

## Size, growth and mortality of riverine golden perch (*Macquaria ambigua*) across a latitudinal gradient

Daniel W. Wright<sup>id</sup> <sup>A,I</sup>, Brenton P. Zampatti<sup>B,H</sup>, Lee J. Baumgartner<sup>C</sup>, Steven Brooks<sup>D</sup>, Gavin L. Butler<sup>E</sup>, David A. Crook<sup>F</sup>, Ben G. Fanson<sup>G</sup>, Wayne Koster<sup>id</sup> <sup>G</sup>, Jarod Lyon<sup>G</sup>, Arron Strawbridge<sup>B</sup>, Zeb Tonkin<sup>G</sup> and Jason D. Thiem<sup>A,C</sup>

<sup>A</sup>Department of Primary Industries, Narrandera Fisheries Centre, PO Box 182, Narrandera, NSW 2700, Australia.

<sup>B</sup>Inland Waters and Catchment Ecology Program, SARDI Aquatic Sciences, PO Box 120, Henley Beach, SA 5022, Australia.

<sup>C</sup>Institute for Land, Water and Society, Charles Sturt University, PO Box 789, Albury, NSW 2640, Australia.

<sup>D</sup>Department of Agriculture and Fisheries, GPO Box 46, Brisbane, Qld 4001, Australia.

<sup>E</sup>Department of Primary Industries, Grafton Fisheries Centre, Private Mail Bag 2, Grafton, NSW 2460, Australia.

<sup>F</sup>Centre for Freshwater Ecosystems, La Trobe University, PO Box 821, Wodonga, Vic. 3689, Australia.

<sup>G</sup>Arthur Rylah Institute for Environmental Research, Department of Environment, Land, Water and Planning, PO Box 137, Heidelberg, Vic. 3084, Australia.

<sup>H</sup>Present address: Commonwealth Scientific and Industrial Research Organisation (CSIRO), Locked Bag 2, Glen Osmond, SA 5064, Australia.

<sup>I</sup>Corresponding author. Email: [daniel.wright@dpi.nsw.gov.au](mailto:daniel.wright@dpi.nsw.gov.au)

**Abstract.** Effective fisheries management requires fish size, growth and mortality information representative of the population and location of interest. Golden perch *Macquaria ambigua* is long lived, potamodromous and widespread in the Murray–Darling Basin (MDB), Australia. Using a sample spanning 13 river systems and 10° of latitude, we examined whether the maximum size of golden perch differed by latitude and whether growth and mortality varied between northern and southern MDB regions. The length, weight and age ranges of golden perch sampled ( $n = 873$ ) were 52–559 mm, 2–3201 g and 0+ to 26+ years respectively, and maximum length and weight were unaffected by latitude. Length and age–length distributions represented by age–length keys varied by region, with greater variability in age-at-length and a larger proportion of smaller individuals in northern MDB rivers, which generally exhibit greater variability in discharge. Growth and mortality rates were similar between regions, and an MDB-wide von Bertalanffy growth model ( $L_{\infty} = 447$ ,  $k = 0.32$  and  $t_0 = -0.51$ ) and instantaneous mortality rate ( $Z = 0.20$ ) best described the data. An MDB-wide length–weight equation also provided the best fit ( $W = 6.76 \times 10^{-6} L^{3.12}$ ). Our data suggest that the MDB can be treated as one management unit in terms of golden perch maximum size, growth and mortality parameters.

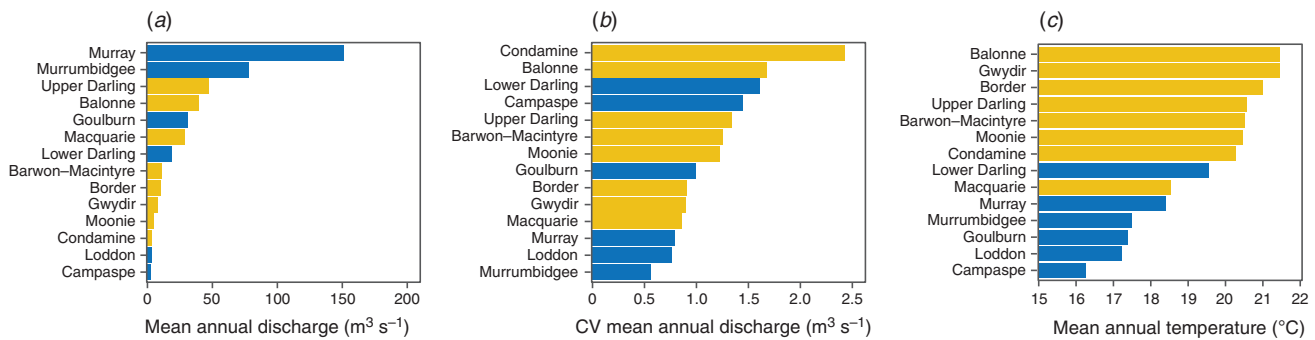
**Additional keywords:** age–length key, Bergmann’s rule, Murray–Darling Basin, otolith aging, von Bertalanffy growth function.

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### Introduction

Robust age and length information, along with derivative growth and mortality rate estimates, are critical to fisheries management (e.g. size-based harvest regulations). The accuracy of this information is improved by using a sample that is representative of the broad population and collected in a standardised manner (e.g.

same sampling method, year and time of year; [Coggins \*et al.\* 2013](#)). For widespread freshwater fish species, geographic span, particularly between latitudes and their associated climates, is a potential source of variation in age–length relationships within populations ([Belk and Houston 2002](#); [Heibo \*et al.\* 2005](#); [Rypel 2014](#)). In this respect, freshwater fishes may adhere to



**Fig. 1.** Magnitudes of (a) mean annual river discharge, (b) the coefficient of variation (CV) of mean annual river discharge and (c) mean annual water temperature from 1993 to 2018 at a single station or multiple stations proximate to or within the range of sampling locations for each river sampled in northern (yellow bars) and southern (blue bars) Murray–Darling Basin regions. Rivers are ranked in descending order of parameter values. Data were used from Stations 405232, 406202, 407202, 410005, 414203, 416001, 4106040, 418004, 421127, 425004, 425005, 417204A, 422201E, 422201F and 422333A (Bureau of Meteorology water data online, see <http://www.bom.gov.au/waterdata>, accessed October 2019).

Bergmann's rule, which predicts maximum fish size across a species range will increase with latitude from the equator towards the poles or at lower temperatures (Angilletta and Dunham 2003). However, many freshwater fishes also follow the converse to this rule (i.e. decreasing maximum size at latitudes towards the poles or lower temperatures) or exhibit no temperature–size relationship (Belk and Houston 2002; Heibo *et al.* 2005; Rypel 2014). Consequently, age–length data should be gathered from a wide geographical and latitudinal area relative to a species range and, where appropriate, sample sizes should be bolstered to explore region-specific growth and mortality rates (e.g. Piddocke *et al.* 2015).

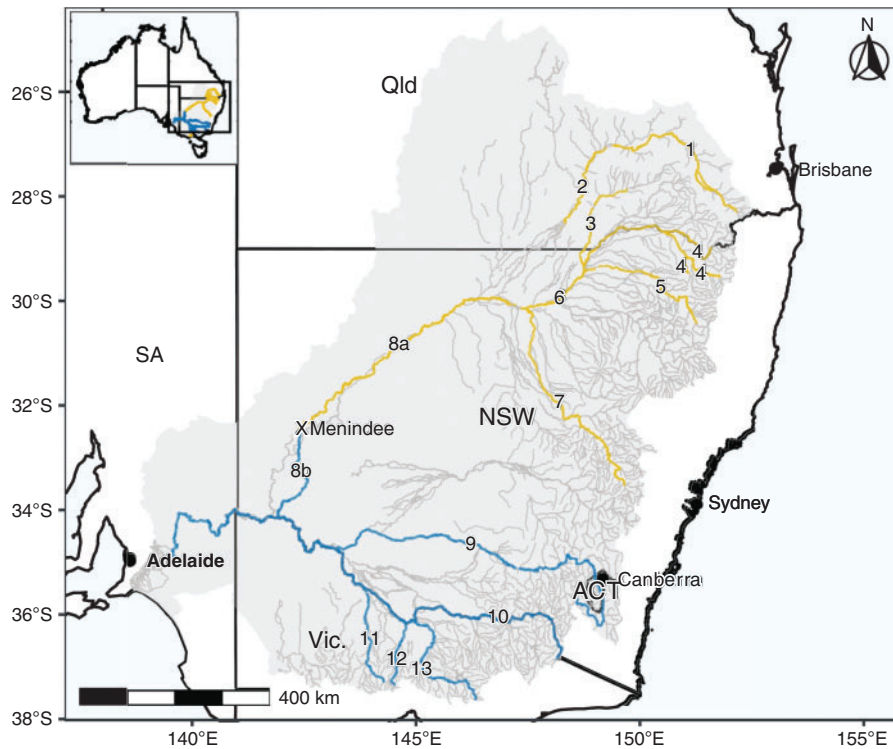
Golden perch *Macquaria ambigua* is a common native species in the Murray–Darling Basin (MDB) of eastern Australia that supports extensive recreational and historical and contemporary commercial fisheries (Kailola *et al.* 1993; Lintermans 2007; Earl 2016). Golden perch are a medium-bodied and long-lived fish, reaching ~550-mm total length (TL) and 27 years of age in river habitats (Mallen-Cooper and Stuart 2003; Forbes *et al.* 2015). Using the life history classifications of Winemiller and Rose (1992), golden perch are considered a periodic species (King *et al.* 2013). Although they may be strongly site attached for long periods (Crook 2004; O'Connor *et al.* 2005), golden perch can also undertake facultative migrations of tens to thousands of river kilometres across a range of river conditions at certain times (Reynolds 1983; Zampatti *et al.* 2018); they are also highly fecund, with females producing >500 000 eggs (Harris and Rowland 1996), and demonstrate episodic recruitment (Zampatti and Leigh 2013). MDB golden perch are genetically distinct from other members of the golden perch species complex that are found in the Lake Eyre, Bulloo and Fitzroy basins (Beheregaray *et al.* 2017). Throughout the MDB, minimal genetic structuring in golden perch reflects their ability to migrate and disperse long distances (Faulks *et al.* 2010; Beheregaray *et al.* 2017; Attard *et al.* 2018). Nevertheless, genetically similar MDB golden perch may still grow and survive at different rates depending on climatic conditions, such as rainfall run-off and temperature, which vary over the broad geographical region of the MDB (Attard *et al.* 2018).

Riverine golden perch age–length relationships have been previously examined, but not from a whole-of-range sample. Anderson *et al.* (1992) assessed the growth rate of golden perch ( $n = 796$ ) in six river systems throughout the southern MDB, although that study was somewhat confounded by including typically larger golden perch from impoundments (Mallen-Cooper and Stuart 2003; Forbes *et al.* 2015). Age–length relationships have also been explored for golden perch in three separate rivers in the northern and southern MDB, although sample sizes were variable ( $n = 39–216$ ; Mallen-Cooper and Stuart 2003). In a comparison of golden perch growth rates between reservoir and river systems, Forbes *et al.* (2015) used large sample sizes ( $n = 540–582$ ) from two rivers in the southern MDB. Nevertheless, suitably robust age–length information, in terms of both sample size and geographical coverage, is lacking. Identifying whether region-specific demographic parameters (e.g. growth and mortality rates) differ is particularly valuable for future monitoring and management efforts because distinct management units may need to be treated differently.

In the present study we used an extensive sample of MDB golden perch collected from 13 river systems throughout their geographic range to examine demographic parameters across a latitudinal gradient. Vital population statistics, including maximum body sizes, growth and mortality rates and length–weight relationships, were developed to provide contemporary whole-of-range information to support and evaluate current and future fisheries management. This was undertaken to: (1) test the hypothesis that, following Bergmann's rule (Angilletta and Dunham 2003), maximum body size would increase with latitude in a southerly direction and lower temperatures (and potentially other correlated environmental variables, such as discharge volume and variability); and (2) determine whether distinct spatial units, such as northern and southern MDB regions, are required in golden perch management due to differing demographic parameters.

## Materials and methods

Between November 2017 and June 2018, 873 golden perch were sampled from 13 river systems throughout the MDB, Australia, spanning 10° of latitude and a range of temperature regimes and hydrological characteristics (Fig. 1, 2). Sampling occurred at



**Fig. 2.** River systems sampled for golden perch within the Murray–Darling Basin (grey shading) in south-east Australia (see inset). The sampled river systems, numbered by increasing latitude, and the number of fish collected were: Condamine (1;  $n = 50$ ), Balonne (2;  $n = 30$ ), Moonie (3;  $n = 56$ ), Border (4;  $n = 54$ ), Gwydir (5;  $n = 65$ ), Barwon–Macintyre (6;  $n = 53$ ), Macquarie (7;  $n = 61$ ), Darling (8a ( $n = 60$ ) and 8b ( $n = 65$ )), Murrumbidgee (9;  $n = 78$ ), Murray (10;  $n = 211$ ), Loddon (11;  $n = 20$ ), Campaspe (12;  $n = 27$ ) and Goulburn (13;  $n = 43$ ). Colour coding represents northern (yellow) and southern (blue) rivers within the basin, separated by a major barrier to fish passage at Menindee (X). Note that golden perch collections from the Darling River in upper (8a) and lower (8b) sections contributed to northern and southern regions respectively. In addition, the upstream section of the Macintyre River and the Severn and Dumaresq rivers here constitute a single border river system (4). NSW, New South Wales; Qld, Queensland; SA, South Australia; Vic., Victoria; ACT, Australian Capital Territory.

multiple sites in eight river systems (including the upper Darling River) in the northern MDB region (latitudinal range from 26.9 to 32.2°S) and in six rivers (including the lower Darling River) in the southern MDB region (latitudinal range from 32.6 to 36.7°S). Southern and northern regions were above and below the Menindee main weir in New South Wales (NSW) on the Darling River (32.4°S, 142.4°E), which was near the latitudinal midpoint of sampling locations and is a semi-impermeable barrier to fish passage separating the two regions. Of the 873 individuals collected, 429 and 444 were from the southern and northern MDB respectively. All individuals were captured using boat-mounted electrofishing (either GPP 5 or 7.5-kW boat-mounted units, Smith-Root, Vancouver, WA, USA) and killed at the point of capture using an anaesthetic overdose (immersion in benzocaine or AQUI-S, AQUI-S New Zealand Ltd, Lower Hutt, New Zealand). Fish were then kept on an ice slurry and were subsequently measured for TL (mm) before sagittal otolith extraction. A subset of individuals ( $n = 863$ ) was also measured for whole weight ( $W$ ; g) at the time of sampling. A further subset ( $n = 704$  individuals) was sexed. Methods previously validated for annual age estimation of golden perch (Stuart 2006) were

used in this study. Whole otoliths were embedded in clear casting resin and, after setting, were hardened for a minimum of 4 h at 40°C in a drying oven fitted with a Eurotherm 3216 controller (Steridium, Gold Coast, Qld, Australia). Using a Isomet low-speed saw (Buehler, Lake Bluff, IL, USA), multiple 400–500  $\mu\text{m}$  transverse sections were cut through the primordium and the sections were then fixed to a microscope slide with super glue (Morison *et al.* 1998). Annulus counts for each fish were performed by two experienced readers (A. Strawbridge and D. Fleer.) under a Wild M3C microscope (Leica Biosystems, Wetzlar, Germany) at a magnification of 16 $\times$ , and age ( $t$ ) was assessed based on the annulus count and edge type in relation to the sampling date of an individual, with a nominal birthdate of 1 October assigned for all individuals (Stuart 2006).

This study was conducted in accordance with animal care and ethics committee guidelines (NSW: NSW Animal Care and Ethics permit 14/10 and Scientific Collection Permit P01/0059(A)-3.0; Queensland: General Fisheries Permit 186281 and Animal Care and Ethics CA 2016/01/938; South Australia: under an exemption (No. 9902132) of Section 115 of the *Fisheries Management Act* 2007 and

**Table 1. Sample sizes, length and age ranges, von Bertalanffy growth model (VBGM) coefficients and instantaneous total mortality rate ( $Z$ ) for sampled golden perch partitioned by region**

VBGM coefficient ranges between upper and lower 95% confidence intervals are provided in parentheses.  $L_{\infty}$ , maximum mean length of an individual;  $k$ , growth coefficient;  $t_0$ , theoretical age at length zero; MDB, Murray–Darling Basin; TL, total length

MDB region (latitudes)	Sample size ( $n$ )	TL (mm)	Age (years)	Weight (g)	$L_{\infty}$ (mm)	$k$ (year <sup>-1</sup> )	$t_0$ (years)	$Z$
North (26.9 to 32.2°S)	429	54–559	0+ to 26+	2–3174	439 (415–442)	0.34 (0.31–0.40)	-0.49 (-0.61 to -0.29)	0.24
South (32.6 to 36.7°S)	444	52–549	0+ to 24+	12–3210	455 (445–465)	0.31 (0.27–0.35)	-0.54 (-0.80 to -0.32)	0.18
Overall (26.9 to 36.7°S)	873	52–559	0+ to 26+	2–3210	447 (440–456)	0.32 (0.30–0.35)	-0.51 (-0.67 to -0.34)	0.20

following the *South Australian Animal Welfare Act* 1985; and Victoria: Victorian Fisheries Research Permit RP827, Fauna and Flora Guarantee Research Permit 10007273, Department of Environment, Land, Water and Planning Animal Ethics 14/04).

### Statistical analysis

Two alternative approaches were used to examine the relationship between age and length of golden perch. First, age–length keys were used to assess age–length distributions; second, von Bertalanffy growth models (VBGMs) were used to investigate growth rates. To predict the proportion of annual ages at each length interval, age–length keys (ALKs) were modelled using multinomial logistic regression model analysis with a maximum of 100 iterations (Ogle 2016). This assigned probabilities to each age–length category, even in the absence of data. Age categories  $\geq 14+$  years were combined because there were few contributing individuals in older age classes. Growth rate estimation, allowing predictions of mean length at age ( $E[L|t]$ ), was examined using the VBGM (von Bertalanffy 1938; Ogle 2016):

$$E[L|t] = L_{\infty} \left( 1 - e^{-k(t-t_0)} \right)$$

where  $L_{\infty}$  is the maximum mean length of an individual,  $k$  is the growth coefficient and  $t_0$  is the theoretical age at length zero.

In addition, we used ages in catch curve linear regression models to predict instantaneous mortality rates ( $Z$ ; Ogle 2016):

$$\log \left( \frac{C_t}{E_t} \right) = \log(qN_0) - Zt$$

where  $N_0$  is starting population size,  $C_t \div E_t$  is catch per unit effort,  $t$  is age and  $q$  is the constant proportion of the population caught. We only included the age classes at and above the maximum catch, which was approximately at 4 years of age (Smith *et al.* 2012; Ogle 2016), to ensure that fish were large enough for capture with the sampling method.

Finally, linear regression models were used to predict length–weight relationships between  $\log_{10}(L)$  and  $\log_{10}(W)$  to assess condition (Ogle 2016).

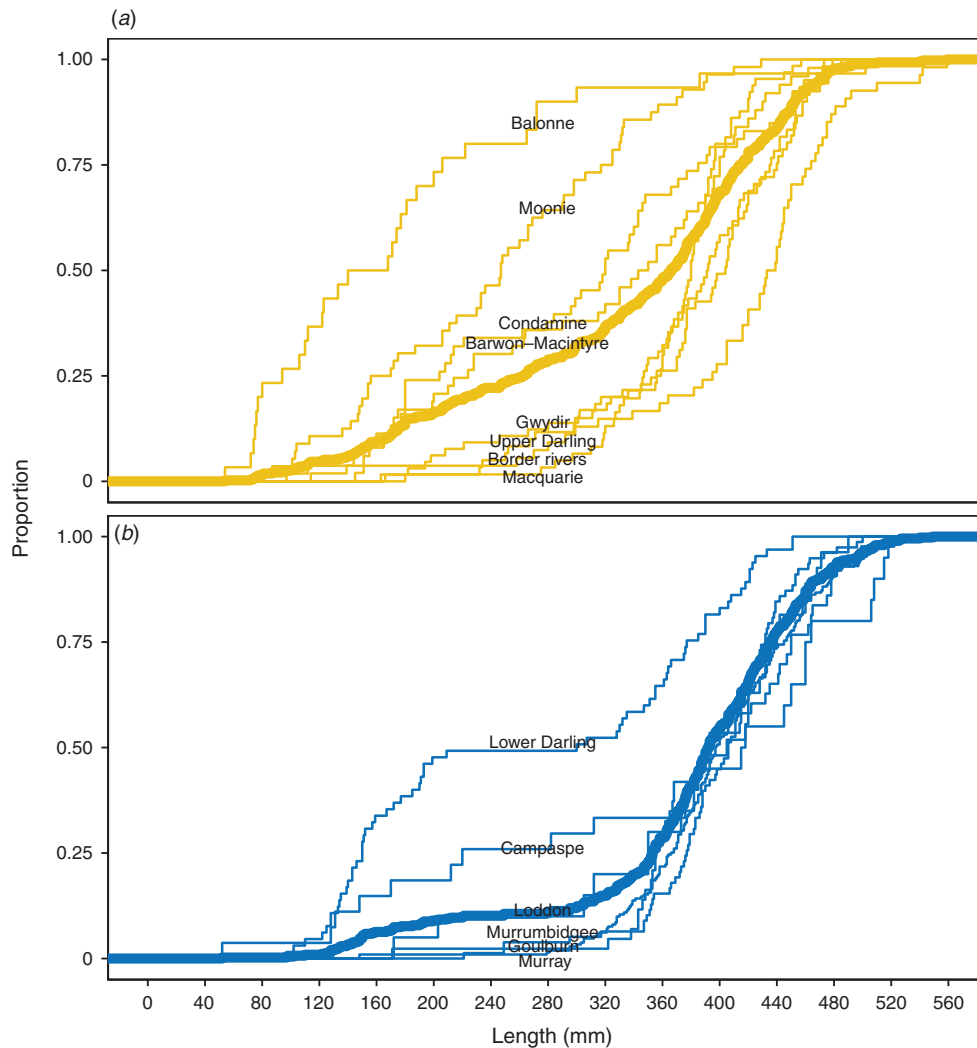
Relationships between maximum body size, in terms of length and log-transformed weight, and latitude at sampling sites with  $\geq 10$  individuals were examined using linear regression model analysis. Length–frequency distributions were compared between northern and southern MDB golden perch by a two-sample Kolmogorov–Smirnov test. Differences in multinomial model ALKs between golden perch captured in the

northern and southern MDB were assessed by a likelihood ratio test. Non-linear model analysis examined differences in VBGMs between golden perch in the northern and southern MDB (nlstools packages, R Foundation for Statistical Computing, Vienna, Austria, see <https://cran.r-project.org/web/packages/nlstools/>, accessed October 2019; Baty *et al.* 2015). In this analysis, we used a model selection approach to compare models from different geographic regions using all combinations of coefficients. Models with all coefficient combinations ( $\{L_{\infty}\}$ ,  $\{k\}$ ,  $\{t_0\}$ ,  $\{L_{\infty}, k\}$ ,  $\{L_{\infty}, t_0\}$ ,  $\{k, t_0\}$ ,  $\{L_{\infty}, k, t_0\}$  and the null model  $\{\Omega\}$ ) were ranked using the Akaike information criterion (AIC), and the model that had the lowest AIC was deemed to have the best fit (after Ogle 2016). Models with a  $\Delta$ AIC of  $< 2$  were also identified as having substantial support (Burnham and Anderson 2002). Linear regression model length–weight relationships were compared between northern and southern MDB golden perch using a Type II analysis of variance (ANOVA) to identify whether condition varied between regions. Concomitantly, to test whether mortality rate differed between regions, we examined differences in linear regression model catch curves predicting instantaneous mortality rates ( $Z$ ) for northern and southern MDB golden perch using a Type II ANOVA. Residual plots and histograms of residuals were used to confirm parametric assumptions were met. All statistical analyses were performed in R (ver. 3.5.2, R Foundation for Statistical Computing, Vienna, Austria, see <http://www.R-project.org/>, accessed March 2019).

### Results

Golden perch lengths and ages were in the range 54–559 mm and 0+ to 26+ years in the northern MDB, and 52–549 mm and 0+ to 24+ years in the southern MDB (Table 1). Weights of sampled golden perch in the northern and southern MDB were in the range 2–3174 and 12–3210 g respectively (Table 1). Sex ratios (males : females) were 1 : 1.8 ( $n = 385$  individuals) and 1 : 2.2 ( $n = 319$  individuals) in the northern and southern regions respectively. Latitude did not affect maximum body size in terms of length ( $R^2 = 0.04$ ,  $F_{1,37} = 1.66$ ,  $P = 0.21$ ) or weight ( $R^2 = 0.07$ ,  $F_{1,37} = 2.80$ ,  $P = 0.10$ ) at sites with  $\geq 10$  individuals (see Fig. S1, available as Supplementary material to this paper).

Length–frequency distributions differed between northern and southern regions of the MDB ( $P < 0.01$ ), with greater proportions of smaller golden perch in the northern MDB (Fig. 3). Differences in age–length composition between regions were identified in ALK comparisons ( $P < 0.01$ ), with golden perch in the northern MDB demonstrating a more variable age–length distribution. For example, an individual of 300 mm in



**Fig. 3.** Cumulative length–frequency distributions for Murray–Darling Basin golden perch in individual (a) northern and (b) southern river systems (thin lines), and within each (a) northern and (b) southern region (thick lines).

length in the northern region would have a  $>0.01$  probability of belonging to an age category between 1+ and 13+ years, whereas an individual of the same length in the southern region would have the same probability of being aged between 2+ and 9+ years (Fig. 4; Tables S1, S2).

Using VBGMs to analyse growth rates, we found the null  $\{\Omega\}$  model without regional differences to be the top-ranked VBGM, using overall coefficients of  $L_{\infty} = 447$ ,  $k = 0.32$  and  $t_0 = -0.51$  (Tables 1, 2; Fig. 5, S2). There was also substantial support ( $\Delta\text{AIC} < 2$ ) for additional models including regional differences in the coefficients (or combinations of coefficients)  $\{L_{\infty}\}$ ,  $\{L_{\infty}, k\}$ ,  $\{L_{\infty}, t_0\}$  and  $\{k\}$  in VBGMs (Table 2). These mostly included  $L_{\infty}$ , which was smaller in the northern MDB (439 mm; 95% confidence interval (CI) 415–442) than in the southern MDB (447 mm; 95% CI 445–465; Tables 1, 2; Fig. 5, S3, S4). Prediction intervals (PI) in  $L_{\infty}$  also varied between the northern (95% PI 372–506) and southern (95% PI 403–506) MDB (Fig. 5, 6).

Slopes of length–weight relationships were not significantly different between the northern and southern regions of the MDB ( $F = 0.68$ ; d.f. = 1, 864;  $P = 0.41$ ), and the overall model  $W = 6.76 \times 10^{-6}L^{3.12}$  explained most of the variation observed ( $r^2 = 0.99$ ; Fig. 6).

Golden perch mortality rates were comparable between the northern and southern MDB ( $F = 3.53$ ; d.f. = 1, 40;  $P = 0.07$ ), and an instantaneous mortality rate  $Z = 0.20$ , translating to an annual mortality rate ( $A$ ) of 18%, was determined for the overall population (Fig. 7).

## Discussion

Rather than following Bergmann's rule, namely that animals reach larger maximum body sizes at latitudes towards the poles or at cooler temperatures (Angilletta and Dunham 2003), golden perch maximum length and weight was not affected by latitude in the present study. These results suggest that golden perch do not have a pronounced body size relationship with latitude or

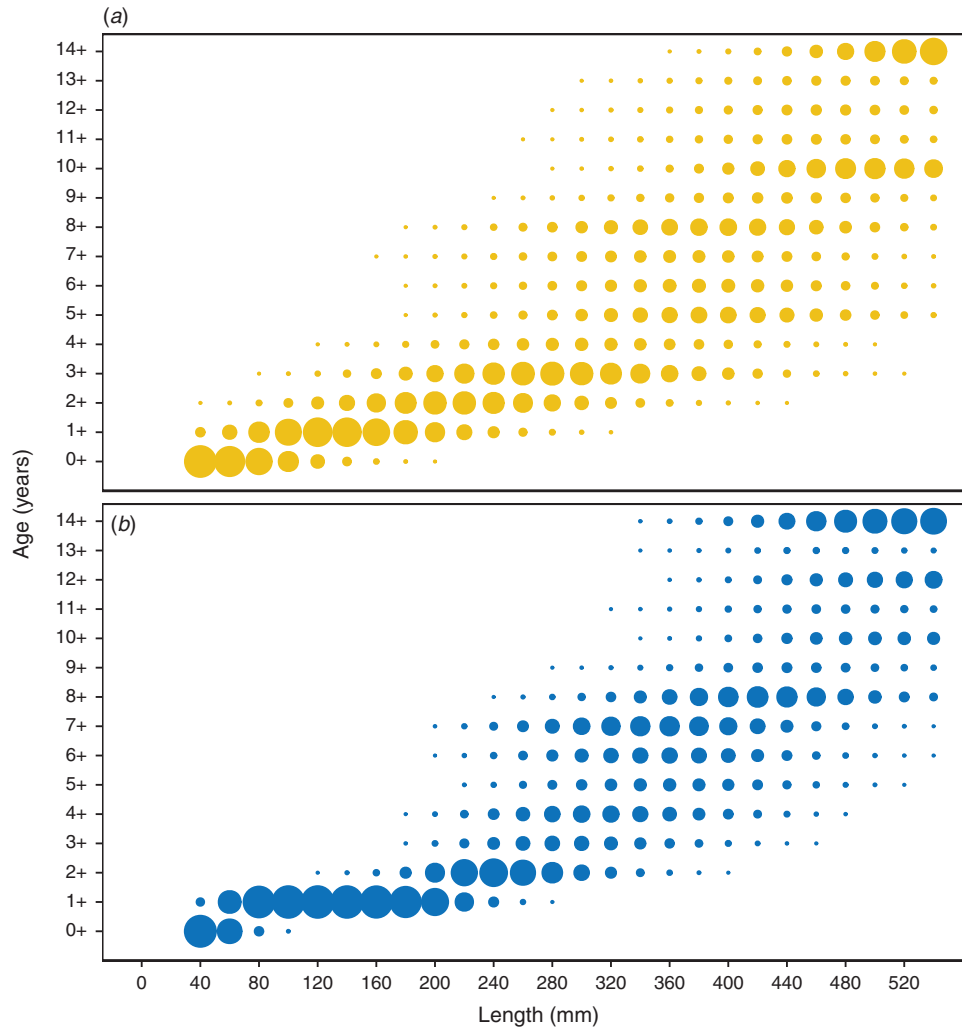


Fig. 4. Modelled age-length keys for Murray-Darling Basin golden perch in (a) northern and (b) southern regions. At each 20-mm interval, the proportion of fish at each age is indicated by bubble size.

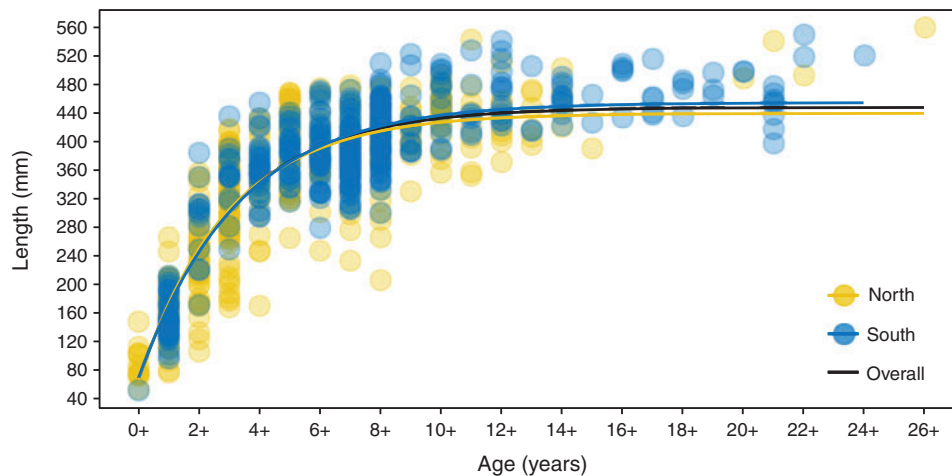
Table 2. Selection of the best model fit for von Bertalanffy growth models (VBGMs) between northern and southern Murray-Darling Basin golden perch

Models with all three VBGM coefficients, two coefficients or one coefficient varying along with the null model were compared, and models with delta Akaike information criterion (AIC) values < 2 for which there was substantial support are in bold.  $L_{\infty}$ , maximum mean length of an individual;  $k$ , growth coefficient;  $t_0$ , theoretical age at length zero;  $\{\Omega\}$ , null model;  $E[L|t]$ , mean length at age

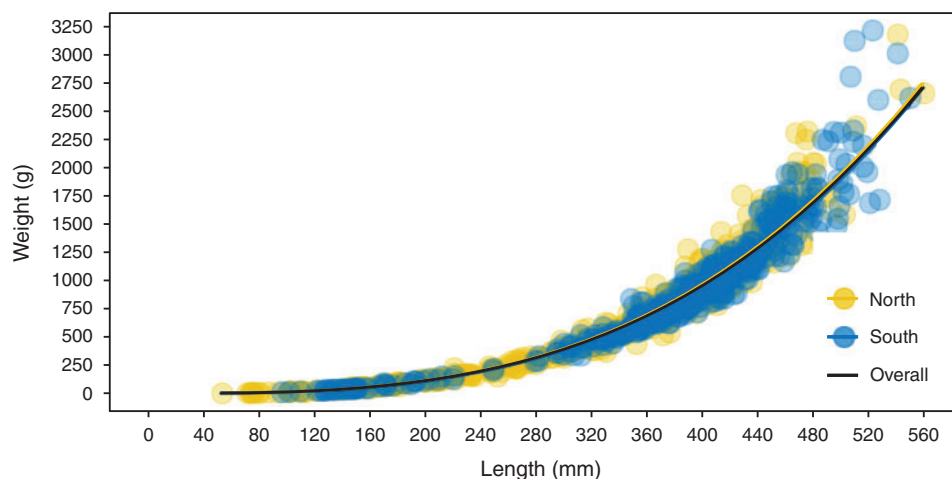
Abbreviation	Model	AIC	$\Delta$ AIC	d.f.
$\{\Omega\}$	$E[L t] = L_{\infty}(1 - e^{-k(t-t_0)})$	<b>9199.3</b>	<b>0</b>	<b>4</b>
$\{L_{\infty}\}$	$E[L t] = L_{\infty}[\text{Region}](1 - e^{-k(t-t_0)})$	<b>9199.7</b>	<b>0.4</b>	<b>5</b>
$\{k\}$	$E[L t] = L_{\infty}(1 - e^{-k[\text{Region}](t-t_0)})$	<b>9201.2</b>	<b>1.9</b>	<b>5</b>
$\{t_0\}$	$E[L t] = L_{\infty}(1 - e^{-k(t-t_0[\text{Region}])})$	9201.3	2	5
$\{L_{\infty}, k\}$	$E[L t] = L_{\infty}[\text{Region}](1 - e^{-k[\text{Region}](t-t_0)})$	<b>9199.9</b>	<b>0.6</b>	<b>6</b>
$\{L_{\infty}, t_0\}$	$E[L t] = L_{\infty}[\text{Region}](1 - e^{-k(t-t_0[\text{Region}])})$	<b>9200.7</b>	<b>1.4</b>	<b>6</b>
$\{k, t_0\}$	$E[L t] = L_{\infty}(1 - e^{-k[\text{Region}](t-t_0[\text{Region}])})$	9202.4	3.1	6
$\{L_{\infty}, k, t_0\}$	$E[L t] = L_{\infty}[\text{Region}](1 - e^{-k[\text{Region}](t-t_0[\text{Region}])})$	9201.9	2.6	7

temperature. There was no conclusive evidence for differences in growth rates, and similar mortality rates and length-weight relationships were also identified for golden perch between the

northern MDB (28 to 32°S) and southern MDB (33 to 37°S) regions. Thus, this study indicates that the MDB may be considered as a single management unit for golden perch from the



**Fig. 5.** Golden perch total lengths plotted against otolith-derived ages in northern and southern (circles) within the Murray–Darling Basin. The lines indicate von Bertalanffy growth model relationships between length and age for northern and southern regions, as well as for all individuals.



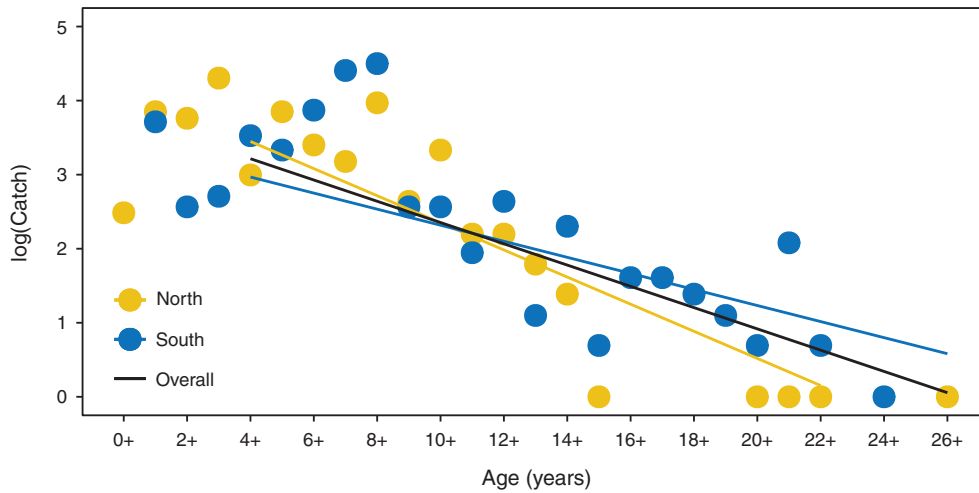
**Fig. 6.** Golden perch weight plotted against total length in northern and southern regions (circles) within the Murray–Darling Basin. The lines indicate modelled length–weight relationships for northern and southern regions, as well as for all individuals.

perspective of maximum size, growth and mortality demographic parameters.

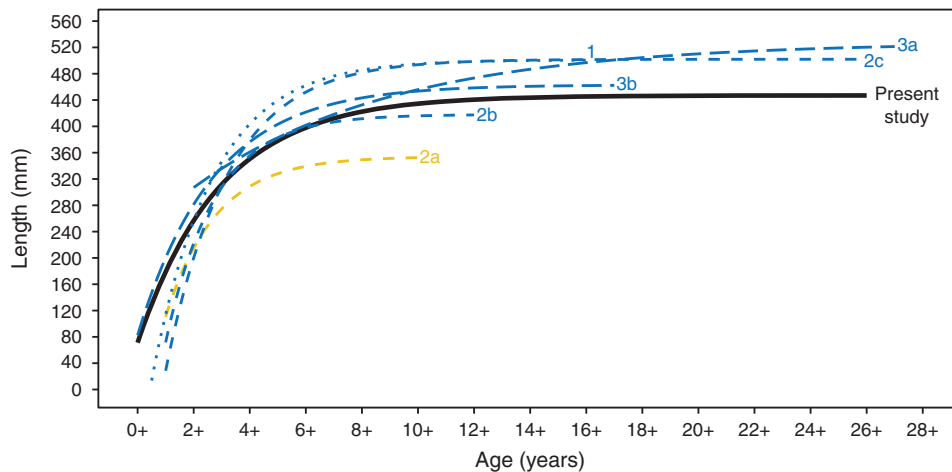
Many freshwater fish species exhibit the converse of Bergmann's rule, particularly warm-water species, which may have reduced growth opportunities and longevity at cooler temperatures (Belk and Houston 2002; Rypel 2014). Golden perch, which could be considered a warm-water or transitional species based on limited knowledge of their thermal preference and tolerance (Lyon *et al.* 2008), do not appear to have a relationship between body size and latitude or temperature. In other taxa, size–latitude relationships are known to weaken for highly mobile species that can evade environmental extremes by repositioning themselves (Ramirez *et al.* 2008). In this respect, golden perch may, at times, be highly mobile, such as during broad-scale flooding when riverine connectivity is high (Reynolds 1983; Zampatti *et al.* 2018). Nevertheless, in the

MDB there may be long periods (years) when low discharge and a multitude of instream barriers prevent large-scale fish movement and the ability to escape environmental extremes (Baumgartner *et al.* 2014). Consequently, it is unlikely that movement is the sole reason that golden perch do not adhere to Bergmann's rule.

Golden perch growth rate, mortality rate and condition (length–weight relationship) were similar between the northern and southern MDB regions. A factor contributing to these similarities was that we sampled many rivers representative of the variety of environmental conditions across regions. This may have masked differences in parameters between regions for specific river types (e.g. perennial or ephemeral). Anthropogenic effects on river systems also conceivably affected golden perch demographic parameters between regions. For example, recreational fishing pressure (Cooke and Cowx 2004) and



**Fig. 7.** Catch curves for Murray–Darling Basin golden perch in northern and southern regions, as well as an overall catch curve.



**Fig. 8.** Golden perch von Bertalanffy growth models (VBGMs) for the entire Murray–Darling Basin (MDB) in this study ( $n = 873$  in 2017–18) and MDB regions or rivers in three previous papers, numbered as follows: 1, VBGM for the southern MDB ( $n = 796$  in 1984–91) in *Anderson et al. (1992)*; 2, VBGM in the Darling (2a;  $n = 39$  in unknown years), Murray (2b;  $n = 216$  in 1990–92) and Murrumbidgee (2c;  $n = 95$  in unknown years) rivers in *Mallen-Cooper and Stuart (2003)*; and 3, VBGM in the Murray (3a;  $n = 582$  in 2009–13) and Murrumbidgee (3b;  $n = 540$  in 2009–13) rivers in *Forbes et al. (2015)*. VBGMs are coloured yellow and blue when developed from fish samples in northern and southern MDB regions respectively and black for this paper using fish samples from throughout the MDB. Note that some individuals from impoundments contributed to the VBGM for the southern MDB reported in *Anderson et al. (1992)*.

human-driven climate and hydrological modifications (*Hobday and Lough 2011; Baumgartner et al. 2014*) may be reducing growth opportunities and survival throughout the MDB. Global warming also has the potential to increase golden perch growth in cooler southern regions (*Morrongiello et al. 2011*) and decrease growth in warmer northern regions, conceivably leading to equivocal growth, survival and condition in both regions. Again, large-scale movement by golden perch, enabling access to favourable environments and between regions, is another potential factor explaining equivalent demographic parameters between regions, but these movements are compromised much of the time by barriers to fish.

We generated a VBGM from a representative, whole-of-range sample of riverine MDB golden perch. This model differs substantially from previous models (*Fig. 8; Table S3*), which can be plausibly explained. First, in contrast to previous studies of individual rivers or regions, we accounted for potential among-river and among-region differences by sampling a range of rivers within the northern and southern MDB. Second, unlike *Anderson et al. (1992)*, we excluded individuals from impoundments, which typically grow faster and larger (*Mallen-Cooper and Stuart 2003; Forbes et al. 2015*). Large sample sizes of particularly small or large individuals can considerably affect growth models (*Lucena and O'Brien 2001*). Third, we used a



standardised boat electrofishing method to collect all samples, rather than multiple sampling methods (e.g. gill-netting, fyke netting, angler captures and specimen donations), which may have altered the age-length structure of samples used between the present and some previous models (Anderson *et al.* 1992). However, we acknowledge that there may be some effect of electrofishing capture probability on the lengths of fish captured (Lyon *et al.* 2014). Fourth, we collected samples within a single year, whereas previous studies have collected samples over multiple years. Despite methodological differences across studies, the lower growth rate in the present study in 2017–18 ( $k = 0.31$ ) and in the study of Forbes *et al.* (2015) in 2009–13 ( $k = 0.14$ – $0.37$ ) compared with those before 2004 ( $k = 0.45$ – $0.57$ ) warrants further investigation into a possible temporal decline in golden perch growth rate (e.g. from global warming or discharge alteration; Hobday and Lough 2011; Morrongiello *et al.* 2011; Tonkin *et al.* 2017; Huss *et al.* 2019).

The oldest golden perch recorded in this study was 26 years, almost equal to the maximum age known for the species of 27 years (Forbes *et al.* 2015). The overall annual mortality rate of 18% observed here includes both recreational fishing and natural sources of mortality. In regions of the MDB, recreational fishing can remove >30% of fish in some size classes each year (Allen *et al.* 2009), and this may have overshadowed patterns in natural mortality within the present study. Interestingly, the overall length-weight relationship ( $W = 6.76 \times 10^{-6}L^{3.12}$ ; e.g. 500-mm fish = 1781 g) determined in this study indicates poorer condition than in 1959–70 ( $W = 7.08 \times 10^{-6}L^{3.12}$ ; e.g. 500-mm fish = 1866 g; Llewellyn 2011) and in 1984–91 ( $W = 3.34 \times 10^{-7}L^{3.66}$ ; e.g. 500-mm fish = 2523 g) (Anderson *et al.* 1992). However, the sample used in Anderson *et al.* (1992) included a significant number of large fish from impoundments and few fish <100 mm, thus potentially biasing their length-weight relationship. Deteriorating golden perch condition over time could again be linked to discharge alterations or global warming.

Length and age-length distributions (from ALKs) differed between northern and southern MDB regions, with greater variability in age-at-length data in the northern MDB. Regional differences in ALKs, but not VBGMs, likely resulted from ALK age-at-length probability matrices being more sensitive to detecting age variation within length classes than VBGMs forcing a single curve or age-at-length prediction through all data (Goodyear 1997). Greater age-at-length variation in the northern MDB may relate to the hydrological diversity of rivers in this region (Thoms and Sheldon 2000). In the ephemeral northern rivers, rewetting and episodic flooding could contribute more to growth (Sternberg *et al.* 2008; Balcombe and Arthington 2009), but low discharge over extended periods of drought may lessen growth (Tonkin *et al.* 2017).

Golden perch appear to have no relationship between maximum size and latitude or temperature. Using a VBGM analysis, an MDB-wide growth model was found to be most appropriate to predict age-at-length for golden perch throughout its range. Nevertheless, ALKs differed between northern and southern regions of the MDB, and in the northern region greater variation in age-at-length data may be associated with increased ephemerality of some rivers. Consequently, to predict age-at-length probabilities for many fish within each length category, region-specific ALKs would be most

appropriate, whereas to predict the age of individual fish within a length category an MDB-wide VBGM would suffice. We advocate that researchers considering fish age-length data similarly examine geographic differences and use both VBGM and ALK approaches to provide end users with case-by-case models. Our data were collected within a single year, and the demographic parameters presented here will vary through time as recruitment and cohort strength vary. Thus, further investigation will be essential to: (1) ensure demographic parameters remain relevant for golden perch management; and (2) identify long-term trends in these parameters and the causal effects. Based on our contemporary results, golden perch size, growth, mortality and condition do not differ between northern and southern MDB regions, and the MDB should be considered as a single management unit for these demographic parameters.

### Conflicts of interest

Lee J. Baumgartner is an Associate Editor of *Marine and Freshwater Research*. Despite this relationship, he did not at any stage have Associate Editor-level access to this manuscript while in peer review, as is the standard practice when handling manuscripts submitted by an editor to this journal. *Marine and Freshwater Research* encourages its editors to publish in the Journal and they are kept totally separate from the decision-making process for their manuscripts. The authors declare that they have no further conflicts of interest.

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