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Manatū Ahu Matua

Stock assessment of hake (Merluccius australis) in the Sub-Antarctic (part of HAK 1) for the 2011-12 fishing year

New Zealand Fisheries Assessment Report 2013/5.
P.L. Horn

ISSN 1179-5352 (online)
ISBN 978-0-478-40511-8 (online)
January 2013


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## EXECUTIVE SUMMARY

## Horn, P.L. (2013). Stock assessment of hake (Merluccius australis) in the Sub-Antarctic (part of HAK 1) for the 2011-12 fishing year. <br> New Zealand Fisheries Assessment Report 2013/5. 52 p.

This report summarises the stock assessment of hake in Quota Management Area HAK 1 south of latitude $46^{\circ} \mathrm{S}$ (the Sub-Antarctic) for the 2011-10 fishing year. An updated Bayesian assessment was conducted using the general-purpose stock assessment program CASAL v2.22. The assessment incorporated all relevant biological parameters, the commercial catch histories, updated CPUE series, and series of proportion-at-age data from the commercial trawl fishery and two research survey series. The analysis includes fishery data up to the end of the 2009-10 fishing year. New model input data and revised catch histories for all three hake stocks (Sub-Antarctic, Chatham Rise, and west coast South Island) are also reported here.

Initial investigations of the available data and preliminary model runs indicated that the sex ratios in the at-age data were inconsistent; the CPUE series was inconsistent with survey biomass estimates; and widely fluctuating estimates of year class strengths throughout the late 1970 s were driven by the error structure applied to the age data acting on sparse data sets. Consequently, a base case model was developed in which sex was excluded, the 1974 to 1979 year class strengths were smoothed, and the CPUE series was omitted.

Three additional models were run as sensitivity analyses. The 'two sex' model included sex in the partition and 'at-age' data were provided by sex. The 'CPUE' model was the same as the single sex model, but it incorporated a trawl fishery CPUE series. The 'estimate $M$ ' model estimated instantaneous natural mortality as a function of age.

The stock status of hake in the Sub-Antarctic is not well known in absolute terms. It appears likely that the stock has declined throughout the 1990s, with the decline driven by poor recruitment rather than fishing pressure. The single sex model indicated that the spawning stock is currently at about $52 \% \mathrm{~B}_{0}$, and that $\mathrm{B}_{0}$ was about 95000 t . The estimate $M$ model produced slightly lower estimates $\left(52 \% \mathrm{~B}_{0}\right.$, and 84000 t$)$, but fitted the available data better than the single sex model. The two sex model was considered undesirable because it was not believed that the inclusion of sex in the partition could logically account for a more than doubling of biomass. The CPUE model produced the most optimistic assessment (current spawning stock of $68 \% \mathrm{~B}_{0}$ ), but the CPUE series was not well fitted and did not capture the decline in biomass during the 1990 s. However, none of the model runs were indicative of current biomass being lower than the target of $40 \% \mathrm{~B}_{0}$, and all projected an increase in biomass over the next five years with future catches equal to those from recent years ( 2300 t ).

The stock is probably being well monitored by the November-December trawl survey series. While the stock status appears to be reasonably well defined, estimates of past and current absolute stock size are very uncertain owing to poor contrast in the relative abundance series. Because of the high uncertainty in estimates of absolute biomass, yield estimates are also very uncertain. However, there are probably no current sustainability issues for this stock.

## 1. INTRODUCTION

This report outlines the stock assessment of hake (Merluccius australis) in the Sub-Antarctic section of Quota Management Area (QMA) HAK 1 (i.e., HAK 1 south of latitude $46^{\circ}$ S), with the inclusion of data up to the end of the 2009-10 fishing year. The current stock hypothesis for New Zealand hake suggests that there are three separate hake stocks (Colman 1998); the west coast South Island stock (WCSI, the area of HAK 7 off the west coast of South Island), the Sub-Antarctic stock (the area of HAK 1 that encompasses the Southern Plateau, the Stewart-Snares shelf, and Puysegur Bank), and the Chatham Rise stock (HAK 4 and the area of HAK 1 on the western Chatham Rise).

The stock assessment of hake in the Sub-Antarctic is presented as a Bayesian assessment implemented as a single stock model using the general-purpose stock assessment program CASAL (Bull et al. 2008). Estimates of the current stock status and projected stock status are provided.

This report fulfils Objective 3 of Project DEE201002HAKA "To update the stock assessment of hake, including biomass estimates and sustainable yields", funded by the Ministry of Fisheries. Revised catch histories are reported here, as are new model input data and research results for all three hake stocks. Although some of these data are not relevant to the assessment reported here, they are included to provide in one place an up-to-date summary of the available knowledge and literature on Merluccius australis in New Zealand waters.

### 1.1 Description of the fishery

Hake are widely distributed through the middle depths of the New Zealand Exclusive Economic Zone (EEZ) mostly south of latitude $40^{\circ} \mathrm{S}$ (Anderson et al. 1998). Adults are mainly distributed in depths from 250 to 800 m although some have been found as deep as 1200 m , while juveniles $(0+$ ) are found in shallower inshore regions under 250 m (Hurst et al. 2000). Hake are taken by large trawlers often as bycatch in fisheries targeting hoki, although target fisheries also exist (Devine 2009). Present management divides the fishery into three main fish stocks: (a) the Challenger QMA (HAK 7), (b) the Southeast (Chatham Rise) QMA (HAK 4), and (c) the remainder of the EEZ comprising the Auckland, Central, Southeast (Coast), Southland, and Sub-Antarctic QMAs (HAK 1). An administrative fish stock exists in the Kermadec QMA (HAK 10) although there are no recorded landings from this area. The hake QMAs are shown in Figure 1.

The largest fishery has been off the west coast of the South Island (HAK 7) with the highest catch ( 17000 t ) recorded in 1977, immediately before the establishment of the EEZ. Currently, the TACC for HAK 7 is the largest, at 7700 t out of a total for the EEZ of 13211 t . The WCSI hake fishery has generally consisted of bycatch in the much larger hoki fishery, but it has undergone a number of changes during the last decade (Devine 2009). These include changes to the TACCs of both hake and hoki, and also changes in fishing practices such as gear used, tow duration, and strategies to limit hake bycatch. In some years, notably in 1992, 1993, 2006, and 2009 there has been a hake target fishery in September after the peak of the hoki fishery is over (Ballara 2012).

On the Chatham Rise and in the Sub-Antarctic, hake have been caught mainly as bycatch by trawlers targeting hoki, although significant targeting occurs in both areas (Devine 2009). Increases in TACCs from 2610 t to 3500 t in HAK 1 and from 1000 t to 3500 t in HAK 4 from the 1991-92 fishing year allowed the fleet to increase the landings of hake from these fish stocks. Reported catches rose over a number of years to the levels of the new TACCs in both HAK 1 and HAK 4, with catches in HAK 1 remaining relatively steady since. The TACC for HAK 1 has increased since then to its current level of 3701 t . Landings from HAK 4 steadily declined from 1998-99 to a low of 811 t in 2002-03, but increased to 2275 t in 2003-04. However, from 2004-05, the TACC for HAK 4 was reduced from 3500 t to 1800 t with an overall TAC of 1818 t . Annual landings have been markedly lower than the new TACC since then. From 1 October 2005 the TACC for HAK 7 was increased to 7700 t with an
overall TAC of 7777 t . This new catch limit was set equal to average annual catches over the previous 12 years, a catch level that is believed to be sustainable in the short term.

Dunn (2003a) found that area misreporting between the WCSI and the Chatham Rise fisheries occurred from 1994-95 to 2000-01. He estimated that between 16 and $23 \%$ ( $700-1000 t$ annually) of WCSI landings were misreported as deriving from Chatham Rise, predominantly in June, July, and September. Levels of misreporting before 1994-95 and after 2000-01, and between WCSI and SubAntarctic, were estimated as negligible, and there is no evidence of significant misreporting since 2001-02 (Devine 2009).


Figure 1: Quota Management Areas (QMAs) HAK 1, 4, 7, and 10; and the west coast South Island (light shading), Chatham Rise (dark shading), and Sub-Antarctic (medium shading) hake stock boundaries assumed in this report.

### 1.2 Literature review

Previous assessments of hake, by fishing year, are as follows: 1991-92 (Colman et al. 1991), 1992-93 (Colman \& Vignaux 1992), 1997-98 (Colman 1997), 1998-99 (Dunn 1998), 1999-2000 (Dunn et al. 2000), 2000-01 (Dunn 2001), 2002-03 (Dunn 2003b), 2003-04 (Dunn 2004a, 2004b), 2004-05 (Dunn et al. 2006), 2005-06 (Dunn 2006), and 2006-07 (Horn \& Dunn 2007). The Bayesian stock assessment software CASAL (Bull et al. 2008) has been used for all assessments since 2002-03. The most recent assessments by stock are: Chatham Rise (Horn \& Francis 2010), Sub-Antarctic (Horn 2008), and WCSI (Horn 2011).

Since 1991, resource surveys have been carried out from R.V. Tangaroa in the Sub-Antarctic in November-December 1991-1993 and 2000-2009, September-October 1992, and April-June 1992,

1993, 1996, and 1998. On Chatham Rise, a consistent time series of resource surveys from Tangaroa has been carried out in January 1992-2011. Appendix A gives more details on these surveys.

Standardised CPUE indices for the Sub-Antarctic, Chatham Rise, and WCSI stocks were updated for the period up to the 2008-09 fishing year (Ballara \& Horn 2011). These update the indices estimated by Phillips \& Livingston (2004), Kendrick (1998), Dunn et al. (2000), Dunn \& Phillips (2006), Devine \& Dunn (2008), and Devine (2010). Indices for the Sub-Antarctic stock only were updated to the 2009-10 fishing year by Ballara (2012). A descriptive analysis of all New Zealand's hake fisheries up to the 2005-06 fishing year was prepared by Devine (2009).

## 2. REVIEW OF THE FISHERY

### 2.1 TACCs, catch, landings, and effort data

Reported catches from 1975 to 1987-88 are shown in Table 1, and reported landings for each QMA since 1983-84 and TACCs since 1986-87 are shown in Table 2. Revised estimates of landings by QMA for 1989-90 to 2009-10 (Table 3) were derived by examining the reported tow-by-tow catches of hake and correcting for possible misreporting, using the method of Dunn (2003a).

Revised landings by stock are given in Table 4. The derivation of the catch from 1974-75 to 1988-89 was described for the Chatham Rise and Sub-Antarctic stocks by Dunn et al. (2000) and for WCSI by Dunn (2004b). Landings since 1989-90 from Chatham Rise and Sub-Antarctic and since 1991-92 for WCSI were obtained from the corrected data used to produce Table 3, but this time summing the landings reported in each of the three shaded areas shown on Figure 1. WCSI revised estimates for 1988-89 to 1990-91 are from Colman \& Vignaux (1992), who estimated the actual hake catch in HAK 7 by multiplying the total hoki catch (which was assumed to be correctly reported by vessels both with and without observers) by the ratio of hake to hoki in the catch of vessels carrying observers. Reported and estimated catches for 1988-89 were respectively 6835 t and 8696 t ; for 1989-90, 4903 t reported and 8741 t estimated; and for 1990-91, 6189 t reported and 8246 t estimated. The catch for 2010-11 from the Sub-Antarctic was assumed based on landings from recent years (Table 4).

Table 1: Reported hake catches (t) from 1975 to 1987-88. Data from 1975 to 1983 from Ministry of Agriculture \& Fisheries (Fisheries); data from 1983-84 to 1985-86 from Fisheries Statistics Unit; data from 1986-87 to 1987-88 from Quota Management System.

|  |  | New Zeala | vessels |  |  | $n$ licens | vessels |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Fishing year | Domestic | Chartered | Total | Japan | Korea | USSR | Total | Total |
| $1975{ }^{1}$ | 0 | 0 | 0 | 382 | 0 | 0 | 382 | 382 |
| $1976{ }^{1}$ | 0 | 0 | 0 | 5474 | 0 | 300 | 5774 | 5774 |
| $1977{ }^{\text { }}$ | 0 | 0 | 0 | 12482 | 5784 | 1200 | 19466 | 19466 |
| 1978-79 ${ }^{2}$ | 0 | 3 | 3 | 398 | 308 | 585 | 1291 | 1294 |
| 1979-80 ${ }^{2}$ | 0 | 5283 | 5283 | 293 | 0 | 134 | 427 | 5710 |
| 1980-81 ${ }^{2}$ |  |  |  | data av | able |  |  |  |
| 1981-82 ${ }^{2}$ | 0 | 3513 | 3513 | 268 | 9 | 44 | 321 | 3834 |
| 1982-83 ${ }^{2}$ | 38 | 2107 | 2145 | 203 | 53 | 0 | 255 | 2400 |
| $1983{ }^{3}$ | 2 | 1006 | 1008 | 382 | 67 | 2 | 451 | 1459 |
| 1983-84 ${ }^{4}$ | 196 | 1212 | 1408 | 522 | 76 | 5 | 603 | 2011 |
| 1984-85 ${ }^{4}$ | 265 | 1318 | 1583 | 400 | 35 | 16 | 451 | 2034 |
| 1985-86 ${ }^{4}$ | 241 | 2104 | 2345 | 465 | 52 | 13 | 530 | 2875 |
| 1986-87 ${ }^{4}$ | 229 | 3666 | 3895 | 234 | 1 | 1 | 236 | 4131 |
| 1987-88 ${ }^{4}$ | 122 | 4334 | 4456 | 231 | 1 | 1 | 233 | 4689 |
| 1. Calendar year |  |  |  |  |  |  |  |  |
| 2. 1 April to 31 March |  |  |  |  |  |  |  |  |
| 3.1 April to 30 September |  |  |  |  |  |  |  |  |
| 4. 1 October to 30 September |  |  |  |  |  |  |  |  |

Table 2: Reported landings ( $t$ ) of hake by QMA from 1983-84 to 2009-10 and actual TACCs ( $\mathbf{t}$ ) for 198687 to 2009-10. Data from 1983-84 to 1985-86 from Fisheries Statistics Unit; data from 1986-87 to 200910 from Quota Management System (- indicates that the data are unavailable).

| QMA | HAK 1 |  | HAK 4 |  | HAK 7 |  | HAK 10 |  | Total |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Landings | TACC | Landings | TACC | Landings | TACC | Landings | TACC | Landings | TACC |
| 1983-84 | 886 | - | 180 | - | 945 | - | 0 | - | 2011 |  |
| 1984-85 | 670 | - | 399 | - | 965 | - | 0 | - | 2034 | - |
| 1985-86 | 1047 | - | 133 | - | 1695 | - | 0 |  | 2875 | - |
| 1986-87 | 1022 | 2500 | 200 | 1000 | 2909 | 3000 | 0 | 10 | 4131 | 6510 |
| 1987-88 | 1381 | 2500 | 288 | 1000 | 3019 | 3000 | 0 | 10 | 4689 | 6510 |
| 1988-89 | 1487 | 2513 | 554 | 1000 | 6835 | 3004 | 0 | 10 | 8876 | 6527 |
| 1989-90 | 2115 | 2610 | 763 | 1000 | 4903 | 3310 | 0 | 10 | 7783 | 6930 |
| 1990-91 | 2603 | 2610 | 743 | 1000 | 6148 | 3310 | 0 | 10 | 9567 | 6930 |
| 1991-92 | 3156 | 3500 | 2013 | 3500 | 3026 | 6770 | 0 | 10 | 8196 | 13780 |
| 1992-93 | 3525 | 3501 | 2546 | 3500 | 7154 | 6835 | 0 | 10 | 13224 | 13846 |
| 1993-94 | 1803 | 3501 | 2587 | 3500 | 2974 | 6835 | 0 | 10 | 7363 | 13847 |
| 1994-95 | 2572 | 3632 | 3369 | 3500 | 8841 | 6855 | 0 | 10 | 14781 | 13997 |
| 1995-96 | 3956 | 3632 | 3465 | 3500 | 8678 | 6855 | 0 | 10 | 16082 | 13997 |
| 1996-97 | 3534 | 3632 | 3524 | 3500 | 6118 | 6855 | 0 | 10 | 13176 | 13997 |
| 1997-98 | 3809 | 3632 | 3523 | 3500 | 7416 | 6855 | 0 | 10 | 14749 | 13997 |
| 1998-99 | 3845 | 3632 | 3324 | 3500 | 8165 | 6855 | 0 | 10 | 15333 | 13997 |
| 1999-00 | 3899 | 3632 | 2803 | 3500 | 6898 | 6855 | 0 | 10 | 13600 | 13997 |
| 2000-01 | 3504 | 3632 | 2472 | 3500 | 8134 | 6855 | 0 | 10 | 14110 | 13997 |
| 2001-02 | 2870 | 3701 | 1424 | 3500 | 7519 | 6855 | 0 | 10 | 11813 | 14066 |
| 2002-03 | 3336 | 3701 | 811 | 3500 | 7433 | 6855 | 0 | 10 | 11581 | 14066 |
| 2003-04 | 3461 | 3701 | 2272 | 3500 | 7943 | 6855 | 0 | 10 | 13686 | 14066 |
| 2004-05 | 4797 | 3701 | 1266 | 1800 | 7316 | 6855 | 0 | 10 | 13377 | 12366 |
| 2005-06 | 2743 | 3701 | 305 | 1800 | 6906 | 7700 | 0 | 10 | 9955 | 13211 |
| 2006-07 | 2025 | 3701 | 900 | 1800 | 7668 | 7700 | 0 | 10 | 10592 | 13211 |
| 2007-08 | 2445 | 3701 | 865 | 1800 | 2620 | 7700 | 0 | 10 | 5930 | 13211 |
| 2008-09 | 3415 | 3701 | 856 | 1800 | 5954 | 7700 | 0 | 10 | 10226 | 13211 |
| 2009-10 | 2156 | 3701 | 208 | 1800 | 2351 | 7700 | 0 | 10 | 4715 | 13211 |

Table 3: Revised landings (t) by QMA 1989-90 to 2009-10 from Ballara (2012).

| Fishing |  |  | QMA | Total |
| :--- | ---: | ---: | ---: | ---: |
| Year | HAK 1 | HAK 4 | HAK 7 |  |
| $1989-90$ | 2115 | 763 | 4903 | 7781 |
| $1990-91$ | 2593 | 726 | 6175 | 9494 |
| $1991-92$ | 3156 | 2013 | 3027 | 8196 |
| $1992-93$ | 3522 | 2546 | 7157 | 13225 |
| $1993-94$ | 1787 | 2587 | 2990 | 7364 |
| $1994-95$ | 2346 | 2896 | 9538 | 14780 |
| $1995-96$ | 3828 | 3070 | 9089 | 15987 |
| $1996-97$ | 3300 | 3190 | 6846 | 13336 |
| $1997-98$ | 3659 | 3239 | 7683 | 14581 |
| $1998-99$ | 3703 | 2740 | 8786 | 15229 |
| $1999-00$ | 3781 | 2756 | 7042 | 13579 |
| $2000-01$ | 3429 | 2321 | 8351 | 14101 |
| $2001-02$ | 2865 | 1420 | 7499 | 11784 |
| $2002-03$ | 3334 | 805 | 7406 | 11545 |
| $2003-04$ | 3455 | 2254 | 7943 | 13652 |
| $2004-05$ | 4795 | 1260 | 7302 | 13357 |
| $2005-06$ | 2742 | 305 | 6897 | 9944 |
| $2006-07$ | 2006 | 900 | 7660 | 10566 |
| $2007-08$ | 2442 | 865 | 2615 | 5922 |
| $2008-09$ | 3409 | 854 | 5945 | 10208 |
| $2009-10$ | 2156 | 208 | 2340 | 4704 |

Table 4: Revised landings (t) from 1974-75 to 2009-10 for the Sub-Antarctic (Sub-A), Chatham Rise (Chat), and west coast South Island (WCSI) stocks. The Sub-Antarctic catch for the most recent year is assumed based on recent catch levels in the fishery.

| Fishing <br> year | Sub-A | Chat | WCSI | Fishing <br> year | Sub-A | Chat | WCSI |
| :--- | ---: | ---: | ---: | :--- | :--- | :--- | ---: |
| $1974-75$ | 120 | 191 | 71 | $1993-94$ | 1453 | 2934 | 2971 |
| $1975-76$ | 281 | 488 | 5005 | $1994-95$ | 1852 | 3387 | 9535 |
| $1976-77$ | 372 | 1288 | 17806 | $1995-96$ | 2873 | 4028 | 9082 |
| $1977-78$ | 762 | 34 | 498 | $1996-97$ | 2262 | 4234 | 6838 |
| $1978-79$ | 364 | 609 | 4737 | $1997-98$ | 2606 | 4252 | 7674 |
| $1979-80$ | 350 | 750 | 3600 | $1998-99$ | 2796 | 3669 | 8742 |
| $1980-81$ | 272 | 997 | 2565 | $1999-00$ | 3020 | 3517 | 7031 |
| $1981-82$ | 179 | 596 | 1625 | $2000-01$ | 2790 | 2962 | 8346 |
| $1982-83$ | 448 | 302 | 745 | $2001-02$ | 2510 | 1770 | 7498 |
| $1983-84$ | 722 | 344 | 945 | $2002-03$ | 2738 | 1401 | 7404 |
| $1984-85$ | 525 | 544 | 965 | $2003-04$ | 3245 | 2465 | 7939 |
| $1985-86$ | 818 | 362 | 1918 | $2004-05$ | 2531 | 3526 | 7298 |
| $1986-87$ | 713 | 509 | 3755 | $2005-06$ | 2557 | 489 | 6892 |
| $1987-88$ | 1095 | 574 | 3009 | $2006-07$ | 1818 | 1081 | 7660 |
| $1988-89$ | 1237 | 804 | 8696 | $2007-08$ | 2202 | 1096 | 2583 |
| $1989-90$ | 1927 | 950 | 8741 | $2008-09$ | 2427 | 1825 | 5912 |
| $1990-91$ | 2370 | 931 | 8246 | $2009-10$ | 1958 | 391 | 2282 |
| $1991-92$ | 2750 | 2418 | 3010 | $2010-11$ | 2000 | - | - |

### 2.2 Other sources of fishing mortality

The recreational fishery for hake is believed to be negligible. The amount of hake caught by Maori is not known, but is believed to be negligible. There is likely to be some mortality associated with escapement from trawl nets, but the level is not known.

## 3. BIOLOGY, STOCK STRUCTURE, AND RESOURCE SURVEYS

### 3.1 Biology

Data collected by observers on commercial trawlers and from resource surveys suggest that there are at least three main spawning areas for hake (Colman 1998). The best known area is off the west coast of the South Island, where the season can extend from June to October, possibly with a peak in September. Spawning also occurs to the west of the Chatham Islands during a prolonged period from at least September to January. Spawning fish have also been recorded occasionally near the Mernoo Bank. Spawning on the Campbell Plateau, primarily to the northeast of the Auckland Islands, may occur from September to February with a peak in September-October. Spawning fish have also been recorded occasionally on the Puysegur Bank, with a seasonality that appears similar to that on the Campbell Plateau (Colman 1998).

Horn (1997) validated the use of otoliths to age hake. New Zealand hake reach a maximum age of at least 25 years. Males, which rarely exceed 100 cm total length, do not grow as large as females, which can grow to 120 cm total length or more. Readings of otoliths from hake have been used as age-length keys to scale up length frequency distributions for hake collected on resource surveys and from commercial fisheries in all three stocks. The resulting age frequency distributions were reported by Horn \& Sutton (2012). The observed proportions-at-age data available from resource surveys of the Sub-Antarctic and Chatham Rise stocks are also shown in Figure 2, and estimated proportions-at-age data from the commercial trawl fisheries are shown for the Sub-Antarctic and west coast South Island in Figure 3a and the Chatham Rise in Figure 3b.

Colman (1998) found that hake reach sexual maturity between 6 and 10 years of age, at total lengths of about $67-75 \mathrm{~cm}$ (males) and $75-85 \mathrm{~cm}$ (females); he concluded that hake reached $50 \%$ maturity at between 6 and 8 years in HAK 1, and $7-8$ years in HAK 4. In assessments before 2005, the maturity ogive for the Chatham Rise and Sub-Antarctic was assumed from a combination of the estimates of Colman (1998) and model fits to the west coast South Island stock presented by Dunn (1998). From 2005 to 2007, maturity ogives for the Chatham Rise and Sub-Antarctic stocks were fitted within the assessment model to data derived from resource survey samples with information on the gonosomatic index, gonad stage, and age (Horn \& Dunn 2007, Horn 2008). Individual hake were classified as either immature or mature at sex and age, where maturity was determined from the gonad stage and gonosomatic index (GSI, the ratio of the gonad weight to body weight). Fish identified as stage 1 were classified as immature. Stage 2 fish were classified as immature or mature depending on the GSI index, using the definitions of Colman (1998) - i.e., classified as immature if GSI was less than 0.005 (males) or GSI less than 0.015 (females), or mature if GSI at least 0.005 (males) or GSI at least 0.015 (females). Fish identified as stages 3-7 were classified as mature. Model fits indicated that Chatham Rise hake reach $50 \%$ maturity at about 6 years for males and 7 years for females, and SubAntarctic hake reach $50 \%$ maturity at 6 years for males and 7.5 years for females (Figure 4). From 2009, fixed ogives (derived from the fitted curves in Figure 4) were used in the assessment models, with values listed in Table 5 . The values for Chatham Rise combined sexes maturity were taken as the mean of the male and female values. In the absence of sufficient data to estimate an ogive for WCSI hake, maturity for this stock was assumed to be the same as for Chatham Rise hake.

Von Bertalanffy parameters were previously estimated using data up to 1997 (Horn 1998). The parameters for all three stocks were updated using all data available at February 2007 (Horn 2008). Plots of the fitted curves on the raw data indicated that the von Bertalanffy model tended to underestimate the age of large fish (Figure 5). Consequently, the growth model of Schnute (1981) was fitted to the data sets (Table 5). This model appeared to better describe the growth of larger hake (Figure 5), and the resulting parameters can be used in the CASAL stock assessment software. Most aged hake have been 3 years or older. However, juvenile hake have been taken in coastal waters on both sides of the South Island and on the Campbell Plateau. It is known that they reach a total length of about $15-20 \mathrm{~cm}$ at 1 year old, and about 35 cm total length at 2 years (Horn 1997).

Estimates of natural mortality $(M)$ and the associated methodology were given by Dunn et al. (2000); $M$ was estimated as $0.18 \mathrm{y}^{-1}$ for females and $0.20 \mathrm{y}^{-1}$ for males. Colman et al. (1991) estimated $M$ as $0.20 \mathrm{y}^{-1}$ for females and $0.22 \mathrm{y}^{-1}$ for males using the maximum age method of Hoenig (1983) (where they defined the maximum ages at which $1 \%$ of the population survives in an unexploited stock as 23 years for females and 21 years for males). These are similar to the values proposed by Horn (1997), who determined the age of hake by counting zones in sectioned otoliths and concluded from that study that it was likely that $M$ was in the range $0.20-0.25 \mathrm{y}^{-1}$.

Dunn et al. (2010) found that the diet of hake on the Chatham Rise was dominated by teleost fishes, in particular Macrouridae. Macrouridae accounted for $44 \%$ of the prey weight and consisted of at least six species, of which javelinfish, Lepidorhynchus denticulatus, was most frequently identified. Hoki were less frequent prey, but being relatively large accounted for $37 \%$ of prey weight. Squids were found in $7 \%$ of the stomachs, and accounted for $5 \%$ of the prey weight. Crustacean prey were predominantly natant decapods, with pasiphaeid prawns occurring in $19 \%$ of the stomachs.

Length-weight relationships for hake from the Sub-Antarctic and Chatham Rise stocks were revised here using all available length-weight data collected during trawl surveys since 1989. Parameters were calculated for males, females, and both sexes combined (Table 5). Sample sizes were large (SubAntarctic: 3182 males, 6010 females; Chatham Rise, 3527 males, 3511 females) and all $r^{2}$ values were greater than 0.96 .

## Sub-Antarctic



Figure 2: Age frequencies by year class and year (symbol area proportional to the proportions-at-age within sampling event) from the Sub-Antarctic (ages 3-20) and Chatham Rise (ages 3-18) resource surveys. Zero values are represented by a dash.

## Sub-Antarctic



Figure 3a: Age frequencies by year class and year (symbol area proportional to the proportions-at-age within sampling event) from commercial trawl fisheries in the Sub-Antarctic (ages 3-20) and west coast South Island (ages 3-18) resource surveys. Zero values are represented by a dash.

Chatham Rise west


Chatham Rise east


Figure 3b: Age frequencies by year class and year (symbol area proportional to the proportions-at-age within sampling event) from commercial trawl fisheries on the western and eastern sections of Chatham Rise (ages 3-18). Zero values are represented by a dash.

Table 5: Estimates of biological parameters for the three hake stocks.

|  | Estimate Source |  |  |
| :---: | :---: | :---: | :---: |
| Natural mortality |  |  |  |
| Males | $M=0.20$ |  | (Dunn et al. 2000) |
| Females | $M=0.18$ |  | (Dunn et al. 2000) |
| Weight $=a \cdot(\text { length })^{b}($ Weight in $t$, length in cm$)$ |  |  |  |
| Sub-Antarctic |  |  |  |
| Males | $a=2.13 \times 10^{-9}$ | $b=3.281$ | (Current study) |
| Females | $a=1.83 \times 10^{-9}$ | $b=3.314$ | (Current study) |
| Both sexes | $a=1.95 \times 10^{-9}$ | $b=3.301$ | (Current study) |
| Chatham Rise |  |  |  |
| Males | $a=2.56 \times 10^{-9}$ | $b=3.228$ | (Current study) |
| Females | $a=1.88 \times 10^{-9}$ | $b=3.305$ | (Current study) |
| Both sexes | $a=2.00 \times 10^{-9}$ | $b=3.288$ | (Current study) |

von Bertalanffy growth parameters
Sub-Antarctic
Males $\quad k=0.295 \quad t_{0}=0.06 \quad L_{\infty}=88.8 \quad($ Horn 2008)
Females $\quad k=0.220 \quad t_{0}=0.01 \quad L_{\infty}=107.3 \quad$ (Horn 2008)

Chatham Rise

$$
\begin{array}{rllll}
\begin{array}{r}
\text { Males } \\
\text { Females }
\end{array} & k=0.330 & t_{0}=0.09 & L_{\infty}=85.3 & (\text { Horn 2008) } \\
& k=0.229 & t_{0}=0.01 & L_{\infty}=106.5 & (\text { Horn 2008) } \\
\text { Males } & k=0.357 & t_{0}=0.11 & L_{\infty}=82.3 & (\text { Horn 2008) } \\
\text { Females } & k=0.280 & t_{0}=0.08 & L_{\infty}=99.6 & (\text { Horn 2008) }
\end{array}
$$

WCSI

Schnute growth parameters ( $\tau_{1}=1$ and $\tau_{2}=20$ for all stocks)
Sub-Antarctic

$$
\begin{array}{rlllll}
\text { Males } & y_{1}=22.3 & y_{2}=89.8 & a=0.249 & b=1.243 \text { (Horn 2008) } \\
\text { Females } & y_{1}=22.9 & y_{2}=109.9 & a=0.147 & b=1.457 \text { (Horn 2008) } \\
\text { Both sexes } & y_{1}=22.8 & y_{2}=101.8 & a=0.179 & b=1.350 \text { (Current study) }
\end{array}
$$

Chatham Rise
Males $\quad y_{1}=24.6 \quad y_{2}=90.1 \quad a=0.184 \quad b=1.742$ (Horn 2008)
Females $y_{1}=24.4 \quad y_{2}=114.5 \quad a=0.098 \quad b=1.764$ (Horn 2008)
Both sexes $y_{1}=24.5 \quad y_{2}=104.8 \quad a=0.131 \quad b=1.700$ (Horn \& Francis 2010)
WCSI

$$
\begin{array}{rrrrrr}
\text { Males } & y_{1}=23.7 & y_{2}=83.9 & a=0.278 & b=1.380 \text { (Horn 2008) } \\
\text { Females } & y_{1}=24.5 & y_{2}=103.6 & a=0.182 & b=1.510 \text { (Horn 2008) } \\
\text { Both sexes } & y_{1}=24.5 & y_{2}=98.5 & a=0.214 & b=1.570 \text { (Horn 2011) }
\end{array}
$$

Maturity ogives (proportion mature at age)

| Age | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Sub-Antarctic |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Males | 0.01 | 0.03 | 0.09 | 0.22 | 0.46 | 0.71 | 0.88 | 0.96 | 0.98 | 0.99 | 1.00 | 1.00 | 1.00 | 1.00 |
| Females | 0.01 | 0.02 | 0.05 | 0.11 | 0.23 | 0.43 | 0.64 | 0.81 | 0.91 | 0.96 | 0.98 | 0.99 | 1.00 | 1.00 |
| Both | 0.01 | 0.02 | 0.07 | 0.16 | 0.29 | 0.57 | 0.76 | 0.88 | 0.94 | 0.97 | 0.99 | 0.99 | 1.00 | 1.00 |
| Chatham | Rise (and assumed for WCSI) |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Males | 0.0 | 0.06 | 0.15 | 0.32 | 0.55 | 0.77 | 0.90 | 0.96 | 0.98 | 0.99 | 1.00 | 1.00 | 1.00 | 1.00 |
| Females | 0.04 | 0.07 | 0.13 | 0.22 | 0.34 | 0.49 | 0.64 | 0.77 | 0.86 | 0.92 | 0.95 | 0.98 | 0.99 | 1.00 |
| Both | 0.03 | 0.06 | 0.14 | 0.27 | 0.45 | 0.63 | 0.77 | 0.86 | 0.92 | 0.96 | 0.98 | 0.99 | 1.00 | 1.00 |


| Miscellaneous parameters |  |
| :--- | :--- |
| Steepness (Beverton \& Holt stock-recruitment relationship) | 0.90 |
| Proportion spawning | 1.0 |
| Proportion of recruits that are male | 0.5 |
| Ageing error c.v. | 0.08 |
| Maximum exploitation rate $\left(U_{\max }\right)$ | 0.7 |



Figure 4: Estimated ogives of proportions mature by age for Sub-Antarctic and Chatham Rise hake males (solid lines) and females (broken lines).


Figure 5: Raw age-length data, by sex, for hake from Chatham Rise (CHAT), west coast South Island (WCSI), and Sub-Antarctic (SUBA), with fitted von Bertalanffy curves (solid lines) and Schnute curves (broken lines).

### 3.2 Stock structure

There are at least three hake spawning areas: off the west coast of the South Island, on the Chatham Rise, and on the Campbell Plateau (Colman 1998). Juvenile hake are found in all three areas, there are differences in size frequency of hake between the west coast and other areas, and differences in
growth parameters between all three areas (Horn 1997). There is reason, therefore, to believe that at least three separate stocks may exist in the EEZ.

Analysis of morphometric data (J.A. Colman, NIWA, unpublished data) showed little difference between hake from the Chatham Rise and from the east coast of the North Island, but highly significant differences between these fish and those from the Sub-Antarctic, Puysegur, and on the west coast. The Puysegur fish are most similar to those from the west coast South Island, although, depending on which variables are used, they cannot always be distinguished from the Sub-Antarctic hake. However, the data are not unequivocal, so the stock affinity is uncertain.

For stock assessment models, the Chatham Rise stock was considered to include the whole of the Chatham Rise (HAK 4 and the western end of the Chatham Rise that forms part of the HAK 1 management area). The Sub-Antarctic stock was considered to contain hake in the remaining Puysegur, Southland, and Sub-Antarctic regions of the HAK 1 management area. The stock areas assumed for this report are shown earlier, in Figure 1.

### 3.3 Resource surveys

In the Sub-Antarctic, three resource surveys were carried out by Tangaroa with the same gear and similar survey designs in November-December 1991, 1992, and 1993, but the series was then terminated as there was evidence that hake, in particular, might be aggregated for spawning at that time of the year and that spawning aggregations had a high probability of being missed during a survey. However, research interest in hoki in the Sub-Antarctic resulted in a return to the NovemberDecember survey annually from 2000 to 2009. Surveys by Tangaroa in April 1992, May 1993, April 1996, and April 1998 formed the basis for a second series, with hake appearing to be more evenly distributed through the survey area at that time of year. A single survey in September 1992 by Tangaroa was also completed. The biomass estimates from the Sub-Antarctic Tangaroa surveys are shown in Figure 6 with further details given in Appendix A. There was some variation in the area surveyed within the November-December and April-May series. Consequently, multiple biomass estimates have been presented in Figure 6 for some of these surveys to show biomass estimates across consistent sets of strata. However, only the longest consistent time series (i.e., November-December 1991 area, April-May 300-800 m strata) are included as assessment model inputs.

Sub-Antarctic surveys were conducted by Shinkai Maru (March-May 1982 and October-November 1983) and Amaltal Explorer (October-November 1989, July-August 1990, and November-December 1990). However, these vessels used different gear and had different performance characteristics (Livingston et al. 2002), so biomass estimates from these surveys cannot be used as part of a consistent time series.

Resource surveys have been carried out at depths of 200-800 m on the Chatham Rise since 1992 by Tangaroa with the same gear and similar survey designs (Figure 7, Appendix A). While the survey designs since 1992 have been similar, there was a reduction in the number of stations surveyed between 1996 and 1999, and some strata in the survey design used between 1996 and 1999 were merged (see Bull \& Bagley 1999). The surveys since 2000 used a revised design, with some strata being split and additional stations added. Since 2000 some of the Tangaroa surveys included deepwater strata (i.e., $800-1300 \mathrm{~m}$ ) on the Chatham Rise. Although only the longest consistent time series (i.e., $200-800 \mathrm{~m}$ strata) is included in the assessment model, additional estimates from surveys fishing the deepwater strata are shown in Figure 7.

Chatham Rise surveys were conducted by Shinkai Maru (March 1983 and June-July 1986) and Amaltal Explorer (November-December 1989). However, these surveys used a range of gear, survey methodologies, and survey designs (Livingston et al. 2002), and cannot be used as part of a consistent time series.

Research surveys of hoki and hake have been conducted periodically off WCSI, but these have been 'one-off' surveys by different vessels (i.e., Shinkai Maru in 1976, James Cook in 1978-79, Wesermünde in 1979, Giljanes in 1990, and Tangaroa in 2000) so any biomass estimates from them are not useful model inputs. It is possible that the 2000 Tangaroa survey (O'Driscoll et al. 2004) may be able to be linked to a trawl and acoustic survey series due to commence off WCSI in winter 2012 to produce a future series. A long-running trawl survey series of inshore waters off WCSI by Kaharoa has not provided a useful index of hake biomass as it surveys no deeper than 400 m (Stevenson \& Hanchet 2000). Age data, and consequent estimates of proportion-at-age, are available for only the 1979 Wesermünde survey; these are incorporated in the WCSI assessment model.


Figure 6: Hake biomass estimates from the Tangaroa surveys of the Sub-Antarctic (1991-2009 including the November-December, April-May, and September series), with approximate 95\% confidence intervals. (See also Appendix A.)


Figure 7: Hake biomass estimates from the Tangaroa (1992-2011 for the January series) of the Chatham Rise, with approximate $95 \%$ confidence intervals. (See also Appendix A.)

### 3.4 Observer age data

### 3.4.1 Chatham Rise

The fishery on the Chatham Rise was stratified using a tree-based regression on mean lengths of hake in tows where observers had measured five or more hake (Horn \& Dunn 2007). The defined strata are shown in Figure 8. Mean fish length tends to increase from west to east, and with increasing depth. Area 404 contains a known spawning ground. Where sufficient data were available, catch-at-age series were developed and selectivity ogives estimated separately for each fishery in the stock assessment model by Horn \& Dunn (2007).


Figure 8: Fishery strata defined for the Chatham Rise hake fishery. Large numbers show longitudes or depths of fishery boundaries; small numbers denote statistical areas. The stratum boundary defined by depth ( 530 m ) is shown only approximately. Isobaths at 1000,500 , and 250 m are also shown.

However, Horn \& Francis (2010) showed that the two western fisheries had similar age-frequency distributions, and the two eastern fisheries were data poor. Consequently, they used two strata, eastern and western, divided at $178.1^{\circ} \mathrm{E}$. Observer data from each fishery stratum were converted into catch-at-age distributions if there were at least 400 length measurements (from western strata) or 320 length measurements (from eastern strata), and the mean weighted c.v. over all age classes was less than $30 \%$. The available data (described by Horn \& Sutton (2012)) are from 1991-92 and 1993-94 to 2009-10. Although the observer length data from each year were partitioned into fisheries (i.e., two strata in each of the two fisheries, as shown in Figure 8), the age data from each year were not (i.e., a single age-length key was constructed for each year and applied to all available sets of length data from that year). Horn \& Dunn (2007) showed that mean age at length did not differ between fisheries, so the use of a single age-length key per year would not bias the age distributions.

### 3.4.2 Sub-Antarctic

The Sub-Antarctic hake observer data were found to be best stratified into the four areas shown in Figure 9 (Horn 2008). Most of the hake target fishing, and most of the catch (average $94 \%$ per year), is associated with the Snares-Pukaki area. Puysegur is the next most important area with about $3 \%$ of the catch. Available observer data are also concentrated in the Snares-Pukaki region, but it is clear that the smaller fisheries (particularly the Campbell Island area) can be over-sampled in most years. Consequently, the Sub-Antarctic observer data are analysed as one major and three very minor fisheries, with a single fishery ogive. However, because of clear differences in mean fish length between the fisheries (Horn 2008), it is important to use the four fishery strata when calculating catch-at-age distributions. Without stratification, the frequent over-sampling in the minor fisheries could strongly bias the catch-at-age distributions. A single age-length key was applied to the scaled lengthfrequency distributions for each fishery to produce the catch-at-age data. Catch-at-age distributions
from the Sub-Antarctic trawl fishery are available from all but three years from 1989-90 to 2009-10 (Horn \& Sutton 2012).


Figure 9: Fishery strata defined for the Sub-Antarctic hake fishery. Large numbers show latitudes or longitudes of fishery boundaries; small numbers denote statistical areas. Isobaths at 1000, 500, and 250 m are also shown.

### 3.4.3 WCSI

The fishery off WCSI was stratified using a tree-based regression on mean lengths of hake in tows where observers had measured five or more hake (Horn \& Dunn 2007). A single catch-at-age distribution was estimated for each year, stratified as shown in Figure 10. Catch-at-age distributions from the WCSI trawl fishery are available from all years from 1989-90 to 2009-10 (Horn \& Sutton 2012).


Figure 10: Fishery strata defined for the WCSI hake fishery. Large numbers show latitudes or depths of fishery boundaries; small numbers denote statistical areas. The stratum boundary defined by depth $(629 \mathrm{~m})$ is shown only approximately. Isobaths at 1000,500 , and 250 m are also shown.

### 3.5 CPUE indices

Standardised CPUE indices were calculated by Ballara \& Horn (2011) from daily processed summary data up the end of the 2008-09 fishing season. Series were produced for the separate eastern and western fisheries on the Chatham Rise, and for all areas combined in each of the Chatham Rise, SubAntarctic, and WCSI stocks.

As only the HAK 1 assessment is being completed under the current project, the Sub-Antarctic CPUE was updated using data to the end of the 2009-10 fishing year by Ballara (2012) (Table 6).

Table 6: Hake CPUE indices (and associated 95\% confidence intervals and c.v.s) for the Sub-Antarctic trawl fishery (from Ballara 2012).

| Year | Index | $95 \%$ CI | c.v. | Year | Index | $95 \% \mathrm{CI}$ | c.v. |
| ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: |
|  |  |  |  |  |  |  |  |
| 1991 | 1.25 | $1.15-1.35$ | 0.04 | 2001 | 1.05 | $0.99-1.11$ | 0.03 |
| 1992 | 1.21 | $1.13-1.29$ | 0.03 | 2002 | 0.95 | $0.90-1.01$ | 0.03 |
| 1993 | 1.18 | $1.11-1.26$ | 0.03 | 2003 | 0.93 | $0.88-0.98$ | 0.03 |
| 1994 | 1.07 | $0.99-1.15$ | 0.04 | 2004 | 1.05 | $0.99-1.11$ | 0.03 |
| 1995 | 0.93 | $0.87-1.00$ | 0.04 | 2005 | 0.81 | $0.75-0.87$ | 0.04 |
| 1996 | 1.17 | $1.08-1.27$ | 0.04 | 2006 | 1.14 | $1.04-1.25$ | 0.05 |
| 1997 | 0.96 | $0.90-1.02$ | 0.03 | 2007 | 1.04 | $0.95-1.13$ | 0.05 |
| 1998 | 0.91 | $0.86-0.96$ | 0.03 | 2008 | 0.90 | $0.83-0.97$ | 0.04 |
| 1999 | 0.86 | $0.81-0.91$ | 0.03 | 2009 | 0.88 | $0.81-0.95$ | 0.04 |
| 2000 | 0.88 | $0.83-0.93$ | 0.03 | 2010 | 1.00 | $0.93-1.09$ | 0.04 |

## 4. MODEL STRUCTURE, INPUTS, AND ESTIMATION

### 4.1 Introduction

An updated assessment of the Sub-Antarctic stock is presented here. The previous assessment of this stock (Horn 2008) partitioned the population into two sexes and age groups $1-30$, with the last age class considered a plus group. The partition included maturity, with ogives estimated within the model. The model's annual cycle began on 1 September and was divided into three steps. The current assessment model used the same annual cycle (Table 7), but maturity ogives were always fixed and maturity was never included in the partition. Note that model references to "year" within this document are labelled as the most recent calendar year, i.e., the year 1 September 1998 to 31 August 1999 is referred to as " 1999 ". Some previous assessments of Sub-Antarctic hake have been based on fishing year, i.e., years starting on 1 October. However, landings peaks tend to occur from September to January (Ballara 2012), so it is logical to include the September catch with landings from the five months following it, rather than with catch taken about seven months previously.

Table 7: Annual cycle of the Sub-Antarctic stock model, showing the processes taking place at each time step, their sequence within each time step, and the available observations. Fishing and natural mortality that occur within a time step occur after all other processes, with half of the natural mortality for that time step occurring before and half after the fishing mortality.

|  |  |  |  |  | Observations |  |
| :--- | :--- | :--- | ---: | :--- | ---: | :--- |
|  | Step | Period | Processes | $M$ <br> fraction $^{1}$ | Age <br> fraction $^{2}$ | Description |

${ }^{1}$ The proportion of natural mortality that was assumed to have occurred in that time step.
${ }^{2}$ The age fraction (used for determining length at age) that was assumed to occur in that time step.
${ }^{3} \% \mathrm{Z}$ is the $\%$ of total mortality in the step that was assumed to have taken place at the time each observation was made.
For all subsequent models, estimates of fixed biological parameters used in the assessments are given in Table 5. A Beverton-Holt stock-recruitment relationship, with steepness 0.9, was assumed. Variability in the Schnute age-length relationship was assumed to be lognormal with a constant c.v. of 0.1. The maximum exploitation rate was assumed to be 0.7 for the stock. The choice of the maximum exploitation rate has the effect of determining the minimum possible virgin biomass allowed by the model, given the observed catch history. This value was set relatively high as there was little external information from which to determine it. The model's annual cycle was as described in Table 7.

Biomass estimates from the resource surveys were used as relative biomass indices, with associated c.v.s estimated from the survey analysis. The survey catchability constant $(q)$ for each survey series was assumed to be constant over all years in that series. Three $q$ values were estimated; one for each of the summer, autumn, and spring survey series. Although the summer and autumn series were believed to have different $q$ values, no information was available to indicate whether the spring survey might have a $q$ similar to either of the other series, so a separate $q$ was estimated for it. Catch-at-age observations were available for each Tangaroa research survey (see Figure 2), from a single Amaltal Explorer survey in November 1989, and from commercial observer data for the fishery. Two selectivity ogives were used for the survey catch-at-age data; one for the summer series (i.e., the Tangaroa and Amaltal Explorer data), and another for the autumn and spring Tangaroa data. There was no information to indicate that the Amaltal Explorer and the Tangaroa in summer had similar selectivities, or that the autumn and spring Tangaroa selectivities were the same. However, it was considered undesirable to estimate selectivity ogives for single surveys (i.e., the Amaltal Explorer and the Tangaroa in spring), so they were analysed as described above. Lognormal errors, with known c.v.s, were assumed for all relative biomass and proportions-at-age observations. Ageing error was
assumed to occur for the observed proportions-at-age data, by assuming a discrete normally distributed error with c.v. 0.08 .

The c.v.s (for observations fitted with lognormal likelihoods) are assumed to have allowed for sampling error only. Additional variance, assumed to arise from differences between model simplifications and real world variation, was added to the sampling variance for all observations in all model runs. For catch-at-age data the additional variance, termed process error, was estimated in MPD runs of each model. Process error of 0.2 was added to all survey biomass indices following the recommendation of Francis et al. (2003). For CPUE indices, process error was fixed at 0.18 so that the final point c.v.s were approximately 0.2 , as recommended by Francis et al. (2001).

Year class strengths were assumed known (and equal to one) for years before 1974 and after 2007, when inadequate or no catch-at-age data were available. Otherwise, year class strengths were estimated under the assumption that the estimates from the model must average one. The Haist parameterisation for year class multipliers is used here (see Bull et al. (2008) for details).

The catch history assumed in all model runs was derived as follows. Using the grooming algorithms of Dunn (2003a), landings of hake reported on TCEPR and CELR forms from 1989-90 to 2009-10 were allocated to month and fishery (based on reported date and location). Annual totals were obtained by summing the monthly totals using a September to August year (for reasons described above). Thus, catch histories for model years 1990 to 2010 were produced for the Sub-Antarctic section of HAK 1 (Table 8). Annual catches before 1990 are as presented in Table 4.

Table 8: Estimated catch (t) from the Sub-Antarctic stock, by fishing year, and model year. Note that from 1989-90 totals by fishing year and model year differ because the September catch has been shifted from the fishing year into the following model year. Model year landings from 2011 are estimated assuming catch patterns similar to the previous year.

| Fishing year | Catch <br> (t) | Model year | Catch <br> (t) | Fishing year | Catch <br> (t) | Model year | Catch <br> (t) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1974-75 | 120 | 1975 | 120 | 1993-94 | 1450 | 1994 | 1596 |
| 1975-76 | 281 | 1976 | 281 | 1994-95 | 1852 | 1995 | 1995 |
| 1976-77 | 372 | 1977 | 372 | 1995-96 | 2870 | 1996 | 2779 |
| 1977-78 | 762 | 1978 | 762 | 1996-97 | 2271 | 1997 | 1915 |
| 1978-79 | 364 | 1979 | 364 | 1997-98 | 2628 | 1998 | 2958 |
| 1979-80 | 350 | 1980 | 350 | 1998-99 | 2802 | 1999 | 2854 |
| 1980-81 | 272 | 1981 | 272 | 1999-00 | 3030 | 2000 | 3108 |
| 1981-82 | 179 | 1982 | 179 | 2000-01 | 2849 | 2001 | 2820 |
| 1982-83 | 448 | 1983 | 448 | 2001-02 | 2512 | 2002 | 2444 |
| 1983-84 | 722 | 1984 | 722 | 2002-03 | 2729 | 2003 | 2777 |
| 1984-85 | 525 | 1985 | 525 | 2003-04 | 3252 | 2004 | 3223 |
| 1985-86 | 818 | 1986 | 818 | 2004-05 | 2528 | 2005 | 2592 |
| 1986-87 | 713 | 1987 | 713 | 2005-06 | 2554 | 2006 | 2541 |
| 1987-88 | 1095 | 1988 | 1095 | 2006-07 | 1815 | 2007 | 1711 |
| 1988-89 | 1237 | 1989 | 1237 | 2007-08 | 2204 | 2008 | 2329 |
| 1989-90 | 1917 | 1990 | 1897 | 2008-09 | 2432 | 2009 | 2446 |
| 1990-91 | 2370 | 1991 | 2381 | 2009-10 | 1958 | 2010 | 1927 |
| 1991-92 | 2743 | 1992 | 2810 | 2010-11 | - | 2011 | 2000 |
| 1992-93 | 3254 | 1993 | 3941 |  |  |  |  |

### 4.2 Prior distributions and penalty functions

The assumed prior distributions used in the assessment are given in Table 9 . The priors for $\mathrm{B}_{0}$ and year class strengths were intended to be relatively uninformed, and had wide bounds. The prior for the survey $q$ was informative and was estimated by assuming that the catchability constant was the product of areal availability, vertical availability, and vulnerability. This same $q$ prior was used in the
previous Sub-Antarctic hake assessment (Horn 2008). A simulation was conducted that estimated a distribution of possible values for the catchability constant by assuming that each of these factors was independent and uniformly distributed. A prior was then determined by assuming that the resulting, sampled, distribution was lognormally distributed. Values assumed for the parameters were areal availability ( $0.50-1.00$ ), vertical availability ( $0.50-1.00$ ), and vulnerability ( $0.01-0.50$ ). The resulting (approximate lognormal) distribution had mean 0.16 and c.v. 0.79 , with bounds assumed to be 0.01 and 0.40 (Figure 11). Priors for all selectivity parameters were assumed to be uniform. The values of survey catchability constants are dependent on the selectivity parameters, and the absolute catchability can be determined by the product of the selectivity by age and sex, and the catchability constant $q$.

Penalty functions were used a) to constrain the model so that any combination of parameters that resulted in a stock size that was so low that the historical catch could not have been taken was strongly penalised, b) to ensure that all estimated year class strengths averaged 1 , and $c$ ) to smooth the year class strengths estimated over the period 1974 to 1979.

Table 9: The priors assumed for estimated parameters. The given parameters for the priors are mean (in natural space) and c.v.

| Parameter description | Distribution |  | eters |  | Bounds |
| :---: | :---: | :---: | :---: | :---: | :---: |
| $B_{0}$ | uniform-log | - | - | 5000 | 350000 |
| Year class strengths | Lognormal | 1.0 | 1.1 | 0.01 | 100 |
| Trawl survey $q^{1}$ | Lognormal | 0.16 | 0.79 | 0.01 | 0.4 |
| CPUE $q$ | uniform-log | - | - | $1 \mathrm{e}-8$ | $1 \mathrm{e}-3$ |
| Selectivities | Uniform | - | - | 0 | $20-200^{2}$ |
| $M\left(x_{0}, y_{0}, y_{1}, y_{2}\right)^{3}$ | Uniform | - | - | $3,0.01,0.01,0.01$ | $15,0.6,1.0,1.0$ |
| ${ }^{1}$ Three trawl survey $q$ values were estimated, but all had the same priors. |  |  |  |  |  |
|  |  |  |  |  |  |
| ${ }^{3}{ }^{3}$ A , age at minimum $M ; y_{0}, M$ at $x_{0} ; y_{1}, M$ at the minimum age in the partition; $y_{2}, M$ at the maximum age in the partite |  |  |  |  |  |



Figure 11: The prior distribution for all three of the survey catchability constants (q), lognormal where $\mu=0.16$, c.v. $=0.79$, and bounds $(0.01,0.40)$.

### 4.3 Developing a 'base' model

Some initial investigations were completed to develop a 'base' model. Model parameters were estimated for final model runs using Bayesian estimation implemented using the CASAL software. However, only the mode of the joint posterior distribution (MPD) was estimated in these initial runs. All runs included survey biomass indices and proportion-at-age data, and the fishery catch-at-age data. Full details of the CASAL algorithms, software, and methods were detailed by Bull et al. (2008).

In developing a base model a series of seven models was considered, with each new model typically differing from previous models in only one key assumption (Table 10).

Table 10: Brief description of the assumptions that differed amongst the eight models that were considered in developing a base case model (see text for more detail). For each model, the underlined assumption is the main one that distinguished it from preceding models.

| Assumption |  |  |  |  | Model number |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 1 | 2 | $3^{1}$ | 4 | $5^{2}$ | 6 | 7 |
| Ageing error assumed | Y | $\underline{\mathrm{N}}$ | Y | Y | Y | Y | Y |
| Smooth 1974-79 year-class strengths | N | N | $\underline{Y}$ | Y | Y | Y | Y |
| All selectivities domed ${ }^{3}$ | Y | Y | Y | N | Y | Y | N |
| Sex in partition and data | Y | Y | Y | Y | $\underline{N}$ | N | N |
| CPUE data used | N | N | N | N | N | Y | N |
| $M$ estimated in model as an ogive | N | N | N | N | N | N | $\underline{Y}$ |
| ${ }^{1}$ Referred to as the 'two sex' model. <br> ${ }^{2}$ Base case model for the assessment. <br> ${ }^{3}$ Logistic selectivities were used for the s | m | a | fish | m |  |  |  |

All models estimated an absolute biomass trajectory, year class strengths from 1974 to 2007, fits to any included relative abundance series, selectivity ogives for the trawl surveys and trawl fishery, and trawl survey catchability coefficients. One model also estimated instantaneous natural mortality.

An initial model (model 1) was set up, partitioning the population into two sexes and age groups $1-$ 30, with the last age class considered a plus group. The partition did not include maturity. The model used six selectivity ogives: male and female survey selectivities for the summer resource survey series, male and female survey selectivities for the autumn and spring resource survey series, and male and female selectivities for the commercial trawl fishery. Male selectivity was estimated relative to female selectivity. Selectivities were assumed constant over all years in the fisheries and the research surveys. All selectivity ogives were estimated using the double-normal parameterisation. Process error for all the catch-at-age series was estimated in this initial MPD model, and these values were used in all subsequent models. The initial model produced the following estimates of process error for the catch-at-age series: summer research survey, 0.3 ; autumn research survey, 0.2 ; spring research survey, 0.1 ; fishery, 0.6 . No CPUE data were incorporated.
$B_{0}$ was 184500 t , and stock status in 2011 was estimated to be $57 \%$ of $B_{0}$. The two survey biomass series with multiple points appeared to be reasonably well fitted, with no obvious trends in the residuals (Figure 12). However, a striking feature of model 1 is that the spawning biomass was estimated to have increased by $56 \%$ in the 1980 s, before the survey series started, and this increase was driven primarily by extremely strong year classes in 1977 and 1980 (Figure 13).


Figure 12: Fits to the summer (thick line, with observed values as solid circles) and autumn (thin line, with observed values as open squares) research survey series.


Figure 13: Estimated year class strengths and spawning stock biomass, from models 1-3.
The age data were examined to see what information existed to indicate that the 1980 and 1977 year classes were particularly strong (Figure 14). There was clear indication from the early parts of the proportion-at-age series (i.e., 1990 and 1992-1994) that the 1980 year class is strong. However, the 1977 year class seldom appears to be strong in any data set; only the 1990 summer and 1993 autumn survey distributions suggest that this year class might be stronger than average. It was suspected that the estimated 1977 strong year class was an artefact, the consequence of a tendency for models which assume ageing error to estimate high variability in year-class strength in periods with few data.

Consequently, two additional models were run to investigate year class strength estimation: in model 2 , the assumption of ageing error was dropped; model 3 retained this assumption but smoothed the year class strengths from 1974 to 1979.

The new models both produced early year class strength estimates that were markedly different from those of model 1, but there was little change in estimated spawning stock biomass or stock status (Figure 13). It appeared very likely that the extreme estimates of year class strength before 1980 (both high and low) are artefacts of the application of ageing error to age classes with few data. All three models produced similar patterns of year class strengths from 1981 to 2008, where the data were more abundant. The two new models were similar over their entire range, and they still provided a clear indication of some stronger than average year classes in the late 1970s.

It was considered desirable to include some ageing error in the assessment model, so the smoothing of early year class strengths was retained for all subsequent models. The effect of including this smoothing was to only slightly degrade the overall fit (Table 11).

Table 11: Negative log likelihood of all data series for models 1 and 3, showing how the smoothing of early year class strengths in the latter model slightly degraded the fits compared to those in the former.

| Data series | Model 1 | Model 3 | Gain |
| :--- | ---: | ---: | ---: |
| Survey biomass (summer) | -10.7 | -10.4 | -0.3 |
| Survey biomass (autumn) | -3.7 | -4.3 | 0.6 |
| Survey biomass (spring) | -1.3 | -1.3 | 0.0 |
| Survey age (summer) | 183.1 | 185.9 | -2.8 |
| Survey age (autumn) | 55.0 | 50.1 | 4.9 |
| Survey age (spring) | 22.7 | 24.7 | -2.0 |
| Fishery age | 162.0 | 161.6 | 0.4 |
| Priors \& penalties | 84.1 | 85.8 | -1.7 |
| Total log likelihood | 491.2 | 492.3 | -1.1 |

## Trawl surveys



Figure 14: Observed (symbols) and estimated (lines, calculated for model 1) proportions-at-age, by year, from the research trawl surveys and commercial fishery. Observed data for the $\mathbf{1 9 8 0}$ year class are represented as open squares, and 1977 year class data as open triangles.

The models investigated so far had selectivity ogives that had been fitted using the double-normal parameterisation. The effects of forcing logistic selectivity ogives for the research biomass surveys were examined in model 4 , with the underlying assumptions being that the surveys comprehensively sample all the adult population. However, the overall fit for this model was much worse than for model 3, particularly for the three series of at-age data where logistic selectivity ogives were applied (Table 12). Consequently, it was concluded that given the currently used constant values for natural mortality rate, catch-at-age data from all sources are much better fitted by double-normal, rather than logistic, ogives. However, the double-normal selectivity ogives were markedly different between sexes (Figure 15), particularly for the summer survey, and this was believed to be unsatisfactory and unrealistic.

Table 12: Negative log likelihood of all data series from models 3 and 4, showing how forcing ogives for the survey and east fishery to be logistic substantially degraded the fit to the corresponding at-age data.

| Data series | Model 3 | Model 4 | Gain |
| :--- | ---: | ---: | ---: |
| Survey biomass (summer) | -10.4 | -11.2 | 0.8 |
| Survey biomass (autumn) | -4.3 | -3.9 | -0.4 |
| Survey biomass (spring) | -1.3 | -1.3 | 0.0 |
| Survey age (summer) | 185.9 | 214.1 | -28.2 |
| Survey age (autumn) | 50.1 | 59.2 | -9.1 |
| Survey age (spring) | 24.7 | 29.4 | -4.7 |
| Fishery age | 161.6 | 161.8 | -0.2 |
| Priors \& penalties | 85.8 | 86.1 | -0.3 |
| Total log likelihood | 492.3 | 534.2 | -41.9 |



Figure 15: MPD estimates of trawl selectivity from model 3, for male (solid line) and female (dashed line) hake.

A likelihood profile for model 3 showed that there were two groups of inputs producing conflicting signals (Figure 16). High values of $\mathrm{B}_{0}$ (greater than 270000 t ) were strongly supported by the three largest sets of at-age data (i.e., the summer survey, the autumn survey, and the trawl fishery) and the summer survey biomass indices. Low values of $\mathrm{B}_{0}$ ( 90000 t or less) were weakly supported by the autumn survey biomass indices and the at-age data from the spring survey. However, biomass levels at the high end of this range would require exceptionally low trawl survey $q$ values (i.e., less than 0.02 ), and would be inconsistent with $B_{0}$ estimates for other New Zealand hake stocks, i.e., 41000 t and 82000 t for the Chatham Rise and west coast South Island stocks, respectively (Horn \& Francis 2010, Horn 2011).

It was also apparent that the sex ratio information for the trawl surveys and commercial fishery were inconsistent. Sex ratios in the surveys have been relatively consistent over time, with perhaps a weak trend of increasing proportions of males over time (Figure 17). However, the samples from the commercial catch are indicative of an increasing proportion of males being removed from the population over time (Figure 17). If proportionally more males have been removed in recent years then the surveys would be expected to show a reduction in the proportion of males over time. It was also found that male proportion-at-age data from the fishery were consistently poorly fitted relative to female data; the
residuals for male data points were, on average, 1.5 times greater than for female data. Similar characteristics were observed in the Chatham Rise and west coast South Island hake fisheries, and the Chatham Rise fishery also exhibited similar conflicts in sex ratio trends. However, for both those stocks, the subsequent modelling problems were alleviated by removing sex from the partition (Horn \& Francis 2010, Horn 2011).


Figure 16: Likelihood profile on $B_{0}$ for model 3, showing both the total likelihood (heavy line) and those for individual data series. Vertical dashed line shows the model estimate of $B_{0}$.


Figure 17: Proportion of male fish recorded in trawl surveys (scaled to survey area) and in the observer length data for the commercial trawl fishery (scaled to total catch by year). Solid lines are linear regressions.

Consequently, a single-sex model was tested. In that model, sex was removed from the partition, lengthweight and Schnute growth parameters were calculated for both sexes combined (see Table 5), $M$ was set at 0.19 (the average of the male and female values), all catch-at-age data were unsexed, and the 1974-79 year class strengths were smoothed as before.

Removing sex from the model substantially reduced the estimate of $B_{0}$ (from 194000 t to 95000 t ) (Figure 18). It had comparatively little effect on the estimated year class strengths, slightly flattening the estimates (Figure 18). Despite the lower $B_{0}$, stock status in 2011 of $51 \%$ of $B_{0}$ was little different to the $54 \% \mathrm{~B}_{0}$ estimated from the two sex model. A period of relatively strong recruitment in the late 1970s was still indicated, resulting in a moderate increase in stock biomass during the 1980s before the start of the survey series. The selectivity ogives are logical, i.e., the two survey ogives are similar and exhibit greater selectivity on young (small) fish than the commercial fishery, as would be expected given the smaller codend mesh in the survey trawl (Figure 19).

A major effect of removing sex from the model was to reduce the degree of conflict between the input data series. All series strongly reject a $\mathrm{B}_{0}$ lower than 60000 t . However, there was little clear support for a distinct value greater than about 90000 t (Figure 20).


Figure 18: Biomass trajectories and estimates of year class strengths from the initial two-sex model (model 3, thick lines) and the subsequent single-sex model (model 5 , thin lines).


Figure 19: Estimated selectivity ogives for the research surveys and the commercial fishery from model 5.


Figure 20: Likelihood profile on $B_{0}$ for model 5, showing both the total likelihood (heavy line) and those for individual data series. Vertical dashed line shows the model estimate of $\mathbf{B}_{0}$.

The impact of adding a trawl fishery CPUE series to the single-sex model was investigated and found to markedly increase the estimate of $\mathrm{B}_{0}$ (from 95000 t to 131000 t ) (Figure 21). It had comparatively little effect on the estimated year class strengths, generally reducing earlier estimates and increasing later ones (Figure 21). It encouraged a flatter biomass trajectory, and consequently a better stock status in 2011 of $67 \%$ of $\mathrm{B}_{0}$ (compared to $51 \% \mathrm{~B}_{0}$ for the single-sex model). However, the initial part of the CPUE series is poorly fitted; it does not show the marked decline in biomass throughout the early 1990s indicated by other model inputs (Figure 22). Consequently, the CPUE also encourages flatter (but not clearly worse) fits to the trawl survey series (Figure 22).


Figure 21: Biomass trajectories and estimates of year class strengths from the single-sex model without CPUE (model 5, solid lines) and with CPUE (model 6, dashed lines).


Figure 22: MPD fits to the CPUE series ('o', observed; ' $e$ ', expected), and fits to the trawl survey series from the single-sex model without CPUE (model 5 , solid lines) and with CPUE (model 6, dashed lines).

The data for this stock comprise a large amount of catch-at-age information so it was considered desirable to investigate the estimation of $M$ in the model. Because $M$ is confounded with selectivity, the model run where $M$ was estimated (model 7) had the selectivity ogive for the trawl fishery estimated using the logistic (rather than double-normal) parameterisation, with the underlying assumption being that the fishery comprehensively samples all the adult population. The effects of forcing logistic selectivity ogives for the surveys were examined above in model 4 , but $M$ was fixed as a constant in that case. In model $7, M$ was estimated using the double exponential parameterisation. It was clear that by allowing the model to estimate an age-dependent $M$, the fits to the summer survey age data could be markedly improved with no significant degradation to the fits to any other input series (Table 13).

Table 13: Negative log likelihood of all data series from models 5 and 7, showing how estimating $M$ as agedependent and forcing the fishery ogive to be logistic substantially improved the fit to the corresponding atage data.

| Data series | Model 5 | Model 7 | Gain |
| :--- | ---: | ---: | ---: |
| Survey biomass (summer) | -10.1 | -9.6 | -0.5 |
| Survey biomass (autumn) | -4.3 | -4.3 | 0.0 |
| Survey biomass (spring) | -1.4 | -1.4 | 0.0 |
| Survey age (summer) | 7.9 | -11.6 | 19.5 |
| Survey age (autumn) | -9.3 | -8.4 | -0.9 |
| Survey age (spring) | 2.6 | 2.8 | -0.2 |
| Fishery age | -5.5 | -6.1 | 0.6 |
| Priors \& penalties | 81.0 | 66.8 | 14.2 |
| Total log likelihood | 60.1 | 28.1 | 32.0 |

The estimated ogive for $M$ was logical with a minimum at age 7 (slightly lower than the estimated age at $100 \%$ maturity of 11 years), and a range from 0.08 to 0.74 (Figure 23). The impact of estimating $M$ was a marked decrease in the estimate of $B_{0}$ (69 $100 t$, down from $95000 t$ in model 5) (Figure 24). It had comparatively little effect on the estimated year class strengths, and also resulted in only slight changes in the fits to the trawl survey series (Figure 24). The biomass trajectory still exhibits a steep decline throughout the 1990 s, resulting in a stock status in 2011 of $43 \%$ of $\mathrm{B}_{0}$ (compared to $51 \% \mathrm{~B}_{0}$ for the single-sex model). This was the most pessimistic of all the models tested.


Figure 23: Estimated $M$ ogive from model 7. The horizontal dashed line indicates the constant $M$ of 0.19 used in all other model runs.


Figure 24: Biomass trajectories, estimates of year class strengths, and fits to the trawl survey series from model 5 (single-sex model, solid lines) and model 7 (estimate $M$ model, dashed lines).

Following the investigations above with MPD model fits it was concluded that the best base case model for MCMC estimation was model 5 (the single sex model). Sensitivity model runs using MCMC estimation were also completed to investigate sex in the partition (model 3, the two sex model), the influence of CPUE (model 6, the CPUE model), and the estimation of $M$ (model 7, the estimate $M$ model).

### 4.4 Model estimation using MCMC

Model parameters were derived using Bayesian estimation implemented using the CASAL software. For final runs, the full posterior distribution was sampled using Monte Carlo Markov Chain (MCMC) methods, based on the Metropolis-Hastings algorithm. MCMCs were estimated using $3 \times 10^{6}$ iterations, a burn-in length of $5 \times 10^{5}$ iterations, and with every $2500^{\text {th }}$ sample kept from the final $2.5 \times 10^{6}$ iterations (i.e., a final sample of length 1000 was taken from the Bayesian posterior). Year class strengths were estimated as in the MPD runs except that values for 2008-10 were no longer fixed at 1 .

## 5. MODEL ESTIMATES

Estimates of biomass were made using the biological parameters (see Table 5) and model input parameters described earlier. MCMC estimates of the posterior distribution were obtained for all four model runs (i.e., single sex, two sex, CPUE, and $M$ estimation), and are presented below. In addition, MCMC estimates of the median posterior and $95 \%$ percentile credible intervals are reported for the key output parameters. The MCMC chains for estimates of $\mathrm{B}_{0}$ and $\mathrm{B}_{2011}$ from the base (single sex) model are not well converged (Figure 25). However, the distributions of estimates of $\mathrm{B}_{0}$ and $\mathrm{B}_{2011}$ (as $\% \mathrm{~B}_{0}$ ) from the single sex model are reasonably consistent between the first, middle, and last thirds of the chain (Figure 25), so convergence is probably adequate for stock-assessment purposes.

The estimated MCMC marginal posterior distributions for selected parameters from the single sex model are shown in Figures 26-30. The selectivity ogives for the trawl survey series and the trawl fishery were all strongly domed (Figure 26). The ogives suggest that hake were fully selected by the fishery by age 10 , and that younger fish were more selected by the trawl surveys (as would be expected given the smaller codend mesh). There is no information outside the model that allows the shape of the estimated selectivity ogives to be verified.

Year class strength estimates were well estimated for all years from 1985 to 2005, and moderately well estimated in all other years (Figure 27). Recall, however, that the estimates from 1974 to 1979 were smoothed. The 1980 year class stands out as being exceptionally strong, but variation in year class strength since then does not appear to be great with virtually all median estimates being between 0.5 and 2.

Estimated biomass for the Sub-Antarctic stock increased throughout the 1980s owing to relatively low catch levels, and the recruitment of stronger than average 1974 to 1980 year classes (Figure 28). Biomass then steadily declined from 1988 to about 2000 owing to higher levels of exploitation (Figure 29) and the recruitment of year classes that were generally of below-average strength. A slower decline continued to about 2009, with a slight increase since then being a consequence of the recruitment of three year classes since 2005 that are estimated to be of above-average strength. Bounds around the biomass estimates are wide, with current stock size being about $52 \%$ of $\mathrm{B}_{0}(95 \%$ credible interval 39-64\%) (see Figure 28 and Table 14). Exploitation rates (catch over vulnerable biomass) were negligible (less than 0.02 ) up to 1990 , and have probably not exceeded $0.1 \mathrm{yr}^{-1}$ in any year since then (Figure 29).

The research survey catchability constants are estimated to be about $3 \%, 4 \%$ and $4 \%$ for the summer, autumn and spring survey series, respectively, suggesting that the absolute catchability of all the surveys is very low, and not very consistent with the prior (Figure 30).


Figure 25: Trace diagnostic plot of the MCMC chains for estimates of $B_{0}$ and $B_{2011}$ for the single sex model run (top and middle panel). MCMC diagnostic plots showing the cumulative frequencies of $B_{0}$ and $\mathbf{B}_{2011}\left(\%_{0}\right)$ for the first (thick line), middle (medium line), and last (thin line) third of the MCMC chain for the single sex model (bottom panel).

Table 14: Bayesian median and $95 \%$ credible intervals of $B_{0}, B_{2011}$, and $B_{2011}$ as a percentage of $B_{0}$ for all model runs.

| Model run | $\mathrm{B}_{0}$ |  |  | $\mathrm{B}_{2011}$ |  | $\mathrm{B}_{2011}\left(\% \mathrm{~B}_{0}\right)$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Single sex | 94150 | (59 220-156 350) | 49590 | (23 860-95 220) |  | (39.0-64.5) |
| Two sex | 192190 | (121 120-295 650) | 106740 | (59 400-178 110) | 54. | (45.9-63.9) |
| CPUE | 135700 | (84 040-227 040) | 91930 | (51 870-166 760) |  | (58.3-77.3) |
| Estimate M | 78240 | (51 810-135 590) | 36170 | (17 820-77 080) | 46. | (32.3-58.6) |



Figure 26: Single sex model - Estimated median selectivity ogive (with 95\% credible intervals shown as dashed lines) for the trawl surveys and the trawl fishery.


Figure 27: Single sex model - Estimated posterior distributions of year class strengths. The dashed horizontal line indicates the year class strength of one. Individual distributions are the marginal posteriors, with horizontal lines indicating the median.


Figure 28: Single sex model - Estimated median trajectories (with $95 \%$ credible intervals shown as dashed lines) for absolute biomass and biomass as a percentage of $\mathbf{B}_{0}$.


Figure 29: Single sex model - Estimated median trajectory of exploitation rate.


Figure 30: Single sex model - Estimated posterior distribution (thin line) and prior (thick line) of survey catchability constants ( $q$ ) for the three resource survey series.

The first sensitivity run investigated the effect of including sex in the partition. As in the MPD run for this model, the selectivity ogives varied markedly between sexes; old females were always much less selected than old males in the surveys and the fishery (Figure 31). Estimated year class strengths were little different to those from the single sex model. Estimates of absolute biomass, both virgin and current, were more than double those of the single sex model, but stock status was virtually identical between the two models (Table 14, Figure 32). The increased biomass results in even smaller estimates of survey catchability constants than in the single sex model, i.e., $2 \%, 2 \%$ and $3 \%$ for the summer, autumn and spring survey series, respectively. These values appear unrealistic and are certainly not consistent with the prior. Exploitation rates (catch over vulnerable biomass) were estimated to have never been greater than $0.05 \mathrm{yr}^{-1}$.


Figure 31: Two sex model - Estimated median selectivity ogive (with $\mathbf{9 5 \%}$ credible intervals shown as dashed lines) for the trawl surveys and the trawl fishery.


Figure 32: Two sex model - Estimated median trajectories (with $\mathbf{9 5 \%}$ credible intervals shown as dashed lines) for absolute biomass and biomass as a percentage of $\mathbf{B}_{\mathbf{0}}$.

A sensitivity run investigating the influence of adding the CPUE series to the single sex model also resulted in higher estimates of absolute biomass (both virgin and current) and also in stock status, i.e., $68 \%$ of $\mathrm{B}_{0}$, compared to $52 \%$ for the single sex model (Table 14). The CPUE series is relatively flat (see Figure 21), so it has the effect of flattening the biomass trajectory (particularly the latter part), and hence, improving current stock status (Figure 33). A consequence of the overall flattening of the biomass trajectory is that the earlier year classes are estimated to be weaker and the later ones are stronger (Figure 34). The selectivity ogives are little different to those estimated in the single sex model. Exploitation rates (catch over vulnerable biomass) were estimated to have never been greater than $0.05 \mathrm{yr}^{-1}$. The estimates of trawl survey catchability constants (i.e., $3 \%, 2 \%$ and $4 \%$ for the summer, autumn and spring survey series, respectively) are very low, and probably unrealistic.


Figure 33: CPUE model - Estimated median trajectories (with 95\% credible intervals shown as dashed lines) for absolute biomass and biomass as a percentage of $\mathbf{B}_{0}$.


Figure 34: CPUE model - Estimated posterior distributions of year class strengths. The dashed horizontal line indicates the year class strength of one. Individual distributions are the marginal posteriors, with horizontal lines indicating the median.

The sensitivity run where an $M$-at-age relationship was estimated was the most pessimistic of the four models investigated. Estimates of absolute biomass were lower than in the single sex model, and current stock status was estimated to be $46 \% \mathrm{~B}_{0}$ (see Table 14). Recall that in this model the selectivity ogive for the trawl fishery was estimated using the logistic (rather than double-normal) parameterisation, with the underlying assumption being that the fishery comprehensively samples all the adult population. The fishery selectivity ogive was tightly defined, while the survey ogives had quite wide bounds (Figure 35). The survey ogives, although estimated using double normal parameterisation, was essentially logistic shaped. Age at full selectivity for the summer survey series was markedly lower than it was in the single sex model, but was essentially unchanged for the autumn survey series and the fishery (see Figure 26).

The estimated year class strengths were little different to those from the single sex model, and the estimated maximum exploitation rate (about $0.08 \mathrm{yr}^{-1}$ ) was only slightly higher. The estimates of trawl survey catchability constants (i.e., medians of $3 \%, 5 \%$ and $5 \%$ for the summer, autumn and spring survey series, respectively) are higher than in any other model run, but still not very consistent with the prior (Figure 36).

The age-based instantaneous natural mortality $(M)$ relationship estimated independent of sex had a minimum of about 0.1 at 6 years, rising to about 0.7 at 30 years, and a relatively narrow $95 \%$ credible interval across ages 6 to 23, i.e., the age classes most abundant in the surveys and the commercial catch (Figure 37). However, the estimation of $M$ will be confounded with the estimation of survey and fishery selectivities, so we cannot be confident that the true relationship has been determined here.


Figure 35: Estimate $M$ model — Estimated median selectivity ogive (with $\mathbf{9 5 \%}$ credible intervals shown as dashed lines) for the trawl surveys and the trawl fishery.


Figure 36: Estimate $M$ model - Estimated posterior distribution (thin line) and prior (thick line) of survey catchability constants ( $q$ ) for the three resource survey series.


Figure 37: Estimate M model - Estimated median age-based natural mortality (M) relationship (with $\mathbf{9 5 \%}$ credible intervals shown as dashed lines) for both sexes combined. The horizontal dashed line is at 0.19 , the value that has been used as a fixed value for $M$ in previous single sex assessments.

### 5.1 Biomass projections

Biomass projections from the four models investigated using MCMC estimation were made assuming future annual catches of $2300 t$ from 2012 to 2016. This catch is the average from 2005 to 2010, and is only slightly lower than the average annual catch since $1990(2500 \mathrm{t})$. There is no specific TACC for the Sub-Antarctic stock. The HAK 1 administrative stock (with a TACC of 3701 t ) includes all of the Sub-Antarctic, the western end of the Chatham Rise, and all around the North Island.

Year class strengths from 2008 onwards were selected randomly from the previously estimated year class strengths from 1974 to 2007.

Projections from all the models suggested that biomass will increase markedly to be between $80 \%$ and $100 \%$ of $B_{0}$ by 2016 (Table 15, Figure 38). The most pessimistic of these models suggests that there is little chance that biomass in 2016 will be lower than $51 \%$ of $\mathrm{B}_{0}$. However, these projections are quite uncertain as indicated by the rapidly spreading confidence intervals after 2011 (Figure 38).


Figure 38: Estimated median trajectories (with $95 \%$ credible intervals shown as dashed lines) for biomass as a percentage of $B_{0}$, projected to 2016 with future catches assumed to be $2300 t$ annually, for all models.

Table 15: Bayesian median and $95 \%$ credible intervals of projected $B_{2016}, B_{2016}$ as a percentage of $B_{0}$, and $\mathbf{B}_{2016} / \mathbf{B}_{2011}$ (\%) for all model runs, with future annual catch equal to the mean from 2005 to 2010.

| Model run | Future catch $(\mathrm{t})$ | $\mathrm{B}_{2016}$ |  | $\mathrm{~B}_{2016}\left(\% \mathrm{~B}_{0}\right)$ |  | $\mathrm{B}_{2016}\left(\mathrm{~B}_{2011}(\%)\right.$ |
| :--- | ---: | ---: | ---: | ---: | ---: | ---: |
|  |  |  |  |  |  |  |
| Single sex | 2300 | $74630(35390-147810)$ |  | $78.4(53.5-110.9)$ | $150(119-200)$ |  |
| Two sex | 2300 |  | $156470(81790-279230)$ |  | $80.2(59.5-108.3)$ | $146(118-192)$ |
| CPUE | 2300 |  | $136200(73690-250450)$ |  | $99.8(78.0-126.3)$ | $147(121-187)$ |
| Estimate $M$ | 2300 | $62080(27760-136220)$ |  | $78.8(51.2-111.6)$ | $169(132-229)$ |  |

### 5.2 Management biomass targets

Probabilities that current and projected biomass will drop below default management reference points from the Harvest Strategy Standard (i.e., target, $40 \% \mathrm{~B}_{0}$; soft limit, $20 \% \mathrm{~B}_{0}$; hard limit, $10 \% \mathrm{~B}_{0}$ ) are shown for all four models in Table 16. It appears extremely unlikely (i.e., less than 1\%) that $\mathrm{B}_{2016}$ will be lower than the soft limit of $20 \% \mathrm{~B}_{0}$, or even the target of $40 \% \mathrm{~B}_{0}$.

Table 16: Probabilities that current ( $B_{2011}$ ) and projected ( $B_{2016}$ ) biomass will be less than $\mathbf{4 0 \%}, \mathbf{2 0 \%}$ or $\mathbf{1 0 \%}$ of $\mathbf{B}_{\mathbf{0}}$. Projected biomass probabilities are presented assuming a future annual catch of $\mathbf{2 3 0 0} \mathbf{t}$.

| Model run | Biomass | Management reference points |  |  |
| :--- | :--- | ---: | ---: | ---: |
|  |  | $40 \% \mathrm{~B}_{0}$ | $20 \% \mathrm{~B}_{0}$ | $10 \% \mathrm{~B}_{0}$ |
| Single sex | $\mathrm{B}_{2011}$ | 0.036 | 0.000 | 0.000 |
|  | $\mathrm{~B}_{2016}, 2300 \mathrm{t}$ catch | 0.003 | 0.000 | 0.000 |
| Two sex | $\mathrm{B}_{2011}$ | 0.000 | 0.000 | 0.000 |
|  | $\mathrm{~B}_{2016}, 2300 \mathrm{t}$ catch | 0.000 | 0.000 | 0.000 |
| CPUE | $\mathrm{B}_{2011}$ | 0.000 | 0.000 | 0.000 |
|  | $\mathrm{~B}_{2016}, 2300 \mathrm{t}$ catch | 0.000 | 0.000 | 0.000 |
| Estimate $M$ | $\mathrm{~B}_{2011}$ | 0.150 | 0.000 | 0.000 |
|  | $\mathrm{~B}_{2016}, 2300 \mathrm{t}$ catch | 0.000 | 0.000 | 0.000 |

## 6. DISCUSSION

This document updates the assessment of the Sub-Antarctic hake last reported by Horn (2008). Past assessments of this stock have produced relatively uncertain results owing to the lack of strong contrast in the available series of relative abundance, i.e., two trawl survey series, and trawl CPUE (Dunn 2006, Horn 2008). This problem still persists in the current assessment, but at least it is indicative of a stock that has not been markedly depleted by exploitation.

An exploratory analysis of the length and sex data collected by observers from the commercial trawl fishery showed that the sex ratios in catches varied markedly between tows, trips, and years. It had also been shown previously that sex ratios varied by location across the Stewart-Snares shelf and Campbell Plateau region (Horn 2008). Similar variations by sex were found in assessments of the Chatham Rise (Horn \& Francis 2010) and west coast South Island (Horn 2011) hake stocks. For those stocks, a single sex model fitted the research biomass series and the commercial fishery proportion-atage data better as it did not have to try and deal with conflicting information about changes in sex ratios over time and area. A similar solution was adopted here for the Sub-Antarctic assessment. As for the Chatham Rise and WCSI assessments, an initial two sex model was markedly more optimistic than the single sex model. However, it is unlikely that sex alone provides sufficient 'logical' information to increase the Sub-Antarctic $\mathrm{B}_{0}$ by about $204 \%$, so the two sex model is considered less desirable than the single sex model at this stage.

The single sex model estimated that the Sub-Antarctic spawning stock is currently at about $52 \% \mathrm{~B}_{0}$, and that continued fishing at recent catch levels is likely to allow stock size to increase by about a half in the next five years. Sensitivity model runs including sex in the partition or allowing the estimation in the model of an age-based relationship for instantaneous natural mortality ( $M$ ) did not markedly alter the estimate of stock status. A sensitivity including a trawl CPUE series did indicate a much more optimistic current stock status. However, none of the model runs were indicative of current biomass being lower than $34 \%$ of $B_{0}$ (i.e., the most pessimistic $95 \%$ credible interval lower bound).

Estimated year class strengths often have quite wide $95 \%$ bounds, particularly at the start and the end of the estimated series. However, the median estimates suggest that variation in year class strength is
not great for this stock; only three of the estimates from 1974 to 2007 are outside the range $0.5-2$ (i.e., 1980 is much higher, and 1986 and 2002 are lower). A similar relatively low level of year class strength variation was estimated for the hake stocks on the Chatham Rise and WCSI (Horn \& Francis 2010, Horn 2011). However, it is not possible to tell whether the low variability in year class strengths is correct (i.e., the actual variability is low) or is a consequence of uninformative data (e.g., the yearclass signal in the at-age data could be poor either because these data are not representative of the catch, or because it is masked by year-to-year variation in selectivity).

The residuals of the MPD fits to the at-age data (Appendix B) show that in the summer trawl survey series observed numbers of 3 and 4 year old fish are, respectively, lower and higher than the estimated values. There are no apparent trends in the residuals for other age classes. The residuals for the fishery at-age data show some clear trends, particularly when sex is included in the partition (Appendix B, Figure B2). But even when sex is removed, residual trends are still apparent (Figure B1). The poor fits are generally associated with fish 6 years or younger, but with a change in the residual trend (from positive to negative) occurring about 2001. This may be indicative of some change in fishing selectivity. However, the assessment model described above used a single selectivity ogive for all years.

The structural assumptions of the model reported here are likely to lead to the Bayesian posteriors of stock status underestimating the true level of uncertainty. The projected stock status relies on adequate estimation of recent recruitment driven by year class strength estimates. The research survey proportions-at-age distributions are collected systematically over time or space, but the fishery proportions-at-age are not. Although the stratification used in the analyses of these data coupled with the removal of sex from the partition is believed to produce reasonable estimates of catch-at-age for the fishery, the projections of future stock status based on these data are likely to underestimate the true level of uncertainty.

Information about absolute stock status of hake in the Sub-Antarctic is probably quite weak owing to the low level of contrast in all of the relative abundance series. Estimates of stock size and projected stock status are strongly influenced by the shape of the selectivity ogives and the way in which $M$ is included in the model. When a constant $M$ is used, the model concluded that very young and very old fish comprise a cryptic biomass that is not selected by the fishing gear (both in the surveys and the fishery). Such a situation where large, old fish are essentially hidden from the fishing gear could occur in trawl surveys if these larger fish spend more time in midwater or on rough ground not able to be bottom trawled. However, the commercial fishery uses midwater trawls and so should have access to biomass 'hidden' from the resource surveys. When an $M$-at-age relationship is estimated (and the resource survey ogives are forced to be logistic), the model concluded that the old fish are not in the catch because they have died, producing a relatively high value of $M$ for old fish, and lower levels of absolute biomass (because the old fish are dead, rather than hidden). The marked improvement in fits to the age data when $M$ is estimated provides support for this model being better than the single sex model with a constant $M$.

Estimates of resource survey catchability ( $q$ s) are very low in all model runs (i.e., 2-5\%), particularly for the summer series, suggesting that the absolute catchability of the Sub-Antarctic trawl surveys is extremely low. It is not known if the catchability of the Sub-Antarctic trawl survey series is as low as estimated by the model, but hake are believed to be relatively more abundant over rough ground (that is likely to be avoided during a trawl survey), and it is known that hake tend to school off the bottom, particularly during their spring-summer spawning season, hence reducing their availability to the bottom trawl. Another plausible explanation for the low values is that there is little contrast in the biomass indices from the Sub-Antarctic trawl survey series, and that the model has little information on which to determine an appropriate "scale" of biomass estimates. Higher estimates of the relativity constant $q$ (although confounded with selectivity) would result in lower current and virgin biomass estimates. A survey $q$ of $10 \%$ was estimated using the same fishing gear to catch hake in a summer Chatham Rise survey (Horn \& Francis 2010). The assessment of that stock, fitting to a trawl biomass series with strong contrast, produced a very believable $\mathrm{B}_{0}$ estimate of 41000 t (Horn \& Francis 2010).

That value, and the $\mathrm{B}_{0}$ of 82000 t estimated for the west coast South Island stock (Horn 2011), by analogy may suggest that the Sub-Antarctic virgin biomass is more likely at the lower end of the range estimated from the four model describe above (i.e., $80000-90000 \mathrm{t}$ ).

The Working Group noted that there was a marked jump in estimated biomass between the 2007 and 2008 summer surveys (i.e., the December 2006 and 2007 summer surveys, see Figure 11), and requested that the possibility of a change in survey selectivity be investigated. An analysis similar to that completed for hoki from the same survey series (Francis 2009) indicated that there appeared to be marked changes in hake catchability between three adjacent survey pairs (i.e., 1991 and 1992, 2006 and 2007, 2008 and 2009) (Figure 39). However, the vertical scale in Figure $39(0.1-4)$ is much narrower than that for hoki ( $0.05-10$, see figure 26 in Francis (2009)), so the case for any maintained step change in survey selectivity is weak. It is possible that hake behaviour differs between years, perhaps related to environmental parameters, resulting in catchability variation.


Figure 39: Testing for step changes in hake catchability. Changes, between pairs of surveys in the subAntarctic summer series, in estimated numbers of hake in selected cohorts: A, surveys in consecutive years and B, surveys two years apart. Each plotted point indicates how the estimated number in a cohort changed between two surveys; the plotting symbol is the age of the cohort in the earlier survey. For example, the bottom right point in panel A shows that the estimated number in the cohort that was aged 14 in the December 2008 survey decreased by about $85 \%$ between the 2008 and 2009 surveys.

The assessment for Sub-Antarctic hake has been updated, and is indicative of a stock that has declined throughout the 1990 s, but is still very likely to be above $50 \%$ of $\mathrm{B}_{0}$. The decline is most likely a result of poor recruitment rather than fishing pressure. An annual catch in the range of 2000-3000 t appears very likely to be sustainable in the medium term at least. The stock is probably being reasonably well monitored by the November-December trawl survey series. While the stock status appears to be reasonably well defined, estimates of past and current absolute stock size are very uncertain owing to a lack of contrast in the relative abundance series. Because of the high uncertainty in estimates of absolute biomass, any yield estimates would also be very uncertain.

Future assessments of this and other hake stocks, consideration should be given to adopting the recommendations of Francis (2011) relating to data weighting. For the Sub-Antarctic assessment the following changes should be considered:

- use multinomial, rather than lognormal, errors for at-age and at-length data;
- use method TA1.8 from Francis (2011) for stage-2 weighting of at-age data;
- use a lowess smoother (or similar) to set a (total) c.v. for CPUE data; and
- avoid the use of conflicting biomass indices in the same model (so the trawl survey biomass indices would be removed from the final 'CPUE' model).


## 7. ACKNOWLEDGMENTS

I thank the many NIWA staff and Ministry of Fisheries observers who were involved in the collection of data at sea, and Chris Francis and members of the Middle Depth Fishery Assessment Working Group for providing useful comments and suggestions on the development of the assessment. Chris Francis produced Figure 39. Chris Francis and Kevin Sullivan provided valuable reviews of this document. The work was funded by the Ministry of Fisheries under project DEE201002HAKA.

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## APPENDIX A: RESOURCE SURVEY BIOMASS INDICES FOR HAKE IN HAK 1 AND HAK 4

Table A1: Biomass indices (t) and coefficients of variation (c.v.) for hake from resource surveys of the Sub-Antarctic. (These estimates assume that the areal availability, vertical availability, and vulnerability are equal to one.)

| Vessel | Date | Trip code | Depth |  | Biomass | c.v. | Reference |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Wesermünde | Mar-May 1979 |  | - | 1 | - | - | Kerstan \& Sahrhage 1980 |
| Wesermünde | Oct-Dec 1979 |  | - | 1 | - | - | Kerstan \& Sahrhage 1980 |
| Shinkai Maru | Mar-Apr 1982 | SHI8201 | 200-800 |  | 6045 | 0.15 | N.W. Bagley, NIWA, pers. comm. |
| Shinkai Maru | Oct-Nov 1983 | SHI8303 | 200-800 |  | 11282 | 0.22 | N.W. Bagley, NIWA, pers. comm. |
| Amaltal Explorer | Oct-Nov 1989 | AEX8902 | 200-800 |  | 2660 | 0.21 | Livingston \& Schofield 1993 |
| Amaltal Explorer | Jul-Aug 1990 | AEX9001 | 300-800 |  | 4343 | 0.19 | Hurst \& Schofield 1995 |
| Amaltal Explorer | Nov-Dec 1990 | AEX9002 | 300-800 |  | 2460 | 0.16 | N.W. Bagley, NIWA, pers. comm. |
| Tangaroa | Nov-Dec 1991 | TAN9105 | Reported | 2 | 5686 | 0.43 | Chatterton \& Hanchet 1994 |
|  |  |  | 300-800 | 3 | 5553 | 0.44 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1991 area | 4 | 5686 | 0.43 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1996 area | 5 | - | - |  |
| Tangaroa | Apr-May 1992 | TAN9204 | Reported | 2 | 5028 | 0.15 | Schofield \& Livingston 1994a |
|  |  |  | 300-800 | 3 | 5028 | 0.15 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1991 area | 4 | - | - |  |
|  |  |  | 1996 area | 5 | - | - |  |
| Tangaroa | Sep-Oct 1992 | TAN9209 | Reported |  | 3762 | 0.15 | Schofield \& Livingston 1994b |
|  |  |  | 300-800 | 3,7 | 3760 | 0.15 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1991 area | 4 | - | - |  |
|  |  |  | 1996 area | 5 | - | - |  |
| Tangaroa | Nov-Dec 1992 | TAN9211 | Reported | 2 | 1944 | 0.12 | Ingerson et al. 1995 |
|  |  |  | 300-800 | 3 | 1822 | 0.12 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1991 area | 4 | 1944 | 0.12 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1996 area | 5 | - | - |  |
| Tangaroa | May-Jun 1993 | TAN9304 ${ }^{6}$ | Reported | 2 | 3602 | 0.14 | Schofield \& Livingston 1994c |
|  |  |  | 300-800 | $\stackrel{ }{ }$ | 3221 | 0.14 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1991 area | 4 | - | - |  |
|  |  |  | 1996 area | 5 | - | - |  |
| Tangaroa | Nov-Dec 1993 | TAN9310 | Reported | 2 | 2572 | 0.12 | Ingerson \& Hanchet 1995 |
|  |  |  | 300-800 | 3 | 2286 | 0.12 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1991 area | 4 | 2567 | 0.12 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1996 area | 5 | - | - |  |

Table A1 ctd.

| Vessel | Date | Trip code | Depth |  | Biomass | c.v. | Reference |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Tangaroa | Mar-Apr 1996 | TAN9605 | Reported | 2 | 3946 | 0.16 | Colman 1996 |
|  |  |  | 300-800 | 3 | 2026 | 0.12 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1991 area | 4 | 2281 | 0.17 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1996 area | 5 | 2825 | 0.12 | O'Driscoll \& Bagley 2001 |
| Tangaroa | Apr-May 1998 | TAN9805 | Reported | 2 | 2554 | 0.18 | Bagley \& McMillan 1999 |
|  |  |  | 300-800 | 3 | 2554 | 0.18 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1991 area | 4 | 2643 | 0.17 | O'Driscoll \& Bagley 2001 |
|  |  |  | 1996 area | 5 | 3898 | 0.16 | O'Driscoll \& Bagley 2001 |
| Tangaroa | Nov-Dec 2000 | TAN0012 | 300-800 | 3 | 2194 | 0.17 | O'Driscoll et al. 2002 |
|  |  |  | 1991 area | 4 | 2657 | 0.16 | O'Driscoll et al. 2002 |
|  |  |  | 1996 area | 5 | 3103 | 0.14 | O'Driscoll et al. 2002 |
| Tangaroa | Nov-Dec 2001 | TAN0118 | 300-800 | 3 | 1831 | 0.24 | O'Driscoll \& Bagley 2003a |
|  |  |  | 1991 area | 4 | 2170 | 0.20 | O'Driscoll \& Bagley 2003a |
|  |  |  | 1996 area | 5 | 2360 | 0.19 | O'Driscoll \& Bagley 2003a |
| Tangaroa | Nov-Dec 2002 | TAN0219 | 300-800 | 3 | 1283 | 0.20 | O'Driscoll \& Bagley 2003b |
|  |  |  | 1991 area | 4 | 1777 | 0.16 | O'Driscoll \& Bagley 2003b |
|  |  |  | 1996 area | 5 | 2037 | 0.16 | O'Driscoll \& Bagley 2003b |
| Tangaroa | Nov-Dec 2003 | TAN0317 | 300-800 | 3 | 1335 | 0.24 | O'Driscoll \& Bagley 2004 |
|  |  |  | 1991 area | 4 | 1672 | 0.23 | O'Driscoll \& Bagley 2004 |
|  |  |  | 1996 area | 7 | 1898 | 0.21 | O'Driscoll \& Bagley 2004 |
| Tangaroa | Nov-Dec 2004 | TAN0414 | 300-800 | 3 | 1250 | 0.27 | O'Driscoll \& Bagley 2006a |
|  |  |  | 1991 area | 4 | 1694 | 0.21 | O'Driscoll \& Bagley 2006a |
|  |  |  | 1996 area | 7 | 1774 | 0.20 | O'Driscoll \& Bagley 2006a |
| Tangaroa | Nov-Dec 2005 | TAN0515 | 300-800 | 3 | 1133 | 0.20 | O'Driscoll \& Bagley 2006b |
|  |  |  | 1991 area | 4 | 1459 | 0.17 | O'Driscoll \& Bagley 2006b |
|  |  |  | 1996 area | 7 | 1624 | 0.17 | O'Driscoll \& Bagley 2006b |
| Tangaroa | Nov-Dec 2006 | TAN0617 | 300-800 | 3 | 998 | 0.22 | O'Driscoll \& Bagley 2008 |
|  |  |  | 1991 area | 4 | 1530 | 0.17 | O'Driscoll \& Bagley 2008 |
|  |  |  | 1996 area | 7 | 1588 | 0.16 | O'Driscoll \& Bagley 2008 |
| Tangaroa | Nov-Dec 2007 | TAN0714 | 300-800 | 3 | 2188 | 0.17 | Bagley et al. 2009 |
|  |  |  | 1991 area | 4 | 2470 | 0.15 | Bagley et al. 2009 |
|  |  |  | 1996 area | 7 | 2622 | 0.15 | Bagley et al. 2009 |
| Tangaroa | Nov-Dec 2008 | TAN0813 | 300-800 | 3 | 1074 | 0.23 | O'Driscoll \& Bagley 2009 |
|  |  |  | 1991 area | 4 | 2162 | 0.17 | O'Driscoll \& Bagley 2009 |
|  |  |  | 1996 area | 7 | 2355 | 0.16 | O'Driscoll \& Bagley 2009 |

## Table A1 ctd

| Vessel | Date | Trip code | Depth | Biomass | c.v. | Reference |
| :--- | ---: | ---: | ---: | ---: | ---: | :--- |
| Tangaroa | Nov-Dec 2009 | TAN0911 | $300-800$ | 3 | 992 | 0.22 |
|  |  |  | Bagley \& O'Driscoll 2012 |  |  |  |
|  |  |  | 1991 area | 4 | 1442 | 0.20 |
| Bagley \& O'Driscoll 2012 |  |  |  |  |  |  |
|  |  | 1996 area |  | 1602 | 0.18 | Bagley \& O'Driscoll 2012 |

1. Although surveys by Wesermünde were carried out in the Sub-Antarctic in 1979, biomass estimates for hake were not calculated
2. The depth range, biomass and c.v. in the original report.
3. The biomass and c.v. calculated from source records using the equivalent 1991 region, but excluding both the $800-1000 \mathrm{~m}$ strata in Puysegur region and the Bounty Platform strata.
4. The biomass and c.v. calculated from source records using the equivalent 1991 region, which includes the $800-1000 \mathrm{~m}$ strata in Puysegur region but excludes the Bounty Platform strata.
5. The biomass and c.v. calculated from source records using the equivalent 1996 region, which includes the $800-1000 \mathrm{~m}$ strata in Puysegur region but excludes the Bounty Platform strata. (The 1996 region added additional $800-1000 \mathrm{~m}$ strata to the north and to the south of the Sub-Antarctic to the 1991 region).
6. Doorspread data not recorded for this survey. Analysis of source data with average of all other survey doorspread estimates resulted in a new estimate of biomass
7. The biomass and c.v. calculated from source records using the equivalent 1996 region, which includes the $800-1000 \mathrm{~m}$ strata in Puysegur region but excludes the Bounty Platform strata. (The 1996 region added additional $800-1000 \mathrm{~m}$ strata to the north and to the south of the Sub-Antarctic to the 1991 region). However, in 2003 , stratum 26 (the most southern $800-1000 \mathrm{~m}$ strata) was not surveyed. In previous years this stratum yielded either a very low or zero hake biomass. The yield in 2003 from stratum 26 was assumed to be zero.

Table A2: Biomass indices (t) and coefficients of variation (c.v.) for hake from resource surveys of the Chatham Rise. (These estimates assume that the areal availability, vertical availability, and vulnerability are equal to one.)

| Vessel | Date | Trip code | Depth |  | Biomass | c.v. | Reference |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Wesermünde | Mar-May 1979 |  | - | 1 | - | - | Kerstan \& Sahrhage 1980 |
| Wesermünde | Oct Dec 1979 |  | - | I | - | - | Kerstan \& Sahrhage 1980 |
| Shinkai Maru | Mar 1983 | SHI8301 | 200-800 |  | 11327 | 0.12 | N.W. Bagley, NIWA, pers. comm. |
| Shinkai Maru | Nov-Dec 1983 | SHI8304 | 200-800 | 2 | 8160 | 0.12 | N.W. Bagley, NIWA, pers. comm. |
| Shinkai Maru | Jul 1986 | SHI8602 | 200-800 |  | 7630 | 0.13 | N.W. Bagley, NIWA, pers. comm. |
| Amaltal Explorer | Nov-Dec 1989 | AEX8903 | 200-800 |  | 3576 | 0.19 | N.W. Bagley, NIWA, pers. comm. |
| Tangaroa | Jan 1992 | TAN9106 | 200-800 |  | 4180 | 0.15 | Horn 1994a |
| Tangaroa | Jan 1993 | TAN9212 | 200-800 |  | 2950 | 0.17 | Horn 1994b |
| Tangaroa | Jan 1994 | TAN9401 | 200-800 |  | 3353 | 0.10 | Schofield \& Horn 1994 |
| Tangaroa | Jan 1995 | TAN9501 | 200-800 |  | 3303 | 0.23 | Schofield \& Livingston 1995 |
| Tangaroa | Jan 1996 | TAN9601 | 200-800 |  | 2457 | 0.13 | Schofield \& Livingston 1996 |
| Tangaroa | Jan 1997 | TAN9701 | 200-800 |  | 2811 | 0.17 | Schofield \& Livingston 1997 |
| Tangaroa | Jan 1998 | TAN9801 | 200-800 |  | 2873 | 0.18 | Bagley \& Hurst 1998 |
| Tangaroa | Jan 1999 | TAN9901 | 200-800 |  | 2302 | 0.12 | Bagley \& Livingston 2000 |
| Tangaroa | Jan 2000 | TAN0001 | 200-800 |  | 2090 | 0.09 | Stevens et al. 2001 |
|  |  |  | 200-1000 |  | 2152 | 0.09 | Stevens et al. 2001 |
| Tangaroa | Jan 2001 | TAN0101 | 200-800 |  | 1589 | 0.13 | Stevens et al. 2002 |
| Tangaroa | Jan 2002 | TAN0201 | 200-800 |  | 1567 | 0.15 | Stevens \& Livingston 2003 |
|  |  |  | 200-1000 |  | 1905 | 0.13 | Stevens \& Livingston 2003 |
| Tangaroa | Jan 2003 | TAN0301 | 200-800 |  | 888 | 0.16 | Livingston et al. 2004 |
| Tangaroa | Jan 2004 | TAN0401 | 200-800 |  | 1547 | 0.17 | Livingston \& Stevens 2005 |
| Tangaroa | Jan 2005 | TAN0501 | 200-800 |  | 1048 | 0.18 | Stevens \& O'Driscoll 2006 |
| Tangaroa | Jan 2006 | TAN0601 | 200-800 |  | 1384 | 0.19 | Stevens \& O'Driscoll 2007 |
| Tangaroa | Jan 2007 | TAN0701 | 200-800 |  | 1824 | 0.12 | Stevens et al. 2008 |
|  |  |  | 200-1000 |  | 1976 | 0.12 | Stevens et al. 2008 |
| Tangaroa | Jan 2008 | TAN0801 | 200-800 |  | 1257 | 0.13 | Stevens et al. 2009a |
|  |  |  | 200-1000 |  | 1323 | 0.13 | Stevens et al. 2009a |
| Tangaroa | Jan 2009 | TAN0901 | 200-800 |  | 2419 | 0.21 | Stevens et al. 2009b |
| Tangaroa | Jan 2010 | TAN1001 | 200-800 |  | 1701 | 0.25 | Stevens et al. 2011 |
|  |  |  | 200-1300 |  | 1862 | 0.25 | Stevens et al. 2011 |
| Tangaroa | Jan 2011 | TAN1101 | 200-800 |  | 1099 | 0.15 | Stevens et al. 2012 |
|  |  |  | 200-1300 |  | 1201 | 0.14 | Stevens et al. 2012 |

1. Although surveys by Wesermünde were carried out on the Chatham Rise in 1979, biomass estimates for hake were not calculated.
2. East of $176^{\circ} \mathrm{E}$ only.

## APPENDIX B: RESIDUALS OF MPD MODEL FITS TO THE CATCH-AT-AGE DATA



Trawl survey (autumn)


Figure B1: Single sex model - MPD residual values for the proportions-at-age data for the SubAntarctic resource survey series and fishery. Symbol area is proportional to the absolute value of the residual, with filled circles indicating positive residuals and open circles indicating negative residuals.


Figure B2: Two sex model - MPD residual values for the proportions-at-age data for the Sub-Antarctic resource survey series and fishery. Symbol area is proportional to the absolute value of the residual, with filled circles indicating positive residuals and open circles indicating negative residuals.


Figure B3: Estimate $M$ model - MPD residual values for the proportions-at-age data for the SubAntarctic resource survey series and fishery. Symbol area is proportional to the absolute value of the residual, with filled circles indicating positive residuals and open circles indicating negative residuals.

