# Bayesian Modelling of Catch in a Northwest Atlantic Fishery 

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#### Abstract

We model daily catches of fishing boats in the Grand Bank fishing grounds. We use data on catches per species for a number of vessels collected by the European Union in the context of the Northwest Atlantic Fisheries Organization. Many variables can be thought to influence the amount caught: a number of ship characteristics (such as the size of the ship, the fishing technique used, the mesh size of the nets, etc.), are obvious candidates, but one can also consider the season or the actual location of the catch. Our database leads to 28 possible regressors (arising from six continuous variables and four categorical variables, whose 22 levels are treated separately), resulting in a set of 177 million possible linear regression models for the log of catch. Zero observations are modelled separately through a probit model. Inference is based on Bayesian model averaging, using a Markov chain Monte Carlo approach. Particular attention is paid to prediction of catch for single and aggregated ships.


Keywords. Bayesian model averaging; Categorical variables; Grand Bank fishery; Predictive inference; Probit model

## 1. Introduction

The mismanagement of the world fisheries is one of the most important global environmental problems that we face today. Nine of the world's 17 major fisheries are in serious decline, and four others are classified as 'commercially depleted' by the Food and Agricultural Organization of the United Nations (Tibbets, 1994).

The Northwest Atlantic Fisheries Organization (NAFO) is one of several international organizations that tries to alleviate overexploitation through voluntary cooperation. It was established in 1978 to contribute to the optimal exploitation and rational use of fisheries resources in the Grand Bank outside Canada's exclusive economic zone (see http:/ / www.nafo.ca for a map of the area covered by the treaty). Countries which are members of the NAFO assign quotas among themselves and grant inspection rights to each other. Three inspection ships -two Canadian and one belonging to the European Union - board vessels of member states and register the information in their logbooks. In addition, ships from signatory countries report (through the so-called "hails") their entry and exit of the different zones of the fishing grounds. Finally, there are two daily flights over the Grand Bank and the Flemish Cap made by inspection airplanes with the purpose of locating and identifying all ships fishing in the area. Boarding ships on high seas to verify catch is expensive and disrupts their operations. Furthermore, ships from non-signatory countries can not be inspected. It then becomes important to construct models that allow for catch prediction and monitoring conditional on the information from aerial sightings and hails, ship characteristics and other variables (such as month of the year). Thus, our aim is to model how all these variables influence catch. This could provide useful information for regulatory measures and guidelines related to issues like mesh size, optimal size of the fleet, etc. More importantly,
it allows to estimate the total amount caught by a group of ships operating in a certain area at a certain time of the year.

The data that we have consist of daily catch (per ship) per species of fish. Since there are many days with zero catch for a given species of fish, our statistical model incorporates a positive probability of zero catch through a probit model. When catch occurs, the log of the quantity caught is modelled through a linear regression structure, where we formally treat the uncertainty concerning the choice of regressors through model averaging in a Bayesian setting using posterior model probabilities as weights. In view of the large number of potential models, we explore the posterior distribution using a Monte Carlo Markov chain over the model space in the spirit of the $\mathrm{MC}^{3}$ methodology of Madigan and York (1995). The Bayesian framework leads to exact small sample results, fully taking both parameter and model uncertainty into account. In the present application we have not used any strong prior information or a formal decision theory framework. Both of these can, however, easily be incorporated into a Bayesian analysis.

The aims of this paper are quite different from those of the large literature in stock assessment, where statistical methods are used in order to assess the size of fish stocks, see, e.g., Hilborn and Walters (1992) for a general introduction and McAllister and Kirkwood (1998) for an overview of Bayesian stock assessment methods. A variety of statistical methods, such as Bayesian statespace models (Millar and Meyer, 2000) and spatial methods (Newman, 1998) has recently been introduced into this literature. There exists, in addition, substantial work on estimation of yeareffects and abundance trends based on modelling catch per hour fished; Quinn and Deriso (1999) provide many examples. In contrast to the above, and like Ferreira and Tusell (1996), our aim is to shed light on how catch can be explained by certain observable characteristics -such as mesh size (Robichaud et al. 1999)—, and provide operational forecasts of commercial landings of various species (Stergiou et al. 1997). It is important to stress that the main aim of our analysis is not necessarily to develop a model that describes the dynamics of fisheries as closely as possible, but rather to provide a framework that can successfully be used for short-term predictions of quantities caught (of a certain species by a certain ship or group of ships) given an easily available information set. This will guide the modelling strategy and the choice of covariates that we will consider.

Section 2 describes the data, while Section 3 introduces the statistical model. The zero observations are treated in Section 4, and the analysis of positive catch is discussed in Section 5. Section 6 focuses on prediction. The empirical results are presented in Section 7 and a final section concludes. Details of the computational implementation are presented in an Appendix.

## 2. The Data

The original data were gathered by the inspection vessel of the European Union operating on the Grand Bank fishery. Inspectors board the fishing boats and record basic characteristics of the ship and the fishing equipment, as well as the quantities caught of different species and where and when this catch was effectuated. They use the ship's logs to collect all the information accumulated since the last time the ship was boarded. All data correspond to 1993 and the first half of 1994, leading to 6,806 observations each corresponding to a particular ship at a given day. In all, there are 59 different ships.

The dependent variable is the live weight of fish caught. Table 1 summarizes the regressors that we consider using. These include four categorical variables: the year when the catch is made ( 2 levels), fishing technique ( 4 levels), zone or division within the fishing grounds ( 4 levels) and month of the year ( 12 levels). In addition, we have four continuous variables, namely mesh size measured in milimeters, length of vessel measured in meters, gross registered tonnage (GRT) and engine kW . See e.g. King (1995, Ch. 2) for a description of fishing gear and methods.

Our dataset also provides the nationality of the ship but we have decided not to consider this variable since one of the purposes of the analysis is to predict catch of ships from non-signatory countries (for which we have no observations). However, we do have a year effect. This is

Table 1. Data Statistics

|  | Regressor | \% Observations |
| :--- | :--- | ---: |
| 1 | Year 1993 | 75.36 |
| 2 | Year 1994 | 24.64 |
| 3 | Drift gillnet | 3.60 |
| 4 | Anchored gillnet | 1.44 |
| 5 | Otter trawl | 79.64 |
| 6 | Otter trawl pair | 15.32 |
| 7 | Zone 3L | 34.64 |
| 8 | Zone 3M | 25.69 |
| 9 | Zone 3N | 35.05 |
| 10 | Zone 3O | 4.62 |
| 11 | January | 4.89 |
| 12 | February | 10.74 |
| 13 | March | 15.05 |
| 14 | April | 12.06 |
| 15 | May | 13.99 |
| 16 | June | 9.48 |
| 17 | July | 7.02 |
| 18 | August | 7.71 |
| 19 | September | 7.98 |
| 20 | October | 7.04 |
| 21 | November | 3.48 |
| 22 | December | 0.56 |
| 23 | Gillnet $\times \log [0.5+$ Mesh size $-\min ($ Mesh size $)]$ |  |
| 24 | Gillnet $\times\{\log [0.5+\text { Mesh size }-\min (\text { Mesh size })]\}^{2}$ |  |
| 25 | Trawl $\times \log [0.5+$ Mesh size $-\min ($ Mesh size $)]$ |  |
| 26 | Trawl $\times \log [0.5+$ Engine kW $-\min ($ Engine kW $)]$ |  |
| 27 | log(Length vessel) |  |
| 28 | log(GRT) |  |


|  | Min | 1st quartile | Median | 3rd quartile | Max |
| :--- | ---: | ---: | ---: | ---: | ---: |
| Mesh size (mm.) gillnetters | 110 | 130 | 140 | 140 | 150 |
| Mesh size (mm.) trawlers | 120 | 120 | 120 | 130 | 150 |
| Engine kW trawlers | 588 | 845 | 1164 | 1470 | 2648 |
| Length vessel (m.) | 29.0 | 42.0 | 47.0 | 61.2 | 84.9 |
| GRT | 252.3 | 376.9 | 664.9 | 970.2 | 2382.0 |

because year class effects are important in fisheries and, from a biological point of view, it would not be sensible to assume equality of catches in, e.g., May 1993 and May 1994. Inevitably, this complicates prediction for years for which no data are available.

The way that mesh size and engine power influence catch is potentially very different for gillnets and otter trawls. Thus, we include these variables in terms of interactions with an indicator variable for the net type used: gillnets (adding drift and anchored) and trawls (both single and paired). In addition, there is prior reason to assume that the effect of mesh size might be nonlinear for gillnets (e.g., catch would decrease if mesh size would either be too large or too small), so we include a quadratic interaction term for this fishing technique. An effect of engine kW on quantity caught is quite plausible for otter trawls (which are towed), but is very unlikely for gillnets (which are passive), so we do not include an interaction term for engine power and gillnets. In order to reduce the collinearity between these interaction terms and fishing techniques, the continuous
variables kW and mesh size are transformed as indicated in Table 1. This substantially increases the spread of the interaction variables and reduces the collinearity in the design matrix. The other continuous variables (length of the vessel and tonnage) are transformed to logarithms in the usual way.

Table 1 indicates the empirical distribution of each of the categorical variables, and quantiles of the continuous variables (before transformation). The data can be obtained from this journal's website. Of course, Table 1 provides only marginal information. Some complementary information is given in Figure 1, where we present bivariate histograms (with lighter shades corresponding to higher relative frequencies) of some combinations of regressors for each of the years in the sample. Levels for the categorical variables are ordered as in Table 1 and continuous variables are categorized into five bins of equal width. From this we note a shift in 1993 from zone $L$ in the period January-May to zone N for the remaining months of the year. The available months of 1994 show a somewhat more even spread over zones $\mathrm{L}, \mathrm{M}$ and N . The month versus mesh size plots are presented for gillnets and trawls separately, which shows that trawlers tend to use smaller mesh than gillnetters. We also see a tendency towards the greater use of small mesh trawl nets (often 120 mm ) in the months April-June 1993. Finally, the length and GRT of the ships are obviously positively correlated as can be seen from the last row of plots.


Fig. 1. Bivariate Greyscale Plots for Regressors
(Lighter shades correspond to higher relative frequencies)

Table 2 lists the five most important species caught in the Grand Bank and has one category for all the other species ("rest"). Every time we observe a ship, we observe its daily catches for all six species. A look at the data tells us that a ship's catch on any given day often does not include all species. In particular, we are going to model the first five species listed in Table 2, for which the
percentage of zeroes in the data (6806 ship-days) ranges from $18.50 \%$ (halibut) to $88.33 \%$ (cod). Thus, this is an important aspect, which, if overlooked, would lead to substantial overestimation of catch. Hence, we shall model zero catches explicitly, by means of a probit model. This feature of the data was not accounted for by Ferreira and Tusell (1996), who analyze the same data set but only take the positive observations into account. Table 2 also lists the fraction of the total live weight that each species constitutes. We will consider separate models for each of the species, to allow for the explanatory variables to affect catch for each species differently.

Table 2. Catch for Different Fish Species

| Sp. | Description | Mean (kg) | Std (kg) | $\%$ zeroes | $\%$ of catch |
| ---: | :--- | ---: | ---: | ---: | ---: |
| 1 | Atlantic cod (Gadus morhua) | 550.20 | 2517.48 | 88.33 | 9.80 |
| 2 | Greenland halibut (Reinhardtius hippoglossoides) | 3503.48 | 3610.55 | 18.50 | 62.40 |
| 3 | Redfish (Sebastes sp.) | 658.51 | 2794.49 | 85.73 | 11.73 |
| 4 | Roundnose grenadier (Coryphaenoides rupestris) | 213.46 | 502.18 | 43.20 | 3.80 |
| 5 | Skate (Raja sp.) | 503.39 | 1661.50 | 55.44 | 8.97 |
| 6 | Rest | 185.23 | 625.76 | 72.27 | 3.30 |

## 3. The statistical model

In this section we outline our statistical model for daily catch of a given species of fish per ship. The observations will be denoted by $s_{i}, i=1, \ldots, n(n=6806)$, and we define $s=\left(s_{1}, \ldots, s_{n}\right)^{\prime}$. Clearly, each of the $n$ observations is non-negative, and a certain number of them, say $Q$, are strictly positive (those that correspond to positive catch). For notational convenience, we shall order the observations so that the first $Q$ observations are positive, whereas the remaining $n-Q$ observations are equal to zero.

As explained in the previous section, it is crucial to take account of the fact that there is a positive probability of zero catch. A natural approach is to use the probit model:

$$
\begin{cases}s_{i}=0 & \text { with probability }\left(x_{i}^{\prime} \gamma\right),  \tag{3.1}\\ s_{i}>0 & \text { with probability } 1-\left(x_{i}^{\prime} \gamma\right),\end{cases}
$$

where (.) denotes the c.d.f. of the standard Normal distribution, the vector $x_{i} \in \Re^{1+k}$ contains the element 1 as well as the explanatory variables presented in Table 1 , and $\gamma \in \Re^{1+k}$ groups the parameters. Each categorical variable is handled through dummies taking the values zero or one, with one level excluded (thus, $k=24$ instead of 28 ). Throughout the paper, the design matrix $X \equiv\left(x_{1}, \ldots, x_{n}\right)^{\prime}$ will be of full column-rank.

If $s_{i}>0$, we further assume a linear regression structure for $y_{i} \equiv \log \left(s_{i}\right)$. This is easier to handle than the probit model, so a more ambitious strategy is feasible. In particular, we will allow for model uncertainty, where each of the potential models considered will assume that

$$
\begin{equation*}
y_{i}=\log \left(s_{i}\right) \text { is distributed as } \operatorname{Normal}\left(\alpha+z_{i}^{\prime} \beta, \sigma^{2}\right), i=1, \ldots, Q, \tag{3.2}
\end{equation*}
$$

and the vector $z_{i}$ corresponds to a subset of the regressors in Table 1. For computational convenience, all the variables are now demeaned, so that each column of the resulting design matrix $Z \equiv\left(z_{1}, \ldots, z_{Q}\right)^{\prime}$ sums to zero. The matrix $Z$ is also of full column-rank. In (3.2), $\alpha \in \Re$ is the intercept, $\sigma^{2}>0$ denotes the sampling variance, whereas the vector $\beta$ groups the regression coefficients.

Note that (3.1) and (3.2) have been modelled entirely separately, using different parameters, and we will also assume prior independence between the parameters in (3.1) and (3.2). This is done partly for pragmatic reasons (as in this case we can conduct posterior inference independently, greatly simplifying the computations), but also because it is not obvious to us that the effects of a given variable on the probability of zero catch and on the actual amount caught (when catch
is positive) should be linked. One might possibly consider sign restrictions for the elements of $\gamma$ in (3.1) and $\beta$ in (3.2). For example, for otter trawls, increasing mesh size could be expected to decrease the amount caught and to increase the probability of zero catch: this would imply a negative component in $\beta$ and a positive component in $\gamma$, but it would not mean that their actual magnitudes are necessarily linked. Thus, such restrictions would not imply that both models should be analysed jointly. Here we have chosen not to impose prior constraints like these, and will instead let the data find the most appropriate parameter ranges. As we do not possess strong prior information, our prior distribution (presented in the following two sections) will generally try to incorporate as little subjective input as possible.

We will use the entire sample to make inference on $\gamma$ and to predict the probability of zero versus positive catch; this analysis only uses the fact whether $s_{i}$ is zero or strictly positive. The actual value of the $Q$ positive observations will be used to conduct inference on $\alpha, \beta$ and $\sigma$, and to predict the amount of catch given that it is positive. The probit model will be examined in Section 4, whereas Section 5 will be devoted to the model for positive catch. Lo et al. (1992) also model zero observations separately from positive ones in the context of analyzing relative fish abundance, using classical statistical procedures and a simple linear probability model for zero observations.

## 4. Analysis of zero observations

In this section, we focus on posterior inference on $\gamma$, the parameter in the probit model in (3.1). We shall complement the latter sampling distribution with the prior

$$
\begin{equation*}
p(\gamma)=f_{N}^{1+k}\left(\gamma \mid 0,\left(h_{0} X^{\prime} X\right)^{-1}\right), \tag{4.1}
\end{equation*}
$$

i.e., $\mathrm{a}(1+k)$-variate Normal distribution with zero mean and covariance matrix $\left(h_{0} X^{\prime} X\right)^{-1}$, where $h_{0}>0$. This corresponds to the $g$-prior introduced in Zellner (1986), and essentially says that the prior precision is a fraction $h_{0}$ of that of the sample. This prior is often used for relatively high-dimensional parameters in the context of a lack of strong prior information, as it typically does not distort the information in the sample. We took zero as the prior mean for $\gamma$, since, from (3.1), $P\left(s_{i}=0 \mid \gamma=0\right)=1 / 2$. For $h_{0}$ we adopt the value $h_{0}=1 / n$, which roughly corresponds to the information in one observation and will easily be dominated in posterior and predictive inference. With the prior in (4.1) the predictive distribution is invariant with respect to the choice of the reference levels for the categorical variables, as is desirable.

Although the posterior distribution corresponding to a sample of $n$ observations from (3.1) and the prior in (4.1) can not be computed analytically, we can use Gibbs sampling (with data augmentation) to approximate $p(\gamma \mid s)$ to any required precision (see Appendix A. 1 for details, and Albert and Chib (1993) for a similar probit analysis).

## 5. Analysis for positive observations

### 5.1. Model specification

We will incorporate model uncertainty in the sense that we allow for any subset of the variables in Table 1 to appear as regressors in (3.2). This means that instead of a single model, we have a set $\mathcal{M}=\left\{M_{j}: j=1, \ldots, J\right\}$, where each model corresponds to a particular choice of regressors.

In order to fully understand our model space $\mathcal{M}$, we need to explain carefully how we deal with categorical variables in this situation. We will treat different levels of a categorical variable separately, so that a model in $\mathcal{M}$ can include or exclude any level with the only restriction that not all levels of a categorical variable can be included in the same model. This gives us extra flexibility with respect to the simpler approach that treats categorical variables as single entities which can only be fully excluded (which means that all levels have exactly the same effect) or included (which implies that all levels have different effects). With our treatment, we also allow for intermediate situations where several levels of a variable have the same effect (and are,
therefore, excluded from the model) whereas other levels (the ones included in the model) have different effects. This is an issue of empirical relevance as the results in Section 7 will illustrate. Our approach implies that we can not fix a reference level, as we want to treat all levels in a symmetric fashion. As an example, consider the categorical variable month of the year, which has twelve levels. If we were to designate, say, December as a reference level we would be able to capture a situation where, e.g., January has the same effect as December (by also excluding January), but not a situation where January has the same effect as, say, February, yet not the same as December. By allowing a free reference level, we can accommodate any combination of levels having the same effect.

With $K$ continuous variables and $R$ categorical variables with $L_{1}, L_{2}, \ldots, L_{R}$ levels, respectively, this strategy implies a model space $\mathcal{M}$ with $J=2^{K} \prod_{r=1}^{R}\left(2^{L_{r}}-1\right)$ elements, which for our application leaves us with $176,904,000$ possible models. We stress that whereas all of the $\tilde{k}=28$ variables in Table 1 will appear in some of the models, the maximum number of regressors that any single model can contain is $k=28-4=24$ (since there are 4 categorical variables). Whenever a model contains all but one levels of a categorical variable, we say that the model is "full" in that categorical variable. Note that models that are full in one or several categorical variables appear with different representations in $\mathcal{M}$, each corresponding to a particular choice of reference level. This feature will be taken into account when setting a prior distribution for the models.

### 5.2. Priors under different models

We now turn to the issue of eliciting priors for the parameters in (3.2) given a particular model $M_{j}$. For these parameters we specify a prior distribution that incorporates minimal prior information while leading to analytical tractability. On the intercept $\alpha$ and the scale parameter $\sigma$, which are present in all the models, we assume the usual non-informative distributions, respectively defined through

$$
\begin{equation*}
p(\alpha) \propto 1 \quad \text { and } \quad p(\sigma) \propto \sigma^{-1} . \tag{5.1}
\end{equation*}
$$

For the vector $\beta_{(j)}$, which groups the relevant regression coefficients under model $M_{j}$, we assume the $g$-type prior

$$
\begin{equation*}
p\left(\beta_{(j)} \mid \sigma, M_{j}\right)=f_{N}^{k_{j}}\left(\beta_{(j)} \mid 0, \sigma^{2}\left(g_{0} Z_{j}^{\prime} Z_{j}\right)^{-1}\right), \tag{5.2}
\end{equation*}
$$

where $k_{j}$ is the number of explanatory variables included in $M_{j}$ and $Z_{j}$ denotes the corresponding design matrix. This prior specification requires minimal judgmental user input, since only the scalar $g_{0}$ is left to be chosen. We shall take $g_{0}=1 / \max \left\{Q, \hat{k}^{2}\right\}$, where $Q$ is the number of positive observations and $\tilde{k}$ is the number of available regressors in Table 1 . This choice is inspired by Fernández et al. (2001a), who find that the use of such a strategy for $g_{0}$ leads to very satisfactory identification of the correct model in simulation exercises, whereas out-of-sample predictive behaviour is also quite good. Besides their empirical simulation justification, they also derive a number of theoretical properties of this prior. Finally, model $M_{j}$ assumes that its excluded explanatory variables do not matter, i.e., that their associated regression coefficients are equal to zero. Now that we have specified the prior distribution, we can immediately conduct Bayesian inference under model $M_{j}$, by combining this distribution with the corresponding sampling model from (3.2). Since this prior distribution resembles a natural-conjugate, computing the posterior and predictive distributions is quite simple, as shall be explained later in the paper.

### 5.3. Model averaging

So far we have considered a single model $M_{j}$ from the space of all possible models $\mathcal{M}$. From a Bayesian perspective, model uncertainty can be treated in a coherent fashion by further specifying a prior distribution $P\left(M_{j}\right)$ on the models. Here we will consider a Uniform distribution on the space of genuinely different models. By this we mean that we take into account that $\mathcal{M}$ contains multiple copies of models which are full in some categorical variable, down-weighting their prior probabilities accordingly. If desired, other prior distributions could be considered with only minor modifications to our framework.

The posterior distribution of a quantity is now given by a mixture of the posterior distributions under each of the models, with mixing probabilities corresponding to the posterior model probabilities. Thus, Bayesian inference provides a coherent framework for treating model uncertainty, leading to an inferential procedure which averages over the inferences resulting from each of the individual models. Madigan and Raftery (1994), Raftery et al. (1997) and Fernández et al. (2001b) find in a series of empirical applications that, in the presence of model uncertainty, Bayesian model averaging leads to the best predictive performance, as measured by a logarithmic scoring rule. In a decision-theory context, mixing over models can be shown to be optimal under predictive squared error loss, provided the set of models considered is exhaustive (Min and Zellner (1993)). We follow this approach and consider model averaging rather than selecting one single model.

Applying Bayes' theorem, the posterior probability of model $M_{j}$ is given by

$$
\begin{equation*}
P\left(M_{j} \mid y\right) \propto l_{y}\left(M_{j}\right) P\left(M_{j}\right), \tag{5.3}
\end{equation*}
$$

where $P\left(M_{j}\right)$ is the prior probability and $l_{y}\left(M_{j}\right)$ the marginal likelihood of model $M_{j}$. The latter is obtained from (3.2), integrating out the parameters with their prior distribution described in Section 5.2. It is easy to show that $l_{y}\left(M_{j}\right)$ is finite if and only if the sample $y=\left(y_{1}, \ldots, y_{Q}\right)^{\prime}$ contains at least two different observations. This condition will be both necessary and sufficient for posterior and predictive inference throughout the paper.

Although we can derive an explicit expression for $l_{y}\left(M_{j}\right)$ (see (A.1) in Appendix A.2), direct computation of the posterior probability in (5.3) is very difficult due to the large number of models in $\mathcal{M}$ (approximately 177 million in our application). Therefore, we shall approximate the posterior distribution of the models via simulation, using a Markov chain Monte Carlo (MCMC) sampler on the model space $\mathcal{M}$. Appendix A. 2 provides more details on the particular sampler we have adopted, which is of the Metropolis-Hastings type. In case we have no categorical variables, the sampler essentially simplifies to the $\mathrm{MC}^{3}$ method of Madigan and York (1995) also used in Raftery et al. (1997).

### 5.4. Inference on regression coefficients

We now consider inference on a linear combination $b^{\prime} \beta \equiv \sum_{l=1}^{\tilde{k}} b_{l} \beta_{l}$ of the elements of the $\tilde{k}$ dimensional regression vector $\beta$, where $\tilde{k}=28$, corresponding to all variables in Table 1 . To do this, we need to apply the model averaging ideas explained in the previous subsection. Under model $M_{j}, b^{\prime} \beta$ takes the value zero if none of the regressors corresponding to a non-zero element of $b$ is included in $M_{j}$, and has a Student- $t$ distribution otherwise. The exact form of the posterior distribution of $b^{\prime} \beta$ is:

1. With probability $p \equiv \sum_{j: B_{j} b=\mathbf{0}} P\left(M_{j} \mid y\right)$,

$$
\begin{equation*}
b^{\prime} \beta=0 \tag{5.4}
\end{equation*}
$$

2. With probability $1-p, b^{\prime} \beta$ has density

$$
\begin{equation*}
\frac{1}{1-p} \sum_{j: B_{j} b \neq \mathbf{0}} f_{S}\left(b^{\prime} \beta \mid Q-1, \frac{b^{\prime} B_{j}^{\prime}\left(Z_{j}^{\prime} Z_{j}\right)^{-1} Z_{j}^{\prime} y}{g_{0}+1}, \frac{Q-1}{G_{j}} \frac{g_{0}+1}{b^{\prime} B_{j}^{\prime}\left(Z_{j}^{\prime} Z_{j}\right)^{-1} B_{j} b}\right) P\left(M_{j} \mid y\right), \tag{5.5}
\end{equation*}
$$

where $B_{j}$ is the relevant selection matrix under model $M_{j}$ in the sense that $\beta_{(j)}=B_{j} \beta$, with $\beta_{(j)}$ corresponding to the regressors included in $M_{j}, \mathbf{0}$ is a vector of zeroes of the appropriate dimension, and $f_{S}(x \mid \nu, m, a)$ denotes the p.d.f. of a Student- $t$ distribution with $\nu$ degrees of freedom, location $m$ (the mean if $\nu>1$ ) and precision $a$ (with variance $\nu /\{(\nu-2) a\}$ provided $\nu>2$ ). Finally, $G_{j}$ is defined in (A.2) in Appendix A.2. From (5.4) - (5.5) it is clear that, once we have run the Markov chain on $\mathcal{M}$ to compute $P\left(M_{j} \mid y\right)$, we can obtain the distribution of $b^{\prime} \beta$ analytically.

## 6. Prediction

We now focus on forecasting the value of a new observable, say $s_{f}$, given a vector of explanatory variables and the observed sample $s$. Our forecast for $s_{f}$ will be based on the out-of-sample predictive distribution, which is obtained from (3.1)-(3.2) after integrating out all the parameters and all possible models using their respective posterior distributions. From (3.1) - (3.2) it is immediate that the predictive distribution for $s_{f}$ will be a mixture of a point mass at zero and a continuous distribution. In particular, we have:
1.

$$
\begin{equation*}
s_{f}=0 \tag{6.1}
\end{equation*}
$$

with probability

$$
\begin{equation*}
\omega_{f} \equiv \int\left(x_{f}^{\prime} \gamma\right) p(\gamma \mid s) d \gamma, \tag{6.2}
\end{equation*}
$$

where $x_{f} \in \Re^{1+k}$ contains the element one and the explanatory variables for $s_{f}$. The integral in (6.2) can be calculated by averaging ( $x_{f}^{\prime} \gamma$ ) over the draws of $\gamma$ generated through the Gibbs sampler in Appendix A.1.
2. With probability

$$
\begin{equation*}
1-\omega_{f}, \tag{6.3}
\end{equation*}
$$

$s_{f}>0$ and it has p.d.f.

$$
\begin{align*}
p\left(s_{f} \mid y\right)=\frac{1}{s_{f}} \sum_{j=1}^{J} f_{S}( & \log \left(s_{f}\right) \mid Q-1, \bar{y}+z_{f(j)}^{\prime} \frac{\left(Z_{j}^{\prime} Z_{j}\right)^{-1}}{g_{0}+1} Z_{j}^{\prime} y,  \tag{6.4}\\
& \left.\frac{Q-1}{G_{j}}\left\{\frac{Q+1}{Q}+z_{f(j)}^{\prime} \frac{\left(Z_{j}^{\prime} Z_{j}\right)^{-1}}{g_{0}+1} z_{f(j)}\right\}^{-1}\right) P\left(M_{j} \mid y\right),
\end{align*}
$$

where $z_{f(j)}$ is the $k_{j}$-dimensional vector that contains the explanatory variables (demeaned as indicated after (3.2)) relevant under model $M_{j}$.

In a practical context, we may be interested in predicting the aggregate catch of a group of ships during a certain spell of time. This means that we focus on the predictive distribution of $s_{\text {sum }} \equiv \sum_{i=1}^{I} s_{f_{i}}$ rather than considering one single observable $s_{f}$ as was the case above. The predictive distribution of $s_{\text {sum }}$ is computed by averaging its sampling distribution over parameters and models using the relevant posterior distributions. It is clear from (3.1) that in the sampling $s_{\text {sum }}$ is zero with probability $\omega(\gamma) \equiv \prod_{i=1}^{I}\left(x_{f_{i}}^{\prime} \gamma\right)$ (where $x_{f_{i}} \in \Re^{1+k}$ corresponds to the explanatory variables for $s_{f_{i}}$ ), and has some p.d.f. with probability $1-\omega(\gamma)$. This means that we forecast:

1. $s_{\text {sum }}=0$, with probability $\omega_{\text {sum }} \equiv \int \omega(\gamma) p(\gamma \mid s) d \gamma$, which, as before, we compute by averaging $\omega(\gamma)$ over the Gibbs draws of $\gamma$.
2. With probability $1-\omega_{\text {sum }}, s_{\text {sum }}>0$ and has a predictive distribution given through some p.d.f. on $(0, \infty)$. Although an explicit expression for the latter p.d.f. is complicated to derive, we can approximate this distribution via simulation drawing a set of values from (3.1)-(3.2) where the parameters are, in turn, drawn from the posterior distribution (taking model averaging into account).

## 7. Discussion of results

### 7.1. Computational issues and model probabilities

Most of the discussion in this subsection will focus on the Markov chain on model space, since it is the most computationally demanding aspect of our model. In the interest of the practical importance of this methodology, and to enhance its appeal to applied researchers, we have made
particular efforts to create an efficient set of programs that can deal with problems of empirical relevance. The programs are coded in Fortran-77 and make efficient use of CPU-time, e.g., through storing results for already visited models in stacks (saving recalculations when a model is revisited by the chain). As a consequence, e.g., the entire single-ship analysis presented in the sequel takes between 1 and 3 hours (depending on the species) on a 200 MHz PowerPC-based desktop computer. The source code is available from this journal's website.

Throughout, we shall split the available observations into a subsample used for posterior inference (the "estimation subsample") and the remaining observations, which will be used for comparison with the predictive distribution (the "prediction subsample"). Observations are randomly assigned to the estimation subsample with probability 0.75 and the resulting number of observations in this subsample is $n=5087$ with $Q$ in Table 3 indicating the number of positive observations in this subsample. The total number of regressors is $\tilde{k}=28$ (all those in Table 1) for halibut, redfish and grenadier. For cod, $\tilde{k}=26$ because there are no catches in November or December. For skate, $\tilde{k}=27$ as we leave out the quadratic interaction term between mesh size and gillnets to avoid collinearity problems. Thus, we obtain $Q>\tilde{k}^{2}$ for halibut, grenadier and skate, which leads to choosing $g_{0}=1 / Q$ in the prior in (5.2), whereas for cod and redfish we choose $g_{0}=1 / \tilde{k}^{2}$.

Table 3. Monte Carlo Performance and Posterior Probabilities

|  | Species |  |  |  |  |
| :--- | ---: | ---: | ---: | ---: | ---: |
|  | cod |  |  |  |  |
| Number of Observations $Q$ | halibut | redfish | grenadier | skate |  |
| Number of Retained Drawings | 583 | 4161 | 727 | 2891 | 2256 |
| Number of Discarded Drawings | $1,000,000$ | 500,000 | $1,000,000$ | 500,000 | $2,000,000$ |
| Number of Models Visited | 500,000 | 100,000 | 500,000 | 100,000 | $1,000,000$ |
| Number of Non-equivalent Models Visited | 32,739 | 1906 | 18,264 | 2840 | 5202 |
| Wind. Est. and Emp. Freq. Correlation Coeff. | 24,229 | 485 | 15,940 | 1766 | 3266 |
| Weighted Average $q$ | 0.9890 | 0.9782 | 0.9919 | 0.9909 | 0.9659 |
| Post. Prob. Covered by Chain | 0.1602 | 0.0554 | 0.2834 | 0.0776 | 0.0392 |
| Post. Prob. of Best Model | 0.8811 | 0.9438 | 0.9530 | 0.9695 | 0.9988 |
| Number of Models Required for 90\% Post. Prob. | 0.0335 | 0.0510 | 0.0791 | 0.1019 | 0.0883 |
| Post. Prob. of Stepwise Model | 4022 | 144 | 1494 | 156 | 235 |

The Markov chain used for computing posterior model probabilities is described in Appendix A.2. Table 3 lists the number of retained drawings and the initial number of discarded draws (the "burn-in"), as well as the total number of visited models. We consider several strategies for assessing the convergence of this chain. Since the marginal likelihood for model $M_{j}, l_{y}\left(M_{j}\right)$, can be calculated explicitly, we will apply the formula in (5.3) to compute posterior probabilities on the basis of the models visited by the Markov chain (instead of using the empirical frequencies of visiting each model). This idea, called "window estimation" by Clyde et al. (1996), implies that the computed posterior odds (ratios of posterior probabilities) between any two models visited are the actual posterior odds. From Table 3 we see that the correlation coefficient between the posterior probabilities of all visited models computed on the basis of empirical frequencies and window estimation is always above 0.96 . This provides an indication of convergence of the chain.

A second diagnostic of convergence is based on the fact that models that are full in one or more categorical variables have exactly equivalent counterparts in the model space (that only differ in the chosen levels of the categorical variables for which they are full). Asymptotically, such equivalent models are visited equally often, which suggests looking at

$$
q \equiv \frac{\max _{i}\left\{\mathrm{freq}_{i}\right\}-\min _{i}\left\{\mathrm{freq}_{i}\right\}}{\sum_{i} \operatorname{freq}_{i}},
$$

where $\mathrm{freq}_{i}$ is the number of times the chain visits representation $i$ of the same model. Clearly, $q \in[0,1]$ with $q=0$ the best result and $q=1$ the worst, indicating that only one of the equivalent representations was visited. Table 3 reports a weighted average of the $q$ values, with weights proportional to the posterior probabilities of each model representation. The reported $q$ values are all reasonably small. Table 3 also lists the number of truly different models that were visited.

A third measure of convergence is provided by an estimate of the total posterior model probability covered by the chain following George and McCulloch (1997). This estimate is based on comparing visit frequencies and the aggregate marginal likelihood for a predetermined subset of models. Table 3 presents this estimate for the various species, which is never below $88 \%$ (and typically well above $90 \%$ ).

All diagnostics indicate that convergence is never a problem, which was corroborated by the fact that other independent runs started from randomly chosen models led to virtually identical results.

The chains visit a relatively small number of models: except for cod always less (and usually much less) than one model in every 9600 . Throughout, the acceptance probability of proposals in the MCMC algorithm is in between $6 \%$ and $18 \%$. The best model (the model with the highest posterior mass) contains in between 7 (cod) and 18 (halibut) regressors, and often receives quite a substantial posterior probability, but never so large that model averaging becomes unnecessary. The number of highest probability models that is needed to cover $90 \%$ of the total visited posterior mass (also presented in Table 3) gives a further indication of the substantial spread of the posterior mass in model space.

Marginal posterior inclusion probabilities of the different regressors ( $1-p$ with $p$ obtained from (5.4)), are given in Table 4. Clearly, the visited models for halibut are always full in the variables year and zone (which means that each of the two years has probability $1-(1 / 2)=0.5$ of inclusion and each of the four zones has probability $1-(1 / 4)=0.75$ of inclusion). Note the large differences in the posterior probabilities of inclusion across the various species, which supports our decision to model each species separately.

Convergence of the Gibbs sampler for the probit model was assessed by monitoring the posterior moments of $\gamma$ in different runs of various lengths. Retaining 20,000 draws after a burn-in of 5,000 was found to be more than sufficient.

### 7.2. Posterior results

Here we present some posterior results for the regression coefficients in $\beta$ and $\gamma$, limiting ourselves to some illustrative findings.

We recall that all available regressors in Table 1 are used for the probit model, where we exclude a reference level (arbitrarily chosen as year 1994, otter trawl by pair, zone 30 and December) for each categorical variable. Since the elements in $\gamma$ are not directly interpretable, we present posterior results for transformations with a clear interpretation. For the categorical regressors, we compute the difference in the probability of zero catch between a category and its reference case -e.g., year 1993 versus year 1994-, when all other explanatory variables are evaluated at typical values. Thus, for categorical variables we compute $\left(\bar{x}_{c}^{\prime} \gamma\right)-\left(\bar{x}_{r}^{\prime} \gamma\right)$, where $\bar{x}_{c}$ and $\bar{x}_{r}$ are vectors of "typical" values, identical except for the relevant categorical variable. For these typical values we take the modal level for categorical variables and median values for continuous variables. We shall consider two sets of values throughout: one corresponding to a typical gillnetter (taking modes and medians over the gillnet observations, and taking anchored gillnet as the reference level for fishing technique) and one corresponding to a typical trawler. For the continuous variables, we consider the derivative of the probability of zero catch with respect to the logarithm of the continuous variable. This gives us the (local) effect on the probability of zero catch of a proportionate change in the underlying continuous variable. As with the categorical variables, this effect will be evaluated at typical values for all regressors.

Since all these measures (called "effect" in the sequel) are functions of $\gamma$, we can compute their

Table 4. Marginal Posterior Inclusion Probabilities of Regressors

|  | Species |  |  |  |  |
| :--- | :---: | :---: | :---: | :---: | :---: |
|  | cod | halibut | redfish | grenadier | skate |
| Year 1993 | 0.49 | 0.50 | 0.03 | 0.49 | 0.41 |
| Year 1994 | 0.49 | 0.50 | 0.03 | 0.49 | 0.41 |
| Drift gillnet | 0.12 | 0.01 | 0.15 | 0.96 | 0.06 |
| Anchored gillnet | 0.14 | 0.04 | 0.16 | 0.05 | 0.06 |
| Otter trawl | 0.93 | 0.22 | 0.23 | 0.54 | 0.03 |
| Otter trawl pair | 0.21 | 1.00 | 0.56 | 0.47 | 0.03 |
| Zone 3L | 0.12 | 0.75 | 0.02 | 0.12 | 0.99 |
| Zone 3M | 0.85 | 0.75 | 0.99 | 0.82 | 0.16 |
| Zone 3N | 0.20 | 0.75 | 0.02 | 0.97 | 0.85 |
| Zone 3O | 0.12 | 0.75 | 0.99 | 0.04 | 0.02 |
| January | 0.03 | 0.99 | 0.05 | 0.05 | 0.89 |
| February | 0.04 | 1.00 | 0.87 | 1.00 | 0.23 |
| March | 0.05 | 0.88 | 0.90 | 1.00 | 0.23 |
| April | 0.16 | 0.89 | 0.99 | 1.00 | 0.23 |
| May | 0.75 | 0.88 | 1.00 | 1.00 | 0.49 |
| June | 0.10 | 0.37 | 0.10 | 1.00 | 0.23 |
| July | 0.04 | 0.15 | 0.36 | 0.02 | 0.79 |
| August | 0.03 | 0.16 | 0.30 | 0.02 | 0.80 |
| September | 0.08 | 0.97 | 0.91 | 0.02 | 0.80 |
| October | 0.10 | 0.99 | 0.15 | 0.05 | 0.81 |
| November | - | 0.99 | 0.10 | 0.02 | 0.88 |
| December | - | 0.04 | 0.17 | 0.02 | 0.02 |
| Gillnet $\times \mathrm{f}($ mesh size) | 0.19 | 1.00 | 0.08 | 0.06 | 0.98 |
| Gillnet $\times[f(\text { mesh size })]^{2}$ | 0.13 | 0.94 | 0.10 | 0.09 | - |
| Trawl $\times \mathrm{f}($ mesh size $)$ | 0.58 | 0.03 | 0.97 | 1.00 | 1.00 |
| Trawl $\times \mathrm{f}($ engine kW) | 0.13 | 1.00 | 0.92 | 0.23 | 1.00 |
| log(Length vessel) | 1.00 | 1.00 | 0.11 | 0.02 | 1.00 |
| log(GRT) | 1.00 | 1.00 | 0.15 | 1.00 | 0.60 |
|  | transformation indicated in Table 10 |  |  |  |  |

full posterior distributions. Table 5 presents the posterior mean and standard deviation of the effects of all relevant variables for both typical ships considered. We only present results for halibut and redfish, which are the most important species in terms of live weight caught. In addition, halibut is the species with the lowest proportion of zero catch ( $18.5 \%$ ), while redfish has one of the highest proportions of zeroes ( $85.7 \%$ ).

From Table 5 we see that the regressors can have a large effect on the probability of zero catch, and that the effect is rather specific to the species considered. In view of the decline of the Grand Bank fisheries at the time the data were collected, we could have expected the year to have a large effect. However, only for grenadier (not presented in Table 5) have we found a substantially lower probability of positive catch in 1994. For the other species the difference is small. We now briefly discuss some results for halibut, and merely note that the findings for redfish are often very different, as can be seen directly from Table 5. The probability of catching halibut with a gillnet is higher with a drift gillnet than an anchored one (which serves as the reference case for computing the effects for gillnetters), and a single otter trawl does better than a trawl by pair. As far as location of catch is concerned, the probability of catching halibut is lowest in the reference zone 30, and highest in zone 3L. The time of the year also has a substantial effect: December is the worst month of the year, whereas March and April seem best. Increasing the mesh size of a

Table 5. Posterior Moments of Some Effects in Probit

|  | Species |  |  |  |
| :--- | :---: | :---: | :---: | :---: |
|  | halibut |  | redfish |  |
|  | typical gillnet | typical trawl | typical gillnet | typical trawl |
| Year 1993 | $-0.00(0.04)$ | $-0.00(0.00)$ | $0.02(0.01)$ | $0.01(0.01)$ |
| Drift gillnet | $-0.16(0.09)$ | - | $-0.58(0.08)$ | - |
| Otter trawl | - | $-0.39(0.03)$ | - | $-0.01(0.01)$ |
| zone 3L | $-0.61(0.05)$ | $-0.49(0.06)$ | $0.47(0.06)$ | $0.28(0.05)$ |
| zone 3M | $-0.22(0.04)$ | $-0.21(0.05)$ | $0.03(0.02)$ | $0.07(0.04)$ |
| zone 3N | $-0.48(0.04)$ | $-0.41(0.05)$ | $0.52(0.06)$ | $0.29(0.05)$ |
| January | $-0.31(0.13)$ | $-0.08(0.05)$ | $0.25(0.10)$ | $0.10(0.05)$ |
| February | $-0.40(0.12)$ | $-0.09(0.05)$ | $0.04(0.02)$ | $0.08(0.05)$ |
| March | $-0.52(0.12)$ | $-0.10(0.05)$ | $0.03(0.02)$ | $0.08(0.05)$ |
| April | $-0.55(0.12)$ | $-0.10(0.05)$ | $0.03(0.01)$ | $0.08(0.05)$ |
| May | $-0.33(0.12)$ | $-0.08(0.05)$ | $0.03(0.02)$ | $0.08(0.05)$ |
| June | $-0.21(0.12)$ | $-0.07(0.05)$ | $0.09(0.03)$ | $0.09(0.05)$ |
| July | $-0.25(0.12)$ | $-0.08(0.05)$ | $0.01(0.01)$ | $0.05(0.06)$ |
| August | $-0.31(0.12)$ | $-0.08(0.05)$ | $0.00(0.01)$ | $0.02(0.06)$ |
| September | $-0.35(0.12)$ | $-0.09(0.05)$ | $0.00(0.01)$ | $0.03(0.06)$ |
| October | $-0.34(0.12)$ | $-0.09(0.05)$ | $0.00(0.01)$ | $0.04(0.05)$ |
| November | $-0.38(0.12)$ | $-0.09(0.05)$ | $0.01(0.01)$ | $0.06(0.06)$ |
| mesh size | $-1.49(0.67)$ | $-0.05(0.04)$ | $-0.88(0.28)$ | $-0.08(0.04)$ |
| engine kW | - | $-0.03(0.01)$ | - | $0.01(0.00)$ |
| length vessel | $-1.19(0.15)$ | $-0.10(0.03)$ | $0.15(0.08)$ | $0.04(0.02)$ |
| GRT | $0.81(0.08)$ | $0.07(0.02)$ | $-0.19(0.06)$ | $-0.05(0.01)$ |

(Entries are posterior means with standard deviations in parentheses.)
gillnet in a neighbourhood of the median value ( 140 mm ) has a positive effect on the probability of catch: locally increasing mesh size by $1 \%$ increases the probability of catching halibut by $0.002-$ 0.028 . The local effect of changes to mesh size for a typical trawler, however, is much smaller. This illustrates the importance of treating gillnets and otter trawls separately. The engine power of ships with trawl gear does not seem to play a substantial role either, although more power is consistently associated with higher probability of catch. Finally, longer vessels tend to have a lower probability of zero catch, but the latter is partly offset by the opposite effect of GRT.

Let us now focus on results for the continuous part, modelled as in Section 5. The coefficient $\beta_{l}$ corresponding to a categorical variable has the following interpretation: $\exp \left(\beta_{l}\right)$ is the ratio between the median catch with the corresponding dummy equal to one and the median catch in case this dummy is zero. If a continuous regressor is the logarithm of a variable (length and GRT), then the corresponding regression coefficient $\beta_{l}$ is unequivocally interpreted as an elasticity (i.e., it approximately reflects the relative percentage change in median catch as a consequence of a $1 \%$ relative change in the original untransformed continuous regressor). For the interactions with trawls, to which the more complicated transformation indicated in Table 1 was applied, the elasticity of median catch with respect to that regressor is given by $\beta_{l}$ times a positive factor (which depends on where we evaluate the elasticity). For the gillnet mesh interaction, the elasticity is a linear combination of both the intervening components of $\beta$.

The $\tilde{k}$-dimensional ( $\tilde{k}=28$ for most species) regression vector $\beta$ has a rather complicated posterior distribution, which is a mixture of point masses at zero and continuous parts. It is therefore quite challenging to present this distribution in an easily interpretable format. In the sequel, we shall illustrate some aspects of this posterior distribution for halibut. Again, results vary considerably across species.

Figures 2-4 present, for a number of selected linear combinations of the components of $\beta, b^{\prime} \beta$, the posterior p.d.f. in (5.5) for halibut. In addition, the gauge on top (black shading) indicates the posterior probability that $b^{\prime} \beta \neq 0$. The vertical lines presented in some of these graphs relate to the classical estimate and $90 \%$ confidence interval obtained from a stepwise regression technique, as explained and discussed later in Subsection 7.4.


Fig. 2. Halibut: Year and zone.
Figure 2 focuses on the elements of $\beta$ corresponding to year and zone. From Table 4 we note that all visited models are "full" in these two categorical variables (i.e. $L_{r}-1$ out of the $L_{r}$ possible levels are always included in the model). This induces $L_{r}-1$ modes in the marginal posterior p.d.f. for the regression coefficients, where every mode corresponds to a different level being excluded (and, thus, treated as the reference level). For year we have $L_{r}=2$ possible levels, leading to unimodal distributions which indicate that 1993 is clearly a better year than 1994. For zone we have $L_{r}=4$ levels and we observe the expected $L_{r}-1=3$ modes. From the relative locations of the modes, it is easy to derive that, for example, the three modes for zone 3 L correspond to taking zone $3 \mathrm{~N}, 3 \mathrm{M}$ and 3 O (from left to right) as reference levels. There is a clear ranking in that zone 3 O is the worst, followed by $3 \mathrm{M}, 3 \mathrm{~L}$ and 3 N , in that order. The difference between zones 3 L and 3 N is not very large (about 0.15 between the modes, or a factor of 1.16 between median catch), which accounts for the apparent bimodality of the p.d.f. corresponding to zone 3O. The latter zone is the zone with by far the least observations, leading to Student- $t$ distributions with large spread in (5.5), which means the modes corresponding to reference cases 3 N and 3L can no longer be separately identified in the figure. In a case such as this, where models are full in a categorical variable, it does not matter which level is taken as a reference level (since all levels are always identified as being different), and we could equivalently fix the reference level and present conditional results instead of the marginal ones given here. For example, if we give results for zone conditioned on the reference level zone 30 , only the extreme right modes appear for the other zones. However, when more than one level at a time is excluded from visited models (as is usually the case), we need the extra flexibility provided by our framework where reference levels are not fixed in advance.

In general, one should aim to present results for quantities that have the same meaning regardless of the choice of reference levels. An interesting way to present regression coefficients of categorical variables is in the form of centred coefficients, that is, $\delta_{l} \equiv \beta_{l}-\left(\sum_{i=1}^{L_{r}} \beta_{i}\right) / L_{r}$ $\left(l=1, \ldots, L_{r}\right)$, for a categorical variable with $L_{r}$ levels and original coefficients $\left(\beta_{1}, \ldots, \beta_{L_{r}}\right)$. Clearly, $\sum_{l=1}^{L_{r}} \delta_{l}=0$ and $\delta_{l}$ indicates the difference between level $l$ and the average, so its meaning
is not dependent on any particular choice of reference level. Figure 3 presents the marginal posterior distributions of the centred coefficients associated with each zone. The ranking of zones mentioned above is now immediately obvious from Figure 3.


Fig. 3. Halibut: Zone with centering.
The effects of the fishing techniques and their interactions with mesh size and engine kW are examined in Figure 4. From Table 4, we see that the categorical variable corresponding to fishing technique (with $L_{r}=4$ levels) is not fully represented in every model. Some levels (the gillnet techniques) are almost never included and otter trawl by pair is always included. Thus, otter trawl by pair is never treated as a reference level (indicating it is quite different from the other levels) whereas often more than one of the other levels are excluded (and thus treated as equal). The fact that models now exclude either one, two or three of these levels at the same time creates more possibilities for modes in the marginal distributions of the associated regression coefficients, and interpretation becomes much harder. Note, however, that now we would lose flexibility if we fixed a reference level (e.g., if we had chosen otter trawl by pair as the reference level, we could not have accommodated the situation described above, where trawl by pair is different from all the others and some of the other levels are equal).

Evaluating the relative merits of the fishing techniques is complicated by the presence of interactions with mesh size and kW . Therefore, Figure 4 presents the posterior distribution of the differences between the regression coefficients associated with drift and anchored gillnet (which are equally affected by the interactions) and also between those for otter trawl and otter trawl by pair. These are interpretable quantities (logs of median catch ratios), and reveal little difference between both gillnets, whereas single trawls tend to do better than trawls by pair. To get a rough idea of the overall effects of the different fishing methods, we can consider the configuration of the best model (the model with highest posterior probability), which includes trawl by pair as the only technique and all interactions except for trawl with mesh size. On the basis of the posterior mode of the included regression coefficients for this model, and evaluating the effect at median values for the continuous regressors, we obtain the following ranking from better to worse: otter trawls, trawls by pair (median catch about $58 \%$ of otter trawls) and both gillnets (median catch about $10 \%$ of otter trawls). These numbers are roughly consistent with the observed values (which are, of course, affected by other factors as well). Figure 4 also graphs the difference in log median catch for two gillnet mesh sizes, suggesting higher median catch for 140 mm mesh (median and $3^{\text {rd }}$ quartile from Table 1) than for 130 mm ( $1^{\text {st }}$ quartile). Finally, for trawls, mesh size is almost never included in the model, whereas engine kW has a positive effect on median positive catch of halibut.

To economize on space, we have not shown the posterior density functions of the regression coefficients of the months or the size variables. The main messages here are that the months January until May have a positive effect, whereas July until November lead to lower median catch of halibut. Finally, length has a positive effect and GRT a negative effect. From Figure 1, we know that both variables are strongly positively correlated and, on balance, the effect of size on the median catch of halibut will be quite small.


Fig. 4. Halibut: Fishing techniques and their interactions.

### 7.3. Predictive results

On the basis of the posterior results partially described above, we shall now predict observations in the subsample that was not used for posterior inference.

First of all, let us examine how well we predict the probability of zero catch. For every observation in the prediction subsample, we compute $\omega_{f}=P\left(s_{f}=0 \mid s\right)$ as in (6.2). An interesting check on the adequacy of our probit model is then to compare these predictive probabilities with the actual occurrences of zero or positive catch. Table 6 presents the means and standard deviations of $\omega_{f}$ computed over the zero and the positive observations in the prediction sample. Clearly, $\omega_{f}$ tends to take much higher values for those observations that turn out to be zero, indicating that the probit model does far better than simply assuming that the probability of zero catch is constant across observations.

Table 6. Predictive Zero Catch Probability

|  | Species |  |  |  |  |
| :--- | :---: | :---: | :---: | :---: | :---: |
|  | cod | halibut | redfish | grenadier | skate |
| Zero observations | $0.94(0.14)$ | $0.59(0.30)$ | $0.92(0.13)$ | $0.63(0.30)$ | $0.66(0.20)$ |
| Positive observations | $0.45(0.21)$ | $0.09(0.14)$ | $0.45(0.28)$ | $0.29(0.16)$ | $0.45(0.19)$ |

( Entries are means with standard deviations in parentheses.)
Let us now use the predictive results for the continuous part in (6.4) to assess the predictive adequacy of the modelling of positive observations. For all the positive observations in the prediction subsample we record in which percentile of the continuous part of the predictive distributions (using the corresponding values of the regressors) the actual observations fall. Contrasting predictive quantiles with empirical ones leads to a $\mathrm{Q}-\mathrm{Q}$ plot that indicates how well the model (estimated on the basis of the estimation subsample) fits the data in the prediction subsample. As the assignment of observations to either subsample is random, we would expect such plots to be a good measure of model accuracy. Figure 5 presents these $\mathrm{Q}-\mathrm{Q}$ plots for all five species, indicating that model fit is always quite good.

For illustration, we now show some predictive distributions for particular observations in the prediction subsample. Figure 6 graphs the predictive p.d.f.'s of the nonzero catch of halibut for observations $i=196(270 \mathrm{~kg})$ and $i=1189(3600 \mathrm{~kg})$-as in (6.4). From the probit analysis, the probability of zero catch for observation 196 is 0.43 , while observation 1189 has only a 0.03 probability of being equal to zero. The rather different predictive distributions illustrate the importance of the fishing gear. The main difference between the observations is that 196 corresponds to a vessel using drift gillnets whereas 1189 is with an otter trawl. In both cases, the actual catch (indicated by a dashed vertical line in Figure 6) is quite compatible with the predictive distributions.

For policy purposes, it might be interesting to predict not the catch of one single ship, but the


Fig. 5. Q-Q Plots for Predictions of Individual Observations.


Fig. 6. Halibut: Predictive Densities and Actual Observed Values.
aggregate catch of a number of ships, that are known to be in a certain area of the Grand Bank at a certain time of the year. If we group the data into clusters of 5 ship-days, we can analyze how such predictions, based on the estimation subsample, compare with the actual retained clusters. Clusters of ships that are in the same zone on the same day are likely to be of most interest for practically relevant predictions. In order to mimic such clusters, we have sorted the prediction subsample by year, day and zone (in that order) and selected clusters of 5 consecutive observations from that ordering. The predictive distribution, computed as described at the end of Section 6, leads to $\mathrm{Q}-\mathrm{Q}$ plots (not shown) that indicate adequate predictions for clusters. Figure 7 presents some individual cluster predictives for the nonzero catch of halibut, redfish and grenadier. The probability of zero aggregate catch varies dramatically across these clusters: from less than $10^{-6} \%$ for halibut to $0.91 \%$ for grenadier and $58.8 \%$ for redfish. Again, the quantities caught are well matched by the corresponding predictive distributions. These predictive densities immediately lead to probability statements, e.g., about a fleet of certain characteristics exceeding a certain catch, which could straightforwardly be used in a decision theory context.


Fig. 7. Predictive Densities and Actual Observed Values for Clusters.

### 7.4. Classical methods

In a classical statistical framework, posterior model probabilities are not readily available and, usually, a particular model is selected instead of averaging over models. Given the substantial spread of the posterior mass over the models in $\mathcal{M}$ (see Table 3), that does not seem an adequate strategy for the analysis of these data.

Nevertheless, if we wish to use classical methods for variable selection, a popular technique is stepwise regression. Table 3 records the posterior probabilities of the models selected using forward selection and backward elimination as in Lo et al. (1992). Here we base the choice of reference levels for the categorical variables on the best model: we can choose any reference level for those categorical variables in which the best model is full and for the other categorical variables we choose from the levels that are excluded in the best model the one with lowest posterior inclusion probability (see Table 4).

The models chosen by this stepwise regression technique have in between 6 (cod) and 21 (halibut) variables. For grenadier and skate, this method identifies the important variables reasonably well: no variables with posterior probabilities over 0.8 are left out and only one regressor with posterior inclusion probability under 0.2 is selected (for skate). Accordingly, the posterior probability of the stepwise model is relatively high for these species (see Table 3). For halibut and redfish the performance of stepwise regression is much less in line with the posterior inclusion probabilities. For cod there is an even larger conflict between the stepwise model and the posterior inclusion probabilities, and, as a consequence, the stepwise model picks up virtually no posterior mass.

The classical $90 \%$ confidence intervals corresponding to the models selected by stepwise regression are indicated in Figure 2 and the last two plots of Figure 4 by dotted vertical lines. The estimated value is indicated by a dashed line and a single dash-dot line at zero represents exclusion of the corresponding regressor. Even though some confidence intervals roughly contain $90 \%$ of the posterior mass, they can be quite different from the corresponding Bayesian credible intervals.

## 8. Concluding remarks

In this paper we have outlined the modelling of daily live weight catch of different species of fish in the Grand Bank fishery. An important feature of the data is the fact that on most days not all species are caught by a certain ship. Thus, modelling of these implicit zero observations is crucial. The latter was done through a probit model. For the positive observations, we have used a Lognormal regression model, where we allow for any combination of regressors from a set of different explanatory variables. We deal with model uncertainty through Bayesian model averaging. Many of the regressors are categorical variables, and we pay particular attention to the treatment of categorical variables in a model uncertainty context. In particular, we allow for any combination of levels to be included in the models, as long as no categorical variable appears with all possible levels (thus, no reference level is fixed in advance). In order to deal with the resulting 177 million possible models, we apply an MCMC algorithm, based on the MetropolisHastings sampler to generate a Markov chain of drawings in this large model space. Throughout, we use a carefully chosen prior distribution which also takes into account that models that are full in categorical variables have equivalent counterparts (corresponding to different choices for the excluded level of these categorical variables), and we examine posterior and predictive inference. The former can be instrumental in policy decisions regarding the effect of certain ship characteristics or regulations concerning, e.g., mesh size or fishing techniques. The latter is required if we wish to predict catch per species from easily obtained information regarding the presence of vessels with known characteristics in a certain area at a certain time, rather than having to board these vessels (which is much more costly and altogether impossible for ships from countries outside NAFO). The methods would also be useful for estimating total catch by area when misreporting and black landings are common. Bayesian model averaging naturally takes into account all uncertainty concerning parameter values as well as model uncertainty. Thus, realistic predictions can be made for one or more ship-days, duly taking into account the ships' characteristics, location, month as well as parameter and model uncertainty. Using the efficient code, new data can easily be processed and posterior and predictive inference can be conducted without excessive computational requirements. We find that the proposed model fits our data relatively well, and that results differ crucially between species.

There are a number of ways in which the model used here could be extended. A possible elaboration would be to include random ship effects -i.e., ship-specific intercepts- in either the discrete or the continuous part of the model. That could pick up certain quality aspects of the vessels, not captured in the regressors. A potential interpretation of such individual effects would be as skill of the captain of the vessel, which was equated with technical efficiency in a stochastic frontier model by Kirkley et al. (1998). Barring rather restrictive forms for the distribution of the random effects, this would result in substantial complications: for example, our computations for the continuous part rely on the fact that the marginal likelihood for each model can be computed analytically. We have also avoided including dynamic effects into the model; such effects might provide a "closer fit", but are not in line with the aim of providing easily computed operational predictions on the basis of available information (which typically does not include a recent history of quantities caught by a cluster of ships considered). In addition, their inclusion would be at the cost of adding to the theoretical and computational complexity of the model. Also, it might be a useful exercise to examine the effects of allowing for heteroskedasticity in the error term of (3.2) by making $\sigma$ depend on, e.g., the size of the ship. Of course, both theory and practical implementation would become more cumbersome as a consequence (unless such dependence would be fixed, rather than estimated from the data). Finally, again at a considerable cost in terms of added complexity, one might propose a multivariate model for all species with correlated error terms.

## Appendix A: Samplers

## A.1. Gibbs sampler for probit model

We introduce independent latent variables $m_{i}(i=1, \ldots, n)$, with $m_{i}$ distributed as $\operatorname{Normal}\left(x_{i}^{\prime} \gamma, 1\right)$. From (3.1), it is immediate that $s_{i}=0$ is equivalent to $m_{i}>0$, whereas $s_{i}>0$ is equivalent to $m_{i}<0$. The posterior distribution is, therefore,

$$
p(\gamma \mid s)=p\left(\gamma \mid m_{i}<0 \text { for } i=1, \ldots, Q ; m_{i}>0 \text { for } i=Q+1, \ldots, n\right) .
$$

A Gibbs sampler, augmenting with $m \equiv\left(m_{1}, \ldots, m_{n}\right)^{\prime}$, consists of drawing from:

$$
\begin{aligned}
& p(\gamma \mid m, s)=p(\gamma \mid m)=f_{N}^{1+k}\left(\gamma \mid\left\{\left(1+h_{0}\right) X^{\prime} X\right\}^{-1} X^{\prime} m,\left\{\left(1+h_{0}\right) X^{\prime} X\right\}^{-1}\right), \text { and } \\
& p(m \mid \gamma, s) \propto\left\{\prod_{i=1}^{Q} f_{N}^{1}\left(m_{i} \mid x_{i}^{\prime} \gamma, 1\right) I_{\left(m_{i}<0\right)}\right\}\left\{\prod_{i=Q+1}^{n} f_{N}^{1}\left(m_{i} \mid x_{i}^{\prime} \gamma, 1\right) I_{\left(m_{i}>0\right)}\right\} .
\end{aligned}
$$

## A.2. MCMC sampler on model space

Suppose the chain is currently at $M_{s}$, which has $k_{s}$ continuous regressors and $n_{r}$ levels for categorical variable $r$ (where $0 \leq k_{s} \leq K, r=1, \ldots, R$ and $n_{r} \in\left\{0,1, \ldots, L_{r}-1\right\}$ ). Suppose that there are $f_{s}$ full categorical variables, $c_{1}, \ldots, c_{f_{s}}$, -i.e., $n_{c_{1}}=L_{c_{1}}-1, \ldots, n_{c_{f_{s}}}=L_{c_{s}}-1$. The number of regressors in $M_{s}$ is then $N_{s}=k_{s}+n_{1}+\ldots+n_{R}$, whereas the maximum amount of regressors in any model is $N_{t o t}=K+L_{1}+\ldots+L_{R}-R$. The Metropolis-Hastings algorithm proceeds along the following steps:
[S1] Propose a new model $M_{c a n}$ in several stages. First propose $N_{c a n}$ :

$$
N_{\text {can }}=\left\{\begin{array}{lll}
N_{s}+1 & \text { with probability } & \left(N_{\text {tot }}-N_{s}\right) / N_{\text {tot }} \\
N_{s}-1 & \text { with probability } & N_{s} / N_{\text {tot }}
\end{array}\right.
$$

Now propose $M_{c a n}$ conditionally on the drawn value of $N_{c a n}$ :
(a) If $N_{c a n}=N_{s}+1$ : sample $M_{c a n}$ by uniformly adding one regressor to $M_{s}$, excluding levels of categorical variables in which $M_{s}$ is already full. We can choose from $N_{t o t}-N_{s}+R-f_{s}$ variables, so the probability of adding each is $1 /\left(N_{t o t}-N_{s}+R-f_{s}\right)$. Define $T_{\text {can }, s}=$ $\left(N_{\text {tot }}-N_{s}+R-f_{s}\right) /\left(N_{t o t}-N_{s}\right)$. Proceed to [S2].
(b) If $N_{\text {can }}=N_{s}-1$ : uniformly drop one regressor from $M_{s}$ to form $M_{\text {can }}$; each choice has probability $1 / N_{s}$. Define $T_{\text {can }, s}=\left(N_{t o t}-N_{c a n}\right) /\left(N_{t o t}-N_{c a n}+R-f_{c a n}\right)$. Proceed to [S2].
[S2] Compute:

- $B_{c a n, s}=l_{y}\left(M_{c a n}\right) / l_{y}\left(M_{s}\right)$, where

$$
\begin{equation*}
l_{y}\left(M_{j}\right) \propto\left(\frac{g_{0}}{g_{0}+1}\right)^{k_{j} / 2} G_{j}^{-(Q-1) / 2}, \tag{A.1}
\end{equation*}
$$

with

$$
\begin{equation*}
G_{j}=\frac{1}{g_{0}+1} y^{\prime} M_{W_{j}} y+\frac{g_{0}}{g_{0}+1}\left(y-\bar{y} \iota_{Q}\right)^{\prime}\left(y-\bar{y} \iota_{Q}\right), \tag{A.2}
\end{equation*}
$$

where $\iota_{Q}$ is the $Q$-dimensional vector of ones, $\bar{y}=\iota_{Q}^{\prime} y / Q, W_{j}=\left(\iota_{Q}: Z_{j}\right)$ and $M_{W_{j}}=I_{Q}-$ $W_{j}\left(W_{j}^{\prime} W_{j}\right)^{-1} W_{j}^{\prime}$.

- $L_{c a n, s}=\left(\prod_{i=1}^{f_{s}} L_{c_{i}}\right) /\left(\prod_{i=1}^{f_{c a n}} L_{c_{i}}\right)$, with $f_{\text {can }}$ denoting the number of full categorical variables in $M_{c a n}$.
[S3] With probability $q=\min \left(1, B_{c a n, s} L_{c a n, s} T_{c a n, s}\right)$ the chain moves to $M_{c a n}$, whereas with probability $1-q$ it stays at $M_{s}$.
[S4] Record the new state of the chain (be it $M_{c a n}$ or $M_{s}$ ) after uniformly redrawing the reference level for each of the full categorical variables.

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