

Direct evidence of the impact of longline fishery on mortality in the Black-footed Albatross *Phoebastria nigripes*

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Summary

The incidental bycatch of seabirds in longline fisheries is one of the most striking examples of diffuse and incidental impact of human activities on vertebrate species. While there are various types of evidence of a strong impact of longline fisheries on seabirds, in particular albatrosses, the incidental bycatch mortality has never been directly linked to estimates of bycatch derived from on-board surveys. We develop a capture-recapture analysis which relates the annual probability of survival in the Black-Footed Albatross to the estimated absolute bycatch. By converting the absolute bycatch into an estimate relative to population size and using the theory of exploited populations, we show that survival probability decreases linearly with bycatch, and that the bycatch is underestimated by at least 50%, confirming suspicions based on reviews of the bycatch survey procedures, and raising further concerns about the impact of longline fisheries on seabirds.

Introduction

While the ecological impact of diffuse anthropogenic factors has been long recognised in ecosystem ecology (e.g. Crowe *et al.* 2000), it is only comparatively recently that population conservation issues were explicitly looked at from this point of view. A number of human activities impact on animal populations at rates that are so low or in ways that are so indirect that they have for a long time implicitly been considered as innocuous or sustainable, often even going unnoticed. In fact, the impact is often translated as a rate relative to the human activity concerned, while the correct reference is the target size, i.e. the animal population.

The incidental bycatch of seabirds by fishing devices is one of the most striking examples and is seen as a major threat to some species, especially albatrosses (Croxall *et al.* 2012) although information about the population level effect is lacking. For longline fishing, the rate of bird capture per hook is quite low, often as low as one bird caught per 50,000 to 200,000 hooks set (e.g. Gilman *et al.* 2003, Lewison and Crowder 2003) although values up to 0.5 to 1 bird per 1,000 hooks set have been recorded (Ryan *et al.* 2002, Lewison and Crowder 2003). Such low rates give at first sight a feeling of innocuousness. However, the development of longline fisheries has led in some oceans to several million hooks being set every day (Lewison and Crowder 2003). The absolute numbers caught resulting from the combination of a low rate per hook set and a huge number of hooks set are far from negligible, with estimates reaching 25,000 birds per year in the Laysan Albatross *Phoebastria immutabilis* in the late 1980s (Naughton *et al.* 2007), and probably more than 300,000 seabirds per year at a global scale (Anderson *et al.* 2011). Soon after the first concerns emerged, various approaches provided evidence of a strong impact on some populations, in particular of albatrosses (e.g. Brothers 1991, Weimerskirch *et al.* 1997).

Deciphering the effects of longline fishery bycatch on seabirds has been the subject of intense activity over the last 20 years, from analysing bird behaviour to assessing the consequences on populations (see e.g. Lewison *et al.* 2004, Arnold *et al.* 2006), from estimating of the numbers caught by various types of fishing activities (synthesised by Anderson *et al.* 2011) to developing and assessing the effectiveness of various mitigation measures (Brothers *et al.* 1999, Gilman *et al.* 2003).

After early studies that discussed the estimated bycatch relative to the estimated population size, usually expressed as a number of breeding pairs (e.g. Ryan *et al.* 2002), investigations of population level consequences became rapidly more and more sophisticated. The degree of sustainability of the estimated bycatch levels was approached with the help of population projection models (Weimerskirch *et al.* 1997, Lewison and Crowder 2003, Arnold *et al.* 2006.). Capture-mark-recapture (CMR) analyses went one step further by providing evidence for a drop in survival in correlation with bycatch (Weimerskirch *et al.* 1997).

In fact, even at a low rate, albatrosses are literally fished, and the population problem is thus an exploited population problem, even if exploitation is incidental. Considering the problem of seabird bycatch, as well as other conservation issues (Reynolds *et al.* 2001), as an exploited population problem brings decisive advantages (Lebreton 2005). First, the theory of exploited populations, e.g. with measures of Catch per Unit of Effort (CPUE) (Haddon 2001) relates the effect of exploitation to its intensity. Véran *et al.* (2007) used such an approach to show that survival in the Black-footed Albatross *Phoebastria nigripes* (hereafter BFAL) was directly related to covariates reflecting the intensity of fishing activity. Second, the theory of exploited populations (TEP) also predicts that survival should decrease in an approximately linear fashion with the proportion exploited (for a review see Lebreton 2005). This prediction has been used in the context of conservation and management of animal species for quarry species such as ducks (Anderson and Burnham 1976, Nichols 1991) and geese (Gauthier *et al.* 2001). The various estimates of bycatch available for seabirds have never been used to directly model and predict the change induced on survival, although doing so would bring direct evidence for the effect of bycatch, help assess its biological significance, and thus improve our ability to decipher the complexities of this low rate exploitation.

The purpose of this paper is to relate survival in the BFAL with annual estimates of total bycatch made available by Naughton *et al.* (2007). We use modern capture-recapture methodology, taking advantage of the detailed investigation by Véran *et al.* (2007), simply using a CMR model with a built-in linear regression (Lebreton *et al.* 1992) constraining the survival probability to be a linear function of the estimated bycatch.

We take into account and discuss the unavoidable uncertainties and biases in bycatch estimates. On the one hand, the uncertainties in the covariate tend, as in all regression models, to bias the slope toward 0 (Barker *et al.* 2002), in an “error-in variable” problem. On the other hand, bycatch estimates tend without any doubt to be underestimated (Brothers *et al.* 1999, Brothers 2008, Anderson *et al.* 2011). First, various direct biases in the survey of birds hauled on board exist (Brothers 2008); second, the number really caught should include birds killed by the line but not hauled on board, named for short “drop-off”, that may reach 50% of the total kill (Brothers 2008, Brothers *et al.* 2011). The latter part is the exact equivalent of the crippling loss of quarry species, consisting of animals killed by hunting activity but not retrieved by hunters, and thus not included in hunting bag surveys (e.g. Van Dyke 1981). Finally, the extrapolation to obtain overall figures is inherently fragile, as this extrapolation inevitably encompasses fleets submitted to looser regulations than those having accepted observers on board (Anderson *et al.* 2011). We develop predictions on the effect of these various uncertainties and biases after a description of the methods and data. We expect, by using a more directly functional covariate and a wider time window than Véran *et al.* (2007), to obtain stronger evidence for a bycatch-survival relationship in the BFAL, and more broadly, to contribute to the toolbox for understanding the effects of diffuse man-induced mortality in animal populations.

Material and methods

Data

The Black-Footed Albatross is a near-endemic of the Hawaiian archipelago. Concerns for this rare species (around 60,000 breeding pairs) have been reviewed by Cousins and Cooper (2000) and Naughton *et al.* (2007). The species is classified as “Endangered” by IUCN. We use CMR data already used by Véran *et al.* (2007). The data consist of individual histories of 13,854 BFAL banded as chicks on Tern island from 1980 to 2004, and recaptured as breeders from 1997 to 2004 (see Véran *et al.* 2007 for further details).

Capture-recapture analysis

The data were analysed using multistate capture-recapture recruitment models (see e.g. Lebreton *et al.* 2003), with two states, ‘Breeder’ and ‘Non-Breeder’. Individuals were marked as chicks in the ‘Non-Breeder’ state. The transition from ‘Non Breeder’ to ‘Breeder’ started at age 5. Individuals were never re-observed as Non-Breeders, i.e. the state Non-Breeder was not observable. Various constraints in relation to goodness-of-fit, parameter identifiability, and biologically relevant questions were considered, leading to several different models that were all fitted using E-Surge (Choquet *et al.* 2009a).

The goodness-of-fit of an initial model was assessed based on the adult part of the CMR histories (i.e. reduced to breeders, following Crespin *et al.* 2006, Lebreton *et al.* 2009), using program U-CARE (Choquet *et al.* 2009b). Following Crespin *et al.* (2006), the initial model considered time and age dependence in recruitment (with stabilisation at age 10, i.e. nearly all the information up to the first recapture as a breeder was used in this model to estimate recruitment), time dependence and two age classes for survival.

An estimate of an overdispersion coefficient reflecting heterogeneity in the data was derived from the goodness-of-fit statistics.

The basic parameters in our models were:

- First year survival probability
- After-first year survival probability
- Age-specific probabilities of transition from Non-Breeder to Breeder (at ages 5, 6, and 7+); the constancy of the recruitment probability from age 7 onwards together with the assumption of the same survival for breeders and non breeders of the same age are sufficient to ensure the statistical identifiability of all parameters (Gimenez *et al.* 2003, Lebreton *et al.* 2003).
- Probabilities of recapture of breeders, that were non null only for occasions 17 to 25, i.e. years 1997 to 2004 (as all recaptures took place from November to January, these calendar years are those of the corresponding November and December months).

As there were no recaptures of marked individuals between the chick stage and age 5, only a compound immature survival probability from the chick stage to age 5 could indeed be estimated. As a first approximation, we assumed that survival probability was independent of age from age 1 onwards, whether the bird was a breeder or not. As this after-first year survival probability could be estimated from recaptures of breeders, one could in turn derive from the compound immature survival probability an estimate of the probability of survival from the chick stage to age 1. As this first year survival probability was estimated in an indirect fashion and concerned birds that were at sea for less than a full year, we expected it to be less sensitive to bycatch than the after-first year survival probability.

Bycatch as a covariate

According to TEP, when a proportion h of a population or a population subset such as a particular age-class with baseline survival probability S_0 is harvested, the probability of survival should

approximately vary as $S = S_o(1 - h)$ unless there is some form of compensation, that may lead to $S = S_o(1 - bh)$ with $b < 1$ (see review in Lebreton 2005).

Véran *et al.* (2007) started from this basic relationship and investigated the relationship between survival probabilities and covariates reflecting fishing effort F , to which h is presumably proportional, as $h = gF$, leading thus to $S = S_o(1 - gF)$. To protect the regression against the use of too many variables that would have biased statistical tests, they first reduced eight fishing effort covariates to three principal components, and then showed that the survival probability of adult albatross was significantly related to the second principal component (PC₂), with a test level corrected for multiple testing.

We here take advantage of the recent availability of annual estimates of the total BFAL bycatch (Figure 4 in Naughton *et al.* 2007). We sampled their figure to obtain the values in Table 1 and we used these values as a covariate to model survival probabilities.

Our analysis differs from that of Véran *et al.* (2007) in the following respects:

- 1) Thanks to the availability of the bycatch estimate we were able to use more data over more years. We used a recruitment model with 13,854 individuals from ringing as a chick to recaptures as breeders over 25 years (1980–2004) while Véran *et al.* (2007) used the first capture as breeder as the initial marking and used 2,046 capture histories of breeders over the 12 years (1992–2003) for which measures of fishing effort were available.
- 2) As a side-effect, we were unable to use the sophisticated treatments of heterogeneity used by Véran *et al.* (2007) and we expect a higher overdispersion coefficient. The comparison between three types of models by Véran *et al.* (2007) shows that the estimates are not very sensitive to the treatment of heterogeneity. A higher overdispersion coefficient will induce a decrease in power, i.e. will be conservative relative to bycatch assessment.
- 3) Based on TEP, we will be able to discuss the slope value, as in the absence of bias and compensation it is expected to be equal to 1.

Predictions on the effect of bycatch

The absolute bycatch H can be converted into a relative estimate h by dividing H by an estimate of total population size N , as $h = H/N$. The ratio Population size / Number of breeding pairs in the BFAL is, as in all albatross species, close to 5 before the annual birth pulse (Cousins and Cooper 2000), a value derived from the stable population structure in a matrix model. With an average population size near 60,000 pairs, we used $N = 300,000$. The effect of potential bias and uncertainties at this stage is discussed below.

As mentioned earlier, one may ideally expect a slope b equal to 1 in $S = S_o(1 - bh)$. It could be modified by compensation, a phenomenon of direct biological interest, and by various artefacts. While compensation is not expected to be strong in such a long-lived species (Gauthier *et al.* 2001, Lebreton 2005), two main artefacts exist:

Table 1. Black-footed Albatross absolute bycatch estimates (denoted as H') obtained by sampling Figure 4 of Naughton *et al.* (2007).

Year	1981	1982	1983	1984	1985	1986	1987	1988
Absolute bycatch estimate H'	11328	11653	12900	13333	14146	15447	15230	16152
Year	1989	1990	1991	1992	1993	1994	1995	1996
Absolute bycatch estimate H'	13713	10407	12304	10190	8726	7913	6938	6396
Year	1997	1998	1999	2000	2001	2002	2003	2004
Absolute bycatch estimate H'	5420	5474	5962	6016	4607	5203	5366	5420

- Bias in N and H , the latter estimated as H' , inducing a bias in h . With a multiplicative expression of the bias, $h' = dh$ in which h is the true relative bycatch, the relationship $S = S_o(1 - bh) = S_o(1 - b'h') = S_o(1 - b'dh)$ shows that the slope expected when using $h' = dh$ as a covariate, is $b' = b/d$. If the bycatch proportion is underestimated, i.e. $d < 1$, then $b' > b$. The potential bias in h results first from underestimation of H by H' , a well known problem whose components are: drop-off, extrapolation to periods without observers on decks and to fleets without observers. Most experts (e.g. Brothers *et al.* 2011, Moore and Zydalis 2008) seem convinced each of these two main types of bias may be in the order of several times 10%. Hence, a value of d as low as 0.5 seems quite likely. The BFAL population at sea includes immature birds and is thus larger than that deduced from the number of breeding pairs, which estimates population size just before the annual birth pulse, i.e. after some mortality of these immature birds. Similarly the factor 5 retained for deriving an estimate of total population size N from the number of breeding pairs tends to underestimate total population size, compared with the value 6 proposed by Dillingham and Fletcher (2011). Our choices tend thus to increase the estimated relative bycatch h' and are thus conservative relative to the sources of underestimation.
- Uncertainty in h' , inducing an "error-in-variable" problem resulting in an "attenuation" of the estimated slope towards 0. The estimated values H' in Table 1, scaled down by a factor 10^{-5} have an estimated mean of 0.0975 and variance of 0.001435. In the figure of Naughton *et al.* (2007), these estimates come with confidence intervals that can be grossly converted into a coefficient of variation close to 20%. The sampling variance (still for rescaled values) can thus be grossly estimated as $s^2 \approx 0.2^2 \times 0.0975^2 = 0.000380$. From $\text{var}(H') = \text{var}(H) + s^2 = \sigma^2 + s^2$, and $\text{var}(H') \approx 0.001435$ we deduce $\sigma^2 \approx 0.001058$. The error-in-variable issue is expected to "attenuate" the slope from 1 to $\frac{\sigma^2}{\sigma^2 + s^2} \approx 0.735$. The scaling of H' to h' by dividing by an estimated population size and the sampling of H' from Figure 3 in Naughton *et al.* (2007) potentially contribute also to the attenuation but were considered as second-order effects relative to the effect of uncertainty in the estimated absolute bycatch. For 1988, the only year for which a number is given by Naughton *et al.* (2007), our sampled value, 16,152, differs only by 0.4% from the actual one, 16,215. As all captures and recaptures took place between November and January, the use of calendar year bycatch estimates brings almost no further approximation or uncertainty.

For the sake of simplicity, we modelled the survival probabilities as functions of the estimated absolute bycatch H' , as $S = S_o(1 - eH')$, and converted to results relative to the estimated relative bycatch $h' = H'/N$, as $S_o(1 - b'H'/N)$, by using the relationship $b' = \hat{e}N$.

Results

There was strong evidence for heterogeneity with an overall goodness-of-fit statistic equal to 88.846, for 24 df. The various components gave the following ratios χ^2/df : Test3.SR 27.777/6 = 4.63, Test3.SM 45.7534/9 = 5.08, Test2.CT: 14.8714/5 = 2.97, Test2.CL: 0.4441/4 = 0.11. We attribute the higher values for Test3 components to differences in recapture rates of new breeders and experienced ones as a result of the temporary emigration demonstrated by Véran *et al.* (2007). We used for QAIC based model selection an overdispersion coefficient based on the overall statistic $\hat{c} = 88.246/24 = 3.7019$. As expected, this coefficient was higher than that obtained by Véran *et al.* (2007), equal to 3.02 for a Cormack-Jolly-Seber survival model.

The first five models (Table 2) started from the initial model directly linked to the goodness-of-fit tests and explored a few reduced structures. We considered departure and arrival state dependence combined with time dependence for recapture probability, i.e. this starting point was for breeders the Jolly-Move (JMV) model (with rank = 89 for 169 parameters). The goodness-of-fit results above apply to this model as explained in the Methods section.

Table 2. Initial multistate capture-recapture models for the Black-Footed Albatross data.

Recruitment (starts at age 5)	Recapture	Survival		Deviance	rank	QAIC
		1 st Year	After 1 st year			
Age 10 x time	state x time (JMV)	time	time	20695.892	89	5768.614
Age 10 x time	time (CAS)	time	time	20700.856	83	5757.954
Age 10	Time	time	time	21004.878	40	5754.080
Age 7	Time	time	time	21011.416	37	5749.847
Age 7	Time	Age+time log scale		21043.180	36	5756.427

Recapture: JMV = Jolly-Move model with dependence of recapture probability on time x state of origin x state of arrival; CAS = Conditional Arnason Schwarz model with dependence of recapture probability on time x state of arrival reducing to time dependence as recaptures occur only in state “Breeder”. Recruitment: The age of stabilisation of the recruitment is given, with or without time effect depending on the model. The model in non-Italic bold is the umbrella model. Goodness-of-fit analysis of this provides an estimated overdispersion coefficient equal to 3.7019. Further analyses proceed from the model in Italic bold with minimal QAIC.

Applying the estimated overdispersion coefficient ($\hat{c} = 3.7019$) lead to QAIC = 5768.614. The QAIC decreased when the capture probability was made dependent only on the arrival state, i.e. in the CAS counterpart of the umbrella model (QAIC = 5757.954), and when considering constancy over time in recruitment (QAIC = 5754.080). Recruitment rate could then be stabilised at age 7 (QAIC = 5749.847). This model was still clearly over-parameterised with 37 separately identifiable parameters only for a total of 59 parameters, as with few years of recapture all time-dependent survival probabilities cannot be separately estimated. As the effort in recapture was known to have varied over the years, no attempt was made to make the probability of recapture constant.

Hence, in relation to our main objective, further models (Table 3) examined various relevant structures for first year and after-first-year survival probabilities: time dependence, constancy, or linear dependence on estimated bycatch H' . While the model with constant survival did better than the time-dependent one, models with covariates did uniformly better than those with either constant or time-dependent survival. The model with different slopes for H' for first year and after first year survival in the relation $S = S_0(1 - e^{H'})$ reduced to the model with constant 1st year survival as the estimated slope for 1st year survival was 0, on the boundary of the admissible range. This model (QAIC = 5732.937) was preferable to the model with the same slope for first year and after first year survival (QAIC = 5733.437).

The estimated bycatch H' explained 59.9 % of the difference in deviance between the time dependent and constant models for after first year survival (under constant first year survival), a highly significant value ($t(19) = 5.46, P = 1.4 \cdot 10^{-5}$, one-tailed test accounting for overdispersion, Lebreton *et al.* 2011). The difference in deviance between the time-dependent model and the covariate model (21119.588-21073.588 = 45.995, for a difference of 33-14 = 19 in model

Table 3. Survival structure of the capture-recapture model for the Black-footed Albatross data. The absolute bycatch H' is used to model survival probabilities according to the theory of exploited populations as $S = S_0(1 - b \times H')$

1 st year survival	After 1 st year survival	Deviance	Rank	QAIC
Constant	Covariate H'	21119.083	14	5732.937
Covariate H' , same slope b		21120.957	14	5733.437
Time	Constant	21026.425	32	5743.901
Constant	Constant	21186.907	13	5749.252
Time	Time	21011.416	37	5749.847
Constant	Time	21073.588	33	5758.755

rank) did not provide evidence for any unexplained variation in survival over time when compared to the overdispersion coefficient ($F(19,24) = (45.595/19)/3.7019 = 0.6482, P = 0.831$).

The estimate of the first year survival probability was 0.6536 ± 0.0512 (95 % CI: 0.5532, 0.7540). The estimate of the baseline after first year survival (S_0) was 0.9647 ± 0.0199 (95 % CI: 0.9258, 1.0036). The estimate of the slope of the after first year survival relative to the estimated absolute bycatch H' was $1.14770 \cdot 10^{-5} \pm 0.24699 \cdot 10^{-5}$ (95 % CI: $0.64399 \cdot 10^{-5}, 1.65140 \cdot 10^{-5}$). The estimated recruitment probabilities (Pr (Non Breeder \rightarrow Breeder / age)) were as follows: age 5: 0.0216 ± 0.0064 ; age 6: 0.2904 ± 0.0255 ; age 7 and later: 0.5868 ± 0.0417 . These estimates are quite sensible, with more than 95 % of individuals (if alive) recruiting between ages 6 and 10, and comparable to recruitment patterns in other albatross species (e.g. Alderman *et al.* 2011).

The baseline after 1st year survival, $0.9647 (\pm 0.0199)$ was quite comparable to those obtained in albatross populations considered as not submitted to bycatch (Véran *et al.* 2007; see also Alderman *et al.* 2011). For the years 1997–2002, we obtained an average after-first-year survival equal to 0.904, reasonably comparable to the values 0.920 and 0.917 obtained by Véran *et al.* (2007) based on a model with an unobservable state to account for temporary emigration and a standard Cormack-Jolly-Seber model, respectively. The after-first-year estimated survival probability averaged over the 24 years, 0.858, is much lower as it covers the period with the higher estimated bycatch. It is 11.0% lower than the baseline survival. As all after-first-year age classes are concerned, this percentage translates into a percentage change on the population growth rate through the elasticity of the after-first year survival, $1-1/T$ (Lebreton and Clobert 1991), where T , the generation time is approximately equal to 25 (Cousins and Cooper 2000). The resulting percent change in growth rate is 10.6%. The maximum growth rate of BFAL has been estimated to 1.059 by Niel and Lebreton (2005). Decreasing by 10.9 % the maximum growth rate of 1.059 proposed by Niel and Lebreton, one obtains fair evidence for decline, at an estimated rate slightly higher than 5 % a year. Under the baseline survival probabilities, i.e. in the absence of bycatch, if each BFAL female were raising 0.8 chicks to fledging each year from age six onwards, the resulting growth rate would be 9.38% a year. Even in this optimistic and unrealistic scenario, the population would thus be unable to sustain the bycatch.

The estimated slope can then be analysed with respect to the theory of exploited population. To this effect, we converted the estimated slope \hat{e} relative to the absolute bycatch H into an estimated slope \hat{b}' relative to the estimated relative bycatch h' using the conversion $\hat{b}' = \hat{e}N$ (Table 4). The resulting slope estimate, 3.4431, seriously departs from 1, even more if accounting for the expected level of attenuation. Similar results were obtained under the second best model. The dominant source for such a difference can only be in the underestimation of bycatch, even if one might

Table 4. Estimated slopes of the linear relationship between survival and bycatch expressed in different fashions in the Black-Footed Albatross, from which one deduces minimal, “average” and maximal value of the multiplicative bias of bycatch. Even the value corresponding to the minimal bias, 0.4142, points to a strong underestimation of the bycatch, by at least 50%.

Covariate	Quantity	Estimate	Lower bound 95% CI	Higher bound 95% CI
Estimated absolute bycatch H'	Estimated Slope \hat{e} in $S = S_0(1 - eH')$ (MLE from CMR model)	$1.14770 \cdot 10^{-5}$	$0.64399 \cdot 10^{-5}$	$1.65140 \cdot 10^{-5}$
Estimated Relative bycatch $h' = H'/N$ under $N = 300,000$	Estimated Slope \hat{b}' in $S = S_0(1 - b'h')$ based on $\hat{b}' = \hat{e}N$	3.4431	1.93197	4.9542
Relative bycatch h assuming 30 % attenuation (error-in-variable)	Estimated Slope \hat{b} in $S = S_0(1 - bh)$	4.9187	2.7600	7.0774
Relative bycatch h	Multiplicative bias factor d	0.2033	0.3623	0.1413

imagine a slight effect of heterogeneity. From the further calculations in Table 4, converting in a similar fashion the boundaries of the confidence interval of the estimated slope and exploring the effect of attenuation, it seems unavoidable to conclude that the bycatch estimate is underestimated by at least 50%.

Discussion

Our study first confirms how critical is the bycatch problem for the BFAL, in two ways: first, a direct linear effect of bycatch on survival (explaining nearly 60% of the variation in after-first-year survival, a value certainly biased down by the uncertainty on the bycatch), second, a strong effect size. This strong effect size is apparent both in a strong drop in survival relative to the baseline survival, and a high slope estimate that provides fair evidence for a strong underestimation of bycatch. The slope value does not point to any form of compensation, as generally expected for long-lived species (Gauthier *et al.* 2001).

As there are few years of recapture, one should understand the relationship as follows:

For the years 1997–2004 (years 17 to 24), the drop in the after-first-year survival probability from a plausible baseline value around 0.96 to around 0.92 among 300,000 after-first year individuals at risk of by catch implies a bycatch around $0.04 \times 300\,000 = 12,000$ individuals, while the mean estimated value is 5,555. Using a less conservative estimate of total population size would only tend to increase this figure, which by itself already points to the underestimation of the bycatch. The figure of 12,000 individuals exactly matches an early warning by E. Melvin (pers. comm. in Niel and Lebreton 2005).

Then the rest of the data are, for the years prior to this period, characterised by high bycatch levels, compatible with even lower survival probabilities. The lack of evidence for a variation of first-year survival with bycatch estimate is easily explainable:

- The first year birds fledge around July (Cooper and Cousins 2000) and are thus at risk for only six months;
- In a recruitment model such as ours, what is actually estimated is the survival from fledging to the age of first reproduction at the population level, i.e. five years. The first year survival is estimated indirectly by applying from age 2 to 5 the after-first-year survival estimated from older birds, in an assumption that cannot be tested. This parameterisation makes it possible to manipulate annual survival probabilities that can be related to annual bycatch through the TEP, but dilutes the potential effect of bycatch on first year survival.

The overall immature survival, and in our case the first year survival deduced from it, might also be modified by natal dispersal. However the high first year survival probability estimate rather points to strong philopatry, as is generally the case among albatrosses (Milot *et al.* 2008).

More broadly, our survival results are based on a few years of data and do not provide by themselves unequivocal evidence. However, a) the strength of the relationship, b) the robustness with respect to heterogeneity of the various models used by Véran *et al.* (2007) and in this paper, c) the sensible baseline survival estimates, and d) the similar average survival estimates in the last six years between our model and those by Véran *et al.* (2007) all make strong sense.

Our results match and confirm what is known or suspected for bycatch underestimation (Brothers 2008, Brothers *et al.* 2011), as a result of the following biases:

- 1) The fleets on which observation can take place are likely the more ethical ones and a simple extrapolation may induce underestimation;
- 2) The observer not being continuously on deck; there is certainly some under-declaration when they are absent;
- 3) Drop-off and crippling loss of injured birds are not accounted for in these estimates as only birds hauled on board are counted. Brothers (2008) and Brothers *et al.* (2011) estimate that only 50% of seabirds caught are hauled on board.

It seems thus that an underestimation by at least 50% should be the rule: our results bring independent confirmation for such an order of magnitude.

It seems that the strong effect of bycatch for which we got evidence did not translate into what is known of population size through monitoring of colonies, as the observed decline over the period 1990–2005 (Naughton *et al.* 2007) has not reached a rate of 5% a year. Our results do not point to an imbalance of bycatch towards younger age classes, which could have explained a dampening of the bycatch effect at the population level. Delayed recruitment and a possible compensation in recruitment might explain the apparently limited impact on the number of breeders of the estimated change in survival. A more detailed assessment through a formal population projection model would be needed (see Véran and Lebreton 2008).

Coming back to bycatch as a population exploitation problem, several authors (Wade 1998, Niel and Lebreton 2005) recommended including a safety margin when calculating sustainable bycatch. Our results on the underestimation of bycatch provide clear confirmation of the need for such a margin when assessing sustainability.

Finally, we wish to insist on the fact that the discussion on the underestimation of bycatch should not be taken as a criticism of the observation programmes that produce such estimates. On the contrary, it is clear that each piece in the puzzle suffers from its own shortcomings, but is essential in assembling a coherent image. We recommend that integrated monitoring programmes, combining marked individuals, bycatch assessment and population counts are developed to help produce a coherent image through integrative tools such as those brought by the theory of exploited populations.

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