

Does diet in Celtic Sea fishes reflect prey availability?

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Feeding preferences of Celtic Sea fishes were investigated using a database of stomach content records, collected between 1977 and 1994. The diet of cod *Gadus morhua*, hake *Merluccius merluccius*, megrim *Lepidorhombus whiffiagonis*, whiting *Merlangius merlangus* and saithe *Pollachius virens* changed markedly as the animals grew larger, and although large predators generally chose larger bodied prey, the variability of prey sizes consumed also increased. Large predators continued to select small, low value, benthic prey (e.g. *Callionymus* spp. and *Trisopterus* spp.) which were easier to catch, rather than larger, more energy lucrative pelagic prey (e.g. mackerel *Scomber scombrus*), even though these pelagic prey-fishes were nearly always available and were often very abundant. Stock estimates of the International Council for the Exploration of the Sea and U.K. groundfish survey catches were used as indices of prey abundance. Blue-whiting *Micromesistius poutassou* and other small pelagic fishes (*Argentina* spp. and clupeoids) were identified as being particularly important, and were consumed by some predators more often than would be expected given the abundance of these prey in the environment. There was no evidence for density-dependent feeding by predators on mackerel and only hake exhibited density-dependent feeding on horse-mackerel. Hake, cod and megrim consumed more blue-whiting when this prey was at higher abundance in the environment. In choosing what prey to consume, predators must balance costs and benefits, considering the quality of prey and the energy expended during search, capture and handling.

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INTRODUCTION

The importance of predation as a regulatory process in marine systems has been well documented and piscivorous fishes are known to have a dramatic influence on population and community level dynamics (Juanes *et al.*, 2002). Although a relatively large amount is known about whom eats who in marine systems, virtually nothing is known of the dynamics, and specifically how diets of piscivorous fishes relate to changes in the abundance of their prey (Greenstreet *et al.*, 1998).

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Most fishes are selective foragers; they prefer to feed on some prey types but not on others (Mittelbach, 2002). Much of the theoretical development of foraging theory has revolved around trying to explain why predators choose the items they do and whether a predator should theoretically choose to pursue a particular prey item that it has encountered (Mittelbach, 2002). Ecologists have used optimization criteria to address this question, arguing that natural selection should result in predator behaviours that maximize the rate of energy gain, which is a component of fitness. Charnov (1976) developed one of the first optimal diet models, and three basic predictions stem from this work: (1) predators should prefer prey that yield more energy per unit handling time; (2) as abundance of higher value prey increases in the environment, lower value prey should be dropped from the diet and predators should become more selective; (3) foragers should obey a quantitative threshold rule for when specific prey types should be included or excluded from an optimal diet (Stephens & Krebs, 1986; Sih & Christensen, 2001).

Prey encounter rate is dependent on the abundance of that prey in the environment but, in times when a focal prey is not available, predators can satisfy their nutritional and energy requirements, to some extent through adjustments in selection for prey quality. Prey fishes can exhibit a 10-fold difference in lipid content and a five-fold difference in energy density (Anthony *et al.*, 2000). Thus as prey vary in abundance, resource value, catchability and handling time, predators must trade-off costs and benefits to optimize their own survival and reproductive fitness (Stephens & Krebs, 1986).

As originally formulated, the optimal foraging model of Charnov (1976) considered only diet choice within a homogeneous patch. If the forager moves to another patch the model should be freshly applied and this means that such models cannot be tested by simply looking at stomach contents data and taking overall averages (Stephens & Krebs, 1986). For example, a predator might forage in a part of the environment where low-quality prey items are disproportionately common, because the abundance of these items might compensate for their low food value. A forager choosing such a patch would appear to take too many low-ranking foods and too few high-ranking types compared to an idealized forager for which model parameters would be calculated as if they foraged over the whole environment. In the present study, the validity of optimal foraging theory is not tested using stomach content data, rather it is used as a framework for discussing the trade-offs which consumers face in real systems.

The body mass of many piscivorous fish species can increase by five orders of magnitude during their lives, and a fast-growing animal may begin life as a prey item, only to become the main predator on the same group of species within 1 year (Jennings *et al.*, 2002). Clearly size is an important determinant of fish diets and many authors (Scharf *et al.*, 2000) have attempted to relate physical attributes of the predator (*e.g.* gape size and length) to the prey found in fish stomachs.

Over the past 30 years a considerable amount of stomach sampling effort has been expended in the North Sea (Pope, 1991), but there has been little concerted research elsewhere in the north-east Atlantic (*e.g.* the Celtic Sea). Marked changes are known to have occurred in the Celtic Sea ecosystem in recent

years (Pinnegar *et al.*, 2002), and given such changes it might be expected that the diet of predatory species will have varied in response (Greenstreet *et al.*, 1998).

The main aims of the present study were to: (1) explore how the diet of Celtic Sea fishes change with body-size, (2) examine how the size range of the targeted prey change with increasing body-size and (3) explore whether predators select prey in accordance with availability in the environment.

METHODS

THE CELTIC SEA

The Celtic Sea is an area of continental shelf bordered by Ireland in the north, the U.K. in the east and the Bay of Biscay (47° N) in the south (Fig. 1). It supports a diverse range of fishing fleets (*métiers*), characterized by the use of different fishing gear types and different target species (Marchal & Horwood, 1996). Recent expansion of the Celtic Sea fisheries has prompted concern about the present and future state of fish stocks, the scale of fishery discards and possible implications for ecosystem functioning as a whole (Pinnegar *et al.*, 2002).

STOMACH-CONTENT DATABASE

Fishes were sampled by U.K. and French researchers between 1977 and 1994. A total of 26539 prey items were recorded from 66 predator species (18129 stomachs).

Samples from the U.K. were taken as part of annual research cruises aboard the vessels *Cirolana*, *Clione* and *Scotia*. Location, predator total length (L_T), stomach fullness (on a 10 point scale), prey identity, number and digestion state (on a four point scale), were recorded along with the number of empty stomachs. Where possible, prey L_T was recorded.

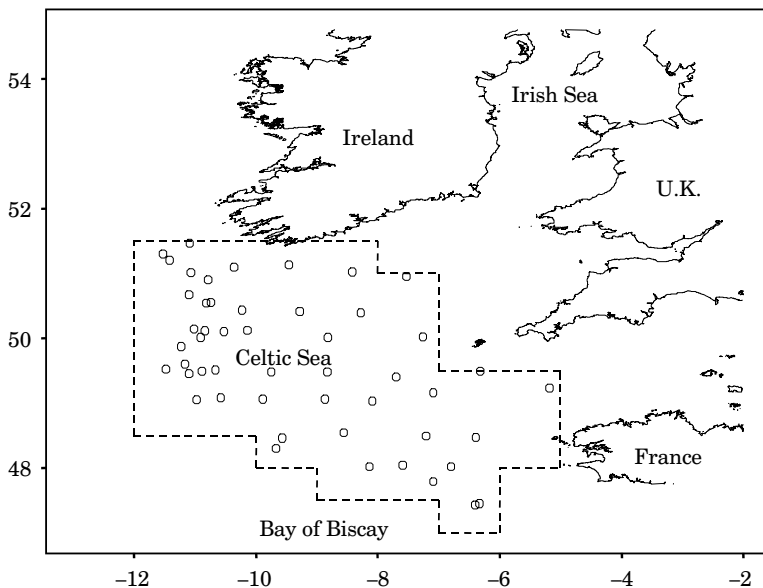


FIG. 1. Area covered by this study, with U.K. spring-sampling sites (○).

French samples were collected aboard commercial trawlers (Agora, Elsinor, Galaxie, Madiana, Melodie, Opera, Peoria, Symphonie and Valérieane) during routine fishing operations in the Celtic Sea. These data have been largely documented by Du Buit (1982, 1992, 1995, 1996) but not in the context of prey availability. Predator L_T , prey identity, prey abundance, prey mass (g), and digestion state (on a five point scale) were recorded. No data were available relating to non-identifiable prey remains or empty stomachs, also there was no detailed information linking samples to particular geographic locations. Prey length data only existed for megrim *Lepidorhombus whiffiagonis* (Walbaum), whiting *Merlangius merlangus* (L.) and saithe *Pollachius virens* (L.).

Of the 66 fish species for which stomach-content data existed, the number of records was greatest for cod *Gadus morhua* (L.), whiting, megrim, hake *Merluccius merluccius* (L.), haddock *Melanogrammus aeglefinus* (L.), and saithe. Initial analyses were carried out to determine which of these predators fed mainly on fishes (in terms of total numbers of prey items). It was determined that haddock consume very little fish material (<2% of all items) and thus this species was excluded from all further analyses.

The temporal coverage of the available data is detailed in the Appendix. Sampling varied greatly from year to year, with the largest number of stomachs collected in 1984, 1985 and 1991. Stomachs were collected from commercial vessels (1977–1988) throughout the calendar year although sampling was particularly intensive in May, June and October. Stomachs were collected from survey vessels (1990–1994) in March and April each year.

PREY ABUNDANCE

Mackerel *Scomber scombrus* (L.), horse-mackerel *Trachurus trachurus* (L.) and blue-whiting *Micromesistius poutassou* (Risso) were identified as being important fish prey on the basis of their occurrence in the stomachs of cod, whiting, megrim, hake and saithe. Stocks of these pelagic prey species are assessed on an annual basis by the International Council for the Exploration of the Sea (ICES), and thus population estimates were available for comparison with fish-prey numbers from stomach contents (available *v.* consumed prey). Population numbers (in millions) were extracted from two reports (WGMHSA, 2002; WGNPBW, 2002). The geographic units considered by ICES, extend beyond the Celtic Sea, thus it was necessary to assume that any temporal patterns apparent at the whole stock level would also be reflected at the Celtic Sea sub-stock level. The abundance of each pelagic species was compared with the stock size of the other two species using a non-parametric Kendall tau test (Conover, 1980).

Abundance estimates for all fish-prey species were available from 1982–2003 through the annual ground-fish survey of the Centre for Environment, Fisheries & Aquaculture Science (CEFAS). Samples are collected in March and April each year using a Portuguese high headline trawl (PHHT), hauled by the research vessel Cirolana. The distance travelled by the vessel whilst hauling, together with the gear geometry (monitored using acoustic transponders) were used to calculate 'swept area' and consequently fish numbers per unit area. Only haul-stations from the central Celtic Sea were included in the analyses (Fig. 1), and the estimated fish density (numbers km^{-2}) was raised to the level of this whole central zone (124505 km^2).

PREY LENGTH

To estimate changes in median, 'minimum' and 'maximum' prey size with increasing predator L_T , quantile regression techniques were used (Scharf *et al.*, 1998, 2000). In order to carry out quantile regression, the BLOSSOM software developed and described by Cade & Richards (1996) and Cade *et al.* (1999) was utilised.

Correlation analyses were performed among 10% quantile, median (50% quantile) and 90% quantile slope estimates to determine whether increases in median prey size were primarily a result of changes in 'minimum' or 'maximum' prey sizes.

PREY-PREFERENCE

Many prey-selection indices have been proposed, and different indices seem appropriate for answering different ecological questions (Pearre, 1982; Confer & Moore, 1987). In the present study, the index of Chesson (1978, 1983) was used (α_a) based on the constant preference coefficient discussed by O'Neill (1969). For a two prey system:

$$\alpha_a = r_a p_a^{-1} \left[\sum_{i=1}^2 r_i p_i^{-1} \right]^{-1} = e a_d (d a_c)^{-1} [e d^{-1} (a_d a_c^{-1} + b_d b_c^{-1})]^{-1}$$

where a_d is the number of prey animals of species a in the predator's diet, b_d is the number of all other prey animals in the diet, a_c is the number of prey animals of species a in the environment, b_c is the number of all other prey animals in the environment, d is the total number of all animals in the diet, e is the total number of all animals in the environment, r_a is the proportion of prey species a in the diet and P_a is the proportion in the environment. The notation used by Pearre (1982) is used throughout.

α_a (also known as the 'standardized forage ratio' s_i) has become popular because of its use in food-web modelling packages (Christensen *et al.*, 2000). The standardized forage ratio as presented, ranges between 0 and 1, with $\alpha_a = 0$ representing complete avoidance and $\alpha_a = 1$ exclusive feeding. The index is independent of prey availability, and in form is broadly similar to the index (β_{Na}) proposed by Manly *et al.* (1972).

Here, α_a was calculated on the basis of all available French and English stomach data, irrespective of year. Small sample sizes (Appendix) precluded the calculation of meaningful indices on an annual basis. α_a was calculated for a 'portfolio' of seven ($m=7$) fish-prey species (*Argentina* spp., *Callionymus* spp., *M. poutassou*, *S. scombrus*, *T. trachurus*, *Trisopterus* spp. and clupeidae) since these were found to be the most commonly observed fishes in the stomachs of cod, hake, whiting, megrim and saithe. Some of these prey 'species' are actually groups of closely related animals (*e.g.* clupeidae, *Trisopterus* spp.) which could not be distinguished in stomach contents. In the present case $\alpha_a = 0.143$ ($\alpha_a = m^{-1}$) denotes 'random-feeding', *i.e.* that the particular prey is taken by the predator in exactly the same proportions as in the environment. Thus $\alpha_a = 0-0.143$ denotes 'negative selection' and $0.143-1$ denotes 'positive selection' of a particular prey type.

PREFERENCES V. CHANGING PREY AVAILABILITY

Fish-prey abundance, characterized on the basis of U.K.-survey data or from ICES stock assessments, were compared with the proportion of identifiable fish prey for each year that stomachs were collected. Tests for significance were based on Kendall's tau (Conover, 1980). The rank-based Kendall tau test is a non-parametric method which is particularly robust to outliers. In addition, in order to obtain robust correlation coefficients, 20% trimming was employed, whereby 20% of observations were removed, starting with the most extreme using a routine within the statistical package S-Plus (Insightful, 2001).

RESULTS

CHANGES IN PREDATOR DIET WITH FISH LENGTH

The diet of all species were found to change markedly with increasing predator L_T (Fig. 2). In cod, whiting and hake there was a marked transition from consumption of crustaceans in smaller predators to a higher proportion of fishes in the diet of larger animals. Fishes never represented >40% of the diet in cod, and polychaetes always represented a small but consistent proportion [Fig. 2(d)].

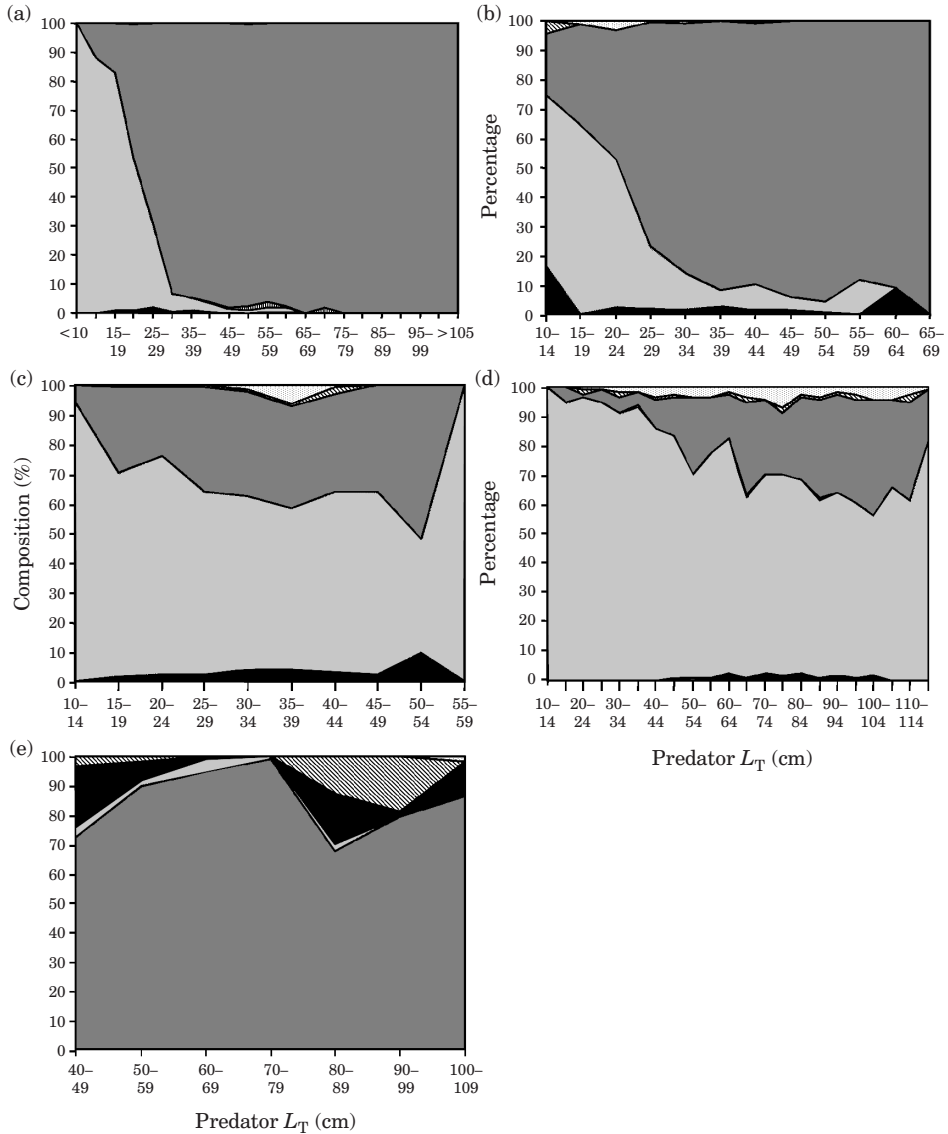


FIG. 2. Changes in the composition of Celtic Sea fish stomach contents (■, fishes □, crustacea; ■, cephalopods; □, polychaetes; ▨, molluscs; ▩, Echinoderms; □, cnidaria) with increasing predator body length, based on the number of total prey items. (a) Hake, (b) whiting, (c) megrim, (d) cod and (e) saithe.

Unfortunately, no diet data were available for saithe <40 cm [Fig. 2(e)], possibly because of distributional differences between adult and juvenile animals in the Celtic Sea. Diets of large saithe were overwhelmingly dominated by fish prey, but with some cephalopod and pteropod molluscs. Megrim appeared to consume similar proportions of crustaceans and fishes throughout their lives, whilst cephalopods also provided a small but consistent contribution [Fig. 2(c)].

Based on the U.K. data it was possible to estimate that 64.7% of all megrim, 11.7% of hake, 20.4% of whiting, 19.4% of saithe and 6.1% of cod stomachs were empty upon examination.

PREDATOR LENGTH *V.* PREY LENGTH

The range of absolute prey sizes eaten expanded with increasing L_T for all five predators (Fig. 3), thus resulting (particularly in megrim) in highly heteroscedastic error distributions. Upper and lower limits of these relationships (90% and 10% quantiles) changed at different rates (Table I), and in every case the

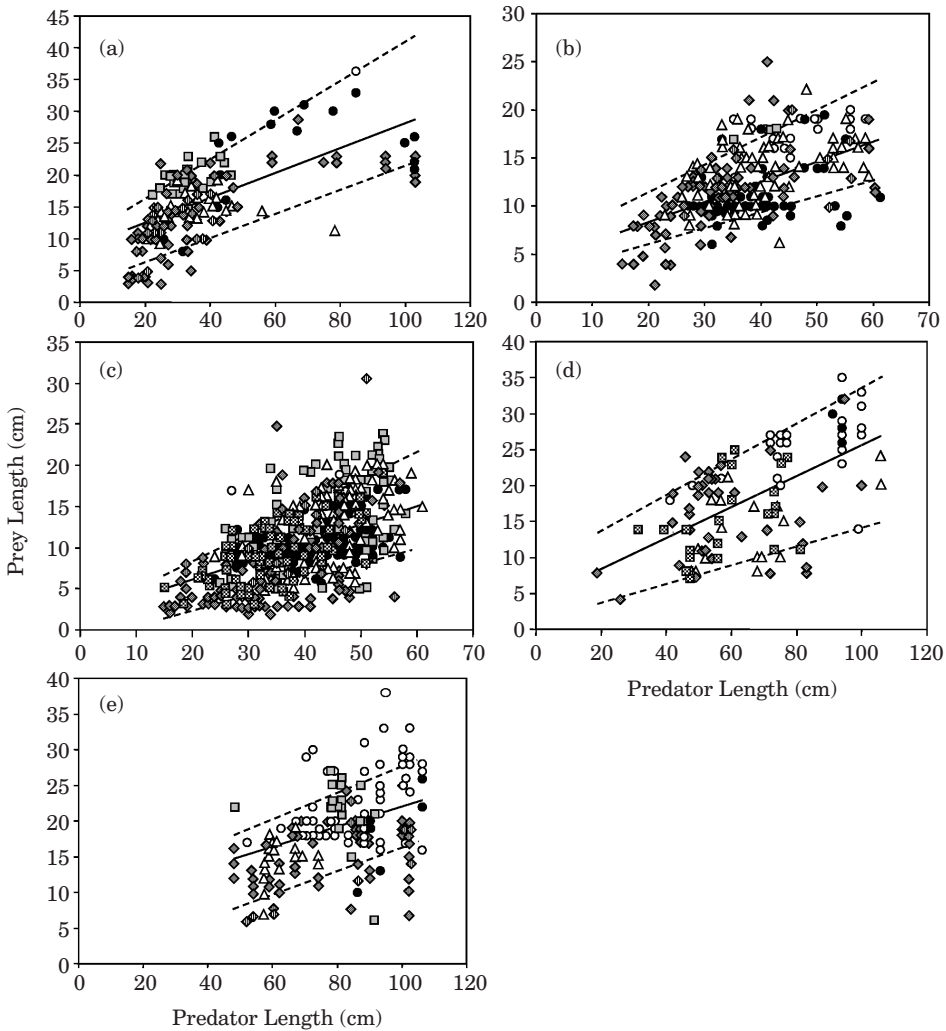


FIG. 3. Lengths of predator and prey (fishes: \blacklozenge , *Argentina* spp.; \blacksquare , *Callionymus* spp.; \blacklozenge , Clupeidae; \blacksquare , *Micromesistius poutassou*; \odot , *Scomber scombrus*; \bullet , *Trachurus trachurus*; \triangle , *Trisopterus* spp.; \blacklozenge , other fishes) in the stomachs of (a) hake, (b) whiting, (c) megrim, (d) cod and (e) saithe. The median (50%; —) and minimum (10%) and maximum (90%; ---) are indicated.

TABLE I. Regression equations (generated using quantile regression techniques) relating median, maximum (90% quantile) and minimum (10% quantile) prey length (in cm) to predator length. Only fish prey were considered here

Predator species	Median (50% quantile)	90% quantile	10% quantile	Mean \pm S.D. predator: prey size ratio	<i>n</i>
Hake	$y = 0.20x + 8.51$	$y = 0.31x + 10.04$	$y = 0.19x + 2.51$	2.60 ± 1.14	267
Megrim	$y = 0.22x + 1.67$	$y = 0.33x + 1.67$	$y = 0.19x - 1.31$	4.29 ± 1.95	690
Cod	$y = 0.24x + 3.44$	$y = 0.25x + 8.50$	$y = 0.13x + 1.23$	4.07 ± 1.71	110
Whiting	$y = 0.21x + 4.21$	$y = 0.29x + 5.71$	$y = 0.17x + 2.67$	3.18 ± 0.99	277
Saithe	$y = 0.14x + 7.71$	$y = 0.18x + 9.21$	$y = 0.17x - 0.33$	4.57 ± 1.61	294

slope of the relationship between predator L_T and maximum (90% quantile) prey size was greater than the slope of the relationship with minimum prey size (10% quantile) (Table I). This indicated that large predators continued to consume disproportionately small prey, throughout their lives.

Overall, for each predator, prey size increased as predator L_T increased, and at different size thresholds predators targeted different prey types (Fig. 3). For hake [Fig. 3(a)] horse-mackerel and clupeoids [sardine *Sardina pilchardus* (Walbaum)] were targeted by larger predators (>50 cm), whilst blue-whiting, *Argentina* spp., *Trisopterus* spp., and small clupeoids [mostly sprat *Sprattus sprattus* (L.)] were exploited by smaller predators (<50 cm).

Whiting continued to consume a wide diversity of fish prey throughout their lives [Fig. 3(b)], although apparently targeting more clupeoids when they were smaller (<40 cm) and only eating mackerel when they themselves were >40 cm and the mackerel were 15–20 cm.

Megrim consumed a wide diversity and size-range of prey [Fig. 3(c)], including many dragonettes (*Callionymus* spp.) particularly when predators were <40 cm, and *Trisopterus* spp. was eaten by larger predators. Blue-whiting and *Argentina* spp. were consumed over the whole predator length range, but were supplemented with 'other' fishes (e.g. gobies and small flatfishes) in smaller predators.

The data for cod [Fig. 3(d)] were relatively sparse (only 110 data points), but it would seem that mackerel (>20 cm L_T) were selected by large predators (>70 cm), whilst dragonettes and *Trisopterus* spp. were favoured by predators <80 cm. Clupeoids (18–25 cm) were only targeted by cod of 45–60 cm.

Data were only available for large saithe (40–110 cm), which selected mackerel >15 cm long and clupeoids and *Trisopterus* spp. of 6–20 cm. Predators of 80–95 cm also targeted blue-whiting [Fig. 3(e)].

The average prey: predator length ratio was lowest for hake and whiting (Table I). The lower the ratio, the larger the prey (on average) chosen by the predator. Thus saithe, megrim and cod tended to choose smaller prey relative to their own size, than did hake and whiting. There was no significant correlation between the slopes of the 90% quantile relationship and the slopes of the median (50% quantile), across the five predators examined ($r = 0.560$). Similarly there was no significant correlation between the slopes of the 10% quantile line and that of the median ($r = 0.394$).

TABLE II. Standardized forage ratio, α_a . Preference in relation to a 'portfolio' of seven key-prey types (irrespective of year). The standardized forage ratio as presented, ranges between 0 and 1, with $\alpha_a=0$ representing complete avoidance and $\alpha_a=1$ exclusive feeding. $\alpha_a=0.143$ denotes 'random-feeding'

Predator	Prey						
	<i>Argentina</i> spp.	<i>Callionymus</i> spp.	Clupeoids	Blue- whiting	Mackerel	Horse- mackerel	<i>Trisopterus</i> spp.
Cod	0.012	0.963	0.009	0.002	0.001	0.000	0.013
Hake	0.400	0.053	0.249	0.166	0.019	0.005	0.108
Megrim	0.076	0.855	0.039	0.007	0.000	0.000	0.023
Saithe	0.471	0.000	0.479	0.016	0.016	0.000	0.019
Whiting	0.136	0.106	0.630	0.012	0.003	0.002	0.111

PREFERENCE V. PREY AVAILABILITY

In terms of Chesson's standardized forage ratio α_a (Table II), relative to a portfolio of the seven key fish prey; and assuming that anything greater than 0.143 ($\alpha_a = k^{-1}$) represents positive selection, then mackerel, horse-mackerel and *Trisopterus* spp. were never positively selected. Cod and megrim showed a particular preference for dragonettes (*i.e.* small benthic fishes), whiting chose clupeoids, saithe chose argentinines and clupeoids, whilst hake exhibited a particular preference for all types of small pelagic prey (*i.e.* blue-whiting, argentinines and clupeoids). Pearre (1982) has demonstrated that α_a estimates are not sensitive to the absolute count of potential prey items in the environment, however α_a is very sensitive to counts or proportions assumed from the stomach contents data.

Correlation analyses of ICES stock numbers for mackerel, horse-mackerel and blue-whiting using Kendall's tau test, revealed no significant relationship between horse-mackerel and mackerel stocks ($r=0.341$, $P=0.056$). Similarly there was no significant correlation between blue-whiting and mackerel stocks ($r=-0.127$, $P=0.076$). Blue-whiting and horse-mackerel numbers were positively related ($r=0.657$, $P=0.056$), illustrating that horse-mackerel and blue-whiting stocks have exhibited broadly similar temporal dynamics over the past 22 years, whilst temporal patterns were very different in the mackerel stock.

When the proportion of mackerel, horse-mackerel and blue-whiting in predator stomachs was plotted against the availability of these prey as defined by ICES stock assessments, there were many positive trends, demonstrating that predators do select certain preys in proportion to their availability in the environment. However, Kendall's tau indicated that the only significant or near significant correlations were for megrim and hake feeding on blue-whiting ($P=0.003$ and 0.05 respectively). These data contained many outliers and much variability, largely related to the very limited number of stomachs sampled in some years (Appendix). Many of these outliers were effectively removed by the 20% 'trimming' procedure employed prior to the application of Kendall's tau test.

When the proportion of a particular prey represented in stomachs was plotted against the 'availability', as determined by CEFAS spring survey data

(1982–1994), there were again many positive relationships. Because of the marked variability in the data, few of these proved to be statistically significant. Despite the relatively large number of whiting stomachs sampled, there were no instances where the proportion in the stomach and availability in the environment were significantly correlated. Megrim appeared to consume significantly more dragonettes and gobies in years when these prey were more abundant ($P=0.036$ and 0.009 respectively). Cod consumed more blue-whiting when these fish were abundant ($P=0.043$) and hake chose more horse-mackerel and blue-whiting in years when these prey were more abundant ($P=0.053$ and 0.053).

In the analyses conducted here, only positive correlations were tested for. There was a clear relationship between consumption of *Trisopterus* spp. by cod, and the abundance of *Trisopterus* spp. in the environment. Because this was a strong negative relationship, however, the test for a positive correlation was rejected ($P=0.958$).

DISCUSSION

Examination of stomach contents remains one of the few tools available for understanding the linkages which occur within natural ecosystems. Stomach content analyses, however, have a number of limitations (Deb, 1997). For example, they tend to provide mere ‘snapshots’ of diets at particular points in time and space, and results depend extensively on the number of samples collected and the subjectivity and taxonomic knowledge of the investigator. Rarely are all links which occur in natural systems adequately quantified using stomach content data (Cohen & Newman, 1988) and for piscivorous species, an additional problem may stem from the fact that animals frequently regurgitate food upon capture (Bowman, 1986). This might explain the relatively high proportion of apparently empty stomachs observed here for megrim, whiting and saithe.

PREY SIZE

The analyses presented here demonstrated that for several predator species the proportion of the diet represented by fishes greatly increased as the animals became larger. For cod however, the proportion represented by invertebrates remained relatively high, even in large animals and this was unlike the pattern which has been observed in the North Sea, where the fraction of invertebrates in cod diets decrease with increasing L_T (J. Hislop, P. J. Bromley, N. Daan, H. Gislason, H. J. L. Heessen, A. P. Robb, D. Skagen, H. Sparholt, & A. Temming, pers. Comm). For cod in the Irish Sea, the decapod *Nephrops norvegicus* is a very important prey (Armstrong, 1982), and diets-at-length seem on the whole, to be broadly similar to those observed in the Celtic Sea.

Many studies have demonstrated that fishes of varying taxonomic groups have an ‘optimal’ prey size, which is selected if the predator is given a choice (Hart & Connellan, 1984). Fishes tend to grow faster when feeding on their ‘optimal’ prey, and models based on particulate feeding fishes suggest that the optimal prey size should be the largest size that a predator can handle. In the present study it was demonstrated that at various size thresholds (of both the

predator and prey), the feeding preference of the predator changes. Thus for example, only above a predator size of 40 cm and a prey size of 15 cm, do whiting start to consume mackerel. The existence of these apparent size thresholds might be seen as supporting one of the predictions of optimal foraging theory (OFT), that 'foraging should obey a quantitative threshold rule for when specific prey should be included or excluded from the diet' (Sih & Christensen, 2001). In each case, however the predator size and prey size distributions were highly asymmetric and predators did not always select the largest prey possible, which would presumably provide the best return for each feeding event. Large predators continued to eat small prey, which could be construed as being against one of the other key principles of OFT, that 'predators should prefer prey that yield more energy per unit handling time' (Scharf *et al.*, 2000). In a recent study by Floeter & Temming (2003) it was demonstrated that >75% of fishes found in the stomachs of North Sea cod originated from the least preferred (*i.e.* the smallest) quantile of the prey size range. As a consequence, these authors suggested that relative prey abundance rather than prey size preference was by far the dominant force determining composition of fish diet.

Many factors can influence the size and type of prey chosen by a predator (Juanes *et al.*, 2002). These include the effort required to find, pursue and handle the prey, the morphology or visual acuity of the predator and the behaviour and habitat use of the prey (which in-turn affect the encounter rate). In the present study, the observation that small prey were retained in the diets of larger predators was not unusual. Scharf *et al.* (2000) demonstrated similar patterns in 18 species of marine fishes (including some of the same species listed here). These authors hypothesized that the combination of high relative abundance and high capture probability for small (often benthic) prey, relative to large (often pelagic) prey, may lead to consistently high vulnerability to predation for small fishes. Schoener (1979) explored the relationships between prey L_T , profitability and distance at which the prey are first encountered. Schoener (1979) noted that the most profitable prey L_T becomes larger as the encounter distance (and hence the difficulty in pursuing and capturing the prey) increases (Stephens & Krebs, 1986).

PREY TYPE AND AVAILABILITY

In balancing costs and benefits, optimally foraging predators can adjust their preferences to take into account prey quality (Anthony *et al.*, 2000). A diet high in lipid provides sufficient metabolizable energy for maintenance, so dietary protein can be allocated to tissue synthesis and growth. Pelagic fishes tend to have a higher lipid content than do demersal or benthic species, hence the observed inclusion of small benthic fishes (*e.g.* the dragonettes, *Trisopterus* spp. and gobies 'positively selected' by cod), may represent a trade-off between the low-value nature of the prey, and the energy saved in pursuit and capture. Interestingly, smaller pelagic species (*e.g.* blue-whiting and argentines), tend to be more lipid-rich than larger pelagic species (*e.g.* mackerel) (Van Pelt *et al.*, 1997). As a consequence it might be expected that predators would exhibit a greater preference for these 'small pelagics' when they are available. The α_a values estimated here seem to suggest that hake and whiting did prefer

small pelagic prey (*Aregentina* spp., clupeoids and blue-whiting), and there was very little evidence to suggest that either mackerel or horse-mackerel (*i.e.* larger pelagics) were preferentially selected by any of the five predators considered.

LIMITATIONS OF THE DATA

Caution must be exercised in interpreting preference indices, in this case α_a , because the U.K. survey data, used as an estimate of numbers of prey in the environment (*i.e.* availability), do not necessarily represent a true picture of the ecosystem. All trawl gears are selective and the choice of sampling method used can greatly affect the perception of the structure and dynamics of the ecological community in a given area (V. Trenkel, unpubl. data). The U.K. spring groundfish survey (using a modified Portuguese high-headline otter trawl) probably under-estimates the abundance of small benthic fishes, even though it is equipped with bobbins, rubber disks and tickler chains on the groundrope, aimed at specifically increasing the capture efficiency for such species (Engås & Godø, 1989). A small experimental beam trawl survey carried out in the same area caught many more gobies, dragonettes and flatfishes per unit area, than did the standard U.K. sampling gear (cod-end mesh 20 mm) (J. Ellis, unpubl. data). The overall implications for prey preference indices of not adequately sampling small benthic fishes, would be to apparently inflate the suggested preference for these rarer species, at the expense of large-pelagics (mackerel, horse-mackerel, blue-whiting) which dominate in the survey. The U.K. survey was originally established specifically to investigate the distribution and biology of mackerel (Warnes & Jones, 1995); it was only widened to cover other more benthic species at a later stage.

In the present paper the proportion of blue-whiting, mackerel and horse-mackerel in the diet of predators were compared with their abundance, as determined by ICES stock assessments. This was completely independent of the U.K. spring groundfish survey and its associated problems, yet there was again no evidence for density-dependent feeding by predators on either mackerel or horse-mackerel. From these analyses blue-whiting emerged as an important prey for megrim and hake, with consumption relative to other prey animals, increasing in years when blue-whiting stocks were high. The importance of blue-whiting for predators in the Celtic Sea and Bay of Biscay has also been noted by Du Buit (1982).

The number of significant correlations observed between prey abundance and the proportion in the predator stomachs, was relatively low. Thus to answer the question posed in the title of this research paper, it might be concluded that the diet of Celtic Sea fishes does not generally reflect prey availability. If many such relationships had been found to exist, however, this would have inferred that piscivorous predators are indiscriminate opportunists, consuming whatever are available and abundant at the time. Such an observation would clearly be inconsistent with the idea that animals evolve to occupy ecological niches, thereby avoiding interspecific competition for resources. The results seem to infer that each predator species exhibits some flexibility in feeding preferences, but they each have a different 'portfolio' of suitable prey types (*i.e.* they are not indiscriminate), and they respond to changes of relative abundance of prey within their particular portfolio.

It should be acknowledged that the data available have many shortcomings, notably that 33% of stomach observations came from a single year (1984) and that trawl survey gears do not give a true picture of fish abundance in every case. Also it is important to recognize that with weak data some wrong conclusions are probably made, but this remains the first study to look at predator–prey relationships in the Celtic Sea.

Any relationships between predator preferences and prey availability apparent from the data, should be considered as being of the upmost importance since this could mean that fishing pressure exerted on a prey species (*e.g.* blue whiting) might have a significant indirect impact on predator stocks (*e.g.* hake and cod) or *vice versa* (Pinnegar *et al.*, 2000). It is becoming increasingly apparent that individual fish stocks cannot be managed in isolation and fishery managers are now being required to take into account wider ecological considerations (Botsford *et al.*, 1997). Incorporation of complex feeding behaviour, such as that described here, into fisheries models is crucial if realistic fishery predictions are to be made in the future and fish stocks managed on a sustainable basis (Magnusson & Palsson, 1991).

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APPENDIX. Number of predator stomachs containing identifiable fish prey

Year	Predator Species				
	Cod	Hake	Megrim	Saithe	Whiting
1977	5	19	17	0	45
1978	0	25	16	3	64
1979	12	4	8	16	2
1981	33	57	22	35	38
1982	43	25	12	0	0
1983	3	202	223	0	66
1984	669	187	146	0	627
1985	46	365	317	42	69
1986	0	126	54	0	53
1987	7	5	1	13	0
1988	0	59	29	0	0
1991	61	232	106	186	92
1992	26	48	43	6	65
1993	43	28	93	17	32
1994	1	3	11	0	0
Total	949	1385	1098	318	1153