INTRODUCTION

Marine mammals that occupy high trophic positions in marine ecosystems can have important top-down effects on those ecosystems (Bowen 1997). As the apex (non-human) marine predator, the killer whale *Orcinus orca* is capable of preying on a great variety of species and has a remarkably diverse diet. The list of prey taken by this cosmopolitan predator includes more than 120 species of fishes, cephalopods, sea turtles, sea birds, mustelids, pinnipeds and cetaceans (Martinez & Klingenhammer 1970, Jefferson et al. 1991, Matkin & Saulitis 1994, Fertl et al. 1996, Similä et al. 1996, Ford et al. 1998). With such a diversity of prey types, killer whales can potentially influence ecosystem structure and function at a variety of levels. For example, it has recently been proposed that predation by killer whales caused the depletion of several marine mammal populations in the North Pacific in the post-whaling era (Springer et al. 2003). A variety of arguments have been raised against this hypothesis (e.g. DeMaster et al. 2006, Mizroch & Rice 2006), which emphasise how much uncertainty exists regarding the dynamics of killer whales and their prey. Developing an understanding of the factors involved in prey selection, as well as the extent of foraging specialisation or flexibility, will be needed if we are to better understand the roles played by killer whales in marine ecosystems. Such knowledge is also important for conservation of both killer whale populations and their prey.

Although a generalist as a species, different killer whale populations can have strikingly divergent foraging specialisations. For example, in coastal waters of the NE Pacific Ocean, 2 sympatric, genetically-distinct killer
whale ecotypes, so-called ‘residents’ and ‘transients’, feed almost exclusively on fish and marine mammal prey, respectively (Bigg et al. 1990, Baird & Dill 1995, Ford et al. 1998, 2000, Hoelzel et al. 1998, Ford & Ellis 1999, Barrett-Lennard 2000). Killer whale populations in other regions, such as Norway, Argentina and Antarctica, also specialise on particular prey types, and often employ elaborate foraging tactics in order to do so (Lopez & Lopez 1985, Similä & Ugarte 1993, Baird 2000, Pitman & Ensor 2003). Such specialisations appear to represent behavioural traditions that are passed across generations by social learning (Guinet & Bouvier 1995, Ford et al. 1998, Saulitis et al. 2000), and have been described as cultures (Rendell & Whitehead 2001).

Behavioural traditions may determine the overall type of prey that is acceptable to a particular killer whale population and the foraging tactics employed for prey capture, but other factors are likely to play important proximate roles in prey selection. Prey choice by predators is influenced by rates of encounters with a prey species and its profitability, which is determined by the prey item’s net energy value and the amount of time needed to catch and handle it (Stephens & Krebs 1986, Scheel 1993, Bowen et al. 2002). Factors that are important in prey choice in killer whale populations are poorly known, but long-term studies in the NE Pacific have provided some insights. Mammal-hunting transient killer whales in this region prey on at least 9 species of marine mammals, including seals, sea lions, porpoises, dolphins and baleen whales, but appear to prefer small species such as harbour seals Phoca vitulina and harbour porpoises Phocoena phocoena, which are common year-round in the whales’ range (Haley 1986). These species are relatively easy to capture and kill, and have a low probability of causing injury to the attacking whales (Baird & Dill 1995, Ford et al. 1998, 2005). Sympatric, fish-eating resident killer whales show movement patterns in nearshore waters that are closely associated with high densities of migrating salmon (Heimlich-Boran 1986, Guinet 1990, Nichol & Shackleton 1996). Nichol & Shackleton (1996) found positive correlations between the seasonal occurrence of resident killer whale populations and 3 of the most abundant salmonid species, pink salmon Oncorhynchus gorbuscha, sockeye salmon O. nerka and chum salmon O. keta, off NE Vancouver Island, British Columbia. Ford et al. (1998) confirmed that salmonids were the predominant food of resident killer whales in these waters by identifying prey species from scales and tissue fragments collected from kill sites. An unexpected finding, however, was that sockeye, pink and chum salmon, despite being correlated with the occurrence of killer whales in this area, formed the minority of prey samples. Instead, chinook salmon O. tshawytscha, one of the least common salmonids during the summer migration period, represented almost two-thirds of prey items identified.

The disproportionate occurrence of chinook salmon in prey samples relative to its low abundance led Ford et al. (1998) to suggest that resident killer whales may forage selectively for this species over other available salmonids. Chinook is the largest salmonid and has a relatively high lipid content, features that may be desirable for foraging killer whales. However, Ford et al. (1998) also raised concerns that potential biases in their prey sampling method may have led to over-representation of chinook and under-representation of other species in samples from predation events. Chief among these was the possibility that chinook, being larger than other salmonids, may be broken up by whales prior to being eaten, thus shedding more scales than smaller salmonids in the process (Ford et al. 1998). However, without knowledge of the details of prey handling and consumption of salmonid and other fish species by resident killer whales, it was not possible to evaluate the significance, if any, of this potential bias. It was concluded that resident whales may have a preference for chinook, but the extent of their selectivity for this species remained uncertain (Ford et al. 1998). These potential biases as well as small sample sizes also prevented Ford et al. (1998) from examining potential differences in prey selection among different resident communities or social groups, as suggested by Nichol and Shackleton (1996), or by different sex or age classes, as suggested by Bain (1989).

In order to address these and other gaps in the current knowledge of resident killer whale diet and prey selection, we undertook field studies of foraging behaviour and feeding by resident killer whales from 1997 to 2005 to build upon the data presented in Ford et al. (1998). In particular, field efforts from 2003 to 2005 were focused on documenting the detailed aspects of prey capture and handling to assess the validity of using prey fragment sampling to interpret dietary preferences. In this report, we present new information on the frequent occurrence of cooperative foraging and prey sharing in resident killer whales, and the implications of this behaviour with respect to the use of prey fragments as indicators of prey selection. Extensive sampling of feeding events also allowed us to evaluate prey selection quantitatively with regard to prey species availability, as well as by geographical location, group membership, and age and sex class of foraging whales.

MATERIALS AND METHODS

Study area and population. Field studies on the life history, social organisation, acoustic behaviour, and population genetics of killer whales in British Colum-
bia have been undertaken since 1973 (Bigg 1982, Bigg et al. 1987, 1990, Olesiuk et al. 1990, Ford 1989, Ford et al. 1998, 2000, Barrett-Lennard 2000). These long-term studies have relied extensively on the photographic identification of individuals from natural markings (Bigg et al. 1987, Ford et al. 2000). In these waters, 2 communities of resident killer whales, ‘northern residents’ and ‘southern residents’ can be found in all months of the year, but are observed mostly during May to November. The northern resident community is found typically from mid Vancouver Island to SE Alaska, and the southern resident community off the southern half of Vancouver Island and in the inland waters of Washington State. Whales from the 2 communities have not been seen to associate, despite extensive overlap in their ranges (Ford et al. 2000). Large aggregations of resident killer whales can be found in certain coastal locations during summer. The whales greatly reduce their use of these locations in winter and spring, and their range during this period is poorly known (Ford et al. 2000, Wiles 2004). The northern and southern resident communities contained 219 and 87 individuals, respectively, in 2004 (authors’ unpubl. data, K. Balcomb, Center for Whale Research, Friday Harbor, Washington, unpubl. data).

The basic social unit of resident killer whales is the ‘matriline’, which consists of individuals that are closely related by matrilineal descent. Matrilines generally contain an old female, or matriarch, and 1 to 3 generations of her descendents of both sexes. Dispersal of individuals from the matriline is extremely rare (Ford et al. 2000). Matrilines are comprised of an average of 6 members (±0.59 SE, range 1 to 26, n = 50). Resident killer whales typically travel in ‘pods’, which consist of related matrilines that spend the majority of their time together (Bigg et al. 1990). Although some pods originally described in the 1970s and 1980s have maintained their stability, others have split in recent years (Ford et al. 2000). ‘Clans’ are comprised of pods and matrilines that have descended from a common matrilineal ancestor and have a unique set of shared dialects. The northern resident community consists of 3 clans, A, G, and R, while the southern resident community is made up of a single clan, J. Members of the northern resident clans frequently associate with one another.

Field effort and procedures. Data on predation by resident killer whales in British Columbia have been collected each year since 1974. Data collected from 1974 to 2002 consisted mostly of opportunistic observations of feeding events and collection of prey fragments from the vicinity of kills. Effort varied widely according to changing research objectives, but predation studies were given higher priority after 1990 (Ford et al. 1998, 2000). The results of these studies up to 1996 were reported by Ford et al. (1998), and some of these data are included in the present study. In 2003 to 2005, field studies were dedicated to systematically documenting foraging behaviour and collecting predation data, in addition to conducting the annual census of individuals by photo-identification (Bigg et al. 1987, Ford et al. 2000). A total of 152 field days were devoted to these objectives in 2003 to 2005.

Field studies from 1974 to 2002 were conducted using a variety of vessels from 5 to 20 m in length. In 2003 to 2005, dedicated focal-individual and focal-group studies were undertaken mainly from a 10 m long power vessel. When whales were encountered, individuals were observed visually or photographed to determine the identity of matrilines present. Photographic identification procedures are described in Bigg et al. (1987) and Ford et al. (2000). Once the identity of killer whales present in the encounter was established, effort was directed to documenting foraging behaviour and collecting scales and tissue fragments from prey killed during feeding events. The activity state of the whales was determined from surfacing and dispersion patterns (see Ford 1989 for definitions of activity states). When foraging, whale groups typically spread out over several square kilometres, with individuals and subgroups swimming and diving independently but travelling generally in the same direction. Surfacing whales were observed by eye or binoculars for signs of prey pursuit or capture. When apparent feeding was observed, the site of the kill was approached quickly, while taking care to avoid disturbing the whales, in order to determine identity of the individual(s) involved and to search for prey fragments in the water. Whether or not feeding was confirmed, the individual or subgroup was then followed at distances of 50 to 150 m to document subsequent feeding events. Focal individuals and subgroups (Altmann 1974, Mann 1999) that were actively feeding were followed for as long as the activity continued or until focal animals joined other groups and could no longer be followed individually.

The behaviour of focal individuals and subgroups was monitored closely and constantly during feeding sessions. Particular attention was given to direction of travel, regularity of dive durations, and extent of subgroup cohesion, as changes in these variables often signalled a feeding event. Individuals or subgroups suspected to have captured a prey item were approached to within 20 m to observe prey handling and consumption. To collect evidence of feeding, the surfacing locations of the feeding whale or subgroup were also examined for prey fragments at the surface or in the water column. The principal observer, who was also the boat operator, was situated approximately 4 m above the water surface on the flying bridge of the research vessel. This position afforded a high-angle view
into the water as the boat was manoeuvred. A second observer stood on the vessel’s bow, holding a fine-mesh dip net (mesh size approximately 1 mm) with 4 m telescoping handle, and also searched for fragments. When fish scales or pieces of tissue were seen, the boat was immediately stopped and the net was deployed to retrieve the fragments. Fragments were collected mostly at depths of 0 to 2 m, but occasionally as deep as 3 to 4 m in calm conditions with good water clarity. Generally, only a subsample of the many scales and tissue fragments seen in the water were collected from each kill. Rain, winds greater than 10 knots, and high water turbidity reduced the success rate of fragment location and collection. Prey fragments and scales were stored in 10 ml vials containing 95% ethanol. Until 2004, no effort was made to collect tissue samples from fish kills when scales were available for collection, but both were collected systematically in that year and in 2005. The date, time, and geographical position (from a GPS instrument) of the feeding event was recorded, as well as the identity of the individual making the kill and other whales involved in the prey capture or consumption. If individuals could not be identified, their age/sex class was noted whenever possible.

**Prey species identification and ageing.** Many species of fishes are readily identifiable at a distance by an experienced observer, but salmon species can be difficult to distinguish without close examination. Although Ford et al. (1998) included salmonid identifications based on field observations, in the current analyses we included only positive species identifications of salmonids based on scales or tissue samples to eliminate this potential source of error. Fish scales were analysed by the Fish Ageing Laboratory at the Pacific Biological Station (Department of Fisheries and Oceans, Nanaimo, British Columbia, Canada) to determine species identity and age. Age was designated using the European method, whereby years in freshwater after hatching and years in marine water are identified and separated by a decimal point (Groot & Margolis 1991), and age class was assigned according to a standard 1 January birth date. Age class was thus calculated by summing the freshwater and marine years of the European age and adding 1.

Scales that could not be positively identified to species and tissue samples collected from feeding events were submitted to the Molecular Genetics Laboratory at the Pacific Biological Station for species identification using analysis of microsatellite DNA. The methodology of these analyses is described in Withler et al. (2004).

**General analyses.** Statistical analyses were conducted using SPSS Version 11.0. The significance of differences between 2 group means was tested using Mann-Whitney U-tests, and among different frequencies of occurrence by chi-squared ($\chi^2$) goodness-of-fit tests. The standard error (SE) of the mean is given as a measure of variability.

**RESULTS**

A total of 529 feeding events were observed during 206 encounters with resident killer whales between 1974 and 2005. Of these, 340 (63%) were recorded during dedicated feeding studies in 2003 to 2005. Prey samples (tissue or fish scales) were recovered from 449 feeding events (85%), while the remaining 15% were documented by observation only (Table 1). A mean of 4.8 scales per feeding event ($\pm$0.24 SE, range 1 to 28) were collected from 428 of the 529 feeding events (81%). During 2004 to 2005, when tissue as well as scales were collected systematically when both were available, tissue fragments were retrieved from 115 of 249 (46%) feeding events.

Feeding events were documented from May to December. A total of 463 (87.5%) feeding events involved northern residents, and 66 (12.5%) events involved southern residents. All 4 resident clans and all but 1 of the 19 resident pods in the northern and southern communities are represented in this data set (the exception is W1 pod; Ford et al. 2000). Samples were collected from most regions of the coast, but two-thirds came from waters off NE Vancouver Island, an important core area for northern residents (Fig. 1, Table 2; see also Ford et al. 2000). Kills made during a total of 274 feeding events could be attributed to whales of known age or sex class. The monthly distribution of these is shown in Table 3.

**Foraging behaviour and prey fragment sampling**

Dedicated studies of foraging behaviour of resident killer whales in 2003 to 2005 resulted in sampling 331 feeding events on 60 d. Within this period, focal subgroups or individuals were followed and observed for a total of 61.7 h during 34 feeding sessions, which we defined as the interval between the first and last feeding events in a series by that individual or subgroup.
Feeding sessions by focal individuals and subgroups lasted an average of 1.9 h (±0.29 SE, range 0.4 to 9.25 h). Of the total 331 feeding events, 170 were documented during these 34 sessions, with an average of 5 feeding events per session (±0.47 SE, range 2 to 15).

Intervals between feeding events ranged from 2 to 120 min, with an average of 25.8 min (±1.84 SE, n = 136 intervals). Almost one-third of feeding events in a session were 10 min or less apart. Based on the distinct behavioural cues associated with prey capture and consumption, we believe that few feeding events were missed during feeding sessions by focal groups or individuals.

An average of 4.5 matrilines were present during encounters in which feeding behaviour was documented (±0.37 SE, range 1 to 12 matrilines, n = 60 encounters). Subgroups, which usually consisted of complete matrilines or partial matrilines comprised of mothers and their young offspring, often foraged 200 m or more apart. Adult males usually foraged independently or in association with their mother, especially in cases where the mother had no juvenile offspring. Whales often foraged close to shorelines, especially in the deep, narrow channels and straits frequented by resident killer whales during summer and autumn.

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Feeding sessions by focal individuals and subgroups lasted an average of 1.9 h (±0.29 SE, range 0.4 to 9.25 h). Of the total 331 feeding events, 170 were documented during these 34 sessions, with an average of 5 feeding events per session (±0.47 SE, range 2 to 15).

Adult males usually foraged further offshore than subgroups. Individuals and subgroups foraging nearshore followed the coastline closely, often within 50 m of shore. Whales foraging offshore often swam in a zigzag pattern rather than in a straight line along a channel.
Signs of prey pursuit and capture were at times very conspicuous. When an appropriate prey item was detected, an individual would break suddenly into a high-speed chase that continued for 10 to 30 s, or (rarely) up to 3 min. Chases were directional and non-directional, the latter accompanied by fast turns and rolls at the surface. Chases often took place along steep shorelines, with whales swimming at high speed within a few metres of the rocks. Although vigorous chases were obvious indicators of predation, more often signs of feeding were quite subtle, and close attention to several behavioural cues was necessary in order to detect them. For example, a change in the otherwise consistent swimming speed and direction of foraging whales often indicated that a pursuit was underway. An unusually long 5 to 7 min dive following an extended series of regular, 2 to 3 min dives was also a good indication of prey pursuit and possible capture.

Whale interactions during feeding events indicated that the majority of prey items were shared by 2 or more individuals. Typically, a whale that made a kill was joined by others, and the group would swim together for 2 to 3 surfacings before splitting up once again. Individuals converged on the successful whale from as far as 400 m, although more often joining whales were within 100 to 200 m when the kill took place. On other occasions, several whales were involved in the pursuit and would work cooperatively to take prey that had sought refuge in crevices along rocky shorelines or in kelp beds. After making the kill, members of the group joined at the surface and swam together for several surfacings. Inspection of the site at which whales joined in such situations invariably revealed fish scales or pieces of tissue in the water. Often, as the whales swam together after joining, a trail of prey fragments was left in the water, indicating that the prey item was being torn up along the way.

Close observations of prey handling and consumption provided strong evidence that sharing was taking place in such circumstances, and that intentional provisioning of other whales was also frequently involved.

Table 3. *Orcinus orca*. Monthly distribution (for 1974 to 2005) of sampled feeding events where age/sex class of whale making kill was determined; n = 274 kills

<table>
<thead>
<tr>
<th>Month</th>
<th>Adult male</th>
<th>Adult female</th>
<th>Juvenile</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>May</td>
<td>8</td>
<td>5</td>
<td>1</td>
<td>14</td>
</tr>
<tr>
<td>Jun</td>
<td>7</td>
<td>2</td>
<td>1</td>
<td>10</td>
</tr>
<tr>
<td>Jul</td>
<td>16</td>
<td>26</td>
<td>25</td>
<td>67</td>
</tr>
<tr>
<td>Aug</td>
<td>19</td>
<td>71</td>
<td>31</td>
<td>121</td>
</tr>
<tr>
<td>Sep</td>
<td>13</td>
<td>3</td>
<td>6</td>
<td>22</td>
</tr>
<tr>
<td>Oct</td>
<td>14</td>
<td>15</td>
<td>11</td>
<td>40</td>
</tr>
<tr>
<td>Total</td>
<td>77</td>
<td>122</td>
<td>75</td>
<td>274</td>
</tr>
</tbody>
</table>

Table 4. *Orcinus orca*. Frequency of prey sharing in feeding events by resident killer whales where age and/or sex class of individuals making kills could be determined. Juvenile whales were <14 yr old

<table>
<thead>
<tr>
<th>Sharing</th>
<th>Adult male</th>
<th>Adult female</th>
<th>Juvenile</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
<td>9</td>
<td>97</td>
<td>36</td>
<td>142</td>
</tr>
<tr>
<td>No</td>
<td>44</td>
<td>4</td>
<td>9</td>
<td>57</td>
</tr>
<tr>
<td>Total</td>
<td>53</td>
<td>101</td>
<td>45</td>
<td>199</td>
</tr>
</tbody>
</table>

On numerous occasions, members of a subgroup milled at the surface while one of the group was underwater on a long dive. Upon surfacing with prey, the whale was seen to carry the fish in the direction of the milling individuals. Scales and/or tissue were found where the whales joined. On other occasions, a whale was observed to surface with prey and carry it for 3 to 5 shallow dives and surfacings while another whale swam quickly in its direction. Although small numbers of scales were often seen in the water in the trail of the whale carrying the fish, much larger numbers of scales and pieces of tissue were observed at the site of joining, indicating that prey consumption was delayed until the whales were together.

Observations during 235 feeding events provided sufficient evidence to judge with reasonable confidence whether or not sharing took place. In 57 of these feeding events (24%) there was no indication of any sharing. In the other 178 cases (76%), sharing was either clearly evident or strongly suspected. Between 2 and 6 whales (including the individual making the kill) were involved in shared feeding events, though it was generally not possible to determine how many individuals actually took part in prey consumption. Whales involved in shared feeding events typically belonged to the same matriline. The frequency of sharing by members of different age and sex classes of whales differed significantly ($\chi^2 = 30.8$, df = 2, $p < 0.001$; Table 4), with adult males sharing significantly fewer of their kills (17%) than expected compared to adult females (96%) and juveniles (80%), (these were not distinguished by sex).

**Prey species and age composition**

All 529 feeding events involved fishes, at least 96% of which were salmonids (Table 2). We could not identify 10 samples (2.5%) to species, and some of these may also have been salmonids. The only non-salmonids identified were 3 Pacific herring *Clupea pallasi*, 1 sablefish *Anoplopoma fimbria*, 1 yelloweye rockfish *Sebastes ruberrimus*, 1 quillback rockfish *S. maliger*, and 1 Pacific halibut *Hippocampus stenolepis*. 
Of the 7 species of Pacific salmon (Oncorhynchus spp.) found in the study area, 6 were represented in feeding samples (the exception was the cutthroat trout O. clarki). Species identity was determined for 432 salmonids, 20 by DNA analysis and the remainder by scale analysis. The frequency of occurrence of these species in whale kills is shown by region in Table 2. Chinook salmon was by far the predominant salmonid observed, representing 71.5% of salmonid kills identified to species. The second most important salmonid was chum, at 22.7% of samples. Coho, pink, sockeye and steelhead together represented less than 6% of the salmonids identified. Ages were determined for 344 salmonid samples (Table 5).

Chinook was the principal species taken by resident killer whales in all regions of the coast (Table 2). It was the most common species in feeding samples from both northern and southern communities, as well as from each of the resident clans (Table 6). Chinook represented the majority of salmonid samples collected from resident killer whales during May to August (Table 7). However, chinook salmon was the predominant species identified from feeding events in September to October. Pink salmon occurred in small numbers in July to September samples, as did coho salmon during July to October. Southern residents were responsible for the single sockeye salmon sample, collected in July, and the 2 steelhead samples, collected in November and December.

### Prey selection as a function of whale age/sex class

Adult male, adult female and juvenile (<14 yr old) resident killer whales all preyed on chinook salmon more than on any other species (Table 8). Chum and coho salmon were also taken by each sex and age category, although the proportion of chum in salmonid kills by adult males was significantly greater than in kills by adult females and juveniles ($\chi^2 = 10.4$, df = 1, $p < 0.01$). This difference may be due to a seasonal bias in sampling of kills by different sexes. A greater proportion of kills by males (35% of total samples from males; Table 3) than by females (15% of total, Table 3) were sampled during September and October, when chum salmon was the predominant prey species (Table 7). All 11 pink salmon kills by identified whales were by juveniles. In fact, 6 of the 11 pink salmon kills were made during a single 1.5 h long feeding session by a 1 yr old calf. Because pink salmon are the smallest of the Pacific salmonids (Quinn 2005), we examined the age distribution of chinook salmon kills to determine if younger (and thus smaller) salmon were taken more frequently by young whales than by adults. The mass of chinook salmon increases dramatically with increasing age, from a mean of 1.1 kg in 2 yr old fish to over 13 kg in 5 to 6 yr old fish (Table 9).

### Table 5. Orcinus Orca. Ages of 344 salmonids killed by resident killer whales. (\text{Age} = 1.0 \times \text{Age}_\text{freshwater} + 1.0 \times \text{Age}_\text{salt water} + 1.0) \begin{table}
<table>
<thead>
<tr>
<th>Species</th>
<th>n</th>
<th>Age class</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0.1</td>
</tr>
<tr>
<td>Chinook</td>
<td>236</td>
<td>1</td>
</tr>
<tr>
<td>Coho</td>
<td>9</td>
<td>57</td>
</tr>
<tr>
<td>Chum</td>
<td>87</td>
<td>8</td>
</tr>
<tr>
<td>Pink</td>
<td>12</td>
<td></td>
</tr>
</tbody>
</table>
\end{table}

### Table 6. Orcinus Orca. Salmonid species sampled from feeding events by members of different resident clans. (\text{Clan A, G, R, J}) \begin{table}
<table>
<thead>
<tr>
<th>Species</th>
<th>Clan</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A</td>
<td>G</td>
</tr>
<tr>
<td>Chinook</td>
<td>178</td>
<td>70</td>
</tr>
<tr>
<td>Chum</td>
<td>64</td>
<td>24</td>
</tr>
<tr>
<td>Coho</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>Pink</td>
<td>9</td>
<td>2</td>
</tr>
<tr>
<td>Sockeye</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Steelhead</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>258</td>
<td>96</td>
</tr>
</tbody>
</table>
\end{table}

### Table 7. Orcinus Orca. Salmonid species sampled from resident killer whale feeding events by month; n = 432 kills. \begin{table}
<table>
<thead>
<tr>
<th>Month</th>
<th>Species</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Chinook</td>
<td>Chum</td>
</tr>
<tr>
<td>May</td>
<td>22</td>
<td>0</td>
</tr>
<tr>
<td>Jun</td>
<td>34</td>
<td>12</td>
</tr>
<tr>
<td>Jul</td>
<td>94</td>
<td>13</td>
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<td>Aug</td>
<td>139</td>
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<td>Sep</td>
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<td>0</td>
</tr>
<tr>
<td>Dec</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>309</td>
<td>98</td>
</tr>
</tbody>
</table>
\end{table}
Although the smallest chinook (2 and 3 yr olds) were taken more often by juvenile than by adult whales (Fig. 2), the overall trend was not significant (\(U\)-test = 1688, \(p = 0.11\)). There was also no significant difference (\(U\)-test = 1377.5, \(p = 0.13\)) between the mean age of chinook taken by adult males (4.50 yr ± 0.13 SE, \(n = 38\)) and by adult females (4.26 yr ± 0.08 SE, \(n = 86\)).

Prey species and sizes in shared and non-shared kills

Of the 6 salmonid species taken by resident killer whales, 4 were identified from both shared and non-shared feeding events (Table 10). The great majority (84%) of chinook salmon tended to be shared, while a significantly lower proportion of chum salmon were shared (55%; \(\chi^2 = 4.47, df = 1, p < 0.05\)). Although chinook are often larger than chum salmon (Healey 1986), larger size appears not to be the reason for the greater incidence of sharing of chinook. Chum salmon formed a higher proportion of the prey samples from adult males (4.50 yr ± 0.13 SE, \(n = 38\)) and by adult females (4.26 yr ± 0.08 SE, \(n = 86\)).

Prey selection versus availability

To assess the extent to which foraging resident killer whales select for particular species or sizes of salmonids, the species and age composition of kills sampled off NE Vancouver Island was compared to salmonid availability. The relative abundance of salmonids was determined from catch statistics resulting from Fisheries and Oceans Canada seine test fisheries (Data available from www-ops2.pac.dfo-mpo.gc.ca/xnet/content/salmon/testfish/default.htm), which were undertaken concurrently with and in close proximity to our sampling of killer whale feeding events in the western Johnstone Strait area, off NE Vancouver Island (see Fig. 1). These data provide a reliable index of the relative availability of different salmonid species to the whales foraging in that area during particular periods of the season. Fig. 4 illustrates the species composition of salmonids caught in test fisheries and by resident killer whales during 3 periods in July to October 2004. From 15 July to 15 August, the test catches

![Table 8. *Orcinus Orca*. Salmonid species sampled from feeding events as a function of age and sex class of resident whales; \(n = 274\) kills. Specific names of prey in Table 2](image)

<table>
<thead>
<tr>
<th>Species</th>
<th>Age/sex class</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Adult male</td>
<td>Adult female</td>
</tr>
<tr>
<td>Chinook</td>
<td>47</td>
<td>104</td>
</tr>
<tr>
<td>Chum</td>
<td>26</td>
<td>16</td>
</tr>
<tr>
<td>Coho</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>Pink</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>77</td>
<td>122</td>
</tr>
</tbody>
</table>

![Table 9. *Oncorhynchus tshawytscha*. Mean fork lengths and mass (±SE) of chinook salmon at different ages. Data from seine caught fish in Mark Recovery Program of Fisheries and Oceans Canada (Kuhn 1988)](image)

<table>
<thead>
<tr>
<th>Age (yr)</th>
<th>Length (mm)</th>
<th>Mean mass (kg)</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>425 ± 1.19</td>
<td>1.1 ± 0.01</td>
<td>3072</td>
</tr>
<tr>
<td>3</td>
<td>581 ± 2.14</td>
<td>3.1 ± 0.04</td>
<td>3206</td>
</tr>
<tr>
<td>4</td>
<td>808 ± 3.43</td>
<td>8.5 ± 0.11</td>
<td>917</td>
</tr>
<tr>
<td>5</td>
<td>939 ± 4.21</td>
<td>13.3 ± 0.20</td>
<td>426</td>
</tr>
<tr>
<td>6</td>
<td>961 ± 15.0</td>
<td>13.7 ± 0.72</td>
<td>37</td>
</tr>
</tbody>
</table>

![Table 10. *Orcinus Orca*. Frequency of prey sharing by resident killer whales in feeding events where prey was identified to species. \(n = 229\) kills. Specific names of prey in Table 2](image)

<table>
<thead>
<tr>
<th>Sharing</th>
<th>Prey species</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Chinook</td>
<td>Chum</td>
</tr>
<tr>
<td>Yes</td>
<td>141</td>
<td>29</td>
</tr>
<tr>
<td>No</td>
<td>27</td>
<td>24</td>
</tr>
<tr>
<td>Total</td>
<td>168</td>
<td>53</td>
</tr>
</tbody>
</table>
were dominated by migrating sockeye and pink salmon, which together comprised over 90% of salmon sampled. During this period, 59 salmonid kills by resident killer whales in the area were documented, 95% of which were chinook and 5% coho (Fig. 4A). These 2 species each represented less than 1% of the available salmonids in the area. During 22 to 31 August 2004, the abundance of chinook increased to approximately 6% of the available salmonids, and represented 97% of kills during this period (Fig. 4B). A substantial change in the relative abundance of both salmonids and kills by killer whales took place between 12 and 18 October 2004 (Fig. 4C). Autumn-migrating chum salmon comprised 97% of the test fishery catch during this period, and chum was the predominant salmonid taken by feeding killer whales (90% of kills). Chinook still represented 5% of kills during this period despite being extremely scarce in test catches (1 chinook in 57,435 salmonids sampled).

To evaluate whether resident killer whales preyed equally on all sizes of their preferred prey species, the age distribution of chinook salmon taken by whales was compared to the age distribution of chinook available to them for waters off NE Vancouver Island in 2000 to 2004. The relative abundance of chinook age classes was determined from estimates developed by the Pacific Salmon Commission Joint Chinook Technical Committee1. Although killer whales took all 5 yr classes available to them (Fig. 5), the frequency distributions were significantly different, with killer whales taking fewer young chinook and more older chinook than expected based on the proportions of those ages available. The mean age of chinook taken by whales was 4.20 yr (±0.06 SE, n = 124), significantly older than the mean of 3.52 yr (±0.03 SE, n = 976,212) for available chinook (U-test = 33057, p < 0.001).

**Fig. 3.** *Oncorhynchus tshawytscha*. Age distribution of chinook salmon taken in shared (n = 117 kills) and non-shared (n = 22 kills) feeding events by *Orcinus Orca*.

**Fig. 4.** Species composition of salmonids caught in seine test fisheries (open bars) and by resident killer whales (shaded bars) in western Johnstone Strait area, NE Vancouver Island from (A) 15 July to 15 August, (B) 22 to 31 August and (C) 12 to 18 October 2004. Samples sizes were (A) 69,847 (test fishery) and 59 (whale kills), (B) 1548 (test fishery) and 30 (whale kills), and (C) 57,435 (test fishery) and 41 (whale kills). Specific names of prey in Table 2.

**DISCUSSION**

Assessing the selectivity of a predator requires information on both the predator’s diet and the relative availability of its various prey types. Although such information is often readily available for terrestrial predators (e.g. Scheel 1993, Karanth & Sunquist 1995), it can be very difficult to obtain for fish-eating marine mammals. The diet of aquatic predators must usually be determined by indirect means, such as from stomach content, faecal, or fatty acid analyses, and the
abundance of prey species is seldom known unless those species are assessed for commercial fisheries (Bowen and Siniff 1999). Studies on foraging in pinnipeds have revealed prey selection based on size, energy density or profitability (Tollit et al. 1997, Lawson et al. 1998, Bowen et al. 2002), but no comparable analyses have until now been undertaken for a cetacean. Our study is the first to analyse a large number of feeding events involving a variety of prey species positively identified either by direct observation or from prey fragments, and to compare predation rates with availability of prey species and age classes determined from concurrent abundance assessments. Our findings indicate that, for a significant part of the year, resident killer whales fed on certain prey species at rates far out of proportion to their relative availability compared to other alternative prey. This strong selectivity is likely to have important influences on the whales’ foraging tactics and seasonal movements, as well as on their social structure and behaviour.

Prey fragments as indicators of diet

The reliability of our data on prey selection by resident killer whales is dependent on all prey species having a similar—ideally equal—probability of being sampled from kill sites. Concerns about potential biases of the prey fragment sampling technique that have been raised include (1) the possibility that large fishes, such as chinook, are more subject to being broken up prior to consumption and are thus more likely to shed scales or tissue than smaller fishes, and (2) the possibility that whales foraging at depth are less likely to bring prey to the surface prior to consumption than prey captured in the upper portion of the water column (Ford et al. 1998, Baird 2000, Baird et al. 2005). Our findings reported here suggest that neither of these potential biases is significant enough to alter the patterns of prey selection we describe, particularly with respect to salmonid prey.

Most feeding events, especially those by female and juvenile whales, involved the transport of the prey item to the surface, where it was broken up for sharing or provisioning. Chinook, chum, coho and pink salmon, which differ widely in average size (Healey 1986, Groot & Margolis 1991), were all shared, suggesting that salmonids are shared regardless of prey size. Also, the age distribution of chinook salmon taken in shared and non-shared feeding events did not differ significantly, providing further evidence that sharing is independent of prey size. Thus, we conclude that salmonid prey is typically brought to the surface and torn apart for social feeding, rather than to facilitate the consumption of large prey items. Adult male killer whales, which shared the minority of their prey, brought a range of salmonid species and sizes to the surface to be consumed alone. Although it is not clear why adult males do this, we suspect that consumption of prey at the surface is routine in these whales.

Although sharing of salmonid prey was noted earlier in our opportunistic studies of resident killer whale predation (Ford et al. 1998), the extent of this behaviour was not apparent until close focal individual and subgroup observations were undertaken during feeding sessions. Signs of prey capture, handling and consumption are subtle and easily overlooked, and consistent detection of these cues requires considerable experience. Given this difficulty, it is not surprising that previous descriptions of the foraging behaviour of resident killer whales, which were generally based on less field effort and/or on opportunistically-collected samples, have not reported food sharing (Jacobsen 1986, Heinlisch-Boran 1988, Felleman et al. 1991, Hoelzel 1993, Nichol & Shackleton 1996, Saulitis et al. 2000).

There is no evidence that proximity to the surface differentially affected the likelihood of salmonid prey species being represented in feeding samples. Fish tracking studies off NE Vancouver Island, where the majority of our salmonid prey samples were collected, revealed that chinook travelled at mean depths of 25 to 64 m during the day compared to 14.9 m in sockeye salmon (Candy & Quinn 1999). Despite being deeper and much less abundant than sockeye in this area during July and August (Fig. 5), chinook salmon was by far the predominant species observed in feeding events, and no sockeye salmon were found in prey samples. For these same reasons, prey fragment collection from feeding events should reveal kills of demersal, non-salmonid fish species as well as salmonids, and we be-
lieve that the scarcity of non-salmonids in our samples is a true reflection of their minor role in the diet of resident killer whales, at least during the seasons covered by our observations. For example, lingcod *Ophiodon elongatus*, found in stomach remains of a stranded resident whale (Ford et al. 1998), reach similar sizes to chinook salmon and are most abundant at depths of 10 to 100 m, which overlap the preferred depths of chinook salmon (Cass et al. 1990, Candy and Quinn 1999). Lingcod have extremely small scales that are unlikely to be shed or recovered, but fish species identification is not reliant on scales alone. Tissue samples, from which species identity can be readily determined using molecular techniques, were collected from almost half of the feeding events sampled during 2003 to 2005. Of 21 prey items identified solely by DNA analysis of tissue fragments, only 1, a sablefish, was a non-salmonid. We thus conclude that any negative bias in representation of bottomfishes in our samples is unlikely to be significant. Studies of diving behaviour in southern resident killer whales found most activity to be concentrated in the upper portion of the water column (Baird et al. 2005), which would be expected of a predator focused on salmonid rather than demersal prey.

**Prey selection**

During the months of May to October, the main period of our field studies, salmonids are clearly the preferred prey type of resident killer whales, representing over 96% of identified prey. The only non-salmonids found were a Pacific halibut, 2 rockfishes, a sablefish and 3 herring. Although the halibut and sablefish were eaten, both rockfish species were abandoned by the whales after being partially consumed. We suspect that rockfishes may be an undesirable prey type due to their prominent dorsal spines, as this part of the fishes’ body was discarded. No rockfish remains have been found in stomach contents of stranded resident killer whales (Ford et al. 1998). It seems unlikely that the herring were targeted prey items, since herring scales were collected only during feeding sessions involving chinook prey. As chinook feed extensively on herring (Healey 1991), it is likely that herring scales were released when whales killed and broke chinook apart or were left in the water after chinook predation on herring.

Chinook and chum salmon together represented 94% of salmonids identified from kills by resident whales. Chinook is the predominant species taken during May to August. This would be expected for May and June, since other salmonids are uncommon in nearshore waters during these months (Groot & Margolis 1991, Quinn 2005). However, chinook remains the predominant prey species in July and August, when migrating sockeye and pink salmon form the overwhelming majority of salmonids available to the whales (our Fig. 4; Groot & Margolis 1991, Nichol & Shackelton 1996, Wydoski & Whitney 2003, Quinn 2005). With only a single sockeye kill sample collected, it is clear that despite its abundance during the summer migratory period, this species is rare in the diet of resident killer whales. Pink salmon were also very uncommon in our samples, and the species does not appear to be a significant prey item. Coho salmon are relatively uncommon throughout the region (Groot and Margolis 1991, Wydoski and Whitney 2003, Riddell 2004, Quinn 2005), and were consistently represented in small numbers in killer whale feeding events. Early summer runs of chum salmon occur during June and July, particularly on the northern coast of British Columbia (Riddell 2004), and this species represented a significant component of the killer whale diet in this period and region. During September and October, the diet of northern resident killer whales shifts to predominantly chum salmon, although chinook are also taken frequently. This period coincides with the autumn migration of chum salmon through the area, which starts in mid-September, peaks in mid-October, and is over by late October (Ryall et al. 1999).

Comparisons of prey selection by resident killer whale communities and clans revealed few differences. Northern and southern resident communities both fed predominantly on chinook. The greater proportion of chum in the samples of northern resident A and G clans probably resulted from their presence off NE Vancouver Island during September and October 2003 and 2004, when intensive sampling was undertaken. Little sampling of feeding events by R and J clans during autumn has yet been undertaken. It appears likely that southern resident whales also target chum salmon in addition to chinook in the autumn, as their movement into waters of Puget Sound in late October and November coincides with migratory aggregations of these species (Osborne 1999). The single sockeye and 2 steelhead salmon kills were observed only in southern residents, but this might be due to chance, as a result of such small sample sizes. Different age and sex classes of resident killer whales did show some differences in salmonid prey composition. Kills of pink salmon were made primarily by juvenile whales, and such predation of these small salmonids may represent a form of play or practice (Jacobsen 1986, Ford et al. 1998). Chinook, chum and coho were taken by both adults and juveniles and by both sexes, but a higher proportion of chum salmon kills were recorded for adult males than for adult females. As noted previously, this difference may be due to a bias towards disproportionately greater sampling of kills by males in September and October,
when chum salmon was the predominant prey species. Bain (1989) suggested that the larger body size of males may allow them to dive more deeply and take larger prey than females. Baird et al. (2005) observed male southern residents to dive deeper more frequently than adult females. However, we observed no difference in the mean ages, and thus general sizes, of chinook taken by the 2 sexes.

Important factors influencing prey selection by predators include size, energy density, availability, and catchability of prey (Stephens & Krebs 1986, Scheel 1993, Lawson et al. 1998, Bowen et al. 2002). Resident killer whales probably prefer chinook and chum salmon over other salmonids primarily because of their large size. Chinook salmon are the largest of the salmonids, and can attain masses of >25 kg (Healey 1991, Quinn 2005). Most chinook taken by killer whales were 4 to 6 yr old, representing mean masses of >8 kg. Chum salmon are smaller, having mean masses of 5.0 to 7.5 kg (Ricker 1980, Salo 1991). Both species are significantly larger than coho (mean mass = 2.95 kg; Sandercock 1991), sockeye (mean mass = 2.73 kg; Burgner 1991) and pink salmon (mean mass = 1.7 to 2.4 kg; Heard 1991). Northern resident killer whales foraging off NE Vancouver Island selected chinook that were older on average than would be predicted from the age distribution of available chinook, indicating a preference for larger-sized fish of this species. It is interesting, however, that the whales also took significant numbers of 3 yr old chinook, which are similar in size to the far more abundant but rarely selected sockeye and pink salmon. It may be that the relatively high lipid content of chinook compared to other salmonids (Stansby 1976, Healey 1986, Winship & Trites 2003) is another factor in the whales’ preference for this species. However, chum salmon is a significant prey species, despite having lower average lipid content than other salmonids (Stansby 1976), suggesting that size may be a more important influence than lipid content in determining salmonid preferences.

Evidence from other regions also suggests that both prey size and lipid content may influence prey choice by fish-eating killer whales. Saulitis et al. (2000) found that resident killer whales in Prince William Sound, Alaska, appeared to prey selectively on coho salmon, which are larger and have higher lipid content than the far more abundant pink salmon that were available to the whales during their study. Chinook are extremely rare in Prince William Sound during July and August, when prey sampling was conducted by Saulitis et al. (2000), which probably explains why few of this species were found in their samples. These whales prey extensively on chinook at other times and in other areas where this prey species is present (C. Matkin, North Gulf Oceanic Society, Homer, Alaska, pers. comm.). Killer whales in Prince William Sound and in the Bering Sea are also known to remove and eat fishes from longline fishing gear, but they do so selectively, according to energy value and size. Pacific halibut, sablefish and Greenland turbot Reinhardtius hippoglossoides are among the favoured species, and the whales take the largest individuals of these species from fishing lines while ignoring other species such as Pacific cod Gadus macrocephalus, walleye pollock Theragra chalcogramma, and rockfish Sebastes spp. (Matkin & Saulitis 1994, Yano & Dahlheim 1995). Favoured species have higher average lipid content and energy densities than species that are shunned (Stansby 1976, Winship & Trites 2003).

Although size and lipid content may be important factors in prey selection by resident killer whales, it is not clear why so few sockeye and pink salmon are taken given their considerable seasonal abundance. During the peak of their migration through whale foraging areas, from mid-July to mid-September, sockeye and pink salmon combined outnumber chinook by as many as 500 to 1 (DFO seine test fisheries data, see ‘Results’). We believe that it is a combination of the chinook salmon’s large size, high lipid content, and year-round availability that makes them the salmonid of choice for resident whales in all seasons, even when alternative species are available in greater abundance during brief migratory pulses.

Unlike most salmonids, chinook are available to resident killer whales in nearshore waters of the region throughout the year. The ‘ocean-type’ chinook ranges over continental shelf waters throughout the marine portion of its life cycle (Healey 1991). The timing of migration to spawning rivers is highly variable in chinook, with different populations of chinook entering freshwater from mid-spring through autumn (Healey 1991). Migrating chinook also tend to travel through coastal waters at slow rates of speed compared to other salmonids (Candy & Quinn 1999). Sockeye, chum and pink salmon, on the other hand, have oceanic distributions most of their lives and only transit coastal regions briefly while en route to spawning rivers (Groot & Margolis 1991). The oceanic range of these salmonids is vast (Groot & Margolis 1991), with fishes distributed widely at densities that are probably too low for whales to effectively utilise. Although resident killer whales range extensively along the coast, there is no evidence that they undertake long distance movements to offshore areas (Ford et al. 2000, Wiles 2004). Sockeye, chum, and pink salmon, the 3 most abundant salmonids in the North Pacific, are thus only available for predation by resident killer whales for a small portion of the year compared to chinook salmon.

Killer whale populations tend to have specialised foraging tactics to hunt their preferred prey (Lopez &
Lopez 1985, Guinet 1992, Barrett-Lennard et al. 1996, Baird 2000, Saulitis et al. 2000, Pitman & Ensor 2003), and this is likely to be the case for resident killer whales also. Resident killer whales may be particularly skilled at finding and capturing chinook, which tend to travel more individually, at greater depths and closer to shore than smaller, schooling salmonids (Groot & Margolis 1991, Candy & Quinn 1999). The echolocation signals of resident killer whales are well suited for the detection of echoes from individual chinook at ranges of 100 m or more (Au et al. 2004). Efficient predation of smaller, schooling salmonids such as pink and sockeye salmon would probably require specialised tactics such as the ‘carousel’ technique used by killer whales to feed on herring in northern Norway (Similä & Ugarte 1993). Such coordinated foraging behaviour has not been observed in resident killer whales (Heimlich-Boran 1988, Ford 1989). It is noteworthy that resident killer whales do not target the large seasonal concentrations of spawning herring that occur in coastal waters throughout their range.

The known distribution patterns of resident killer whales are consistent with what would be expected of a predator focused on chinook salmon. From May to early July, northern residents are found mostly along the coasts of the northern mainland of British Columbia and the Queen Charlotte Islands, where they congregate in areas of high chinook density and feed primarily on this species (Ford et al. 2000, authors’ unpubl. data). This period coincides with the earliest of chinook runs in the region (Riddell 2004). Early-summer-run chum salmon are also found there during this time, although they are preyed upon to a lesser extent than chinook. By mid-July, the whales begin to occur regularly off NE Vancouver Island, concurrently with an increase in chinook abundance in the area and the arrival of migrating sockeye and pink salmon. From mid-July to early September, when sockeye and pink are transiting these waters, chinook salmon are also available at relatively high densities. Although the absolute abundance of chinook is far lower than that of these smaller species, there are probably sufficient numbers available to meet the needs of resident whales without their having to switch to pink or sockeye prey. By October, autumn-migrating chum salmon become the predominant salmonid available and the principle prey of northern residents. Chinook salmon appear to still be taken preferentially during this period, since this species occurred more often in prey samples than would be predicted from its low relative abundance. The movements of southern residents to foraging areas off southern Vancouver Island during June to September also coincide with increased availability of chinook salmon, and in October and November with chum salmon (Osborne 1999).

A significant gap in our knowledge of the feeding ecology of resident killer whales is their diet during November to April. Chinook salmon remain available during these months in the inshore summer and autumn concentration areas of resident killer whales, but mostly at low densities (Osborne 1999; B. Riddell, Pacific Biological Station, Nanaimo, pers. comm.). Whales disperse from these areas during winter and spring, but their whereabouts are for the most part unknown (Ford et al. 2000, Wiles 2004). Their prey may shift in winter and early spring to include more non-salmonid fishes, but there is little evidence to determine the extent to which their diet may change. The stomach of a northern resident whale that died in late November off NE Vancouver Island contained remains of chinook salmon and a variety of demersal fish species, including lingcod, sablefish, and greenling (*Hexagrammos* sp.; Ford et al. 1998). It is probable that the resident killer whales’ preference for chinook continues throughout the winter, and that they travel more widely over remote parts of the coast in pursuit of this species. Future studies are needed to locate resident whales during these months, and to determine whether chinook salmon is indeed their prey of choice throughout the year.

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**LITERATURE CITED**


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