tethering technique with Age-0 cod collected by seine from nearby sites. We used 2 species of cod (58% Atlantic cod *Gadus morhua* and 42% Greenland cod *G. ogac*) for tethering. These 2 species co-occur in the nearshore (Methven et al. 2001) at similar sizes within the first year of life. Susceptibility to predation on tether lines is similar for these 2 species (B. Laurel & R. Gregory unpubl. data).

The tethering apparatus consisted of 2 sections, a 2 m 11.2 kg test vertical float line and a 0.5 m 0.9 kg test horizontal leader line. The float line was kept vertical in the water column using a 564 g anchor and a cork float 1 m from the bottom. A white float at the surface marked the location of the tether line and facilitated retrieval. The horizontal leader line was attached to a #7 barrel swivel on the vertical float line 75 cm from the bottom substrate. Age-0 cod were tethered to the leader line via a #16 hook inserted through the caudal peduncle. The leader line allowed cod prey a 0.5 m radius of movement and access to the eelgrass. A full description of the tethering apparatus and technique is described in Linehan et al. (2001).

Each tether line was deployed from a small aluminum boat (4.2 m) at midday, left undisturbed for a period of 15 min and finally retrieved to determine the incidence of predation. Water clarity was adequate to insure that tethered fish were not lost while being lowered to the bottom. Predation was classified as a 'hook loss', 'fish loss', 'caught predator' or 'predator attack' upon retrieval of the tether line. The absence of predation was recorded if a tethered fish was retrieved alive without any sign of predation, e.g. punctured skin, etc. All predators captured on tether lines were identified, measured to the nearest 0.5 cm standard length (SL) and released.

Our experimental design consisted of setting tethers at the center of each patch (0.32, 1.1, 5.5, 11 and 22 m<sup>2</sup>) and on 6 neighboring areas of unvegetated sand about 3 m from the nearest patch edges. Tethering in patch and unvegetated areas was repeated 6 times during 3 tethering sessions in each year for a total of 1152 tether sets: [6 unvegetated areas + (5 artificial eelgrass patches × 2 patch replicates)] × 6 rounds ×

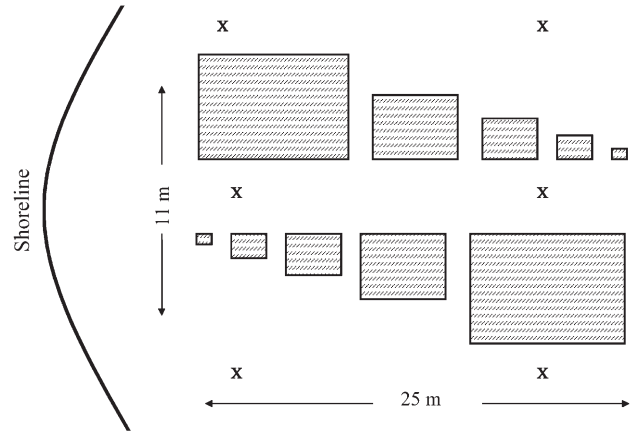


Fig. 2. Experimental layout of artificial eelgrass patches relative to shore at Canning's Cove and Mt. Stamford. Tethers are set at the center of each patch and 6 locations outside of patches (indicated by X)

2 sites × 3 sessions × 2 yr. Tethering sessions were conducted at ca. 2 wk intervals beginning early September of both years and corresponded with concurrent surveys of predator abundance. Tethered prey throughout the time period of the experiment ranged in size from 65 to 106 mm SL. The range in size represents growth in the available *Gadus* prey from early September to mid-October of each year. However, variation in prey size was substantively less within individual tethering sessions, i.e. maximum ± 11 mm SL.

**Predator distribution.** Predator abundance was estimated at 2 wk intervals using a 25 m Danish bag seine (19 mm mesh) deployed 55 m offshore from a boat and hauled to shore by 2 individuals standing 16 m apart. The seine sampled 880 m<sup>2</sup> of bottom habitat up to 2 m in the water column (for a description see Methven & Schneider 1998). Captured fish were transferred to holding containers with seawater. Piscivorous fish captured included Greenland cod, Atlantic cod, cunner *Tautoglabrus adspersus*, sculpin *Myoxocephalus* spp. and white hake *Urophycis tenuis*. Potential predators were identified and measured (±5 mm SL) and released the same day to their respective sites.

Small-scale, localized predator distribution was measured via snorkeling. Snorkeling surveys were always conducted 3 to 4 d before or after seining activity to minimize the effects of net disturbance. Snorkelers swam directly over patches, recording the abundance and type of fish predators within 0.5 m of each patch. Large patches (11 and 22 m<sup>2</sup>) exceeded the viewing capacity of a single snorkeler, so these patches were assessed by 2 snorkelers swimming in parallel. Snorkelers swam with delineated poles (2.82 m each) to minimize overlap of survey areas of the large patches. Water clarity throughout the survey was suffi-

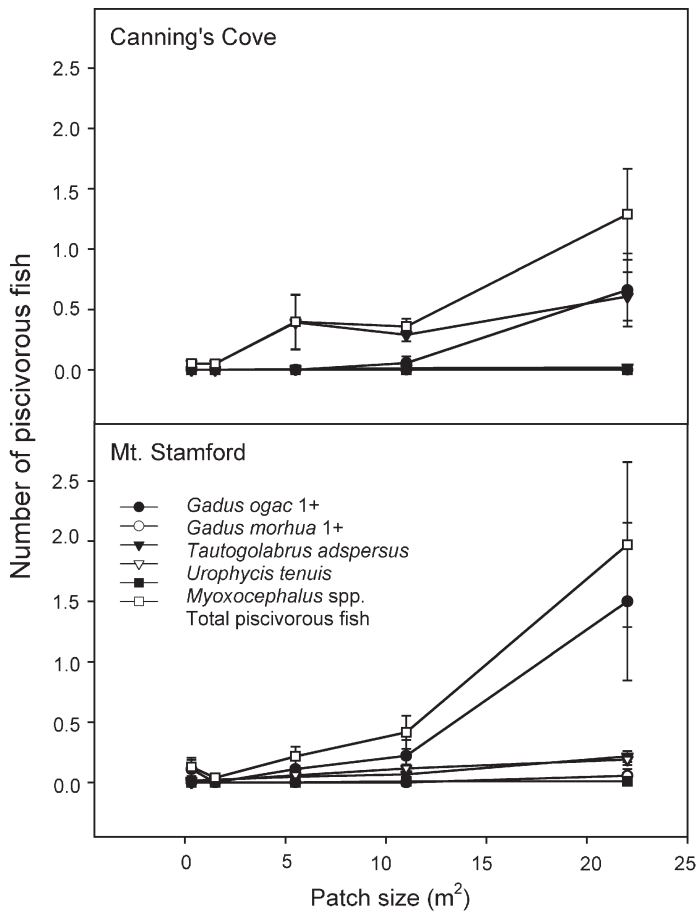


Fig. 3. Mean abundance of predators (*Gadus ogac*, *G. morhua*, *Tautogolabrus adspersus*, *Myoxocephalus* spp. and *Urophycis tenuis*) censused via snorkeling at Canning's Cove and Mt. Stamford on various sized patches of simulated eelgrass. Each point represents a mean of 9 snorkel surveys of 2 replicate patches  $\pm 1$  SE

ciently greater (i.e. 5 to 6 m) than the maximum area censused by a single snorkeler, i.e. 2.82 m. Experimental sites were visually censused in this fashion between August 10 and October 12, 2000 for a total of 9 surveys. The survey was not performed in 1999 for logistical reasons.

Additional control snorkeling surveys were conducted at 2 natural eelgrass meadows of similar depth during the same time period (n = 9). Eelgrass at the control sites was continuous (i.e. non-patchy) over the range of the survey area, but only plot areas similar to those at experimental sites were surveyed. This was accomplished by deploying 15 m transect lines perpendicular to shore at each control site. The vertical dimension and spacing of patches was marked on each transect line with pink surveyor's tape, while the horizontal dimension of patches was estimated by snorkelers swimming with delineated plastic poles over the

transect lines. Snorkelers swam in parallel once over both transect lines at each site, so that only areas falling within the delineated transect lines were surveyed (80 m<sup>2</sup>).

Because of the difficulty in estimating fish size under water, potential predators within patches were size-corrected by integrating seine and snorkel census data using the following equation:

$$P_{tot} = \sum P_{sn} P_{se}$$

where  $P_{tot}$  is the total potential predators in the patch,  $P_{sn}$  is the total predators identified from snorkel surveys and  $P_{se}$  is the percentage of predators caught in the seine greater than 2x the mean tethered prey size (mm SL). From previous cod studies (Bogstad et al. 1994, Grant & Brown 1998b, Linehan et al. 2001) we know that predators greater than 2x the prey size have the potential of capturing and consuming juvenile cod.

**Data analysis.** Tethering data were analyzed using an analysis of variance assuming a binomial error structure on logarithically transformed data (log link;

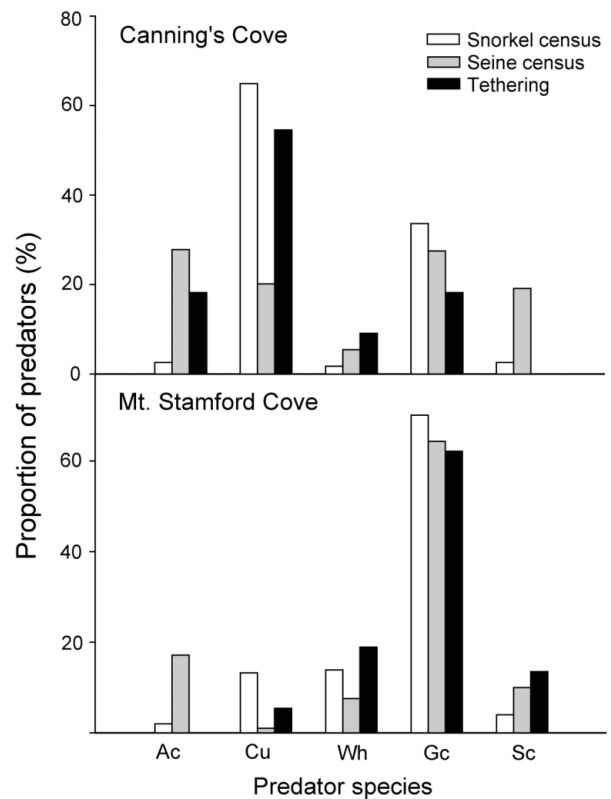


Fig. 4. Proportion of Age-0 cod predators determined by snorkeling, seining and tethering at Canning's Cove and Mt. Stamford in 2000. Proportions are based on 4 beach seine hauls, 288 tether sets and 9 snorkel surveys per site. Greenland cod *Gadus ogac* (Gc); Atlantic cod *Gadus morhua* (Ac); Cunnners *Tautogolabrus adspersus* (Cu); sculpin *Myoxocephalus* spp. (Sc); white hake *Urophycis tenuis* (Wh)

proc GENMOD; SAS Release 6.03). 'Year', 'sampling period' and 'site' were used as explanatory variables. Significant interactions were found between year and site variables, so patches were analyzed separately in 1999 and 2000 within sites. The model assumed a uniform predator distribution; therefore, only patches with similar mean predator density (i.e.  $\pm 2$  SE) were analyzed. In this way, the effects of habitat structure on predation rates were measured independently of the predator distribution.

## RESULTS

A total of 1152 tethering sets were conducted in both years at our 2 sites. Thirty-six sets were omitted from the analysis because lines were either tangled or fish were dead upon retrieval. The remaining 1116 tethering sets were considered successful, of which 12% (138 incidences) resulted in some form of predation. Predation events included 'caught predator' (48%), 'hook gone' (6%), 'fish gone' (22%) and 'predator attack' (24%). Predators retrieved on tether lines included Greenland cod ( $18.7 \pm 3.9$  cm SL), Atlantic cod ( $23.6 \pm 4.7$  cm SL), cunner ( $15.8 \pm 5.9$  cm SL), sculpin ( $17.0 \pm 5.8$  cm SL) and hake ( $12.6 \pm 0.5$  cm SL). Higher overall predation was observed at MS (16%) than CC (9%).

Predator abundance increased with increasing patch size at both sites as expected (Fig. 3). However, the estimated composition of predators differed between sites as well as by census methods (Fig. 4). For example, cunners were found in greater abundance at CC, whereas hake were more numerous at MS. However, snorkel and seine surveys indicated that total predator abundance was higher at MS compared to CC. Census discrepancies included Atlantic cod and sculpins. Both were rarely seen over patches during snorkeling surveys but were often caught in seines and on tether lines. In contrast, cunners were often observed in artificial eelgrass patches yet comprised a smaller portion of total seine catch. However, seine, snorkel and tether data all indicated that *Gadus ogac* was an abundant predator at both sites (Fig. 4).

Distribution of predators among patches was similar between sites (Fig. 5). At MS, patch sizes of 0.3 to 11 m<sup>2</sup> supported similar predator densities ( $0.037 \pm 0.006$  pred. m<sup>-2</sup>). In contrast, predator density in the largest patch (22 m<sup>2</sup>) was 2.3 times higher ( $0.086 \pm 0.011$  pred. m<sup>-2</sup>) than in the smaller patches. A similar distribution was observed in patches at CC; predator densities in small patch sizes were nearly 50% ( $0.030 \pm 0.005$  pred. m<sup>-2</sup>) of those in the largest patch size ( $0.059 \pm 0.008$  pred. m<sup>-2</sup>). Therefore, predator density was similar in all patches except the largest (22 m<sup>2</sup>) during

snorkeling surveys. Differences in 1+ yr old Greenland cod abundance were responsible for this trend (Fig. 6). Although Greenland cod were evenly distributed within natural eelgrass meadows, they were disproportionately distributed among artificial eelgrass patches. Cunners and hake did not deviate consistently from expected trends at either eelgrass meadow or patch sites.

Predation rates increased with diminishing patch size at both sites during each year (Table 1, Fig. 7) when patches had similar predator density. Data collected from the largest patches (22 m<sup>2</sup>) were not included in the analysis because GENMOD assumes uniform predator distribution; only patch sizes 1 to 4 could meet these assumptions. In 2000, there was a significant effect of sampling date on predation rates at both sites; however, no interaction was detected between sampling date and patch (Table 1). Highest predation occurred outside the patches at rates similar to those found over the smallest patches (Fig. 7). Lowest predation was observed in the largest patches analyzed, i.e. 11 m<sup>2</sup>.

## DISCUSSION

Structural complexity provides refuge to juvenile fish in freshwater (e.g. Crowder & Cooper 1982, Savino & Stein 1982, Werner et al. 1983a,b, Gotceitas & Colgan 1989) and marine (e.g. Sale 1991) systems over a wide range of latitudes (Heck & Wilson 1987). Numerous studies have shown that juvenile Atlantic cod use

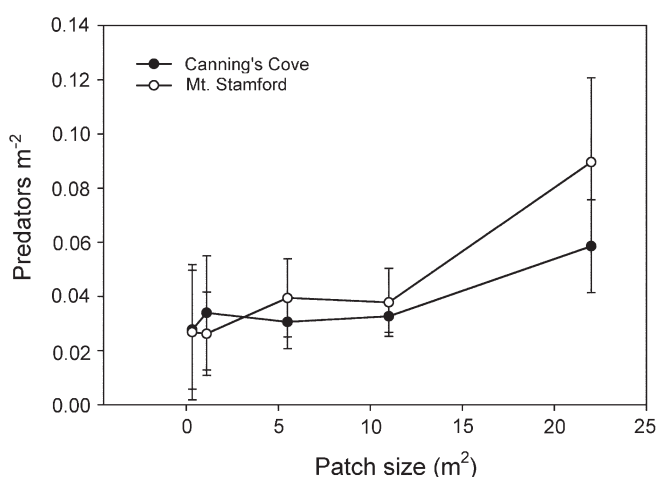


Fig. 5. Total density of piscivorous predators (Greenland cod *Gadus ogac*, Atlantic cod *Gadus morhua*, cunners *Tautoglabrus adspersus*, *Myoxocephalus* spp. and white hake *Urophycis tenuis*) in various sized patches of simulated eelgrass at Canning's Cove and Mt. Stamford. Values represent a mean of 9 snorkel surveys of 2 replicate patches  $\pm 1$  SE



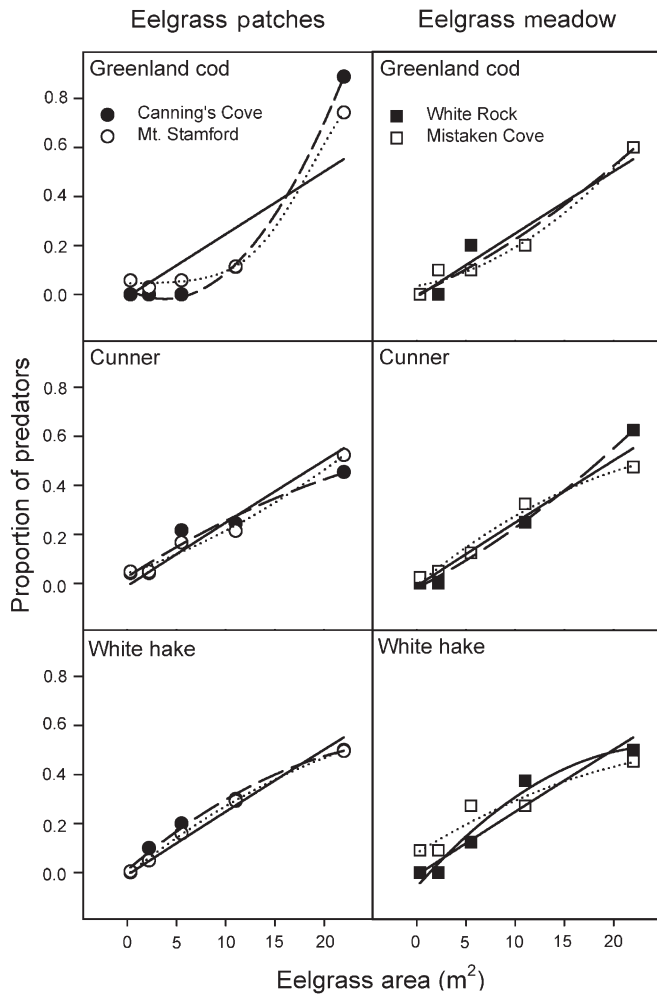


Fig. 6. Proportion of common piscivorous predators (Greenland cod *Gadus ogac*, cunners *Tautoglabrus adspersus* and white hake *Urophycis tenuis*) distributed over a range of artificial eelgrass patch sizes and equivalent areas at natural eelgrass meadows. Expected lines are calculated assuming a linear distribution of fish with area. Actual distributions are plotted with a best-fit quadratic equation to illustrate trends deviating from the expected linear distribution

complex habitats to reduce predation (e.g. Gotceitas & Brown 1993, Gotceitas et al. 1995, 1997, Gregory & Anderson 1997, Lindholm et al. 1999, 2001). Structurally complex habitats impair the visual and swimming capabilities of predators (Savino & Stein 1982), which in turn can reduce the effectiveness of encountering, attacking and capturing prey (Tallmark & Evans 1986). Our results were consistent with these findings. Predation rates of juvenile cod were highest over areas without structural complexity, i.e. unvegetated sand. However, we also show that the areal extent of surrounding refuge habitat, along with predator density, also influence the predation risk of small fish.

Age-0 juvenile cod were subjected to highest predation risk in smallest patches when predator density was uniform across all patch sizes. Fragmented habitat patches are considered risky because: (1) they have high perimeter to area ratios; and (2) the interior habitat is close to the edge and therefore, accessible to hunting piscivores. While these mechanisms have been explored in terrestrial systems (Brittingham & Temple 1983, Andren et al. 1985, Andren 1992, Donovan et al. 1997), marine examples have been restricted to a number of invertebrate species (Irlandi 1997, Bologna & Heck 1999, Peterson et al. 2001). Our results show that predation risk increases for fish in fragmented environments. Our analysis controlled for differential predator density and we suggest that the patch characteristics are directly responsible for the trend observed. We believe that risk was higher in small patches because of the associated changes in a predator's foraging ability, and not simply predator abundance.

Predator density was not uniform among all habitat patches. The largest seagrass patches (22 m<sup>2</sup>) attracted higher densities of piscivorous fish. Consequently, predation risk was higher in these patches than could have been predicted from habitat characteristics alone, i.e. patch size. Predator density and predation rates are positively correlated in studies of juvenile cod (Linehan et al. 2001, Gregory et al. unpubl. data). However, the non-linear distribution of predators with habitat area we observed in our study was unexpected. The

Table 1. GENMOD analysis of predation rates of tethered *Gadus* spp. on varying sampling dates, and artificial eelgrass patches as a function of year (1999 or 2000) and sampling location: Mt. Stamford (MS); Canning's Cove (CC)

Site	Year	Source of variation	df	Deviance	$\chi^2$	p
MS						
1999	Date	Date	2	35.82	1.24	0.5379
		Patch	1	8.06	27.76	<0.0001
		Patch × Date	2	7.34	0.72	0.6990
		Intercept	207	37.06		
2000	Date	Date	2	30.63	11.98	0.0025
		Patch	1	21.04	9.59	0.0020
		Patch × Date	2	16.20	4.84	0.0890
		Intercept	276	42.61		
CC						
1999	Date	Date	2	30.90	5.95	0.0510
		Patch	1	24.08	14.82	0.0090
		Patch × Date	2	9.70	4.38	0.0810
		Intercept	238	36.85		
2000	Date	Date	2	42.95	15.27	0.0005
		Patch	1	15.51	27.44	<0.0001
		Patch × Date	2	14.37	1.13	0.5681
		Intercept	278	58.22		

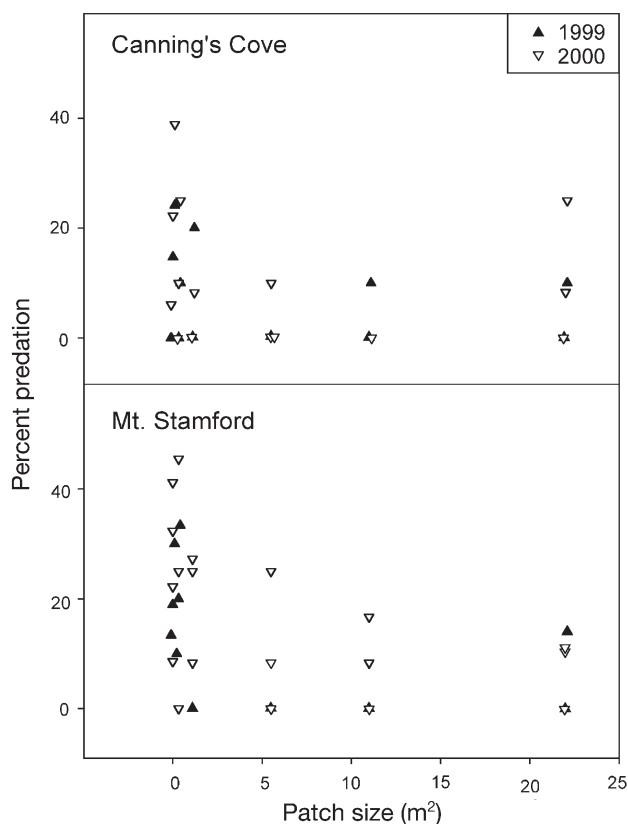


Fig. 7. Percent predation of juvenile cod *Gadus* spp. tethered in unvegetated areas ( $0 \text{ m}^2$ ) and artificial eelgrass patches ( $0.32, 1.1, 5.5, 11$  and  $22 \text{ m}^2$ ) at Canning's Cove and Mt. Stamford between 1999 and 2000. Values consist of 12 to 36 tether sets and are staggered  $\pm 0.1 \text{ m}$  to avoid overlap

higher predator densities in the large patches were due to increased abundance of Age-1+ Greenland cod, a species common in nearshore areas around Newfoundland (Methven et al. 2001) and a known predator of Age-0 cod (Linehan et al. 2001). The distribution of Greenland cod at the control sites (eelgrass meadows) was linear, indicating that this result was not due to overall patch layout, small changes in depth, distance from shore or area surveyed. Rather, it appears that changes in Greenland cod density were a function of eelgrass patch size.

Nonlinear associations between habitat and organisms are not unique to our study; however, the causal mechanisms are uncertain. Species-area curves typically increase with patch size (MacArthur & Wilson 1967) and for some species, density has been related to habitat area. For example, European jays *Garrulus glandarius* L. and ravens *Corvus corax* L. are commonly found in large wooded areas, yet are entirely absent from smaller forest fragments (Andren 1992). Similar findings have been reported for other bird species (Stoufer & Bierregaard 1995) and many species of

invertebrates (Kareiva 1985, Margules et al. 1994, Ingham & Samways 1996, Golden & Crist 1999), including infaunal macroinvertebrate assemblages in eelgrass (Bowden et al. 2001). Small, fragmented patches may not have suitable food or niche space (Galli et al. 1976, Martin 1981) or refuge characteristics (Martin 1988) that large patches offer. Eelgrass is known to both concentrate prey (Connolly 1997) and provide shelter to piscivorous fish from predators (Orth et al. 1984). Therefore, predatory Greenland cod may use large patches of eelgrass to balance feeding success with their risk of being eaten. Our results suggest that predation risk changes with patch size for small cod ( $<10 \text{ cm}$ ); however, whether such an explanation applies to the distribution of larger fish (e.g. Greenland cod) is unknown.

We detected no interaction between date and patch size, suggesting that trends in predator distribution and risk were maintained within the temporal scale of our study. However, diel, monthly, seasonal and annual changes in piscivorous predators are variable in nearshore areas around Newfoundland (Clark & Green 1990, Linehan et al. 2001, Methven et al. 2001). Therefore, observed trends of predator distribution in our study may not uphold at finer (e.g. diel) or broader (e.g. seasonal) temporal scales. Furthermore, such trends will not necessarily result in predictable changes in predation risk for juvenile cod. For example, Linehan et al. (2001) demonstrated a decrease in predation risk at night on Age-0 juvenile cod, despite increased density of larger conspecifics. Similarly, seasonal differences in temperature can affect the metabolic demands of piscivorous fish and consequently, risk to their prey species. Overwintering predation in many aquatic systems is therefore considered negligible (e.g. Keast 1978). We restricted our study to the timescale August to October, a time of year when juvenile cod predation risk is high (Linehan et al. 2001, Gregory et al. unpubl. data) and effects of predator distribution on the risk of juvenile cod to predation would be highest.

Inconsistencies in predator density estimates between our seine, snorkeling and tethering techniques are likely methodological artifacts. We suspect that seining more accurately samples predators than visual census. Previous work has demonstrated that demersal fish are caught by our seine technique with 95% efficiency (Gotceitas et al. 1997), whereas visual methods can underestimate density due to diver avoidance and fish crypsis (Keats 1990). We estimated density within individual eelgrass patches visually because the seine was not suited to sampling at small spatial scales (ca.  $1$  to  $25 \text{ m}^2$ ). The visual census more closely matched seine estimates for site-attached piscivores (e.g. Greenland cod, hake, cunners) that

remained associated with eelgrass patches when approached. However, some fish did not associate with patches at these small scales. For example, we routinely observed Age-1+ Atlantic cod within close proximity to patches, but we did not include these fish in our abundance estimates as they were 'outside' the patch area. Consequently, Atlantic cod were underestimated in snorkel surveys relative to seine catches. Some piscivores may have also been underestimated by our tethering technique due to differential foraging capabilities. For example, cunners were rarely caught on tether lines at our sites, yet they were relatively abundant in the areas. Cunners are considered to be omnivorous, feeding preferentially on mollusks and amphipods rather than fish (Scott & Scott 1988). Prey selection is known to be gape-limited (e.g. Schmitt & Holbrook 1984) and cunners, with their small mouths, may have been restricted in their ability to handle our tethered prey in an equal manner to other similar-sized predators, e.g. Greenland cod or sculpins. However, we have observed *in situ* foraging attempts of large cunners (ca. 25+ cm) on Age-0 cod in Newman Sound.

There are limitations of the tethering technique as a means of measuring predation risk in the field. Tethering studies are numerous, including a wide variety of marine and freshwater organism such as decapod crustacea (Heck & Wilson 1987, Eggleston et al. 1990, Wilson et al. 1990), bivalves (e.g. Arsenault & Himmelman 1996), freshwater fish (Gregory & Levings 1998, Post et al. 1998) and marine fish (Curran & Able 1998, Linehan et al. 2001). The technique has its detractors (e.g. Halpin 2000, Kneib & Scheele 2000, Peterson & Black 1994) who have suggested that artifacts confound estimates of predation risk. As a technique to measure relative predation risk between similar habitats, such artifacts are unlikely to confound our interpretations (see Linehan et al. 2001).

In summary, our study is the first to measure the effects of habitat patch size on predation rates in a marine fish species. Eelgrass afforded greater protection to juvenile cod than unvegetated habitat, but both habitat patch size and predator distribution also contributed to differential rates of predation. Small eelgrass patches were riskier to juvenile cod than large patches when predators were evenly distributed; however, risk often increased in large patches due to higher predator densities. These results stress the importance of considering both habitat patch dimension as well as predator distribution and abundance when evaluating habitat quality for small fish.

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