INTRODUCTION

Humans perturb natural habitats in many ways, often degrading habitat quality to the point where native wildlife is unable to persist (Powers & Jetz, 2019; Sala et al., 2000). However, anthropogenic activities create opportunities as well as costs. One common modification is to create sites with an increased availability of food (resource subsidy) but with a higher risk of predation. For example, agricultural fields attract rats to feast on the crops, but at a greater risk of predation by avian predators (Labuschagne et al., 2016). In many cases, the attributes of the newly created habitat patch differ considerably from that of the ecosystem that they replaced, such that native biota adapted to the previous habitat type will be poorly suited to exploiting the new opportunity. As a result, severe habitat disturbance typically eliminates a high proportion of taxa found in an area (Brooks et al., 2002).

Interestingly, some native taxa (as well as many invasive taxa) thrive in the newly created habitat type (Gangoso et al., 2013). That success is unsurprising for invasive species, many of which exploit anthropogenically modified habitats in their native range and thus do not face the challenge of a niche shift after translocation (Chapple et al., 2011). However, the situation with native taxa is different. The mechanisms that allow that flexibility deserve scrutiny, if we are to understand why anthropogenic activities benefit some native species while disadvantaging others. In particular, which attributes allow a population to exploit a newly created niche, whose use is best-suited to individuals with phenotypic traits different from those required to exploit the pre-existing habitat?
One key to success may be phenotypic diversity within the local population. If a population consists of individuals of a wide range of morphologies (e.g., body sizes) and behavioral types (e.g., along the bold-shy continuum), then some individuals within that population may be well suited to the new opportunity that has arisen. Put simply, ecologically relevant variation among individuals within a population may confer an ecological breadth that pre-adapts that population to using a novel anthropogenically created habitat type. Why should such variation exist, if it does not adapt individuals to local ecological conditions? One possibility is that the resource hotspot advantages a trait that previously conferred an ecological disadvantage and was maintained only through intense sexual selection. For example, suppose that large body size and bold behavior in males enhance mating opportunities through increased success in male–male combat bouts (e.g., large male American rubyspot damselflies outcompete smaller males for territories and thus increase access to females: Serrano-Meneses et al., 2007). That situation may result in males evolving to grow to a size at which their maintenance energy requirements can barely be met; that is, the optimal body size from an energetic perspective is smaller than that favored by sexual selection. For example, reproductive success correlates with size in male marine iguanas, but large males suffer the largest mortality when environmental conditions decline (Wikelski & Trillmich, 1997). In such a system, the creation of a novel habitat type that is rich in food resources and most effectively utilized by a large animal might alleviate the energetic disadvantages of large size. If so, a population with a broad range of body sizes, previously maintained by sexual as well as natural selection, might flourish in a newly created resource hotspot.

Our study species—a large species of varanid (monitor) lizard in eastern Australia—is ideal for investigating this situation. These large (to >2 m) lizards are widely distributed through woodland habitats, and campgrounds scattered within that woodland. Attracted by resource subsidies from campers and picnickers, these lizards (Varanus varius, “lace monitors” or “goannas” in local parlance) sometimes attain high densities in campgrounds and refuse dumps, and individuals sampled within these highly disturbed open habitats differ in morphology from conspecifics in adjacent bushland (Amir, 2018; Jessop et al., 2012). Sexual size dimorphism in this species is extreme, with adult males averaging around 2.4 times heavier than adult females (5,320 g vs. 2,225 g; Carter, 1992; Guarino, 2002; Kirshner, 2007). In exceptional cases, males attain >14 kg (>6.2 times the average mass of females: Weavers, 1988). That huge body size likely reflects sexually selected advantages to larger males in wrestling matches for dominance and access to matings (Darwin, 1871; Frýdlová & Frynta, 2010). However, a population of lace monitors also contains much smaller individuals; hatchlings (snout–vent length [SVL] 120 mm, 23 g) are ecologically independent and tend to use different habitats and prey items than do adult conspecifics (King & Green, 1999). Thus, the provision of open resource-rich habitats potentially offers opportunities to goannas of a wide range of sizes. We predicted (based on the ideas above, plus previous studies: Amir, 2018; Jessop et al., 2012) that campground-dwelling individuals would be larger and also bolder than conspecifics in adjacent bushland regions. Because larger lizards experience high mortality with the invasion of toxic cane toads, Rhinella marina (Jolly et al., 2015), we also predicted that the arrival of cane toads (which are most abundant in disturbed habitats: González-Bernal et al., 2016) would lead to a reduction in mean body sizes and favor monitor lizards that were less bold.

In summary, then, we have a reptilian predator (the lace monitor) and a toxic invader (the cane toad) in a habitat matrix that includes patches of high resource availability (campgrounds) surrounded by large expanses of habitat lacking a resource subsidy (bushland). Surveys of campgrounds and bushland in sites with versus without cane toads allowed us to address three questions: (a) do lizards in campgrounds versus bushland differ in size and boldness?; (b) are habitat-associated differences in boldness a secondary consequence of differences in mean body size?; and (c) is the distribution of phenotypic traits across that habitat dichotomy affected by the arrival of an invasive species?

2 | MATERIALS AND METHODS

2.1 | Study species

Lace monitors are very large (to 14 kg, >2 m) semi-arboreal lizards with an extensive distribution across eastern Australia (Cogger, 2014; Weavers, 1988). Keystone predators and dietary generalists, lace monitors flexibly exploit seasonal resources, and key in on anthropogenic food subsidies (Jessop et al., 2012; Figure 1a). The longterm numerical impact of toads on lace monitors appears to be minor (especially, compared with impacts on other varanid species: Doody et al., 2009; Pettit et al., In review). Native to Central and South America, cane toads (Rhinella marina) were introduced into northeast Australian sugar cane fields in 1935 in an unsuccessful attempt to control insect pests (Zug & Zug, 1979). Once established, toads rapidly spread west through the wet-dry tropics of Queensland, the Northern Territory and Western Australia (Phillips et al., 2006, 2007; Urban et al., 2007), and slowly extended south along coastal regions of Queensland and New South Wales (Seabrook, 1991). The southern invasion front has expanded through a series of inadvertent and intentional introductions as well as slow-moving range expansion (van Beurden & Grigg, 1980; Seabrook, 1991). Cane toads have had disastrous impacts on many Australian taxa, especially large-bodied anuraphagous predators that are fatally poisoned if they ingest a toad (Smith & Phillips, 2006).

2.2 | Study area

We worked at 27 sites along the east coast of Australia (Figure 1b). Each site consisted of a campground and an adjacent 5 km road transect through native bushland. We estimated population sizes of lizards from 10 standardized visual encounter surveys per site (details below). Morphological estimates, habitat characteristics, and
FIGURE 1  A lace monitor (Varanus varius) (a) foraging at a campsite fire pit. Locations of field sites (b) within and beyond the current range of cane toads (Rhinella marina) along the east coast of Australia. Filled squares depict toad-free sites, whereas open triangles show sites that have been colonized by cane toads.
behavioral data were gathered from lizards during these surveys and from opportunistic observations. Seven of our sites lacked cane toads, whereas the other 20 had been invaded 1–80 years previously.

2.3 | Survey methodology

Surveys took place between October 2017 and April 2018 on days with maximum air temperature >23°C. Those five sampling days were split across a minimum of two sampling periods, stratified over time to minimize seasonal biases. Groups of one to three sites were surveyed concurrently for logistical reasons, with surveys of grouped sites, and sites within those groups, randomly ordered to further minimize sampling biases among sites with respect to season and time-of-day.

Diurnal surveys for lizards were conducted twice daily between 0800 and 1700 hr AEST. Each visual survey totaled 1-hr duration, consisting of a 15-min visual encounter survey on foot in and around the campground, and 45 min of observation by a single observer from a slow-moving vehicle (20–40 km/hr) along the 5 km road transect through bushland (see Jolly et al., 2016 for detailed transect survey methods). Nocturnal spotlight surveys were performed between 1900 and 2300 hr to confirm the presence or absence of toads via visual confirmation or calling males.

2.4 | Population sizes of lace monitors

To estimate the relative abundance of lace monitors in campgrounds and bushland at toad-absent and toad-present areas, we used mean sightings per 30 min from our 10 visual encounter surveys per habitat per site to account for differential survey effort between habitats (15 min in campgrounds, 45 min in bushland per survey).

2.5 | Body sizes of lace monitors

A single observer (LP) estimated the SVL of 247 lizards to the nearest 5 cm (Jolly et al., 2016; Lambert et al., 2012). We validated our size estimates for 65 goannas by placing a tape measure next to placid individuals (N = 3) or by using ImageJ (Ver. 2.0) to compare standardized photographs of the lateral side of a goanna versus a tape measure placed in the same location and orientation (N = 62). Our visual estimates accorded closely with these more precise measurements (ANOVA $R^2 = 0.89, F_{1.63} = 491.01, p < .0001$).

2.6 | Behavioral measures

To measure the behavioral attributes of wild reptiles, researchers must adapt their scoring systems to the ecology of the species in question (Byrnes et al., 2016; Ward-Fear et al., 2018). The four behavioral measures listed below are appropriate for assaying lace monitors along a shyness/boldness axis (see Results). We attempted to take data only once from each individual but animals were not marked and thus, some individuals may have been assayed more than once.

2.7 | Vegetation cover when sighted

An animal’s behavioral phenotype can influence how it selects microhabitats, with shyer individuals found in microhabitats that facilitate cryptis (Ward-Fear et al., 2019). Accordingly, we assessed the percentage of vegetation cover above 30 cm within a 5 m radius surrounding each lizard when first observed (n = 244). We then examined associations between vegetation density and lizard behavior in our two habitat contexts (see analysis for details).

2.8 | Flight-initiation distance

Flight-initiation distance (FID) is the distance at which an animal begins to flee from an approaching threat (Cooper & Frederick, 2007), with animals that initiate flight from further away considered more risk-averse. This offers a robust estimate of risk-taking behavior in many taxa (Cooper et al., 2015; Samia et al., 2016), even if individuals are only measured once (Putman et al., 2020). We scored flight-initiation distance (FID) for 231 encounters with lace monitors. As soon as the lizard was sighted, the observer began walking toward the animal at 1 m/s. When the animal responded by fleeing, ascending a tree, or displaying aggression, we placed a mark on the ground and then placed a second mark at the location from which the animal responded. We recorded locations of the two marks using Sightings (v1.1 for iPhone) mobile application software and quantified linear distances between marks to determine the FID. Lizards that allowed approach to within an arm’s length were assigned an FID score of 0. We excluded any lizards that fled at a distance greater than 60 m, due to the uncertainty of assigning the flee response specifically to the approach of the observer.

2.9 | Primary flee response

We scored the behavioral responses of 231 lizards to the approach of an observer as: (a) defend: raising the forebody off the ground, tail-flicking, and/or hissing, (b) no response (FID = 0), (c) walk, (d) run, or (e) climb. We classified the first three behaviors as “weak flee responses” (i.e., bold behaviors) and the last two behaviors as “strong flee responses” (i.e., shy behaviors).

2.10 | Proportion that climbed a tree

Lace monitors often climb trees to escape terrestrial threats (Webb, 1994). Following the approach of the observer for the flight-initiation distance trial, we scored whether or not the lizard fled up a tree (n = 230). We considered animals that climbed trees to be more risk-averse.
2.10.1 | Analyses

Although we were unable to repeatedly assay individuals in a range of ecological contexts to test for behavioral syndromes (Sih et al., 2004), our sampling protocol allows us to test for correlated behaviors of individuals, as expected if those behaviors are driven by underlying syndromes (behavioral axes). To explore these relationships, we used nonparametric Spearman analyses to test for correlations between three behavioral variables: (a) association with dense vegetation (% within 5 m radius), (b) flight-initiation distance, and (c) strength of the flee response.

Next, we tested for associations between lizards that displayed a strong or weak flight response and the habitat in which they were found. We ran a full factorial mixed model with the flight response (categorical: strong flight response or weak flight response), habitat (bushland or campground) and their interaction as predictor variables, and the vegetation density (% cover) surrounding the lizard was seen as a continuous response variable.

We then ran separate models on our six dependent variables (population size, body size, ground cover, flight-initiation distance, flee response strength, and proportion of lizards that climbed trees) with habitat (categorical: bushland or campground) as the independent variable. Following significant results, we next ran additional mixed models with habitat as a factor, body size (continuous) as a covariate, and their interaction. Finally, we ran full factorial models with factors of habitat (campground vs. bushland) and toad presence (present or absent), and their interaction. Survey day was included as a continuous fixed effect in all models (except for population size analyses, as mean counts were derived from across all survey days), and site was added as a random effect in all models.

Mixed model analyses were run using the GLIMMIX procedure in SAS 9.4 (SAS Institute, Cary NC). Analyses of continuous dependent variables (counts, body size, ground cover when sighted, flight-initiation distance) used a normal distribution and identity link function. Analyses with a binary dependent variable (proportion that displayed a strong flee response, proportion that climbed trees) were run with a binomial distribution and a logit link function. We visually checked data for normality, and equality of variances was assessed using the Levene test. We applied transformations to data to improve equality of variances where appropriate. We accepted deviations from normality when samples sizes were large (Sokal & Rohlf, 1995).

3 | RESULTS

3.1 | Correlations between behavioral variables

The three behaviors that we scored were positively correlated (all Spearman’s tests \( p < .0021 \)). That is, lizards that were spotted in denser vegetation displayed a stronger flee response that was initiated at a greater distance from the observer. These positive correlations support our designations of “bold” or “shy” behaviors across individuals. Additionally, lizards that displayed a strong flight

![FIGURE 2](image-url) The vegetation density (% cover) in which a lace monitor (Varanus varius) was seen as a function of the habitat type (bushland or campground) and the strength of the lizards flight response to an approaching observer

| TABLE 1 | Results from models exploring the attributes of lace monitors (Varanus varius) in bushland (BL) or campgrounds (CG) |
|---|---|---|---|---|---|
| Dependent variable | Factors | \( F \) and \( df \) | \( p \) value | Effect direction |
| Population size | Habitat | \( F_{1, 26} = 12.73 \) | .0014 | CG > BL |
| Body size | Habitat | \( F_{1, 222} = 18.27 \) | <.0001 | CG > BL |
| Ground cover | Habitat | \( F_{1, 219} = 12.72 \) | .0004 | BL > CG |
| Flight-initiation distance | Habitat | \( F_{1, 206} = 48.67 \) | <.0001 | BL > CG |
| Flee response strength | Habitat | \( F_{1, 206} = 12.31 \) | .0006 | BL > CG |
| Proportion that climbed trees | Habitat | \( F_{1, 205} = 3.54 \) | .0612 |

Note: Boldface font indicates significant differences between habitat types.

response (running away or climbing a tree in response to an approaching observer) were more likely to be found in dense vegetation \( (F_{1,204} = 4.16, p = .043; \) Figure 2), regardless of being in bushland or campground (habitat \( \times \) flee response interaction \( F_{1,204} = 0.03, p = .86 \)). The main effect of habitat on flight response was also significant, with bushland lizards fleeing from further away \( (F_{1,204} = 5.14, p = .025) \).

3.2 | Overall comparisons between lizards in campgrounds versus bushland

Lace monitors were larger and more abundant in campgrounds than in bushland habitats (Table 1, Figure 3a,b). Despite the more open conditions in campgrounds, lizards in these disturbed areas allowed
closer approach before fleeing from an observer and were more likely to ignore the observer or even to exhibit both agonistic behavior toward him rather than fleeing (Table 1, Figure 3c,d,e). Lizards from habitat types were equally likely to climb a tree in response to the approaching observer (Table 1, Figure 3f).

3.3 | Are behavioral differences between habitats due to differences in mean body size?

There was no significant relationship between the body size of a lizard and the density of vegetation cover in which it was first seen,
In both bushland and campgrounds, smaller lizards were more likely to flee (run or climb), whereas larger lizards walked away slowly or allowed the observer to approach (Table 2). Thus, the difference in mean body sizes between campground and bushland lizards was

TABLE 2 Results of statistical models exploring how the behavioral attributes of lace monitors (Varanus varius) differ with habitat type (bushland [BL] and campgrounds [CG]) and body size

| Dependent variable               | Factors                  | F and df | p value | Effect direction |
|----------------------------------|--------------------------|----------|---------|------------------|
| Ground cover                     | Habitat                  | F₁, 216 = 3.08 | .0808   |                  |
|                                  | Size                     | F₁, 216 = 2.17 | .1422   |                  |
|                                  | Habitat × Size           | F₁, 216 = 0.78 | .3793   |                  |
| Flight-initiation distance       | Habitat                  | F₁, 204 = 0.01 | .9164   |                  |
|                                  | Size                     | F₁, 204 = 2.30 | .1313   |                  |
|                                  | Habitat × Size           | F₁, 204 = 3.60 | .0592   |                  |
| Flee response strength           | Habitat                  | F₁, 204 = 0.49 | .4847   |                  |
|                                  | Size                     | F₁, 204 = 4.04 | .0458   |                  |
|                                  | Habitat × Size           | F₁, 204 = 0.00 | .9538   |                  |
| Proportion that climbed trees    | Habitat                  | F₁, 203 = 1.97 | .1620   |                  |
|                                  | Size                     | F₁, 203 = 0.03 | .8585   |                  |
|                                  | Habitat × Size           | F₁, 203 = 3.65 | .0574   |                  |

Note: Boldface font indicates significant main effects and interactions.

aBushland lizards from toad-present areas fled from the furthest away.

TABLE 3 Results from full factorial models exploring the attributes of lace monitors (Varanus varius) in areas differing in habitat (bushland [BL] and campgrounds [CG]) and toad invasion history (toads absent [TA] or toads present [TP])

| Dependent variable               | Factors                  | F and df | p value | Effect direction |
|----------------------------------|--------------------------|----------|---------|------------------|
| Population size                  | Habitat                  | F₁, 21 = 16.25 | .0005   | CG > BL         |
|                                  | Toad status              | F₁, 21 = 4.76 | .0388   | TA > TP         |
|                                  | Habitat × Toad status    | F₁, 21 = 2.76 | .1091   |                  |
| Body size                        | Habitat                  | F₁, 221 = 7.47 | .0068   | CG > BL         |
|                                  | Toad status              | F₁, 221 = 3.80 | .0525   |                  |
|                                  | Habitat × Toad status    | F₁, 221 = 3.84 | .0513   |                  |
| Ground cover                     | Habitat                  | F₁, 218 = 9.84 | .0019   | BL > CG         |
|                                  | Toad status              | F₁, 218 = 0.00 | .9751   |                  |
|                                  | Habitat × Toad status    | F₁, 218 = 0.12 | .7246   |                  |
| Flight-initiation distance       | Habitat                  | F₁, 205 = 15.42 | <.0001 | BL > CG         |
|                                  | Toad status              | F₁, 205 = 1.87 | .1732   |                  |
|                                  | Habitat × Toad status    | F₁, 205 = 10.12 | .0017   |                  |
| Flee response strength           | Habitat                  | F₁, 205 = 4.79 | .0298   | BL > CG         |
|                                  | Toad status              | F₁, 205 = 2.87 | .092    |                  |
|                                  | Habitat × Toad status    | F₁, 205 = 5.08 | .0253   |                  |
| Proportion that climbed trees    | Habitat                  | F₁, 204 = 2.05 | .1537   |                  |
|                                  | Toad status              | F₁, 204 = 0.07 | .7934   |                  |
|                                  | Habitat × Toad status    | F₁, 204 = 0.01 | .9276   |                  |

Note: Boldface font indicates significant main effects and interactions.

aBushland lizards from toad-present areas fled from the furthest away.

bBushland lizards from toad-present areas exhibited the strongest flee response.
3.4 | Does the invasion of cane toads affect lizard behavior?

In areas with cane toads, lace monitors were smaller and less abundant, and more wary (stronger flee response) (Table 3). In areas with toads, bushland lizards fled from further away, whereas approach distance was unchanged for campground conspecifics (interaction toad presence*habitat—Table 3, Figure 4). If we split the analysis into habitat types, bushland goannas became shyer (fled from further away) after toads arrived (square root-transformed: $F_{1,55} = 4.34, p = .041$), whereas flight-initiation distances of campground lizards did not change significantly [log (1 + x) transformed; $F_{1,135} = 1.32, p = .25$].

4 | DISCUSSION

Our surveys show that campgrounds support larger monitor lizards than does adjacent native bushland. Previous studies on varanid lizards have shown similar patterns in body size associated with anthropogenically subsidized environments, that may reflect processes such as higher growth rates due to resource subsidies, or intraspecific competition for access to favored sites (Ardiantonio et al., 2018; Jessop et al., 2012; Jolly et al., 2016). Our data are the first to document habitat-associated differences in abundance, morphology, and behavior of a reptilian predator following a biological invasion. Boldness can be defined as an animal’s propensity to engage in risky behavior (Reale et al., 2007; Putman et al., 2020) and hence can be measured as the strength of an individual’s response to potential threats (Reale et al., 2007). Bolder individuals may be better able to compete for territory or mates (Reaney & Backwell, 2007), or exploit foraging opportunities in open habitats (Ioannou et al., 2008; Short & Petren, 2008). Hence, boldness may confer strong fitness advantages (Smith & Blumstein, 2008), including higher reproductive success (Ariyomo & Watt, 2012; Reale et al., 2009) and survival (Sinn et al., 2014). However, boldness can also confer costs (e.g., increased predation risk), maintaining variation in behavioral phenotypes within a population (Hulthen et al., 2017).

Although our survey methodology did not allow us to test the same individuals for repeatable behavioral responses across contexts (a requirement to identify behavioral syndromes), significant correlations between our three behavioral measures from each individual are consistent with a spectrum of boldness. Animals that demonstrated the strongest flee response also fled from further away, even though they were concealed in vegetation when first seen (consistent with shy behavior; Ward-Fear et al., 2019). At the other end of the spectrum, individuals that we found in relatively open habitats often did not move away from us at all. Hence, these behaviors are consistent with boldness [as documented in birds, (Ducatez et al., 2017), mammals (Reale et al., 2000), reptiles (Putman et al., 2020) and fish (Coleman & Wilson, 1998)].

Lizards from bushland habitats fled from further away and adopted tactics (such as climbing trees) consistent with more wary responses. Smaller individuals were less bold than were larger conspecifics, as found also for prey-handling behavior in this species (Jolly et al., 2016) and for broader dimensions of behavior in another large varanid species from tropical Australia (V. panoptes: Ward-Fear et al., 2018). The behavioral (boldness) difference between campground versus bushland lizards was due, at least partly, to the larger average size of campground animals. Interestingly, the invasion of highly toxic cane toads exacerbated some of these habitat-based divergences between lizards. Below, we explore causal mechanisms underlying these patterns.

Many populations of free-living animals exhibit substantial variation in individual behavior (consistent patterns of behavior = “personality”: Gosling, 2001), in ways that influence the choice of habitats (Holtmann et al., 2017). For example, we might expect bolder individuals to be better-suited to disturbed areas like campgrounds, because a highly wary individual in such a site would be able to maintain activity only during the relatively brief periods when no potential threats were evident (Dammhahn & Almeling, 2012). Another mechanism that might generate a correlation between boldness and use of open habitats is habituation: that is, a lizard in such a habitat becomes accustomed to frequent disturbance and so learns to tolerate the approach of a potential threat (such as a human being) without fleeing. Thus, all else being equal, we might expect to see bolder individuals in more open habitats. An individual’s body size is likely to affect such habitat partitioning, however, because larger size may render an animal less vulnerable to predators (Urban, 2007), more capable of repelling competitors in competition for a resource (Candolin & Voigt, 2001), and may enable it to ingest a wider range of prey types and sizes (Scharf et al., 2000). Also, size covaries with age, such that a larger animal will likely be older, and thus have had more opportunity to learn the location of local resources (shelters, foraging sites) that facilitate exploitation of a disturbed habitat.
Our data suggest that the behavioral divergence between lizards from campgrounds versus bushland is partly driven by body size effects (larger lizards are bolder, and are more often found in campgrounds) but that campground lizards are bolder than bushland lizards even at the same body size. That correlation might reflect either habituation (campground lizards learn to tolerate frequent disturbance by humans: Samia et al., 2015) or a capacity for individuals to select habitats best-suited to their own behavioral phenotypes (shy lizards fail to thrive in campgrounds, because they are unable to access resources when bolder conspecifics are present), or to natural selection (e.g., rates of growth and mortality are higher for shy lizards in bushland and for bold lizards in campgrounds). Our data do not allow us to choose between these possibilities. Studies on captive-raised hatchlings that are the progeny of adults from bushland versus campground habitats would be of great interest.

Frustratingly, we were unable to determine the sex of the lizards that we observed. There are no reliable overt indicators of sex in monitor lizards, even if the animal can be handled, thereby necessitating molecular-genetics techniques to distinguish between males and females (e.g., Halverson & Spelman, 2002; Ward-Fear et al., 2018). This was not possible in our study, for logistical reasons. Nonetheless, the marked sexual dimorphism in body sizes within this species (Carter, 1992; Guarino, 2002; Kirshner, 2007) indicates that most of the largest specimens were males. A detailed analysis of a similar-sized monitor species (V. panoptes) showed that in general, males were bolder than females (Ward-Fear et al., 2018). If the same is true in V. varius, then at least part of the habitat-based divergence in sizes and behavior that we documented may be attributable to sex-specific habitat partitioning (as reported in several other taxa of reptiles: e.g., Delaney & Warner, 2016; Shine & Wall, 2007). Further work to explore this idea, combining molecular sexing with behavioral observations, would be of great interest. Such a study could also explore the idea (see Introduction) that the large body sizes of adult male lace monitors may confer energetic disadvantages in bushland (where prey are relatively scarce) and hence be maintained by sexual selection rather than natural selection.

The habitat-based disparities in body sizes and behaviors of lace monitors are of interest not only in their own right, but also because this species is a keystone predator. Recent research has detected multiple trophic shifts in tropical Australia following decimation of monitor (V. panoptes) populations by the invasive cane toad (Brown et al., 2013; Doody et al., 2015) and has suggested similar shifts following toad-induced mortality of V. varius (Jolly et al., 2015). The higher abundance, larger size, and bolder behavior of V. varius in disturbed (campground) habitats thus may have strong implications for the diverse array of smaller taxa consumed by these giant lizards (Guarino, 2001; Jessop et al., 2010).

A trend for larger, bolder individuals to exploit anthropogenically disturbed habitats may have implications for people as well as for other species, exacerbating conflicts with wildlife. Habituation and boldness of predator species can render them nuisances, leading to dangerous situations for both humans and other animals. For example, bears seeking food are a pervasive threat at campgrounds throughout North America and have caused human fatalities (Rogers, 2011). Coyotes that key in on human subsidies (livestock) demonstrate patterns of bold behavior that also are exhibited by their offspring (via cultural or heritable transmission; Schell et al., 2018). These encounters often lead to culling. Large varanid lizards can inflict serious damage to humans, via aggressive displays (including tail whipping, biting and scratching) and also venom (Fry et al., 2006). The patterns that our study demonstrate are concerning in this respect, and should be considered by managers.

Interestingly, the association between lizard traits and habitat has been reinforced by the recent invasion of cane toads into southeastern Australia. Broadly, we expect mortality due to lethal toxic ingestion of cane toads to fall more heavily on lizards in campgrounds (because toads thrive in disturbed habitats: González-Bernal et al., 2016) and on larger lizards (because they consume larger toads, and thus ingest more toxin; and because larger goannas evaluate prey less carefully prior to ingesting it: Jolly et al., 2016). So, we might expect toad invasion to disproportionately affect campground lizards, removing the largest and boldest individuals. That would tend to reduce the disparity between campgrounds and bushlands in the attributes of goannas. In practice, however, the disparity was maintained: after toads arrived, bushland lizards fled from further away than did conspecifics in the campground (Table 3, Figure 3b). Indeed, bushland goannas became shyer (fled from further away) after toads, whereas flight-initiation distances of campground lizards did not change significantly. This pattern may result from behavior-dependent emigration of lizards from the bushland to the campground after toad-induced reduction of goanna numbers in campgrounds. That is, the boldest lizards from bushland habitats moved into campgrounds to exploit the newly available opportunities, leaving the shyest individuals as the only ones left in the bushland. That pattern results in an overall decrease in the frequency of bold lizards (as seen in the main effect of toad invasion on strength of the flight response), but no shift within campgrounds in this parameter.

Although the specific type of disturbance and resource subsidy exploited by the lace monitors that we studied are the result of anthropogenic disturbance (i.e., campgrounds), the broad geographic range of lace monitors means that similar “resource hotspots” would have been available even before humans colonized eastern Australia. Thus, for example, monitor lizards have been reported to key in on seasonally or stochastically available food supplies such as fish in drying pools (Shine, 1986; Ward-Fear et al., 2020) and the eggs and hatchlings of sea turtles (Lei & Booth, 2017). The massive range in both body sizes and behavioral syndromes within a population of large varanid lizards (Ward-Fear et al., 2018) thus may have allowed these giant reptiles to effectively utilize unpredictable resource-rich patches within a dynamic habitat mosaic. As humans modified the ecosystems, creating even more extreme spatial and temporal variation in prey availability, species like the lace monitor were ideally placed to take advantage of that new opportunity.
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CONFLICT OF INTEREST
None declared.

AUTHOR CONTRIBUTIONS
Lachlan Pettit: Data curation (lead); formal analysis (lead); funding acquisition (supporting); project administration (lead); software (lead); visualization (lead); writing—original draft (lead). Greg P. Brown: Formal analysis (equal). Georgia Ward-Fear: Conceptualization (supporting); funding acquisition (supporting); investigation (supporting); supervision (supporting); writing—review and editing (supporting). Richard Shine: Conceptualization (lead); formal analysis (supporting); funding acquisition (lead); investigation (lead); resources (lead); supervision (lead); writing—review and editing (lead).

ETHICAL APPROVAL
All procedures were approved by the University of Sydney ethics committee (approval 2017/1202) and were carried out in accordance with relevant guidelines and regulations under license from state and federal wildlife agencies.

DATA AVAILABILITY STATEMENT
Data are housed on the Figshare data repository. https://doi.org/10.6084/m9.figshare.13298732.v1 (Pettit et al., 2020).

ORCID
Lachlan Pettit https://orcid.org/0000-0002-2628-1416
Gregory P. Brown https://orcid.org/0000-0002-2924-9040
Georgia Ward-Fear https://orcid.org/0000-0002-4808-1933
Richard Shine https://orcid.org/0000-0001-7529-5657

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