#### Ecological Society of Australia

# Responses of an Australian freshwater turtle to droughtflood cycles along a natural to urban gradient

BRUNO O. FERRONATO,<sup>1</sup>\* JOHN H. ROE<sup>2</sup> AND ARTHUR GEORGES<sup>1</sup>

<sup>1</sup>Institute for Applied Ecology, University of Canberra, Canberra, Australian Capital Territory 2601, Australia (E-mail: brunoferronato@hotmail.com), and <sup>2</sup>Department of Biology, University of North Carolina, Pembroke, North Carolina, USA

**Abstract** Urban areas provide habitat for numerous native species, but life in towns and cities presents many challenges. The effect of climate on the ecology and the behaviour of non-volant vertebrates inhabiting urban habitats have received little attention. In this study, we investigated demography, growth rates, movements and reproduction of a semi-aquatic freshwater turtle, *Chelodina longicollis*, along a natural to urban gradient during a period of relatively high rainfall (2011–2014) and compared this to a previous study in the same system during drought (2006–2007). In addition to changes in rainfall, urbanization increased considerably over the same time period and a pest-exclusion fence was constructed to mitigate against urban hazards encroaching on the adjacent reserve. Turtles grew at similar rates, had similar abundances and sex ratios and had similar reproductive output across the gradient from urban to non-urban sites during the wet period. Despite increasing urbanization, recruitment occurred at all sites and survivorship estimates were similar among sites. Turtles moved among wetlands at high rates and over long distances (6 km), underscoring the importance of movements in urban landscapes. Our results contrast with those for the same system during drought, when turtles were less abundant and grew slower in the nature reserve compared with the urban environment. Our results underscore the strong influence climate can have on population dynamics and resilience of species to changes brought about by urbanization. Further monitoring is required to understand the long-term population responses of long-lived species to drought cycles.

Key words: climate cycles, dispersal, El Niño, longitudinal study, urbanization.

# INTRODUCTION

Urbanization rates are expected to increase worldwide in the next decades (World Resources Institute et al. 1996; Gakenheimer 1999). Growth of cities and the associated urban sprawl encroaches on natural habitats with negative consequences for many native species (McKinney 2002, 2008; Pauchard et al. 2006). An increasing body of work details how vertebrates are impacted by urbanization (Pautasso 2006; McKinney 2008), including mitigation measures to protect them from urban hazards (Clevenger & Waltho 2005; Bond & Jones 2008), and how generalists species can cope with the life in the city (Chace & Walsh 2006; Bateman & Fleming 2012). Less well understood is how climate, and in particular drought cycles, influences behaviour and population responses along the interface between natural or rural and urban habitats, and how interactive and cumulative effects of climate cycles and urban expansion influence vertebrate populations. Climate change predictions are for an increase in the frequency of drought and flood events (Hughes 2003), which in turn increases urgency to improve this

understanding. Vagile vertebrates (e.g. many birds and mammals) can disperse among habitat patches either within or outside the urban area (Takekawa & Beissinger 1989; Goad *et al.* 2014), but less vagile species may be more limited in their responses to spatial and temporal variation in habitat quality resulting from climate cycles.

Freshwater turtles represent a useful model species to investigate the effects of urbanization and the interactions with climate. Populations can be easily and repeatedly sampled within a series of discrete patches (waterbodies) across targeted land-use gradients (Gibbons 1968). Long lifespans also make turtles amenable to capture-mark-recapture studies that span short-term climactic cycles (Congdon et al. 1994; Roe & Georges 2008a,b) and anthropogenic land-use changes (Plummer & Mills 2008; Eskew et al. 2010). Freshwater turtles rely on movements among waterbodies and to nearby terrestrial habitats for nesting and other critical behaviours (Steen et al. 2006; Roe & Georges 2007), making them particularly sensitive to loss of connectivity arising from roads and other aspects of urbanization (Gibbs & Shriver 2002; Marchand et al. 2002; DeCatanzaro & Chow-Fraser 2010; Eskew et al. 2010). Alternatively, some species may benefit from increased productivity of urban waterways, leading to faster growth, higher

<sup>\*</sup>Corresponding author.

Accepted for publication September 2016.

fecundity and ultimately higher population abundances compared with natural populations (Brown *et al.* 1994; Lindeman 1996; Souza & Abe 2000). Such benefits are water-dependent and so under the influence of drought cycles characteristic of the Australian temperate climate.

The Australian eastern long-necked turtle (Chelodina longicollis) is a generalist and opportunistic species with a marked propensity for overland movement (Roe & Georges 2008a; Rees et al. 2009), enabling it to exploit a wide range of temporary and permanent aquatic habitats (Kennett et al. 2009), including urban waterbodies (Rees et al. 2009; Roe et al. 2011; Stokeld et al. 2014; Hamer et al. 2016). Chelodina longicollis appears to be somewhat resilient to at least some degrees of urbanization, with site occupancy and population demographics in urban areas similar to rural and natural areas (Roe et al. 2011; Stokeld et al. 2014; Hamer et al. 2016). In some cases, they have been shown to grow faster, achieve larger body size and reach higher abundances in urban habitats compared with nearby natural areas (Roe et al. 2011). However, the mechanisms involved in such population responses in urban landscapes are not completely understood and may depend on a suite of interacting factors, including climate (Rees et al. 2009; Roe et al. 2011), environmental contamination (Stokeld et al. 2014) and influence of exotic predators (Spencer & Thompson 2005). Most studies examining turtle responses to urbanization, including for C. longicollis, are limited to short-term snapshots of a few consecutive years (Roe et al. 2011; Stokeld et al. 2014), which can lead to an incomplete understanding of temporal trends in urban impacts.

Here, we examine attributes of the population biology, including demography, growth rates, movements and reproduction, of C. longicollis during a period of high rainfall (wet period; 2011-2014) influenced by La Niña, and we compare these with earlier studies in the same system during a period of low rainfall influenced by El Niño (drought; 2006-2007) (Rees et al. 2009; Roe et al. 2011). In conducting a longitudinal study over nearly a decade, we aimed to better understand vertebrate responses to drought cycles within natural-urban gradients. We also examine how turtles respond over time to additional changes in the system with potential importance for population regulation, including expanding urbanization and the implementation of a barrier fence to mitigate urban impacts in neighbouring nature reserves.

#### METHODS

#### Study area

From October 2011 to March 2014, we studied turtle pop-

ulations in 14 water bodies distributed along a gradient

extending from suburb to rural to nature reserve (urban gradient) in the Gungahlin region of the Australian Capital Territory (ACT), southeastern Australia (Fig. 1). The nature reserve site was Mulligans Flat Nature Reserve comprising 791 ha of woodland, grassland, ponds and upper tributaries of Ginninderra Creek. In June 2009, a predator-proof barrier fence was erected, as part of a wildlife restoration project, enclosing 485 ha of the reserve to isolate it from the impacts of encroaching urbanization, exclude invasive species and allow reintroduction of locally extinct native species (Manning *et al.* 2011). We sampled turtles in five wetlands within the enclosure. The nature reserve site was defined here as having a low degree of anthropogenic impact isolated by the fence enclosure.

Five wetlands were sampled in the rural landscape, including two wetlands in the Ginninderra Experimental Station and three wetlands in Goorooyarroo Nature Park. The Ginninderra Experimental Station consists of areas with native grasses and eucalypts, in addition of areas with crops and pastures (Webster & Butler 1976). Goorooyarroo is adjacent to Mulligans Flat Nature Reserve, with similar vegetation species composition, but is not enclosed by the barrier fence. Together the rural sites of Ginninderra Experimental Station and Goorooyarroo were defined as having intermediate degree of anthropogenic impact, including agriculture, low-level urban development and exposure to invasive predators such as the European fox.

Finally, four wetlands were sampled in a suburban site located in the central Gungahlin suburb, including a large reservoir, a golf course pond, a canal and a storm water drainage pond. This area is subject to industrial and residential development, including high road densities and managed urban green spaces such as golf courses, parks, gardens and sport ovals (Rees *et al.* 2009; Roe *et al.* 2011). This site was defined as having high degree of anthropogenic impact, including urbanization and exposure to invasive and domestic predators.

The climate of the ACT is temperate, with a mean annual rainfall of 633 mm (1974–1988, Australian Bureau of Meteorology). Rainfall in southeast Australia is highly variable, with long periods of drought punctuated by flood. The most recent drought occurred from 2001 to 2009, with a yearly below median rainfall of 483 mm year<sup>-1</sup>, mainly influenced by El Niño events (van Dijk *et al.* 2013). There are also periods of elevated rainfall, influenced by La Niña events (Beard *et al.* 2011) such as in 2012 and 2013 with annual rainfall of 694 and 534 mm, respectively (Canberra Airport weather station, Australian Bureau of Meteorology).

#### **Trapping and marking**

Turtles were captured using traps baited with sardines and bovine liver once per month (five consecutive days of trapping per month) from October 2011 to March 2014, excluding months when turtles are inactive (April–August). Two ponds in each of the nature reserve, rural and suburb sites, hereafter referred to as the fixed sites, were sampled monthly to describe reproductive biology of the turtles. Additional ponds were sampled twice per year to increase sample sizes in other demographic analyses (three ponds



**Fig. 1.** Nature reserve, rural and suburban study sites in Gungahlin, northern Canberra, Australian Capital Territory. Study sites were defined by drawing 700-m polygons around sampled ponds and then joining the polygons to delimit each site. The polygons in the nature reserve were contracted to delimit the area only included within pest-fencing. Number 1 denotes the Ginninderra Experimental Station and number 2 the Goorooyarroo Nature Park, both part of the rural sites.

each in the nature reserve and rural sites, and two in the suburb site) and are referred to hereafter as the occasional sites. We used two to six traps per pond depending on pond size. We marked captured turtles with unique codes by notching the shell and measured maximum straight-line carapace length (CL), carapace width (CW), midline plastron length (PL) and plastron width (PW) with callipers ( $\pm 0.1$  mm) and body mass with a scale ( $\pm 5$  g). Turtles with a CL <145 mm were considered juveniles; those for which CL >145 mm were classified as males or females on the basis of external morphological features (see Kennett & Georges 1990). All turtles were released at their point of capture.

#### Anthropogenic impact

Anthropogenic impact was measured by calculating road density (km of road/km<sup>2</sup>) within 700 m of each of the 14 ponds using ArcGIS (version 9.3.1: ESRI 2009). This distance was based on typical movement distances of *C. longicollis* determined from previous studies in the region (Rees *et al.* 2009; Roe *et al.* 2009). If the buffer encompassed areas within the predator-proof fence, the buffer was rearranged following the fence line, as the fence completely impedes turtle movements (Ferronato *et al.* 2014).

#### Primary and secondary productivity

We estimated a surrogate for primary productivity by measuring total phosphorus and nitrogen (TP and TN; ~0.2 L) from water samples in each pond, once per month from December 2012 to February 2013. Water samples were kept on ice in an insulated container during transportation to the lab and analysed using oxidation with K2S2O8 and low-pressure microwave digestion (Maher et al. 2002). We measured secondary productivity as the standing-crop biomass of potential prey items, sampling wetlands once per month from December 2012 to February 2013. On each occasion, we conducted four time-constrained (30 s) searches in each pond by agitating the sediment and searching in the littoral zone around available structures (e.g. rocks, debris, macrophytes) with the use of a  $34 \times 28$  cm dipnet (250 µm mesh; Roe et al. 2011). Samples were preserved in 90% ethanol for later sorting (see Roe et al. 2011). We only considered potential prey items that are known to be eaten by C. longicollis (Georges et al. 1986). We compared primary (TP and TN) and secondary (prey biomass) productivity among study sites using analysis of variance (ANOVA). TP and prey biomass were log10 transformed and TN was square root transformed to meet the assumptions of normality.

#### Reproduction

Adult female turtles were transported to the University of Canberra for radiographic examination (AJEX Meditech Ltd; Model: AJEX160H; settings: 50 kV, 1.20 mAs, 0.02 s, 70 cm high) and then released within 7 days at their point of capture. Egg length (EL) and egg width (EW) were measured with callipers from the X-ray films, and egg volume (EV) was estimated with the formula

$$\mathrm{EV} = \pi * X * Y^2 / 6$$

where X is the EL and Y is the EW (Vanzolini 1977). The proportion of mature females that were gravid was compared among sites with a chi-square contingency analysis. We used analysis of covariance (ANCOVA) to test whether clutch sizes differed among sites, with site as the factor, clutch size as the response variable and CL as a covariate.

#### Growth rates and movements

We calculated growth rates over both long-term (individuals initially encountered in 2006–2007 sampling and recaptured in 2011–2014, spanning both the drought and wet periods) and short-term (individuals initially encountered and recaptured in 2011–2014 sampling, during the wet period only) intervals. We then compared growth rates among turtles from natural, rural and suburban areas considering the long-term and the short-term scenarios. Annual growth was measured as change in CL, divided by the fraction of the 6-month growing season (15 September to 15 March) that had elapsed between captures. We only included individuals in the analysis if they were recaptured in the same study area and if the period between captures spanned at least one-half of a growing season.

Individuals were determined to have grown appreciably only if the growth increment exceeded the accuracy of measurements ( $\pm 0.5$  mm), and the proportion of individuals that grew appreciably was also determined for each study site. Data from individuals that had not grown appreciably were otherwise excluded from analyses of growth rates. The analysis of growth was as in the previous C. longicollis study during drought (2006-2007) to allow comparisons (Roe et al. 2011). The proportion of individuals showing appreciable growth was compared among study areas with a series of chi-square contingency analyses (juveniles and adults separate). Growth rates were compared among sites using ANCOVA, with site as the factor,  $log_{10}$  carapace growth rate as the response variable and initial CL as a covariate. Growth rates analyses were performed for both long-term and short-term recaptures.

We assessed if recaptured individuals had moved among study sites (not including movements between ponds within a study system), considering both long- and short-term recapture intervals. We calculated minimum straight-line distances animals moved with ArcGIS (version 9.3.1: ESRI 2009).

#### **Demographic parameters**

We compared proportion of females, estimated population size, size–frequency distributions, survivorship and recapture probability among study sites. Proportion of females was compared with ANOVA, with site as the factor and proportion of females as the response variable. For estimation of population size, we used the Horvitz-Thompson type estimator (Seber 1982):

$$N=\frac{n}{p}$$

where *N* is the estimated population size, *n* is the number of unique turtle captures in each pond and *p* is the capture probability. Population size was compared with ANCOVA, with site as the factor, estimated population size as the response variable and pond surface area as a covariate. Overall differences in size–frequency distributions among sites were examined with a chi-square test using the PROC FREQ procedure in SAS. We followed the overall test with a series of chi-square tests to examine in which size classes differences occurred. We used the Dunn–Sidak correction to adjust the per comparison significance ( $\alpha < 0.004$ ) to accommodate compounding error that occurs in multiple related comparisons.

Survivorship and capture probability were estimated using Cormack-Jolly-Seber (CJS) open population capture-recapture models in Program MARK. We estimated parameters among groups (adult male, adult and subadult female, and juvenile), sites (nature reserve, rural and suburb) and over time (sampling occasions). We collapsed capture histories into two occasions of approximately equal duration per year (September-December and January-March) owing to the different sampling effort in our fixed and occasional trapping sites. We started with models where survivorship  $(\Phi)$  and capture probability (p) were allowed to vary over time, among groups and among sites. We then fitted a series of reduced parameters models and ranked them based on Akaike's information criterion (AIC). If competing models had AIC values ≤2.0, we considered them as having some support (Lebreton et al. 1992). We assessed the fully saturated model's adequacy to describe the data using a bootstrap goodness-of-fit test with 500 simulations, and an overdispersion parameter  $(\hat{c})$  was derived by dividing the model deviance by the mean of the simulated deviances (Cooch & White 2014). If there was evidence for overdispersion  $(\hat{c} > 1)$ , we adjusted the models with the derived  $\hat{c}$  to improve model fit and calculated a quasi-likelihood estimator, QAIC<sub>c</sub> (Burnham & Anderson 1998). All parameters were estimated using model averaging.

#### Data analysis

Statistical analyses were performed with SPSS (Version 21), Program MARK version 7.1 (White & Burnham 1999)

and SAS Version 9.1 (SAS Institute 1999). The assumptions of normality and homogeneity of variances were checked by analysis of residuals, and when data failed to meet these assumptions, data were transformed to approximate normal distributions and equal variances; otherwise non-parametric tests were used. Statistical significance was accepted at the  $\alpha = 0.05$  level unless specified otherwise.

#### RESULTS

#### Anthropogenic impact

Road density around sampled ponds ranged from 15.7 to 19.3 km/km<sup>2</sup> at the urban site, from 0.0 to  $5.8 \text{ km/km}^2$  at the rural site, and no roads were within 700 m of ponds in the nature reserve.

### Primary and secondary productivity

The nature reserve and rural ponds had TP and TN similar to one another and higher than suburban ponds (TP: ANOVA:  $F_{2,39} = 10.08$ , P < 0.001; TN: ANOVA:  $F_{2,39} = 47.90$ , P < 0.001; Table 1). However, there was no difference in prey biomass among sites (ANOVA:  $F_{2,39} = 0.35$ , P = 0.70; Table 1).

## Reproduction

Shelled eggs were detected only from October to December in each year (Fig. 2). Of 299 adult females captured during the breeding season, only 8.4% were gravid, and the percentage of gravid females did not vary significantly among sites (nature reserve: 10.7%; rural 5.8%; urban: 8.4%) ( $X^2 = 0.70$ , d.f. = 2, P = 0.71). After controlling for carapace length, clutch sizes were similar among study sites (ANCOVA site:  $F_{2,19} = 0.72$ , P = 0.50; CL:  $F_{1,19} = 24.57$ , P < 0.005; Table 2). The interaction between site and CL was not significant (P = 0.60) and was dropped from the analysis.

**Table 1.** Primary and secondary productivity measurements in ponds inhabited by *Chelodina longicollis* among study sites,

 Australian Capital Territory, Australia

	Prir	Primary			
	TP (mg/L)	TN (mg/L)	Prey biomass (g)		
Nature reserve $(n = 15)$ Rural $(n = 15)$ Suburb $(n = 12)$	$\begin{array}{c} 0.11\pm0.01^{\rm A}(0.040.26)\\ 0.08\pm0.01^{\rm A}(0.050.15)\\ 0.05\pm0.01^{\rm B}(0.020.09) \end{array}$	$\begin{array}{c} 1.42 \pm 0.09^{\rm A} \ (0.892.28) \\ 1.47 \pm 0.08^{\rm A} \ (1.032.01) \\ 0.61 \pm 0.03^{\rm B} \ (0.460.76) \end{array}$	$\begin{array}{c} 2.66 \pm 0.39^{A} \; (0.96{-}5.50) \\ 3.35 \pm 0.58^{A} \; (0.27{-}6.58) \\ 2.52 \pm 0.56^{A} \; (0.52{-}6.89) \end{array}$		

Mean, SE; superscripts indicate similarities or differences among study sites within columns.



**Fig. 2.** Gravid females *Chelodina longicollis* inspected through X-ray in nature reserve (a), rural (b) and suburban (c) sites, in Australian Capital Territory, Australia. X-rays performed from October 2011 to March 2012, September 2012 to March 2013 and September 2013 to March 2014.

	CS (n)	EL (mm)	EW (mm)	EV (mm <sup>3</sup> )
Nature reserve	10.8 ± 0.8 (8.0–14.0)	29.9 ± 0.7 (28.1–31.7)	21.1 ± 0.3 (20.1–22.5)	7009 ± 312 (6151-8319)
(n = 7) Rural	$14.2 \pm 0.9 \; (13.017.0)$	$29.7\pm1.4(27.534.0)$	21.1 ± 0.3 (20.1–21.5)	6942 ± 501 (5822–8254)
(n = 4) Suburb	$12.5\pm0.8(8.017.0)$	$29.7\pm0.4(27.432.6)$	$20.7\pm0.3(19.122.7)$	$6713 \pm 207 \; (5762 – 8088)$
(n = 12)				

Table 2. Clutch size and egg measurements of gravid female *Chelodina longicollis* (through X-ray evaluation) from different study sites, Australian Capital Territory, Australia

Mean, SE, range; CS, clutch size; EL, egg length; EW, egg width; EV, egg volume.

**Table 3.** Growth rates of eastern long-necked turtles (*Chelodina longicollis*), and recaptures for juveniles (J) and adults (A), spanning long-term (drought-wet) and short-term (wet) conditions, Australian Capital Territory, Australia

	Group	Recaptures (n)		Percentage growing		Carapace growth rate (mm year <sup>-1</sup> ) <sup>†,‡</sup>	
Period		J	Α	J	Α	Ν	Mean $\pm$ SE (range)
Long-term 2006–2014	Nature reserve Rural	5 5	19 11	100 80	47 73	14 12	$\begin{array}{c} 4.4^{\rm A} \pm 1.0 \; (0.812.4) \\ 3.8^{\rm B} \pm 0.8 \; (0.59.1) \end{array}$
Short-term 2011-2014	Suburb Nature reserve Rural Suburb	3 4 8 11	19 25 3 24	100 100 100 100	74 48 67 75	17 16 10 29	$\begin{array}{l} 4.5^{\rm C} \pm 0.8 \; (0.9{-}13.9) \\ 5.7^{\rm A} \pm 1.7 \; (0.9{-}27.3) \\ 8.0^{\rm A} \pm 3.0 \; (0.8{-}26.8) \\ 7.7^{\rm A} \pm 1.4 \; (0.6{-}28.0) \end{array}$

<sup>†</sup>Based on a growth year spanning the typical activity season (15 September to 15 March). <sup>‡</sup>Carapace growth values not adjusted for carapace length variation (ANCOVA), which if included reveals statistical differences among study sites during the long-term period, but not during the short-term. Superscripts indicate similarities or differences among study sites within columns.

#### Growth and movements

After controlling for CL, there was a difference in turtle growth rates among sites during the long-term interval spanning both dry and wet periods (ANCOVA site:  $F_{2,39} = 12.49$ , P < 0.005; CL:  $F_{1,39} = 95.21$ , P < 0.005), with urban turtles growing fastest, followed by rural, and then nature reserve turtles (Table 3, Fig. 3). But there was no difference in growth rates among study sites during the short-term interval coinciding with the wet period only (ANCOVA site:  $F_{2,51} = 2.22$ , P = 0.12; CL:  $F_{1,51} = 32.49$ , P < 0.005; Table 3, Fig. 3). The percentage of juveniles and adults growing appreciably did not differ among study sites during the long-term (juveniles:  $X^2 = 1.73$ , d.f. = 2, P = 0.42; adults:  $X^2 = 3.37$ , d.f. = 2, P = 0.18) or short-term intervals (juveniles: not computed as growth was a constant; adults:  $X^2 = 3.80$ , d.f. = 2, P = 0.14; Table 3). We recorded long-distance movements of turtles through capturemark-recapture, spanning 2006-2014 (Table 4). Over the short-term interval (2011-2014), none of the turtles was recaptured in different study areas.

#### -

# **Demographic parameters**

We made 782 captures of 655 different turtles. There was no difference in proportion of females (mean, SE, n, range) among sites (nature reserve:  $0.49 \pm 0.18$  (*n* = 5, 0.28–0.72); rural: 0.38 ± 0.14 (n = 4, 0.21-0.54); suburb:  $0.45 \pm 0.12$  (n = 4, 0.12)0.29–0.57)) (ANOVA:  $F_{2,10} = 0.35$ , P = 0.60). Relative population sizes (corrected for variation in capture probability) increased from nature reserve to rural to urban study areas, but after controlling for wetland surface area, population sizes (i.e. densities) did not differ among sites (nature reserve:  $47.6 \pm 17.5$ (n = 5, 14.0-110.6); rural:  $110.2 \pm 36.6$  (n = 5, 14.0-110.6);10.2–224.3); suburb: 156.3  $\pm$  35.3 (*n* = 4, 74.2– 231.0); ANCOVA: site:  $F_{2,10} = 2.19$ , P = 0.16; wetland surface area:  $F_{1,10} = 1.71$ , P = 0.22). The interaction between site and wetland surface area was not significant (P = 0.09) and was removed from the analysis.

Size-frequency distributions differed among sites (overall  $X^2 = 87.2$ , d.f. = 24, P < 0.001), with significant differences within 60.1–75 mm PL ( $X^2 = 12.0$ , d.f. = 2, P < 0.004) and 105.1–120 mm PL size



**Fig. 3.** Relationships of growth and initial carapace length (mm) in *Chelodina longicollis* inhabiting nature reserve (open circles, smaller black dashed line), rural (black filled circles, solid line) and suburban (grey filled circles, larger grey dashed line) habitats, during 2006–2014 period (a) and 2011–2014 (b) period, Australian Capital Territory, Australia.

classes ( $X^2 = 14.1$ , d.f. = 2, P < 0.004), with more individuals in the rural site in both cases, as well as in the 165.1–180 mm PL size class, with more individuals in the rural and urban sites than the nature reserve ( $X^2 = 15.0$ , d.f. = 2, P < 0.004; Fig. 4).

In the capture-mark-recapture analysis, the model with most support had survivorship constant over

time and among groups and sites, and capture probability varying according to site (Tables 5,6). The other competing models had little support according to  $\Delta$  QAIC<sub>c</sub> values (Table 5). Analyses using only the fixed trapping sites supported the same highest ranked model as analyses using both fixed and occasional sampling sites.

# DISCUSSION

Populations of plant and animal species can be expected to wax and wane under the influence of climate, extending their ranges during more favourable periods, contracting during less favourable periods (Davis & Shaw 2001; Huntley et al. 2008). In the context of drought, critical elements in this process are patches of local persistence during the dry periods, and connectivity with suitable habitat during the wet periods. Often species, both terrestrial and aquatic, will contract and expand their distributions along drainage lines (Burridge et al. 2006; Smit et al. 2007), and maintaining connectivity is an important management objective (Pringle 2001). Urbanization has a major influence on whether or not local refugia for many species during drought are maintained and in what condition, and to what degree dispersal of species through the urban environment is prevented or impeded. Our study and those that preceded it (Roe et al. 2011; Ferronato et al. 2014) establish clearly the importance of these factors for the persistence of the eastern long-necked turtle in urban areas within its natural range.

Long-term or repeat studies of turtles inhabiting urban landscapes have focused specifically on demographic parameters and vital rates before and after major habitat alterations (Plummer & Mills 2008; Eskew et al. 2010). While we were not able to examine turtle populations before urban development, our study is unique in that we evaluated turtle responses not only to changes in degree of urban development over time and space but also to other potentially interacting stressors such as climate and invasive predators. Our recent sampling from 2011 to 2014 coincided with higher rainfall (mean 602 mm year $^{-1}$ , La Niña event, Beard et al. 2011) and a marked increase in urbanization compared to an earlier 2006-2007 sampling when the system was in drought (mean 464 mm year<sup>-1</sup>, El Niño period, van Dijk et al. 2013) and in the early stages of urban development (Rees et al. 2009; Roe et al. 2011). In addition, a fence was recently built to exclude non-native animals and isolate the nature reserve from encroaching urban hazards with substantial impact on turtle movements (Ferronato et al. 2014). Such a longitudinal study across a gradient of anthropogenic impact and following such changes allowed us to

		Recaptured in 2011–2014 <sup>†</sup> ( <i>n</i> )				
Group	Trapped in 2006–2007 (n)	NR	R	S	Distance moved (m) Mean $\pm$ SE (range)	
NR	32	_	_	8	$1446.2 \pm 431 \; (540 – 3800)$	
R	17	_	-	_	_	
S	28	3	4	_	3118.8 ± 582 (1575-6020)	

 Table 4.
 Chelodina longicollis movements among study sites based on capture-mark-recapture, Australian Capital Territory,

 Australia
 Capital Territory,

Denotes recaptures outside its original site of capture. NR, nature reserve; R, rural; S, suburb.



Fig. 4. Size-frequency distributions of *Chelodina longicollis* among study sites, Australian Capital Territory, Australia. Asterisk indicated statistical difference.

**Table 5.** Models of survivorship ( $\Phi$ ) and capture probability ( $\rho$ ) of *Chelodina longicollis* over time (twice per year), among sites (nature reserve, rural and suburb) and among groups (adult male, adult and subadult female, and juvenile) in the Australian Capital Territory, Australia, 2011–2014

Model	QAIC <sub>c</sub>	$\Delta \text{ QAIC}_{c}$	Weight	Parameters	Deviance
$\Phi$ (.) $\rho$ (site)	491.7	0.00	0.52	4	115.8
$\Phi$ (site) $\rho$ (.)	493.8	2.03	0.19	4	117.8
$\Phi(.) \rho(.)$	494.7	2.97	0.12	2	122.8
$\Phi$ (group) $\rho$ (.)	495.6	3.81	0.08	4	119.6
$\Phi$ (.) $\rho$ (group)	496.4	4.62	0.05	4	120.4
$\Phi$ (site × group) $\rho$ (.)	498.1	6.37	0.02	10	109.9
$\Phi$ (.) $\rho$ (site $\times$ group)	498.6	6.88	0.02	10	110.4
$\Phi$ (site × group) $\rho$ (site × group)	508.0	16.26	0.00	18	103.1
$\Phi$ (.) $\rho$ (site × group × time)	552.6	60.89	0.00	46	85.9
$\Phi$ (site × group × time) $\rho$ (.)	554.1	62.40	0.00	46	87.5
$\Phi$ (site × group × time) $\rho$ (site × group × time)	641.8	150.06	0.00	90	66.5

Models were compared and ranked with a quasi-likelihood Akaike's information criterion (QAIC<sub>c</sub>) estimator corrected for overdispersion ( $\hat{c} = 1.51$ ).

examine spatial and temporal responses in behaviour, demography and vital rates, yielding insight into the mechanisms related to turtle persistence in urban systems. The evidence of increased urbanization in the system (Gungahlin suburbs) over the last 8 years includes a 79% growth in human population (Australian Bureau of Statistics 2013), a 76% increase in

doi:10.1111/aec.12462

Site	Group	$\Phi$ (bi-annual)	$\Phi$ (annual)	$\rho^{\dagger}$ (bi-annual)
Nature reserve	Male	$0.688 \pm 0.076$	$0.473 \pm 0.005$	$0.117 \pm 0.028$
	Female	$0.702\pm0.074$	$0.492 \pm 0.005$	$0.121 \pm 0.029$
	Juvenile	$0.687 \pm 0.076$	$0.471 \pm 0.005$	$0.118 \pm 0.029$
Rural	Male	$0.648 \pm 0.078$	$0.419 \pm 0.006$	$0.077 \pm 0.022$
	Female	$0.652 \pm 0.077$	$0.425\pm0.005$	$0.079\pm0.022$
	Juvenile	$0.639 \pm 0.079$	$0.408 \pm 0.006$	$0.077 \pm 0.022$
Suburb	Male	$0.671 \pm 0.078$	$0.450\pm0.006$	$0.099 \pm 0.029$
	Female	$0.686 \pm 0.076$	$0.470\pm0.005$	$0.102 \pm 0.029$
	Juvenile	$0.668 \pm 0.079$	$0.446\pm0.006$	$0.101 \pm 0.029$

**Table 6.** Estimates of survivorship  $(\Phi)$  and capture probability  $(\rho)$  for *Chelodina longicollis* among different sites and groups in the Australian Capital Territory, Australia, 2011–2014

<sup>1</sup>Capture probabilities showed differences among sites according to model selection. Parameters were derived as weighted averages based on their quasi-likelihood Akaike's information criterion (QAIC<sub>c</sub>) values, adjusted for model overdispersion. Results expressed in mean  $\pm$  SE.

traffic volume (Territory and Municipal Services, R. Shoukrallah, pers. comm., 2013) and a 130% higher road density compared to the 2006–2007 period (Roe *et al.* 2011). Such an increase in urbanization, especially regarding road density and traffic volume, could pose a threat for turtles with marked propensity for overland movements, although our measures of various behavioural, demographic and population vital rates together with those of earlier studies suggest otherwise (Roe *et al.* 2011). The apparent resilience of *C. longicollis* to such threats is in contrast with demographic responses of several other species to heavy road density and traffic volume elsewhere (Gibbs & Shriver 2002; Marchand & Litvaitis 2004).

Although we considered habitats in the context of being more or less influenced by anthropogenic stressors, we were also interested in understanding potential differences in productivity. Urban areas may have higher productivity and nutrient loads owing to human subsidized resources (DeStefano & DeGraaf 2003; Shochat et al. 2006), which can ultimately influence growth rates and reproductive output in turtles (Brown et al. 1994; Lindeman 1996). This was not the case in our system, where urban ponds had the lowest primary productivity, a result that could have been influenced by differences in the uptake of resources by organisms and variation in the availability of nutrients during sampling (Jones 1984; Müller 2000). However, availability of food resources for turtles was similar among study sites, suggesting that productivity did not differ in ways relevant to turtle population regulation, a finding in agreement with previous estimates of food availability during drought (Roe et al. 2011). The lack of elevated productivity in urban ponds may be related to the lack of sewage contamination in our system, which can be a significant source of nutrient input in urban areas elsewhere (Galbraith et al. 1988; Souza & Abe 2000).

Our results demonstrate the strong influence of drought on growth rates. During drought, turtles in urban areas grew five times faster than those in the nature reserve (Roe et al. 2011), even though ponds were similar in prey biomass when flooded. The authors hypothesized that the constant availability of water extended the activity period (and thus foraging opportunities) for turtles in suburbs, while ponds in the nature reserve dried and most turtles aestivated on land. Our results support this hypothesis, as growth rates were similar between natural and urban ponds during the wet period, when all ponds remained flooded and no turtles were observed in long-term aestivation (Ferronato et al. 2016). Interestingly, long-term growth rates spanning both dry and wet periods remained lowest in nature reserve turtles, indicating a lack of compensatory growth following the return of rainfall. Such responses suggest that the cumulative effects of several wet-dry cycles may further constrain individual growth rates in nature reserve populations relative to their urban counterparts, though we were only able to measure growth responses over one wet-dry cycle. Indeed, growth rates of animals, including C. longicollis, are strongly influenced by rainfall patterns in wet-dry cycles characteristic of much of Australia (Kennett & Georges 1990; Madsen & Shine 2000; Greenville et al. 2013).

Turtles in urban systems may have higher fecundity than in natural settings (Brown *et al.* 1994; Lindeman 1996), but we did not detect variation among sites in any measure of reproductive biology, including reproductive season, clutch sizes or percentage of gravid females. That our measures of fecundity were similar across environments is again likely related to similar food availability among sites. *Chelodina longicollis* can lay up to three clutches in a reproductive season in the Murray River and in Gippsland, Victoria (Parmenter 1985; Kennett *et al.* 2009), but we found no evidence of multiple clutches based on X- ray analysis of females recaptured within a breeding season, a finding corroborated by a previous study in the Canberra region (Vestjens 1969) and likely reflecting climactic constraints on a shortened reproductive season (October to December). Even though we do not have information on reproductive output of *C. longicollis* during drought in our system, females aestivate on land for several months in response to wetland drying (Rees *et al.* 2009) and may cease reproduction during unfavourable drought conditions at other locations in southeastern Australia (Kennett & Georges 1990).

The observation of similar population sizes among study sites at first suggests a different dynamic from the previous drought (2006-2007), where urban turtles were nearly three times more abundant than nature reserve turtles (Roe et al. 2011). However, despite statistical analyses, abundance in nature reserve ponds was still 3.2 and 2.3 times lower than the urban and rural sites, respectively, a difference that could be biologically meaningful but biased by low sample size. For instance, even though the same ponds were sampled in each study, sample sizes in the wet period (2011-2014) were smaller owing to the construction of the fence that required the natural site to be divided into two regions, reducing power in the analysis. While nature reserve turtles resumed growth and reproduction during the recent wet conditions, perhaps not enough time had passed for a population-level response to be realized. Additionally, at the same time that the drought broke, the predator-proof fence was erected, isolating that population and preventing remigrations of individuals that had left for the suburban ponds during drought. Indeed, many more turtles were encountered on the outside of the fence following the return of rainfall, which likely represents individuals attempting to return to the flooded ponds in the nature reserve (Ferronato et al. 2014). Thus, immigration into nature reserve ponds was eliminated, causing both high mortality and forcing them into other ponds (Ferronato et al. 2014, 2016).

The observation of animals in the smaller size classes in all study sites indicates that recruitment has continued despite expanded urbanization. One of the typical challenges facing turtles in urban land-scapes is limited recruitment owing to high predation rates and lack of nesting habitats (Spinks *et al.* 2003; Marchand & Litvaitis 2004). The presence of recruitment across all levels of anthropogenic impact examined here is a signal that some females are still safely reaching nesting areas, eggs are successfully incubated, and some hatchlings are capable of travelling to water. Although survivorship did not vary across study sites based on capture–mark–recapture estimates, a concurrent radiotelemetry study demonstrated that adult female urban turtles had lower

annual survivorship (0.67) compared with females from the nature reserve (1.00, Ferronato et al. 2016), where most mortality of urban turtles was on roads. However, the radiotelemetry study focused only on females, and we identified localized hotspots that could have biased mortality differences in the telemetry owing to small sample sizes (Ferronato et al. 2016). The overall mortality in the broader study area could be diffuse at the meta-population scale, yet still significant on local scales. Regardless, survivorship estimates during the wet period (2011-2014) are especially low for C. longicollis (Roe et al. 2009) and compared to other freshwater turtles (Shine & Iverson 1995), so we question the accuracy of these survival rates. Considering the biology of C. longicollis and its ability for frequent and long-distance inter-wetland movements (Ryan & Burgin 2007; Roe & Georges 2008b; Roe et al. 2009), there is potential for high emigration to ponds outside the sample locations, which would be interpreted as 'mortalities' in our CJS models (Cooch & White 2014). We did not consider using the Robust Design approach to account for emigration, as females were temporarily removed from the populations for X-ray analysis, which would have violated the assumptions of such models.

Previous research has demonstrated the high vagility of C. longicollis, suggesting that single wetlands should not represent the minimum habitat unit harbouring a population (Roe & Georges 2008b; Roe et al. 2009). Rare dispersal events of up to 5.2 km have been described among dune lakes in an undisturbed setting in southeastern Australia (Roe et al. 2009), and it is apparent that such long-distance movements (up to about 6 km) also occur within urbanized regions, suggesting that the turtles behave similarly with regard to inter-wetland movements for dispersal or migration even where roads and other threats are encountered. Such movements are important for maintaining connectivity and gene flow among populations (Hansson 1991; Coulon et al. 2004), in addition to rescue populations from stochastic events such as drought. However, the current design of the pest-fencing surrounding the nature reserve is disrupting this dynamic in response to wet-dry cycles (Rees et al. 2009; Roe et al. 2011; Ferronato et al. 2014), and causing high mortality for turtles attempting to pass (Ferronato et al. 2014). Nevertheless, these long-distance movements demonstrate that our sites do not satisfy the assumption of independence. However, such important behaviour would have not been documented in short-term studies or if we had only sampled wetlands distant from each other (e.g. 10 km apart). Moreover, another limitation in our design is that the construction of the fence enclosure in the nature reserve in 2009 isolated that population from exchange with other nearby ponds.

### CONCLUSIONS

Our study expands the current understanding of vertebrate responses to drought cycles within natural-urban gradients and demonstrates the strong influence of rainfall on population dynamics. Together, our findings of similar vital rates, demography and recruitment in all study sites indicate that C. longicollis is resilient to the levels of urbanization in our system. Perhaps, the ability of C. longicollis to move overland and settle in different habitats (Kennett & Georges 1990; Roe & Georges 2008a,b; Roe et al. 2009), in addition to its opportunistic carnivorous feeding behaviour (Georges et al. 1986) have also contributed to its successful colonization in urban ponds. As long as turtles can travel safely between natural and urban environments, climate cycles similar to those observed in this study should pose no threat to their persistence in such habitats. A major limitation of our study is the lack of adequate replication, with our study system covering only one gradient of urbanization in a single urban environment. However, taken together with studies in urban environments elsewhere, evidence of C. longicollis resilience and establishment across a range of urban settings over broad spatial scales is growing (Rvan & Burgin 2007; Rees et al. 2009; Roe et al. 2011; Stokeld et al. 2014). Finally, even though our repeat study compared population dynamics over an 8-year period, such duration is still relatively short compared with the turtle life spans that may cover several decades (Gibbons 1987), making further monitoring essential for a more complete understanding of turtle population dynamics in urban landscapes.

# ACKNOWLEDGEMENTS

We thank the staff of Gungahlin Lakes Golf Club (S. Dawson, Wade), Ginninderra Experimental Station (P. Dumbar) and Mulligans Flat Nature Reserve (P. Mills, G. Woodbrigde, J. Lawler) for their logistic support and to colleagues who shared their expertise in some topics in this study (A. Lopez-Aldana: stats; E. Harrison: Freshwater lab; B. Maher, F. Krikowa, V. Vysna: Ecochemistry lab; B. Gruber: MARK Program). Research was conducted with the appropriate approvals and permits from the University of Canberra Committee for Ethics in Animal Experimentation and Environment ACT. The study was funded by the Institute for Applied Ecology, the ACT Herpetological Association and the Turtles Australia Inc. BOF was sponsored by an Australian Endeavour International Postgraduate Research Scholarship and a University of Canberra W.J. Weeden Scholarship.

#### REFERENCES

- Australian Bureau of Statistics (2013) Census Quick Stats. [Cited 19 Dec 2013.] Available from URL: http:// www.abs.gov.au/websitedbs/censushome.nsf/home/data?open document&navpos=200.
- Bateman P. W. & Fleming P. A. (2012) Big city life: carnivores in urban environments. J. Zool. 287, 1–23.
- Beard G., Chandler E., Watkins A. B. & Jones D. A. (2011) How does the 2010–2011 La Niña compare with past La Niña events? *Bull. Aust. Meteorol. Oceanogr. Soc.* 24, 17– 20.
- Bond A. R. & Jones D. N. (2008) Temporal trends in use of fauna-friendly underpasses and overpasses. Wildlife Res. 35, 103–12.
- Brown G. P., Bishop C. A. & Brooks R. J. (1994) Growth rate, reproductive output, and temperature selection of snapping turtles in habitats of different productivities. *J. Herpetol.* 28, 405–10.
- Burnham K. P. & Anderson D. R. (1998) Model Selection and Inference: A Practical Information-Theoretic Approach. Springer-Verlag, New York.
- Burridge C. P., Craw D. & Waters J. M. (2006) River capture, range expansion, and cladogenesis: the genetic signature of freshwater vicariance. *Evolution* **60**, 1038–49.
- Chace J. F. & Walsh J. J. (2006) Urban effects on native avifauna: a review. Landscape Urban Plan. 74, 46–69.
- Clevenger A. P. & Waltho N. (2005) Performance indices to identify attributes of highway crossing structures facilitating movement of large mammals. *Biol. Conserv.* 121, 453–64.
- Congdon J. D., Dunham A. E. & Van Loben Sels R. C. (1994) Demographics of common snapping turtles (*Chelydra* serpentina): implications for conservation and management of long-lived organisms. *Am. Zool.* 34, 397–408.
- Cooch E. & White G. (2014) Using MARK a Gentle Introduction, 13th edn. [Cited 10 Oct 2014.] Available from URL: http:// www.phidot.org/software/mark/docs/book/.
- Coulon A., Cosson J. F., Angibault J. M. et al. (2004) Landscape connectivity influences gene flow in a roe deer population inhabiting a fragmented landscape: an individual-based approach. Mol. Ecol. 13, 2841–50.
- Davis M. B. & Shaw R. G. (2001) Range shifts and adaptive responses to quaternary climate change. *Science* **292**, 673–9.
- DeCatanzaro R. & Chow-Fraser P. (2010) Relationship of road density and marsh condition to turtle assemblage characteristics in the Laurentian Great Lakes. *J. Great Lakes Res.* **36**, 357–65.
- DeStefano S. & DeGraaf R. M. (2003) Exploring the ecology of suburban wildlife. *Front. Ecol. Environ.* **1**, 95–101.
- van Dijk A. I. J. M., Beck H. E., Crosbie R. S. et al. (2013) The Millennium Drought in southeastern Australia (2001– 2009): natural and human causes and implications for water resources, ecosystems, economy, and society. Water Resour. Res. 49, 1040–57.
- Eskew E. A., Price S. T. & Dorcas M. E. (2010) Survivorship and population densities of painted turtles (*Chrysemys picta*) in recently modified suburban landscapes. *Chelonian Conserv. Biol.* 9, 244–9.
- ESRI (Environmental System Resource Institute) (2009) ArcMap 9.3.1. ESRI, Redlands.
- Ferronato B. O., Roe J. H. & Georges A. (2014) Reptile bycatch in a pest-exclusion fence established for wildlife reintroductions. J. Nat. Conserv. 22, 577–85.

© 2016 Ecological Society of Australia

- Ferronato B. O., Roe J. H. & Georges A. (2016) Urban hazards: spatial ecology and survivorship of a turtle in an expanding suburban environment. *Urban Ecosyst.* **19**, 415–28.
- Gakenheimer R. (1999) Urban mobility in the developing world. *Transp. Res. Part A Policy Pract.* 33, 671–89.
- Galbraith D. A., Bishop C. A., Brooks R. J., Simser W. L. & Lampman K. P. (1988) Factors affecting the density of populations of common snapping turtles (*Chelydra* serpentina). Can. J. Zool. 66, 1233–40.
- Georges A., Norris R. H. & Wensing L. (1986) Diet of the freshwater turtle *Chelodina longicollis* (Testudines: Chelidae) from the coastal dune lakes of the Jervis Bay Territory. *Aust. Wildlife Res.* 13, 301–8.
- Gibbons J. W. (1968) Population structure and survivorship in the painted turtle, *Chrysemys picta*. *Copeia* **1968**, 260–8.
- Gibbons J. W. (1987) Why do turtles live so long? *Bioscience* 37, 262–9.
- Gibbs J. P. & Shriver W. G. (2002) Estimating the effects of road mortality on turtle populations. *Conserv. Biol.* 16, 1647–52.
- Goad E. H., Pejchar L., Reed S. E. & Knight R. L. (2014) Habitat use by mammals varies along an exurban development gradient in northern Colorado. *Biol. Conserv.* 176, 172–82.
- Greenville A. C., Wardle G. M. & Dickman C. R. (2013) Extreme rainfall events predict irruptions of rat plagues in central Australia. *Austral Ecol.* 38, 754–64.
- Hamer A. J., Harrison L. J. & Stokeld D. (2016) Road density and wetland context alter population structure of a freshwater turtle. *Austral Ecol.* 41, 53–64.
- Hansson L. (1991) Dispersal and connectivity in metapopulations. Biol. J. Linn. Soc. 42, 89–103.
- Hughes L. (2003) Climate change and Australia: trends, projections and impacts. *Austral Ecol.* 28, 423–43.
- Huntley B., Collingham Y. C., Willis S. G. & Green R. E. (2008) Potential impacts of climatic change on European breeding birds. *PLoS One* **3**, e1439.
- Jones R. C. (1984) Application of a primary production model to epiphytic algae in a shallow, eutrophic lake. *Ecology* 65, 1895–903.
- Kennett R. M. & Georges A. (1990) Habitat utilization and its relation to growth and reproduction of the eastern longnecked turtle, *Chelodina longicollis* (Testudinata: Chelidae), from Australia. *Herpetologica* 46, 22–33.
- Kennett R., Roe J., Hodges K. & Georges A. (2009) Chelodina longicollis (Shaw 1784) – eastern long-necked turtle, common long-necked turtle, common snake-necked turtle. Chelon. Res. Monogr. 5, 031.1–8.
- Lebreton J. D., Burnham K. P., Colbert J. & Anderson D. R. (1992) Modelling survival and testing biological hypotheses using marked animals: a unified approach with case studies. *Ecol. Monogr.* **62**, 67–118.
- Lindeman P. V. (1996) Comparative life history of painted turtles (*Chrysemys picta*) in two habitats in the Inland Pacific Northwest. *Copeia* 1996, 114–30.
- Madsen T. & Shine R. (2000) Rain, fish and snakes: climatically driven population dynamics of Arafura filesnakes in tropical Australia. *Oecologia* **124**, 208–15.
- Maher W., Krikowa F., Wruck D., Louie H., Nguyen T. & Huang W. Y. (2002) Determination of total phosphorus and nitrogen in turbid waters by oxidation with alkaline potassium peroxodisulfate and low pressure microwave digestion, autoclave heating or the use of closed vessels in a hot water bath: comparison with Kjeldahl digestion. *Anal. Chim. Acta* 463, 283–93.

- Manning A. D., Wood J. T., Cunningham R. B. *et al.* (2011) Integrating research and restoration: the establishment of a long-term woodland experiment in south-eastern Australia. *Aust. Zool.* 35, 633–48.
- Marchand M. N. & Litvaitis J. A. (2004) Effects of habitat features and landscape composition on the population structure of a common aquatic turtle in a region undergoing rapid development. *Conserv. Biol.* 18, 758–67.
- Marchand M. N., Litvaitis J. A., Maier T. J. & DeGraaf R. M. (2002) Use of artificial nests to investigate predation on freshwater turtle nests. *Wildlife Soc. Bull.* 30, 1092–8.
- McKinney M. L. (2002) Urbanization, biodiversity and conservation. *Bioscience* 52, 883–90.
- McKinney M. L. (2008) Effects of urbanization on species richness: a review of plants and animals. Urban Ecosyst. 11, 161–76.
- Müller U. (2000) Periphytic primary production during spring. A sink or source of oxygen in the littoral zone? *Limnologica* 30, 169–74.
- Parmenter C. J. (1985) Reproduction and survivorship of Chelodina longicollis (Testudinata: Chelidae). In: Biology of Australasian Frogs and Reptiles (eds G. Grigg, R. Shine & H. Ehmann) pp. 53–61. Royal Society of NSW, Sydney.
- Pauchard A., Aguayo M., Peña E. & Urrutia R. (2006) Multiple effects of urbanization on the biodiversity of developing countries: the case of a fast-growing metropolitan area (Concepción, Chile). *Biol. Conserv.* 127, 272–81.
- Pautasso M. (2006) Scale dependence of the correlation between human population presence and vertebrate and plant species richness. *Ecol. Lett.* 10, 16–24.
- Plummer M. V. & Mills N. E. (2008) Structure of an urban population of softshell turtles (*Apalone spinifera*) before and after severe stream alterations. In: Urban Herpetology, Herpetological Conservation, Vol. 3 (eds R. E. Jung & J. C. Mitchell) pp. 95–105. Society for the Study of Amphibian and Reptiles, Salt Lake City.
- Pringle C. M. (2001) Hydrological connectivity and the management of biological reserves: a global perspective. *Ecol. Appl.* 11, 981–98.
- Rees M., Roe J. H. & Georges A. (2009) Life in the suburbs: behavior and survival of a freshwater turtle in response to drought and urbanization. *Biol. Conserv.* **142**, 3172–81.
- Roe J. H. & Georges A. (2007) Heterogeneous wetland complexes, buffer zones, and travel corridors: landscape management for freshwater reptiles. *Biol. Conserv.* 135, 67–76.
- Roe J. H. & Georges A. (2008a) Maintenance of variable responses for coping with wetland drying in freshwater turtles. *Ecology* 89, 485–94.
- Roe J. H. & Georges A. (2008b) Terrestrial activity, movements and spatial ecology of an Australian freshwater turtle, *Chelodina longicollis*, in a temporally dynamic wetland system. *Austral Ecol.* 33, 1045–56.
- Roe J. H., Brinton A. C. & Georges A. (2009) Temporal and spatial variation in landscape connectivity for a freshwater turtle in a temporally dynamic wetland system. *Ecol. Appl.* 19, 1288–99.
- Roe J. H., Rees M. & Georges A. (2011) Suburbs: dangers or drought refugia for freshwater turtle populations? J. Wildlife Manage. 75, 1544–52.
- Ryan M. & Burgin S. (2007) Gone walkabout? Movement of the eastern long-necked turtle *Chelodina longicollis* from farm dams in northwest peri-urban Sydney (Australia). J. *Biol. Res.-Thessalon* 8, 119–27.

- SAS Institute (1999) SAS Statistics. Version 8.2. SAS Institute, Cary.
- Seber G. A. F. (1982) The Estimation of Animal Abundance and Related Parameters, 2nd edn. Grifin, London.
- Shine R. & Iverson J. B. (1995) Patterns of survival, growth and maturation in turtles. *Oikos* 72, 343–8.
- Shochat E., Warren P. S., Stanley H. F., McIntyre N. E. & Hope D. (2006) From patterns to emerging processes in mechanistic urban ecology. *Trends Ecol. Evol.* 21, 186–91.
- Smit I. P. J., Grant C. C. & Devereux B. J. (2007) Do artificial waterholes influence the way herbivores use the landscape? Herbivore distribution patterns around rivers and artificial surface water sources in a large African savanna park. *Biol. Conserv.* 136, 85–99.
- Souza F. L. & Abe A. S. (2000) Feeding ecology, density and biomass of the freshwater turtle, *Phrynops geoffroanus*, inhabiting a polluted urban river in south-eastern Brazil. J. *Zool.* 252, 437–46.
- Spencer R. J. & Thompson M. B. (2005) Experimental analysis of the impact of foxes on freshwater turtle populations. *Conserv. Biol.* **19**, 845–54.
- Spinks P. Q., Pauly G. B., Crayon J. J. & Shaffer H. B. (2003) Survival of the western pond turtle (*Emys marmorata*) in an urban California environment. *Biol. Conserv.* 113, 257–67.

- Steen D. A., Aresco M. J., Beilke S. G. *et al.* (2006) Relative vulnerability of female turtles to road mortality. *Anim. Conserv.* 9, 269–73.
- Stokeld D., Hamer A. J., van der Ree R., Pettigrove V. & Gillespie G. (2014) Factors influencing occurrence of a freshwater turtle in an urban landscape: a resilient species? *Wildlife Res.* 41, 163–71.
- Takekawa J. E. & Beissinger S. R. (1989) Cyclic drought, dispersal, and the conservation of the snail kite in Florida: lessons in critical habitat. *Conserv. Biol.* **3**, 302–11.
- Vanzolini P. E. (1977) A brief biometrical note on the reproductive biology of some South American Podocnemis (Testudines, Pelomedusidae). *Pap. Avuls. Zool.* 31, 73–102.
- Vestjens W. J. M. (1969) Nesting, egg-laying and hatching of the snake-necked tortoise at Canberra, ACT. Aust. Zool. 15, 141–9.
- Webster R. & Butler B. E. (1976) Soil classification and survey studies at Ginninderra. Aust. J. Soil Res. 14, 1–24.
- White G. C. & Burnham K. P. (1999) Program MARK: survival estimation form populations of marked animals. *Bird Study* 46, 120–38.
- World Resources Institute, United Nations Environment Programme, United Nations Development Programme, The World Bank (1996) World Resources 1996–1997. The Urban Environment. Oxford University Press, Oxford.