Age-based demography of humpback grouper *Cromileptes altivelis*: implications for fisheries management and conservation

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ABSTRACT: The humpback grouper *Cromileptes altivelis* is listed as Vulnerable on the IUCN Red List because of concerns about unsustainable levels of harvest. Quantitative estimates of life history parameters are scarce, limiting formal assessment of long-term harvest rates and conservation risk and the design of robust management measures. We provide the first estimates of age-based population parameters for *C. altivelis*, using samples from the Great Barrier Reef (GBR) and Torres Strait, Australia. Population parameters did not differ significantly between regions. The maximum observed age was 19 yr, the von Bertalanffy growth parameters were $K = 0.30$ yr$^{-1}$, $L_\infty = 597$ mm, and total mortality rate, estimated from an age-based catch curve, was 0.26 yr$^{-1}$. Preliminary estimates of natural mortality for the GBR, based on samples from reefs closed to fishing, were 0.23 yr$^{-1}$ (Hoenig regression) and 0.26 yr$^{-1}$ (catch curve), suggesting recent fishing mortality was low. *C. altivelis* was found to be a protogynous hermaphrodite with 50% of individuals being male at length 547 mm and age 9.6 yr. Peak spawning was between October and January. These results contrast with previous perceptions of life history traits of this species and suggest that *C. altivelis* may not be as vulnerable to managed fishing pressure as previously thought. Notwithstanding this, there remains considerable uncertainty in unfished abundance and average recruitment levels for *C. altivelis* populations. We consider a combination of management strategies including size limits, effort controls and no-take areas that are likely to be most effective in minimising the conservation risk for *C. altivelis* populations and provide sustainable yields across the species’ range.

KEY WORDS: Humpback grouper · *Cromileptes altivelis* · Conservation · Fisheries management · Age-based demography · Great Barrier Reef · Torres Strait · Coral reef

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INTRODUCTION

The humpback grouper *Cromileptes altivelis* is an iconic species of coral reef fish distributed throughout tropical East Asia straddling the western margin of the Pacific and eastern margin of the Indian Oceans (Heemstra & Randall 1999). Heemstra & Randall (1999) classified *C. altivelis* as a unique monotypic genus within the family Serranidae, but recent work by Craig & Hastings (2007) has demonstrated that *C. altivelis*
species within the *Epinephelus* genus in a clade with other large species including *E. lanceolatus*, *E. fuscoguttatus* and *E. polyphedakion*. *C. altivelis* is distinctive because of its unusual high dorsal profile, although this feature is also evident in the other serranid genera *Alphistes* and *Dermatolepis*. Juvenile *C. altivelis* are prize targets for the aquarium trade throughout the species’ distribution, and adults are among the most valued products in the Asian live reef fish trade (Sadovy et al. 2008). The current, largely unmanaged, harvests of *C. altivelis* in many south-east Asian and south Pacific waters are generally considered to be unsustainable and pose a serious conservation risk to the species and long-term yields (Sadovy et al. 2008). This concern has been supported by studies documenting large targeted catches of this and other reef species and the infrequent observations of the species in Indonesia and Malaysia, where the species was previously considerably more abundant (Oakley et al. 1999, Sadovy et al. 2003, Pet-Soede et al. 2004). *C. altivelis* was listed as Vulnerable on the IUCN Red List in 2007 (Sadovy et al. 2008) in response to concerns over declines in many populations, the limited understanding of their life history traits, and high exploitation rates in response to demand in the live fish and aquarium trade.

There are no published studies that provide age-based estimates of life-history parameters required to evaluate the likely performance of alternative harvest and conservation strategies for *Cromileptes altivelis*. The limited information available for wild populations suggests that *C. altivelis* is likely to be naturally low in abundance (Sabetian 2003, Pears 2005), slow-growing and long-lived (Oakley et al. 1999, Morris et al. 2000, Sabetian 2003, Teh et al. 2005) relative to common perch species. *C. altivelis* represents a relatively low proportion of catches in commercial reef line fisheries relative to other species (Samoilys et al. 2002, Williams et al. 2008a) despite growing demand and prices as high as $US130 kg⁻¹ in select markets (Sadovy et al. 2003). The combination of low abundance and high value has driven research on aquaculture and hatchery rearing of this species (Mous 2000, Sugama et al. 2003) in an attempt to capitalise on the unmet demand for *C. altivelis* in live reef fish markets. Observations from the aquaculture industry suggest that *C. altivelis* is a protogynous hermaphrodite (Liu & Sadovy de Mitcheson 2009), but this has not been confirmed through histological analysis.

*Cromileptes altivelis* occurs in tropical waters across northern Australia, including the Great Barrier Reef (GBR), the coral reefs of the Torres Strait (TS) and the tropical fringing reefs of the Northern Territory and Western Australia (Heemstra & Randall 1999). The species was regularly taken on the GBR by a multi-

**MATERIALS AND METHODS**

**Sample collection. GBR:** In total, 225 *Cromileptes altivelis* were sampled from 6 reefs at each of 4 latitudes (areas) along the GBR (Lizard Island, Townsville, Mackay and Storm Cay) over 11 yr from 1995 to 2005 during the Effects of Line Fishing (ELF) Experiment catch surveys (Mapstone et al. 2004; Fig. 1). All samples were collected using standardised commercial line fishing gear. Four reefs within each area were closed to fishing for up to 12 yr prior to sampling, while the other 2 reefs had always been open to fishing but there were too few samples collected from open reefs to compare life history parameters between open (n = 57) and closed (n = 167) reefs. Accordingly, it was necessary to pool the data across open and closed reefs and all 4 areas for most analyses, meaning that parameter estimates are likely to be influenced to some
degree by exploitation prior to _C. altivelis_ becoming a no-take species on the GBR and by any latitudinal variation over the southern half of the GBR.

An additional 22 _Cromileptes altivelis_ samples from the GBR were provided by commercial and recreational fishers and recreational spear fishers. These samples were not used for estimates of age and length frequency distributions or to estimate mortality due to the potential biases associated with different fishing gear selectivities. Fork length (FL) of all samples was measured to the nearest mm, and whole wet weight (W) was measured to the nearest 10 g. Sagittal otoliths were removed, cleaned of any residual material and stored dry. Gonads were removed and stored in 10% phosphate buffered formalin.

**TS:** Fifty-seven _Cromileptes altivelis_ were collected from reefs open to fishing in the TS (Fig. 2) during observer surveys on board indigenous and non-indigenous commercial vessels. Five 2 wk observer surveys were completed with the non-indigenous commercial fishers in 2004, while 3 observer surveys were completed with the indigenous commercial fishers in 2005 and 2006 (Williams et al. 2008a). An additional 145 _C. altivelis_ were obtained from reefs open to fishing from March 2004 to November 2005 by monthly purchases from indigenous and non-indigenous commercial operators in the TS. Whole fish were purchased whenever possible to ensure gonad samples were collected, but reproductive information was not available from all samples as some fish had been gutted prior to purchase. FL was measured to the nearest mm, and otoliths and gonads were removed from each individual. Samples were not available from all months in each year due to the opportunistic nature of sample collection.

**Sample processing.** Otoliths were sectioned and read using the same methods applied to other serranids from the GBR and TS (see Ferreira & Russ 1994, Pears et al. 2006, Williams et al. 2008b). Opaque increments were assumed to be formed annually based on the validation of this assumption for a wide range of other serranids (e.g. Ferreira & Russ 1994, Mosse 2001, Pears et al. 2006). All sectioned otoliths were read twice by a single experienced reader, and the count of opaque increments was accepted as the fish’s age in years if the 2 readings were the same. A third count was made by the same reader if the first 2 counts differed, and a match between the third and either of the first 2 counts was accepted as the fish’s age. The median count was assigned as the final age estimate for those otoliths (18%) from which there was no agreement among 3 counts.

Gonads from fish collected from the GBR were removed and preserved at sea in 10% phosphate-buffered formalin. Gonads from fish collected by observers in the TS were frozen immediately after removal and transported to the laboratory to be preserved. Gonads from fish purchased from the TS were removed from frozen fish and preserved in the labora-
tory. All gonads were preserved in 10% phosphate buffered formalin. Each pair of gonads was dried of excess fixative after fixation, and gonad weight was measured to the nearest 0.1 g. Total gonad weight was estimated by multiplying the mass of a single complete lobe by 2 where 1 gonad lobe was damaged during processing. It was assumed that this provided a reasonably accurate measure of gonad weight because the 2 gonad lobes are generally equal in size for other serranid species (Adams 2003).

Histological sections were taken from all gonads following the procedures outlined by Adams (2003). The stage of ovary development was based on the most advanced non-atretic cell type present (West 1990). Additional features used in histological staging included the presence of brown bodies, atretic oocytes, vascularisation, and the relative thickness of the gonad wall, all of which may indicate prior spawning (Sadovy & Shapiro 1987). Ovaries and testes were classified into reproductive developmental stages adapted from Ferreira (1995) and Adams (2003). Females were classified into 4 stages: Immature, Resting, Ripe, and Running Ripe (hydrated). Males were classified into 3 stages: Resting, Ripe, and Spent. Transitional (fish in the process of changing sex from female to male) individuals were also identified using the criteria described by Sadovy & Shapiro (1987).

**Demographic analyses.** The relationship between FL and W was described using a power function of the form

\[ W = a \times FL^b \]  

where \( a \) is the coefficient of the power function and \( b \) is the exponent. This relationship was compared between populations from the GBR and TS using analysis of covariance (ANCOVA) with ln FL as the covariate of ln W.

Length and age frequency distributions of *Cromileptes altivelis* were compared between the GBR and TS using multi-dimensional contingency tables using the same range of length (340 to 660 mm) and age (3 to 14 yr) classes for each region.

The von Bertalanffy growth function (VBGF) was fitted by nonlinear least-squares regression of FL on age of *Cromileptes altivelis* from the GBR and TS. The form of the VBGF used to model length-at-age data was

\[ L_t = L_\infty \left(1 - e^{-K(t - t_0)}\right) \]  

where \( L_t \) is the length at age \( t \), \( L_\infty \) is the mean asymptotic FL, \( K \) is the growth coefficient or rate at which \( L_\infty \) is approached, \( t \) is the age of the fish and \( t_0 \) is the age at which the fish have a theoretical length of 0. This form of the VBGF is most commonly used to describe the growth of teleosts given their very small size at birth compared to other taxa (Haddon 2001). Alternative parameterisations of the VBGF were tested but revealed very small differences in parameter estimates. A constrained VBGF was also fitted to the length-at-age data, where \( t_0 \) was constrained to 0, as a sensitivity analysis, given the lack of smaller, younger fish in the samples.

VBGFs were compared between the GBR and TS using likelihood ratio tests (Kimura 1980). A common range of age classes (3 to 15 yr) was used in each analysis to assure validity of the comparisons (Haddon 2001).

Age-based catch curves (Ricker 1975) were used to estimate the instantaneous rate of total mortality (\( Z \)) for *Cromileptes altivelis* from the GBR and TS. Catch curves are generated by fitting a linear regression to the natural logarithm of the number of individuals of each age class against fish age. The slope of this regression is an estimate of the rate of annual mortality. The use of catch curves is subject to a number of assumptions, including (1) the population is sampled representatively; (2) the regression only includes fish fully recruited to the sampling gear; (3) mortality is constant among age classes and over time; and (4) recruitment is the same each year. Regressions were fitted from age 6 yr (assumed full selectivity) through to the oldest age class that was preceded by no more than 2 consecutive 0 frequencies. Estimates were compared between the GBR and TS using ANCOVA with age as the covariate and data limited to a common age range of 6 to 16 yr. A catch curve was also fitted to age frequency data from reefs closed to fishing on the GBR to provide an estimate of natural mortality (\( M \)) for *C. altivelis* assuming that these reefs had not been subjected to significant levels of infringements. An additional estimate of \( M \) was obtained by fitting the linear equation ln(\( Z \)) = 1.44 – 0.982 × ln(\( t_{\text{max}} \)) developed by Hoenig (1983) to the maximum age observed, \( t_{\text{max}} \) (19 yr), obtained from reefs closed to fishing, and assuming the estimate of \( Z \) approximated \( M \).

**Reproductive biology.** The spawning season for *Cromileptes altivelis* from the GBR and TS was examined using samples collected in different months throughout the year. A gonadosomatic index (GSI = gonad weight/W × 100) was calculated for each sample, providing a relative measure of reproductive stage. The proportion of samples in each mature female and male reproductive stage in each month was plotted for each region to examine the ovarian and testicular development patterns throughout the year and the degree of spawning activity occurring in each month in each region.

A logistic function was fitted to the proportion of males in each length and age class to estimate the length and age at which *Cromileptes altivelis* changes sex. The length and age at sex change was estimated for the GBR and TS using the logistic equation.
\[ P_s = \left(1 + e^{-\ln(\frac{1}{2}) \left( s_{50} - s_{95} \right)} \right)^{-1} \]  

where \( P_s \) is the proportion of males in age or 10 mm length class \( s \), and \( s_{50} \) and \( s_{95} \) are the age or length at which 50 and 95% of the population are males, respectively. Transitional individuals were not included in these analyses, but their presence or absence in each length and age class provided an additional estimate of the length and age range over which sex change occurred. Likelihood ratio tests were used to test for differences in the length and age at sex change between the GBR and TS using a common age or length range for each region to assure validity of the comparisons (Haddon 2001).

The sex assigned from histology for samples collected by researchers during observer surveys in the TS or ELF catch surveys on the GBR was used to calculate the operational sex ratio (mature females to mature males) for *Cromileptes altivelis* from the GBR and TS. Sex ratios were compared between the GBR and TS using a chi-squared contingency test and compared to an expected ratio of 1:1 with a chi-squared goodness of fit test.

**RESULTS**

**Length and age distributions**

The modal length of *Cromileptes altivelis* was the same (500 mm) for the GBR and TS, and a multi-dimensional contingency table analysis indicated that the length distributions did not vary significantly between regions (\( \chi^2 = 23.75, df = 15, p = 0.07 \)). Therefore, length data were pooled across regions to produce a length frequency distribution for the combined data (Fig. 3). The modal age of *C. altivelis* differed between regions and was 6 yr for the GBR and 9 yr for TS. The maximum age from the GBR (19 yr) was slightly older than from TS (16 yr). Despite different modes and maximum age, multi-dimensional contingency table analyses indicated that the age distributions did not differ significantly between the regions (\( \chi^2 = 11.61, df = 11, p = 0.39 \)). Age data were pooled across regions to produce an age frequency distribution for the combined data (Fig. 3).

**Growth and mortality**

The relationship between FL and \( W \) was approximately isometric for *Cromileptes altivelis* from both regions (Table 1) and did not differ significantly between them (ANCOVA, \( F = 0.17, df = 1, p = 0.68 \)), so data were pooled to provide an estimate of the FL–W relationship (Fig. 4, Table 1).

There was substantial variation in the length-at-age of *Cromileptes altivelis* within the GBR and TS. The parameter estimates for the VBGF indicated a greater average maximum size \( (L_\infty) \) and lower growth coefficient \( (K) \) for the GBR than the TS (Table 1) but the likelihood ratio tests indicated that these differences were not significant (\( \chi^2 = 4.81, df = 3, p = 0.19 \)). The unconstrained growth curve for the pooled data across regions was relatively ‘flat’, as few fish smaller than 400 mm FL were sampled (Fig. 5). Therefore, VBGF parameter estimates from the unconstrained fit are likely to be biased, with an overestimate of \( L_\infty \) and underestimate of \( K \) and \( t_0 \). The constrained fit of the VBGF resulted in a lower \( L_\infty \) and higher \( K \) and \( t_0 \) than the unconstrained fit (Fig. 5). It is important to note, however, that the parameter estimates from the constrained fit may also be
biased, resulting in an underestimate of $L_\infty$ and overestimate of $K$. The average lengths at age for 10 wk, 1 and 2 yr old fish, based on data from Liu & Sadovy de Mitcheson (2009), are plotted in Fig. 5. These data demonstrate the likely better fit of the constrained VBGF to early growth of $C. altivelis$.

The estimate of total mortality ($Z$) for $Cromileptes altivelis$ was similar between the TS (0.25 yr$^{-1}$) and the GBR (0.27 yr$^{-1}$; Table 1) and did not differ significantly between the regions (ANOVA, $F = 0.42$, df = 1, $p = 0.68$), so data were pooled across regions to provide an overall estimate of $Z$ (0.26 yr$^{-1}$). The estimate of $M$

### Table 1. $Cromileptes altivelis$. Parameter estimates (±SE) for the length–weight relationship, constrained ($t_0 = 0$) and unconstrained von Bertalanffy growth function (VBGF) and rates of total ($Z$) and natural ($M$) mortality (yr$^{-1}$; estimated from catch curves and the method of Hoenig 1983) from the Great Barrier Reef, Torres Strait and pooled across regions. $a$ and $b$: parameters of the allometric relationship ($W = a \times FL^b$) between fork length ($FL$) and weight ($W$); $L_\infty$: mean asymptotic $FL$ (mm); $K$: growth coefficient (yr$^{-1}$); $t_0$: theoretical age (yr) at length 0 for the VBGF; $n$: number of samples used to estimate each parameter; na: not available.

<table>
<thead>
<tr>
<th>Function</th>
<th>Parameter</th>
<th>Great Barrier Reef</th>
<th>Torres Strait</th>
<th>Pooled</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length–weight relationship</td>
<td>$a$ ($\times 10^{-9}$)</td>
<td>2.85 (1.27)</td>
<td>2.63 (3.22)</td>
<td>3.48 (1.50)</td>
</tr>
<tr>
<td></td>
<td>$b$</td>
<td>3.27 (0.07)</td>
<td>3.29 (0.19)</td>
<td>3.24 (0.07)</td>
</tr>
<tr>
<td></td>
<td>$n$</td>
<td>127</td>
<td>43</td>
<td>170</td>
</tr>
<tr>
<td>VBGF, unconstrained</td>
<td>$L_\infty$</td>
<td>735 (114)</td>
<td>625 (52)</td>
<td>696 (72)</td>
</tr>
<tr>
<td></td>
<td>$K$</td>
<td>0.08 (0.05)</td>
<td>0.13 (0.07)</td>
<td>0.09 (0.04)</td>
</tr>
<tr>
<td></td>
<td>$t_0$</td>
<td>-6.96 (3.97)</td>
<td>-6.79 (4.19)</td>
<td>-8.00 (3.39)</td>
</tr>
<tr>
<td></td>
<td>$n$</td>
<td>180</td>
<td>196</td>
<td>376</td>
</tr>
<tr>
<td>VBGF, constrained ($t_0 = 0$)</td>
<td>$L_\infty$</td>
<td>597 (11)</td>
<td>558 (7)</td>
<td>575 (6)</td>
</tr>
<tr>
<td></td>
<td>$K$</td>
<td>0.30 (0.02)</td>
<td>0.40 (0.03)</td>
<td>0.35 (0.02)</td>
</tr>
<tr>
<td></td>
<td>$t_0$</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total mortality, catch curve</td>
<td>$Z$</td>
<td>0.27 (0.03)</td>
<td>0.25 (0.04)</td>
<td>0.26 (0.03)</td>
</tr>
<tr>
<td></td>
<td>$n$</td>
<td>152</td>
<td>160</td>
<td>310</td>
</tr>
<tr>
<td>Natural mortality, catch curve</td>
<td>$M$</td>
<td>0.26 (0.02)</td>
<td>na</td>
<td>na</td>
</tr>
<tr>
<td></td>
<td>$n$</td>
<td>113</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Natural mortality, Hoenig</td>
<td>$M$</td>
<td>0.23</td>
<td>na</td>
<td>na</td>
</tr>
</tbody>
</table>

**Reproductive biology**

The monthly trends in GSI and mature ovarian and testicular stages indicated that the peak spawning months differed between the GBR and TS. Mean
monthly GSI values for mature females were highest in December–January on the GBR and October–November in the TS (Fig. 6). The mean monthly GSI values for males were more variable than for females but were generally highest from October to March in both regions (Fig. 7).

The monthly trend in the frequency of mature ovarian stages (ovaries in active vitellogenesis) also suggested a different temporal pattern in spawning activity for the GBR and TS (Fig. 6). The majority of spawning activity, indicated by the presence of ripe and running ripe females, occurred in December–January on the GBR, but was protracted across the months October to January in the TS (Fig. 6). The monthly trend in the frequency of testicular stages indicated that males in both regions were capable of spawning over an extended period on either side of the spawning season indicated by the presence of active females (Fig. 7).

There was a large overlap in the length and age of male and female *Cromileptes altivelis* from the GBR and TS, although females generally dominated the smaller length classes and younger age classes and males dominated the larger length classes and older age classes (Fig. 8). Transitional individuals were relatively common on the GBR, but rare in TS. No immature individuals were sampled from either region.
The estimated length and age at which 50 and 95% of individuals changed sex was very similar between the GBR and TS (Table 2). Likelihood ratio tests indicated that the pattern in the length and age at sex change was similar between the GBR and TS (length: $\chi^2 = 0.25, df = 2, p = 0.88$; age: $\chi^2 = 0.26, df = 2, p = 0.88$). Sex ratio for Cromileptes altivelis also did not differ significantly between regions (contingency table, $\chi^2 = 0.42, df = 1, p = 0.52$). Data were pooled across regions, and the sex ratio for the combined data was significantly female-biased (Table 2, goodness of fit test, $\chi^2 = 6.19, df = 1, p = 0.01$).

### DISCUSSION

Our study provides the first age-based estimates of demographic parameters for Cromileptes altivelis and provides a foundation for formal assessments of the effects of fishing and evaluation of the likely performance of alternative management and conservation efforts for this species. The biological characteristics of C. altivelis (moderate longevity, relatively fast initial growth, moderate total mortality rates and early maturity) suggest that this species may not be as vulnerable to managed fishing pressure as previously thought. The apparently low natural levels of abundance of C. altivelis (Sabetian 2003, Pears 2005), however, suggest that reproductive output, recruitment processes, or a combination of both, may be limiting factors for the productivity of C. altivelis populations. In the absence of robust estimates of recruitment and unexploited post-settlement abundance and evidence of rapid depletion of C. altivelis stocks, it would seem prudent to maintain/implement appropriately conservative management measures.

Serranids, including Cromileptes altivelis, have generally been thought to be long-lived fishes (Manooch 1987, Sadovy et al. 2003) but recent studies have demonstrated that longevity within serranids varies greatly from $< 10$ yr to $> 40$ yr (e.g. Ferreira & Russ

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**Table 2.** Cromileptes altivelis. Peak spawning months, sex ratios and parameter estimates (±SE) of age and length-specific sex change from the Great Barrier Reef, Torres Strait and pooled across regions. $L_{50}$ and $L_{95}$: estimated length (mm FL) at which 50 and 95% of the population are males, respectively; $A_{50}$ and $A_{95}$: estimated age (yr) at which 50 and 95% of the population are males, respectively; $n$: number of samples used to estimate each parameter.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Great Barrier Reef</th>
<th>Torres Strait</th>
<th>Pooled</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Dec–Jan</td>
<td>Oct–Jan</td>
<td>Oct–Jan</td>
</tr>
<tr>
<td>Sex ratio (F:M)</td>
<td>1.22:1</td>
<td>1.41:1</td>
<td>1.32:1</td>
</tr>
<tr>
<td>$n$</td>
<td>151</td>
<td>176</td>
<td>327</td>
</tr>
<tr>
<td><strong>Length-specific sex change</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$L_{50}$</td>
<td>549 (8)</td>
<td>545 (8)</td>
<td>547 (8)</td>
</tr>
<tr>
<td>$L_{95}$</td>
<td>715 (24)</td>
<td>715 (27)</td>
<td>713 (22)</td>
</tr>
<tr>
<td>$n$</td>
<td>151</td>
<td>176</td>
<td>327</td>
</tr>
<tr>
<td><strong>Age-specific sex change</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$A_{50}$</td>
<td>9.7 (0.7)</td>
<td>9.5 (0.7)</td>
<td>9.6 (0.5)</td>
</tr>
<tr>
<td>$A_{95}$</td>
<td>17.9 (2.1)</td>
<td>18.0 (2.2)</td>
<td>18.6 (1.6)</td>
</tr>
<tr>
<td>$n$</td>
<td>131</td>
<td>176</td>
<td>307</td>
</tr>
</tbody>
</table>

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**Fig. 8.** Cromileptes altivelis. Proportion of male, transitional and mature females in each (a) size and (b) age class and estimated logistic function (line) fitted to the proportion of males from the Great Barrier Reef and Torres Strait. Parameters of the logistic functions are given in Table 2.
1994, Mosse 2001, Grandcourt 2002, Pears 2005, Pears et al. 2006). *C. altivelis* from the GBR and TS is a moderately long-lived serranid with a life span approaching 20 yr. This is a shorter lifespan than that of other medium to large serranids such as *Epinephelus polyphekadion* and *E. fuscoguttatus* that live around 40 yr (Pears 2005, Pears et al. 2006) and is counter to previous perceptions that this species is particularly long-lived (e.g. Morris et al. 2000). The longevity of *C. altivelis* is similar to the coral trout species *Plectropomus leopardus*, *P. areolatus* and *P. laevis*, the main reef fishes targeted by many fisheries in the Indo-Pacific. These species have a reported longevity between 14 and 18 yr (Ferreira & Russ 1994, Williams et al. 2008b, Heupel et al. in press).

The growth pattern for *Cromileptes altivelis* is similar to many other serranids that are characterised by large variation in size-at-age and reach about 80% of their average maximum size (*L∞*) within the first quarter of their lifespan (e.g. Ferreira & Russ 1994, Mosse 2001, Pears 2005, Pears et al. 2006). Consequently, the relationship between age and size is weak, with the exception of the first or 2 age-classes. As a result, length data alone are of limited value for estimating life history parameters and assessing the state of the stock and potential effects of different harvest strategies. The distribution of length data, however, indicates that fish <430 mm TL were not fully selected by the fishing gear used in this study. Hence, the estimated growth parameters are likely to be biased, relative to the ‘true’ pattern of growth, due to the lack of small fish and likely sampling bias towards faster-growing individuals in the younger age classes. The constrained and unconstrained fits of the VBGF to the size-at-age data provide an indication of the likely extremes of the growth pattern for *C. altivelis*, although the true growth pattern most likely lies somewhere between the 2 fits. Length-at-age data from hatchery reared *C. altivelis* reported by Liu & Sadovy de Mitcheson (2009) indicate the constrained VBGF is more likely to reflect the early pattern of growth of *C. altivelis*, although these data can be expected to be biased by hatchery conditions.

Published estimates of *Z* vary widely among serranids, reflecting the wide range of longevities among species, the difficulty in estimating mortality rates using catch curves and, presumably, the wide variation in fishing mortality among study locations. Estimates of *Z* for *Plectropomus leopardus*, which has a similar longevity to *Cromileptes altivelis* and is the main target species on the GBR and TS, range from 0.12 to 0.68 (Brown et al. 1994, Russ et al. 1998, Mapstone et al. 2004, Bergenius 2007), encompassing the range of estimates for *C. altivelis* in our study. It is difficult to partition estimates of *Z* into its components of natural mortality (*M*) and fishing mortality (*F*), which are necessary to estimate the likely impacts of different levels of exploitation. The estimate of *Z* for *C. altivelis* from the GBR was based primarily on data from reefs closed to fishing for 10 to 12 yr prior to the first sampling in 1995. Excluding the data from open reefs resulted in a very similar estimate of mortality from catch curves (0.26 yr⁻¹) and Hoenig’s (1983) equation, which can be assumed to provide an indication of the likely values of *M*, assuming no infringement and 0 adult movement among reefs. The very similar estimates of *Z* for the GBR and TS (0.25 to 0.27 yr⁻¹) and the negligible difference between the estimates of *M* and *Z* suggest that current (in the TS) or recent past (on the GBR) fishing mortality on these populations is low.

Most serranids are considered to be late maturing, with first maturity (smallest length at which maturation is observed) reached at approximately 50% of average maximum size but ranging between 33 and 74% (Sadovy 1996). The absence of immature *Cromileptes altivelis* in our study means it is difficult to make rigorous comparisons of the maturity schedule for this species to that of other serranids. Liu & Sadovy de Mitcheson (2009) reported the length at first maturity for *C. altivelis* under culture conditions to be 155 mm *FL*. Although these estimates will have been influenced by culture conditions to some degree, they are consistent with our results. The smallest (mature) female *C. altivelis* we sampled was 350 mm *FL*, suggesting that 100% of the population above this size is mature and that the size at first (and 50%) maturity occurs at a smaller size. On the basis of these results, we conclude that *C. altivelis* appears to reach first maturity at approximately 27% (i.e. 155 mm/575 mm) of its average maximum length, which is smaller than for other serranids reported by Sadovy (1996). Information on age at maturity for serranids is more limited. Liu & Sadovy de Mitcheson (2009) estimate the age at first maturity for *C. altivelis* under culture conditions to be 1.6 yr, which is consistent with the youngest (mature) female *C. altivelis* that we sampled at 3 yr of age. The preliminary results suggest that *C. altivelis* matures at approximately 8 to 16% of potential life span (19 yr) and is likely to have a relatively long reproductive life (up to 17 yr). Further sampling of smaller and younger individuals is required to provide more accurate estimates of the size and age maturity schedules for *C. altivelis*.

Observations from underwater visual surveys (Sabetian 2003, Pears 2005) suggest that *Cromileptes altivelis* usually occur as individuals or pairs and do not form large spawning aggregations like many other medium- to large-sized serranids (Sadovy 1996). Spawning behaviour has been reported for a number of serranid species (e.g. Sadovy et al. 1994, Coleman et al. 1996, Samoilys 1997, Sluka 2001, Whaylen et al. 2004, Bergenius 2007), encompassing the range of estimates for *C. altivelis* in our study. It is difficult to partition estimates of *Z* into its components of natural mortality (*M*) and fishing mortality (*F*), which are necessary to estimate the likely impacts of different levels of exploitation. The estimate of *Z* for *C. altivelis* from the GBR was based primarily on data from reefs closed to fishing for 10 to 12 yr prior to the first sampling in 1995. Excluding the data from open reefs resulted in a very similar estimate of mortality from catch curves (0.26 yr⁻¹) and Hoenig’s (1983) equation, which can be assumed to provide an indication of the likely values of *M*, assuming no infringement and 0 adult movement among reefs. The very similar estimates of *Z* for the GBR and TS (0.25 to 0.27 yr⁻¹) and the negligible difference between the estimates of *M* and *Z* suggest that current (in the TS) or recent past (on the GBR) fishing mortality on these populations is low.
Cromileptes altivelis spawning in the wild has not been documented (Morris et al. 2000). Spawning has been observed in captivity at Indonesian aquaculture facilities, where it occurred throughout most of the year (Mous 2000, Sugama et al. 2003), but these observations are from a single equatorial location and are likely to be strongly influenced by culture conditions. Hence, they are unlikely to be representative of patterns in the wild or variation across the species’ range. Peak periods of spawning activity of *C. altivelis* in the TS were identified to occur in the months of October through January, while peak spawning on the GBR was restricted to December and January. This is consistent with general patterns of reef fish reproduction whereby spawning seasons commence earlier and are more protracted at lower than at higher latitudes (e.g. Kokita 2004).

The sexual development pathway for *Cromileptes altivelis* was unconfirmed prior to our study but assumed to be protogyny, as in most other serranids (Sadovy et al. 2008). We have confirmed that *C. altivelis* is a protogynous hermaphrodite through biological analyses that have revealed a bimodal size and age distribution by sex (with males larger and older than females) and transitional individuals, all of which are necessary features to confirm protogyny (Sadovy & Shapiro 1987). *C. altivelis* was found to change sex (on average) at around 95% of the average maximum length (*Lm* from the constrained VBGF fit) and around 50% of the potential maximum age. This is similar to other protogynous serranids in which sex change occurs close to the maximum size (e.g. Pears et al. 2006, Williams et al. 2008b).

Plasticity often exists in the timing of sex change for fish populations in different regions and locations (Adams et al. 2000, Gust 2004, Williams et al. 2006). The large overlap in the size and age of male and female *Cromileptes altivelis* and the presence of transitional individuals over several ages indicates that sex change occurs close to the maximum size (e.g. Pears et al. 2006). Estimates of biological parameters for *Cromileptes altivelis* from the TS were generally similar to those estimated for populations on the GBR, and the combined estimates provide a sound foundation for the future management of *C. altivelis* populations throughout the GBR and TS. The applicability of the biological information to other parts of the species’ range is less clear. Given the wide range of latitude from which our samples came (9.5 to 21°S), we consider that the estimates of population parameters provided are a useful starting place for other regions in the absence of region- or population-specific data. However, we strongly recommend that local estimates of population parameters are obtained as a matter of priority. Given the significant regional and local variation in population parameters that is becoming increasingly apparent for tropical reef fish populations (e.g. Adams et al. 2000, Williams et al. 2003, 2006, Gust 2004, Bergenius 2007) and its potential importance in the dynamics of species with meta-population structures (Kritzer & Davies 2005), estimates from local populations are likely to significantly improve assessments and advice for particular stocks.

*Cromileptes altivelis* is currently a no-take species on the GBR but is still subject to harvest in the TS. Fisheries management measures in the TS are likely to
provide protection from recruitment overfishing and conservation risk and may provide a suitable default for managing harvests in other locations where management measures are currently insufficient or absent. A minimum legal size limit of 450 mm FL was implemented in the TS on the basis that it would allow *C. altivelis* to spawn at least once before becoming vulnerable to harvest by the fishery. Our data indicate that the current minimum legal size will achieve this objective, given that all individuals we sampled were mature and the smallest individual sampled was 350 mm FL. However, it is important in the case of hermaphroditic species to account for sex change when setting size limits to avoid the disproportionate removal of 1 sex (Alonzo & Mangel 2004). Approximately 15% of *C. altivelis* individuals change sex prior to reaching 450 mm FL, which provides some protection of males prior to becoming selected by the gear used in this fishery. The survival of released fish needs to be relatively high for a minimum legal size limit to be effective, however, and, preferably, gear regulations should limit selectivity of individuals below the minimum legal size to minimise the prospect of capture and release. The post-release survival of *C. altivelis* is unknown, but for the current study it is likely to be relatively high as *C. altivelis* is usually taken in relatively shallow water (Heemstra & Randall 1999), reducing the incidence of barotrauma and increasing the probability of survival (Sumpton et al. 2008).

Closure of coral reef fisheries during peak spawning times has been advocated as an important management tool for protecting spawning aggregations and ensuring the sustainability of populations (e.g. Sadovy de Mitcheson et al. 2008). The relative efficacy of spawning closures, however, depends on numerous factors, including the aggregating behaviour of the species, predictability of locating aggregations, increased catchability during spawning times, and the presence of other management measures, such as catch, effort or size limits. *Cromileptes altivelis* is not known to aggregate in particular locations to feed or spawn, unlike many other serranids, and therefore is unlikely to be particularly vulnerable to targeted harvesting while spawning. Consequently, seasonal closures to fishing are unlikely to be a particularly efficient tool to protect *C. altivelis* populations from the effects of fishing. The implementation of no-take areas (areas closed to fishing) is likely to provide greater protection for *C. altivelis* than seasonal closures, in the absence of other management measures. Even so, it will be necessary to also manage the harvest outside closed areas if the objectives of management are to provide for sustainable harvest as well as reduce the conservation risk to the population (Hilborn et al. 2006, Mapstone et al. 2008).

The high demand for *Cromileptes altivelis* for the aquarium trade and live reef fish trade and the relatively low natural abundance of the species recommend a conservative approach to management of fisheries that harvest *C. altivelis*. The closure of some areas to harvest, to conserve a proportion of the spawning stock biomass, in addition to catch, effort and minimum size limits in areas open to fishing would appear to be a practical and appropriate combination of management measures to reduce the conservation risk to the species and improve long-term harvest. In practice, implementing restrictions on harvest, such as size limits and catch limits, is likely to be difficult to achieve in many developing nations, where a large proportion of global harvest of *C. altivelis* is taken and populations are at greatest conservation risk. In this context, no-take areas may be a more effective management tool for reducing the immediate conservation risk to local populations of *C. altivelis* in these regions in the short term.

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