ABSTRACT: Tidal migrations are inherent in the life histories of numerous demersal predators, and both the indirect and direct effects of these forays are important to the structure and function of littoral communities. In coastal estuaries of the Northeastern Pacific, Dungeness crabs *Cancer magister* are abundant and compose a significant portion of estuarine biomass. The nursery role of complex littoral habitats for young-of-the-year (0+) *C. magister* is well documented, yet the ecology of subsequent age classes within coastal estuarine systems, and within littoral areas in particular, remains unclear. The goal of our study was to elucidate habitat use and migratory patterns of subadult *C. magister* (40 to 130 mm carapace width; 1+ and >1+ yr classes) in littoral eelgrass *Zostera marina*, oyster *Crassostrea gigas*, and unstructured littoral habitats (ULH). We employed 3 sampling techniques (trapping, acoustic telemetry, and underwater video) designed to examine various aspects of migratory behavior within Willapa Bay, Washington, a representative coastal estuary. Baited trap surveys revealed that relative catches of subadult *C. magister* are 30 to 50% higher on ULH than eelgrass or oyster beds, and are negatively correlated with catches of another large cancrid crab, *C. productus*. Ultrasonic telemetry observations suggest that subadult *C. magister* making nighttime foraging incursions prefer ULH to other littoral habitats, and underwater video observations show that migrations are influenced by tidal rhythms since movements are correlated with the direction and velocity of current flow in adjacent channels. Our results dramatically alter the perception of *C. magister* as a predominantly sublittoral predator, and underscore the significance of littoral habitats as important foraging areas.

KEY WORDS: *Cancer magister* · Dungeness crab · Habitat · Preference · Intertidal · Foraging · Estuaries · Sublittoral
and form the basis for many fundamental theories in marine ecology. For example, patterns in the vertical distribution of species in littoral areas largely result from the interplay between interspecific competition (Connell 1961, Dayton 1971), pressure from sublittoral predators that access littoral areas during flood tides and strongly influence lower tidal extent of a species’ distribution and morphology (Connell 1970, Paine 1974), and physiological stresses that limit the upper tidal range of a species’ distribution (Foster 1971, Bertness 1981). Crabs are often the most prominent predators visiting littoral areas during flood tides (especially in estuarine systems), and consequently several authors have concluded that predation by migrating crabs is a mechanism that both reduces littoral prey populations and structures littoral benthic communities (Boulding 1984, Gee et al. 1985, Yamada & Boulding 1996).

The Dungeness crab *Cancer magister* is among the most conspicuous benthic predators in coastal waters of the NE Pacific. Throughout its range from Unalaska, Alaska, to Santa Barbara, California (Jensen 1995), the highest densities of *C. magister* occur in estuarine systems, which act as nurseries for young of the year (0+ age class) and subadult crabs (40 to 130 mm carapace width [CW], 1+ and >1+ age classes, Gunderson et al. 1990). Large, mobile subadult *C. magister* compose the majority of crab biomass and often range in density from 600 to 3000 crabs ha⁻¹ in sublittoral channels during low tide (Rooper et al. 2002). Not surprisingly, predation by *C. magister* is often assumed to impact populations of littoral prey (Asson Batres 1986, Iribarne et al. 1995, Byers 2002). However, evidence to support this claim is largely anecdotal, and only one study to date has attempted to examine patterns of intertidal movement by subadult *C. magister*. Stevens et al. (1984) compared crab abundance at one sublittoral and one littoral (at the level of mean lower low water, MLLW) location in Grays Harbor, Washington. The authors found circumstantial evidence that some crabs moved into littoral areas at high tide (i.e. crabs were captured in littoral areas at high tide and were absent at low tide), but no inferences were made concerning the significance of these migrations. Conversely, Holsman et al. (2003) modeled the energetic demand of subadult *C. magister* and available prey reserves in Willapa Bay, Washington, and concluded that crabs are obligated to migrate since sublittoral prey resources are insufficient to support observed abundances of crabs concentrated in shallow channels at low tide. It is estimated that subadult *C. magister* may derive between 80 and 90% of their daily energetic requirements from littoral sources in many areas of the estuary (Holsman et al. 2003).

There is no paucity of data concerning intertidal migrations by crabs in general (e.g. Dare & Edwards 1981, Abello et al. 1991, Chatterton & Williams 1994), yet to date, no study has examined the relative movement of these transient species into a range of distinct littoral habitats. Inherent structural and biotic differences between habitats likely influence patterns of habitat utilization, and clear identification of patterns is essential in order to accurately address ecological questions regarding predator-prey dynamics. Understanding littoral habitat use is particularly crucial for managing this important economic resource. There is currently a marked disparity between ecological studies that focus on predation by *Cancer magister* as important in structuring littoral communities, and management perspectives of life history that regard it as a primarily sublittoral predator.

In the present study we attempt to discern the spatial and temporal patterns of intertidal movement and habitat use by subadult *Cancer magister* in a typical large NE Pacific estuary, Willapa Bay, Washington. The following studies were designed to investigate the relative occurrence of subadult crabs in discrete littoral habitats in order to elucidate patterns of movement and habitat use by subadult *C. magister*. In particular we ask the following questions: (1) Do subadult crabs regularly migrate into littoral habitats on flood tides, (2) are intertidal migrations correlated with tidal range or current velocity, (3) what is the magnitude and extent of crab migrations, and (4) does the relative abundance of crabs vary among areas of complex epibenthic structure (‘on-bottom’ oyster culture and eelgrass beds) and unstructured habitats (bare mud or sand flats)?

**MATERIALS AND METHODS**

Traditional trawling and visual estimation methods were considered unsuitable for this study because of the cryptic nature of *Cancer magister*, the turbidity of mixed estuarine water, and the difficulty of sampling in shallow, highly-structured habitats (i.e. dense eelgrass beds or on-bottom oyster culture) during flood tides (Gibson 2003). Instead we employed 3 sampling techniques: baited trap arrays (box traps), underwater video observations, and acoustic telemetry. Each method varied in scope and resolution and the combination of all 3 allowed us to resolve patterns in habitat use across spatio-temporal scales. Baited trap arrays provide only coarse information on the relative use of habitats, but allow for broad-scale interpretation across a system over multiple years. Conversely, underwater video observations are spatially and temporally limited in scope, yet yield high-resolution observations of intertidal migrations. Acoustic telemetry is intermediate between the other methods, and provides
detailed information about selection and behavior across a range of habitats accessible to the individual crabs under study.

**Study site.** As a representative NE Pacific estuary, Willapa Bay, Washington (Fig. 1), has been the focus of numerous recent ecological studies (e.g. Armstrong et al. 2003, Rooper et al. 2003, Ruesink et al. 2003) and was chosen for our study because of the abundance of subadult crab resident within. Willapa Bay is a large bar-built estuary (31970 ha at mean high water) characterized by vast littoral flats. Semidiurnal tides that range up to 4 m drain over 50% of the shallow bay daily, exposing 17500 ha of tidelands at MLLW (Feldman et al. 2000). During ebb tides, water exits littoral areas through an extensive network of tidal tributaries and secondary side channels, which eventually drain into a deeper primary channel. In general, the abundance of many mobile species, including *Cancer magister*, is highest in secondary channels (Rooper et al. 2002) where gentle channel slopes and numerous tidal tributaries provide access during flood tides to a mosaic of highly productive littoral eelgrass *Zostera marina* beds and unstructured sand and mud habitats.

The estuary also supports substantial commercial aquaculture of Pacific oysters *Crassostrea gigas*, which have largely replaced the native oyster *Ostreola conchaphila*, and approximately 21% of tidelands are actively utilized for oyster production. Oysters are predominately cultured in 1 of 2 ways; cultch (inert oyster shell with newly settled oyster spat attached) may be strung on a line between ~1 m tall stakes (‘stake’ or ‘long line’ culture), or it may be dispersed directly onto littoral flats (‘on-bottom’ culture). The structural complexity of on-bottom culture is unique in the system and ecologically important as refuge for myriad small invertebrates, including 0+ crabs (Fernandez et al. 1993, Eggleston & Armstrong 1995). Some of the highest densities and biomass of a variety of species within the system occur in mature (2 to 3 yr) on-bottom oyster beds (K. K. Holsman unpubl. data).

**Box-trap surveys.** We conducted box-trap surveys at 4 locations across the bay in July 2002 and 5 locations in June 2004 in order to ascertain relative patterns of habitat use by subadult *Cancer magister* across the entire system. Locations were selected that had discrete patches of on-bottom *Crassostrea gigas* culture, *Zostera marina* beds, and unstructured littoral habitats (ULH) adjacent to secondary sublittoral channels. At each location, trap surveys were done on the 3 habitats simultaneously. Fifteen Fukui multi-species marine traps (model FT-100; 60 × 45 × 20 cm; 12 mm mesh) were baited with ~200 g of mackerel and set in standard arrays at each site with traps 50 m apart. In 2002 traps were immersed for 24 h at 4 locations (Fig. 1): Ellen Sands (ES), South Site (SS), Stony Point (SP), and Toke Point (TP). In 2004 soak time was limited to 12 h during nighttime high tides (when we observed the largest catches in our 2002 survey) and trapping occurred at 5 locations; ES, SS, SP, TP, and Nahcotta (NH). Upon retrieval of the traps, all crabs were identified to species, sexed, assessed for limb damage, and their CW was measured to the nearest 0.1 mm. Using the methods outlined by Zar (1999), traps were treated as replicates and differences in mean catch in each year were tested using a 2-way ANOVA with sampling location (ES, SS, SP, TP, and NH in 2004) and habitat type (oyster, eelgrass, and ULH) as factors. The analysis was followed by Tukey’s HSD pair-wise comparisons. A similar analysis was done to determine differences in the CW of crabs captured in 2002 and 2004, except crabs were pooled within habitats at each location. In all analyses, data were transformed in order to meet model assumptions of normality and equality of variance (Zar 1999). Additionally, the curve estimation procedure (SPSS for Windows 9.0.0) was used to explore the relationship between catches of *C. magister* and *C. productus*.

**Underwater video.** All video observations took place between July and September near TP (Fig. 1), where the variety of littoral habitats found within Willapa Bay is well represented. Dare & Edwards (1981) first

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**Fig. 1. Locations of sampling sites (diamonds) in Willapa Bay: Toke Point (TP), Stony Point (SP), Ellen Sands (ES), South Site (SS), and Nahcotta (NH)**
employed underwater video to monitor intertidal migrations of *Carcinus maenas* in Europe, and a modified design was adopted in the present study to observe *Cancer magister* traveling between littoral and sublittoral areas. Fyke (funnel shaped) ‘video gates’ constructed of 13 mm wire or plastic mesh were placed in littoral habitats approximately 10 m from the edge of the sublittoral channel. Gates were constructed of 15 m long panels that were 0.5 m high. Their orientation created two 15 m catchment zones; one facing the channel and the other a mirror image facing the littoral flat. A black and white underwater camera equipped with infrared LEDs (Atlantis Underwater Video Systems, AUC-125C) was mounted in a box at the constriction between the 2 catchment zones and tethered to a video cassette recorder on a nearby support boat. We made continuous recordings of crab activity throughout the observation period, and the net frequency of crabs migrating between sublittoral and littoral areas per 10 min interval was calculated for all video observations in 2002 and 2003. In 2002 the CWs of crabs were estimated as individuals passed beneath the camera and over a 10 cm grid (10 mm resolution) located on the floor of the camera box. In 2002 the gates were deployed on oyster, eelgrass, and ULH simultaneously, and the movement of crabs was recorded from 11:00 h July 25 through 10:00 h July 26. Concurrent measurements of water temperature and salinity were also taken. In 2003, observations were made on a predominately unstructured habitat for 5 consecutive days (August 1 to 6) but due to equipment failure, only observations made during daylight hours were considered. In addition to temperature and salinity, current velocity and direction were monitored using an Aanderaa Doppler Current Sensor (DCS 3900).

The mean CW of crabs migrating during the 2002 video observations were calculated and differences in mean CW were tested by pooling crabs within habitats and applying an ANOVA with habitat type (oyster, eelgrass, and ULH) as a factor. The analysis was followed by Tukey’s HSD pair-wise comparisons. In all analyses, data were transformed in order to meet model assumptions of normality and equality of variance (Zar 1999). To examine whether migrations were correlated with water velocity, the bivariate correlation procedure (SPSS for Windows 9.0.0) was used to calculate Pearson’s correlation coefficient for the net frequency of *Cancer magister* migrating between littoral and sublittoral areas per 10 min interval and the velocity of water movement (cm s$^{-1}$) during ebb and flood tides in 2003.

**Manual acoustic telemetry.** Ultrasonic telemetry has been used successfully to observe the movement patterns of various crab species (e.g. Wolcott 1995, Freire & Gonzalez Gurriaran 1998) and provides a method for observing in situ activity of subadult *Cancer magister* without dramatically altering crab behavior. Spatial and temporal patterns in movement among sublittoral and littoral microhabitats were recorded and the resulting observations were combined with habitat information to examine patterns of habitat selection. Telemetric observations commenced during the evening of August 17, 2002, when 13 male and female crabs measuring 90 to 110 mm CW were collected using baited box traps. Each crab was fitted with a uniquely coded 1.5 g sonotronics® acoustic tag. The tags were placed within a piece of surgical tubing and attached to the carapace of the crab using cyanoacrylate adhesive. Total time of tagging did not exceed 30 min, and all 13 individuals were released simultaneously. A directional hydrophone was used to manually locate each crab, and its geographic position was determined with a handheld global positioning system (GPS; Garmin Map76®).

Sequential observations complicate analysis of habitat use in telemetric studies, especially when subsequent positions are incorrectly treated as independent observations (Swihart & Slade 1985). In order to alleviate problems of serial correlation of observations, many authors have suggested separating observations of an individual’s position by sufficient time for the individual to utilize all potential habitats within their home range (e.g. Arthur et al. 1996). Since *Cancer magister* are rarely observed in littoral habitats at low tide and accordingly must return to sublittoral channels during ebb tides, we determined that separating observations by tidal period (~6 h) would give adequate time for crabs to move to any of the surrounding habitats and thus reduce the potential for serial correlation. Therefore, positions were recorded for all individuals once per tidal maximum or minimum (4 times daily) for 7 d, and then once per maximum high- and minimum low-tide (2 times daily) for 6 d. At this point crabs had emigrated from the study area or the tag batteries had expired.

Geographic habitat information was collected with a handheld GPS by tracing the perimeter of major habitat features, including unstructured habitats, marshes composed of *Spartina alterniflora*, eelgrass beds, sublittoral channels, and on-bottom and stake oyster culture areas. Habitat perimeters were compiled within a geographic information system (GIS) using ESRI ArcView 3.2® software. The positions of individual crabs were overlaid on the habitat GIS layer and queries were conducted to determine temporal and spatial patterns of crab movement. The animal movement extension for ArcView 3.2® (Hooge & Eichenlaub 1997) was used to calculate 95 (‘range’) and 50% (‘kernel’) distribution polygons for all crabs at each of the 4 tides. The net distance between 2 consecutive points was calculated for each observation (‘path
length') and a natural log transformation was used to meet model assumptions of normality and equality of variance. Transformed data were then compared with an ANOVA (Zar 1999) to determine if mean path lengths varied between daytime and nighttime high and low tides. Path length was assumed to be the linear difference between points and, although additional movement may have occurred, we did not record such activity in order to reduce serial correlation of observations.

Habitat selectivity is often determined by computing the ratio of habitat use to availability (Manly et al. 1993), or by using rank methods to compare use to availability (Aebischer et al. 1993). However, these methods are dependent on accurate estimation of available habitat and assume that availability is constant and unchanging. Habitat availability is often difficult to evaluate, and is particularly difficult to determine for migrating Cancer magister since tidal range may restrict or expand access to various littoral areas thus constantly changing the metric of habitat availability. Therefore, to assess habitat selection by subadult C. magister, and to resolve whether particular habitats were preferred or avoided, we employed methods outlined by Arthur et al. (1996) to calculate habitat selectivity when availability changes. Overall habitat selection indices ($\hat{b}_k$) were calculated for each individual crab based on the following equations (modified from Arthur et al. 1996):

$$\hat{b}_k = \frac{\sum_{j=1}^{H} O_{jk}}{\sum_{j=1}^{H} A_{jk} \hat{b}_j} \tag{1}$$

where for each individual $\hat{b}_k$ is the estimated selection index for a particular habitat $k$ from the set $j = 1 \ldots H$, and $H$ is the total number of habitats. $N$ is the number of tides on which the individual was located, $O_{jk}$ is either 0 or 1 and indicates the use of habitat $k$ during tide $t$, and $A_{jk}$ is the proportional availability (i.e. area) of habitat $k$ on tide $t$. $A_j$ is the total availability for the set of all habitats (i.e. total area). Initially the values for $\hat{b}_k$ were set to $1/H$ and used to solve Eq. (1) for $\hat{b}_k$; values for $\hat{b}_j$ were then calculated through numerical iteration such that:

$$\hat{b}_j' = \frac{\hat{b}_k}{\sum \hat{b}_k} \tag{2}$$

until $\sum \hat{b}_k = 1$ and $\hat{b}_j'$ for each habitat.

In this study the proportional availability of a habitat ($A_{jk}$) was calculated as the area of habitat $k$ divided by the sum of the area of all available habitats within the range of the individual during a given tide $t$. Since the range of an individual crab was constantly changing, we defined total available habitat separately for each crab location as a circle centered on the crab’s previous location (See Fig. 2 in Arthur et al. 1996). The radius of the circle was set to 370 m, which was the maximum distance a crab was likely to move in a single tide; only 2 crabs moved further than 370 m on a few occasions (0.4%). In some instances, crab locations could not be obtained for 1 or 2 consecutive tides and so the radius of the circle was set to 640.5 and 875 m, respectively. These distances represented the maximum distance a crab was observed moving when 1 or 2 previous tidal locations were missed, and encompassed 100% of the crab positions in these cases. The selection indices were determined for each individual crab for daytime and nighttime high-tide positions, and habitat preference was indicated by indices greater than the expected index value if habitat use was random ($1/H$, Arthur et al. 1996); this value was 0.2 in the present study.

**RESULTS**

**Box-trap surveys**

The mean number of Cancer magister caught per trap in both the 2002 (24 h) and 2004 (12 h) box trap surveys varied significantly among littoral oyster, eelgrass, and ULH habitats ($F_{2,142} = 60.4$, $p < 0.001$ and $F_{2,206} = 44.1$, $p < 0.001$, respectively), with highest catches occurring in ULH and lowest catches in oyster habitats (Fig. 2). The total catch from traps placed on ULH composed 48% of all C. magister caught in the 2002 trapping survey (6424 crabs total) and 58% of all crabs caught in 2004 (5002 crabs total). Mean catch of C. magister also varied significantly between locations in both 2002 ($F_{3,142} = 66.2$, $p < 0.001$) and 2004 ($F_{3,206} = 9.2$, $p < 0.001$), and was highest at TP and SS locations (Fig. 3). The combined catches from these locations composed 77 and 52% of all crabs captured in 2002 and 2004, respectively.

Baited traps predominately captured subadult Cancer magister (≥1+ yr) with CW ranging from 50 to 165 mm. In 2002 and 2004 the mean size of crabs varied slightly across the bay ($F_{3,6362} = 239.8$, $p < 0.001$ and $F_{4,1653} = 134.9$, $p < 0.001$, respectively) with the smallest crabs found farthest from the mouth and the largest crabs found in the northern part of the bay. The mean size of crabs caught in 2002 also varied significantly among habitats ($F_{2,6362} = 13.1$, $p < 0.001$), with the largest crabs caught in oyster beds and the smallest caught in eelgrass and unstructured habitats (82.4 ± 0.5, 78.6 ± 0.4, and 77.0 ± 0.3 mm CW, respectively). However, in the 2004 trapping survey CW did not differ significantly among habitats ($F_{2,1653} = 1.7$, $p = 0.175$).
Cancer productus was also abundant in the 2002 and 2004 box-trap surveys. Mean catches of Cancer productus varied significantly among locations, with the largest average catches in 2002 and 2004 at ES and SP (Fig. 3) in the northern part of the estuary ($F_{3,134} = 121.9$, $p < 0.001$, and $F_{4,206} = 65.3$, $p < 0.001$, respectively). Mean catches of Cancer productus also varied significantly among habitats in 2002 ($F_{2,134} = 48.7$, $p < 0.001$) and 2004 ($F_{2,206} = 223.2$, $p < 0.001$), with the highest catches in oyster and the lowest catches in ULH (Fig. 2). Catches of Cancer magister were negatively correlated with catches of Cancer productus in baited traps (Pearson’s $r = –0.273$, $p = 0.003$), and the relationship was most pronounced at SP and ES locations where on-bottom oyster culture was 3 yr old and percent shell cover was greater than at other locations. The average size of Cancer productus caught in baited traps did not vary significantly among habitats in 2002 ($F_{2,1376} = 1.6$, $p = 0.21$ and 2004 ($F_{2,521} = 2.5$, $p = 0.02$), or locations in 2002 ($F_{3,1376} = 1.4$, $p = 0.24$). In contrast, the mean size of Cancer productus captured in baited traps in 2004 varied significantly across the bay ($F_{2,521} = 20.0$, $p < 0.001$), with the largest crabs captured in the northern part of the system. Approximately 58% of all Cancer magister and Cancer productus crabs caught in baited traps in 2002 were males, and for both species males were significantly larger than females ($F_{1,6362} = 193.5$, $p < 0.001$ and $F_{1,1376} = 358.6$, $p < 0.001$, respectively).

### Underwater video

The 2002 video observations showed marked temporal patterns in the magnitude of tidal migrations of Cancer magister into and out of all 3 habitats. In total, over 1300 Cancer magister migrated through video gates between sublittoral and littoral areas during the 24 h observation period, despite a missing interval due to equipment failure (21:00 to 05:00 h on the eelgrass site, see Fig. 4). During the period that all 3 cameras were recording movements (11:00 to 09:00 pm), relatively more crabs migrated into the ULH site than the eelgrass (66% of the ULH value) or on-bottom oyster sites (50% of the ULH value). Similarly, migrations out of the ULH site during the ebb tide were nearly twice as large as migrations out of the eelgrass and oyster sites combined. Significantly larger crabs moved into and out of the oyster site ($F_{2,1305} = 4.8$, $p = 0.009$) than migrated through the eelgrass and unstructured sites (83.8 ± 1.8, 77.3 ± 0.7, and 79.1 ± 0.9 mm CW, respectively).

Video observations in 2002 and 2003 revealed that subadult Cancer magister move in accord with tidal periodicity (Figs. 4 & 5, respectively). Peaks in migration between littoral and sublittoral areas occurred during maximum exchanges when current velocity was highest, often just prior to tidal maxima or minima.
In 2003 the magnitude and direction of crab migrations was significantly related to water current speed and direction on both flood ($r^2 = 0.232$, $p = 0.004$) and ebb tides ($r^2 = 0.224$, $p = 0.004$). While crabs moved into littoral flats during daytime and nighttime flood tides, emigration out of littoral areas was most pronounced during the morning ebb following a nighttime high tide (Figs. 4 & 5). The magnitude of this migration decreased over the observation period as the magnitude of tidal exchanges diminished, and the smallest ebb migration was observed on August 5 when the tide no longer dropped below 0 m MLLW (Fig. 5).

**Manual ultrasonic telemetry**

Telemetric observations of subadult *Cancer magister* movement revealed that subadult crabs exhibit diurnal and tidal patterns in habitat use and movement. Male and female crabs showed similar patterns of movement and were generally more active at night, particularly during high tides, as indicated by the differences in average home-range size and maximum distances traveled during daytime high (DH) and nighttime high (NH) tides (Fig. 7, Table 1). In most measures of activity (home range, path length, and rate of movement), crabs were 2- to 3-fold more active during night than day, irrespective of tidal stage (Fig. 7, Table 1). During NH tides crabs expanded their ranges to include portions of the littoral flats (Fig. 7), and in a few cases, individuals repeatedly migrated more than 1.2 km round-trip (Fig. 8). Crabs moved significantly further during NH tides than DH tides ($F_{3,135} = 12.6, p < 0.001$). More than half (65% ± 0.26) of all positions occurred in littoral habitats during NH tides (Fig. 9). *Cancer magister* generally remained in sublittoral areas during daytime tides (DH and DL), and at nighttime low (NL) tides were usually in deeper channels (>2 m), although some individuals occasionally moved into

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**Fig. 4.** *Cancer magister*. Underwater video observations from 2002 of crabs migrating between sublittoral channels and littoral habitats: eelgrass, oyster, and unstructured littoral habitats (ULH). Positive and negative values represent the number of crabs observed moving into and out of littoral habitats, respectively. The solid line shows tidal changes in water depth (m) relative to mean lower low water (MLLW). Diagonal shading shows periods of darkness, and 'ND' indicates periods when the camera was obstructed and observations could not be made.

**Fig. 5.** *Cancer magister*. Daytime underwater video observations of crabs migrating between sublittoral channels and an unstructured littoral habitat (ULH) during a continuous 5 d observation period in 2003. Positive and negative values represent the number of crabs observed moving into and out of the littoral habitat, respectively. The solid line expresses tidal changes in water depth (m) relative to mean lower low water (MLLW); the horizontal dotted line indicates the approximate depth at which the camera is exposed. Dates in top left corner given as mm/dd/yy; times on x-axes are given in 24 h clock format.
sublittoral eelgrass beds (Fig. 7a,c). Analysis of habitat selection revealed that selection indices ($\hat{b}_k$) were greater for sublittoral channels than other habitats during DH tides. However, selection indices during NH tides were greater for sublittoral channels and ULH than for the other remaining habitats (Fig. 10). Additionally, the values for sublittoral habitats and ULH were greater than 0.2, suggesting that subadult crabs selected these habitats during NH tides (Fig. 10).

**DISCUSSION**

A variety of brachyuran crabs undertake intertidal migrations at some point in their life history, including *Callinectes sapidus* (Hamilton 1976), *Cancer productus* (Robles et al. 1989), *C. novaezelandiae* (Chatterton & Williams 1994), *Carcinus maenas* (Dare & Edwards 1981, Hunter & Naylor 1993), and *Liocarcinus holsatus* (Abello et al. 1991). The results of the present study suggest that subadult *Cancer magister* regularly migrate into littoral habitats as well, particularly during nighttime high (NH) tides. These highly directed migrations are often extensive, with some individuals moving up to 600 m into littoral flats (1.2 km round-trip) within a single tide. Regular intertidal migrations to preferred unstructured littoral habitats (ULH) appear to be an integral component in the life-history of subadult *C. magister*, a finding

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<th>Table 1. <em>Cancer magister</em>. Path lengths (PL) or distance traveled h$^{-1}$ (rate) by crabs during nighttime high (NH) and daytime high (DH) tides. Note that values for path length and rate reflect average movement preceding or following a given tidal period</th>
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Fig. 6. *Cancer magister*. Tidal patterns in depth (m) relative to mean lower low water (MLLW), current (cm s$^{-1}$), and frequency of migrating crabs over a 2 d period in 2003; the horizontal dotted line indicates the approximate depth of camera exposure. Diagonal shading indicates periods of darkness when video quality was low and the frequency of crab movement could not be measured accurately. x-axes are given as times in 24 h clock format, and dates given as mm/dd/yy.

Fig. 7. *Cancer magister*. Locations of tagged crabs (filled circles) during the 2 wk 2002 telemetry study at Toke Point in Willapa Bay, Washington, grouped by tidal period: daytime high tides (DH), nighttime high tides (NH), daytime low tides (DL), and nighttime low tides (NL). The 95% habitat-use polygon (dotted outline) indicates the home range of crab movement, and 50% (diagonal shading) polygon represents their home kernel. Values in parentheses represent the area (ha) of the 50 and 95% habitat-use polygons, respectively.

EEL: eelgrass bed, OYS: on-bottom oyster culture, STK: stake oyster culture, SUB: sublittoral, ULH: unstructured littoral habitat
that dramatically alters the perception of the crab as a primarily sublittoral species in estuarine nurseries.

In NE Pacific estuaries, subadult *Cancer magister* are most abundant in secondary side channels (Rooper et al. 2002) that are surrounded by extensive littoral habitats. Results of a bioenergetics model have suggested that littoral prey consumed during intertidal forays provide a critical subsidy for populations of *C. magister*. These are estimated to fulfill 80 to 90% of daily energetic demands, which cannot be satisfied in sublittoral channels, where prey biomass is too low to support crab production and agonistic interactions further reduce foraging time (Holsman et al. 2003). The results of the present study are consistent with model predictions, since both telemetry and underwater video observations showed that crabs regularly migrate into littoral habitats.

Intertidal migrations may allow crabs to increase foraging opportunities and decrease agonistic activity. Previous studies of estuarine systems have found that prey biomass is significantly greater in littoral areas than in sublittoral channels (See Table 1 in Gunderson et al. 1990), and migrating species are thought to benefit from increased foraging opportunities resulting from differences in prey abundance (e.g. Gibson 2003). Stevens et al. (1984) correlated the abundance of subadult *Cancer magister* with intertidal migrations of a common prey species, sand shrimp *Crangon* spp., and similarly during our underwater video observations we witnessed occasional predation events on juvenile English sole *Parophrys vetulus* that were migrating into littoral areas (K. K. Holsman pers. obs.). Ongoing work is examining the gut content and fullness of crabs migrating into and out of various littoral habitats to determine whether increased prey availability and biomass drive intertidal migrations by subadult *C. magister*; preliminary results indicate that gut fullness is greatest for crabs moving out of littoral habitats during ebb tides (K. K. Holsman unpubl. data).
Additionally, subadult *Cancer magister* compete aggressively while foraging, and larger crabs often displace and occasionally injure smaller ones (P. S. McDonald pers. obs.). Sublittoral crowding and agonistic encounters may be reduced as crabs disperse into littoral habitats to feed. Kaiser et al. (1990) found that smaller *Carcinus maenas* forage in littoral habitats at high tide to avoid agonistic encounters, and subadult *C. magister* may similarly limit encounter time with conspecifics and reduce overall competition by migrating with the tide and increasing their foraging area. Further studies should explicitly examine apparent tidal patterns in the behavior of subadult *C. magister*, particularly in relation to prey availability and agonistic activity.

Our telemetry work shows that subadult *Cancer magister* activity may be influenced by circadian periodicity since, despite access at high tide, few tagged individuals moved into littoral areas during daylight hours. Instead, most crabs were relatively sedentary, but activity increased between dusk and dawn when crabs often migrated to littoral habitats or moved within sublittoral channels. Although we were unable to detect clear evidence of circadian rhythms in video observations, and neither of our trapping surveys explicitly compared day and night catches, diel patterns of movement have been observed in *C. magister* from Grays Harbor, Washington (Stevens et al. 1984). Many other crabs are known to have daily rhythms as well, and nocturnal peaks in activity are common in the *Cancer* clade as a whole (see review by Novak 2004).

Regardless of the purpose, movements into littoral habitats are assumed to confer an advantage to those individuals that migrate. However, the benefits may be offset if the energetic cost of migration is too high (Gibson 2003). We suggest that subadult *Cancer magister* may rely on tidal currents to facilitate intertidal migrations. There is a strong correlation between net movement of crabs and water velocity and direction in the 2003 video data, and recordings revealed that migrating crabs augment walking with swimming or passive gliding just above the substrate. These behaviors may also allow *C. magister* to access distant littoral areas more quickly. Considering that subadult crabs can only sustain moderate walking speeds (3.5 m min⁻¹) for short periods of time (McMahon et al. 1979, DeWachter & McMahon 1996), it is likely that the tagged crabs we observed traveling more than 600 m into littoral flats (1.2 km round-trip) utilized strong tidal currents to do so. During flood tides, tagged individuals often moved into littoral areas along one vector, and subsequently followed an inverse trajectory during the ebb tide to return to the same spot in a sublittoral channel. Conceivably, crabs use a combination of learned migratory paths and directional water movement to navigate in and out of littoral habitats; thus, migration patterns are likely influenced by the magnitude and duration of tidal events. The strong tidal currents associated with extreme spring tides may not only act as directional cues for migration, but may also facilitate the movement of crabs into littoral areas, and migrations may be more extensive in areas where currents are strongest.

In addition to the patterns of activity described above, subadult *Cancer magister* exhibit selective use of available habitats. In the present study, crabs utilized ULH more frequently and to a greater extent than highly structured littoral eelgrass or oyster areas, and this pattern was consistent across all 3 methods of observation. Habitat complexity strongly influences predator-prey dynamics by directly reducing predator foraging efficiency (Crowder & Cooper 1982, Fernandez et al. 1993). In NE Pacific estuaries, highly structured on-bottom oyster culture and eelgrass provide important refugia for small prey (Thayer & Phillips 1977, Iribarne et al. 1995, Dumbauld et al. 2000) since the increased physical relief of structurally complex areas results in lower predation rates. The cryptic nature of benthic organisms living in structured habitats likely increases search and handling time and lowers overall foraging efficiency of predators, such as subadult *C. magister*.

Structural complexity may also affect the foraging efficiency of predators by obstructing their movements. Vegetative structure has been shown to impede the movements of *Callinectes sapidus* migrating into littoral areas during flood tides (Kneib 1995), and patch size has been shown to impact abundance of prey by impeding predators (Eggleston et al. 1999, Hovel & Lipcius 2001). Predation impacts are typically higher along the edges of eelgrass or oyster beds than within patch centers (Iribarne et al. 1995, Fagan et al. 1999), a pattern attributed to the inability of many predators to penetrate structured habitats. We observed similar patterns in the present study; traps set at the edge of eelgrass and oyster beds caught more *Cancer magister* than ones placed within the center. Furthermore, when crabs in our telemetry study occasionally penetrated eelgrass beds during high tides, they frequently could not be located at subsequent low tides, and may have been stranded when water receded. Stranding is particularly stressful for larger *C. magister*, which are more sensitive to temperature fluctuations than smaller juveniles (Gutermuth & Armstrong 1989). We suggest that the risk of stranding during low tide coupled with reduced foraging efficiency in complex habitats likely deter most subadult *C. magister* from using eelgrass and oyster beds, despite the high density of potential prey species they support.
Unstructured habitats dominate littoral areas of Willapa Bay and support known prey of Cancer magister (Gunderson et al. 1990, Ferraro & Faith 2004). Prey species often occur at lower densities in ULH than other habitats (Ferraro & Faith 2004), but increased accessibility may offset lower abundances. Subadult C. magister may also benefit from greater mobility in ULH than structurally complex habitats. Littoral prey species are not distributed evenly (e.g. Clark et al. 2000), and foraging success may be improved by increasing search efficiency over large areas. We observed tagged crabs returning to particular areas of ULH on successive tides, which suggests that once a patch is discovered it may be exploited repeatedly. Our results corroborate previous studies that implicate C. magister as an important predator in unstructured littoral areas (Fernandez et al. 1993, Yamada & Boulding 1996, Fernandez 1999).

While on-bottom oyster culture is critical to young-of-the-year Cancer magister (Dumbauld et al. 1993, 2000, Fernandez et al. 1993), subadult crabs in our study utilized these areas far less frequently. This highlights an important ontogenetic shift in habitat use by C. magister, and may partially result from changes in foraging efficiency and physiology with ontogeny. However, interactions with a congener, C. productus, likely contribute as well. It seems that C. productus are resident within oyster beds of Willapa Bay; we never observed these crabs migrating in video recordings, and they were often found buried in littoral areas at low tide, especially in dense beds of mature oysters (K. K. Holsman pers. obs.). Adult C. productus are aggressive predators and competitors, known to capture and consume smaller crabs (Jensen 1995, Hunt & Yamada 2003) and displace other decapods while foraging. The strong inverse relationship between catches of these species in the present study suggests that subadult C. magister avoid areas occupied by C. productus. Subadult crabs may reduce the risk of predation or injury by utilizing ULH, since large C. productus are found at relatively low densities outside structured habitats. Ongoing experiments will elucidate the nature of competitive interactions between these species and the potential for C. productus to exclude C. magister from oyster beds (K. K. Holsman unpubl. data).

The pattern of habitat use described above for Cancer magister highlights the importance of small differences in biotic and abiotic aspects of microhabitats, and implies that factors other than prey density strongly affect habitat use by a migrating predator. The present study also adds to an increasing body of literature that emphasizes the importance of biotic, physical, and chemical cues to patterns of predator movement (Micheli 1997, Finelli et al. 2000, Alofs & Polivka 2004). Powers & Kittinger (2002) showed that flow dynamics strongly affect odor-mediated movements, and that the specific response of predators to changes in water velocity greatly impact predator-prey dynamics. Similarly, factors such as habitat fragmentation, physical structure, competition, and prey distribution strongly influence these interactions (Paine & Levin 1981, Wootton et al. 1996, Eggleston et al. 1998).

We have shown that tidal influence is also critical when examining the behavior of foraging predators. Incorporating these realized patterns of movement into models of habitat selection is necessary to clarify key mechanisms governing the structure of littoral communities in estuarine systems.

Our findings suggest that unstructured habitats may be the primary foraging ground for migrating crabs and are therefore critical to crab production in these systems. However, a variety of human activities, including ship-canal dredging, upstream sediment loading, mariculture practices, and pesticide use, are modifying littoral habitats at an increasing rate (Wainwright et al. 1992, Simenstad & Fresh 1995, Feldman et al. 2000, Dumbauld et al. 2001). These perturbations threaten to impact populations of Cancer magister either directly, through mortality of crabs in littoral habitats during high tides, or indirectly, through changes in the abundance of littoral prey. Littoral areas are also affected by invasions of non-indigenous species. In recent years, smooth cordgrass Spartina alterniflora and Japanese eelgrass Zostera japonica have begun to colonize ULH in several estuaries of the NE Pacific (Posey 1988, Feist & Simenstad 2000), and their impact on populations of C. magister is poorly understood. Widespread alteration of ULH may negatively impact overall production of C. magister, particularly in areas adjacent to subtidal channels. Considering the importance of estuarine populations to the stability of coastal C. magister fisheries (Armstrong et al. 2003), decisions affecting littoral habitats should be approached cautiously. Far less conservation effort and attention have been directed toward ULH, including mud and sandflats, than other critical estuarine habitats such as eelgrass beds, oyster reefs, or emergent marshes. Regional declines in C. magister populations may be related to widespread loss of ULH in historically productive estuaries such as San Francisco Bay, California. Future restoration and conservation efforts should consider the ecosystem value of ULH, which is essential to sustaining populations of C. magister and myriad other species in coastal estuaries.

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