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Determinants of spread in an urban landscape by an introduced lizard

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35 Abstract

36 Context

37 Urban landscapes are a mixture of built structures, human-altered vegetation, and

- 38 remnant semi-natural areas. The spatial arrangement of abiotic and biotic
- 39 conditions resulting from urbanization doubtless influences the establishment and
- 40 spread of non-native species in a city.

41

42 Objectives

43 We investigated the effects of habitat structure, thermal microclimates, and species

44 coexistence on the spread of a non-native lizard (*Anolis cristatellus*) in the Miami

45 metropolitan area of South Florida (USA).

46

47 Methods

We used transect surveys to estimate lizard occurrence and abundance on trees and to measure vegetation characteristics, and we assessed forest cover and impervious surface using GIS. We sampled lizard body temperatures, habitat use, and relative abundance at multiple sites.

52

53 Results

54 At least one of five *Anolis* species occupied 79% of the 1,035 trees surveyed in

- 55 primarily residential areas, and non-native *A. cristatellus* occupied 25% of trees.
- 56 Presence and abundance of *A. cristatellus* were strongly associated with forest
- 57 patches, dense vegetation, and high canopy cover, which produced cooler

58	microclimates suitable for th	his species. Presence of A.	<i>cristatellus</i> was negatively
50	microcimiates suitable for ti	ins species. I reserve of n.	cristatenus was negatively

59 associated with the ecologically similar non-native A. sagrei, resulting in reduced

60 abundance and a shift in perch use of *A. cristatellus*.

61

62 Conclusions

- 63 The limited spread of *A. cristatellus* in Miami over 35 years is due to the patchy, low-
- 64 density distribution of wooded habitat, which limits dispersal by diffusion. The
- 65 presence of congeners may also limit spread. Open habitats some parks, yards
- 66 and roadsides contain few if any *A. cristatellus*, and colonization of isolated forest
- 67 habitat appears to depend on human-mediated dispersal.
- 68

69 Keywords

- 70 Anolis body temperature canopy cover GIS forest cover impervious surface -
- 71 thermal microclimates tree characteristics urban heat island urban vegetation

72

74 Introduction

75 Upon arrival in a city, non-native species encounter a spatially heterogeneous environment that varies in the types and densities of buildings, 76 77 vegetation, infrastructure, and remnant natural areas (Pickett et al. 2001; Cadenasso 78 et al. 2007; Forman 2014). This variability in habitat structure and its spatial 79 patterning will likely influence the ability of invaders to establish and spread within 80 an urban area. For example, exotic grey squirrels in the UK are positively associated 81 with increased canopy cover, larger trees, and the presence of seed-bearing trees as 82 well as supplementary feeders for birds (Bonnington et al. 2014). Thus, the 83 arrangement of vegetation and non-vegetative features within the urban landscape 84 influences where exotics establish and the routes by which they spread. Identifying 85 those features associated with the occurrence of exotic species is important for 86 understanding their current distributions and potential for future spread. 87 As a consequence of habitat structure modification during urbanization, city 88 temperatures can be several degrees higher than surrounding rural areas (Akbari et al. 2001; Arnfield 2003; Chen and Jim 2008; Rizwan et al. 2008). These urban heat 89 90 islands are spatially and temporally heterogeneous (Ramalho and Hobbs 2011). 91 reflecting variation in the matrix of built structures and local vegetation and 92 creating a thermal mosaic (Georgi and Zafiriadis 2006; Hamdi and Schayes 2008; Huang et al. 2008). This variation influences the microclimates available in a city, 93 including air and surface temperatures, relative humidity, solar radiation, and wind 94 95 speed. Thermal microclimates are critically important to ectotherms (e.g., insects, 96 lizards, frogs), which rely on external sources of heat to regulate their body

97 temperatures. Because temperature is fundamentally important for development,
98 growth, survival, and reproduction in ectotherms (Angilletta 2009), organisms
99 living in a city are likely to be sensitive to variation in vegetation and urban features
100 that affect thermal microclimates (Ackley et al. 2015a).

101 In addition to the habitat structure of a city, interactions with potential 102 competitors and predators can influence occurrence and abundance patterns 103 (Shochat et al. 2006; Anderson and Burgin 2008). For example, abundance of a 104 native gecko, *Lepidodactylus lugubris*, throughout the Pacific is strongly influenced 105 by interactions with a competitively superior non-native gecko, Hemidactylus 106 frenatus (Case et al. 1994), which better exploits insect resources concentrated 107 under artificial night lighting (Petren and Case 1996). In general, more ecologically 108 similar species are predicted to have stronger negative effects on each other at local 109 scales through competition (Pianka 1981; Losos 1994). Thus, both biotic and 110 abiotic factors may influence the establishment, spread, and ultimately the 111 distribution of non-native species in a city.

112 In this study, we investigate the effects of habitat structure, thermal 113 microclimates, and species interactions on the spread of introduced *Anolis* lizards in 114 the Miami metropolitan area. Our study group, Anolis lizards (or anoles), comprise a 115 species-rich genus of small, insectivorous, diurnal lizards found in the Neotropics from the southeastern United States to South America including Caribbean islands 116 (Losos 2009). Many Anolis species occupy both natural and human-modified areas 117 in their native and non-native ranges (Perry et al. 2008; Irschick et al. 2005; 118 119 Marnocha et al. 2011; Kolbe et al. 2015). There are nine Anolis species established

120 in Miami, only one of which — *A. carolinensis* — is native to the U.S. (Lever 2003; 121 Kolbe et al. 2007; Kraus 2009). Four species have very restricted distributions (A. 122 chlorocyanus, A. cybotes, A. garmani and A. porcatus), two are distributed throughout 123 the Miami area (A. distichus and A. equestris), and one is found throughout Florida, 124 the Gulf Coast, and southern Georgia and South Carolina (A. sagrei). In contrast to 125 these either very restricted or widespread species, an eighth non-native species, A. 126 *cristatellus* (Fig. 1, inset), is expanding its distribution in Miami, but is not vet 127 ubiquitous. We can therefore identify factors related to its current distribution and 128 predict whether future spread in urban areas is likely. 129 Anolis lizards have a number of advantages for this study. First, anoles in 130 Miami are conspicuous, easy to detect, and sufficiently different in ecology and 131 morphology to accurately identify to species when present. Second, the 132 introduction history of A. cristatellus in Miami is well studied with two independent 133 introductions from its native range in Puerto Rico (Kolbe et al. 2007). Third, the 134 other four Anolis species that co-occur with A. cristatellus in Miami — A. carolinensis, 135 A. distichus, A. equestris and A. sagrei — were all present prior to its introduction in 136 the mid-1970s. These species span a range of ecological similarity; specifically, A. 137 *sagrei* and *A. distichus* typically perch lower to the ground on tree trunks, similar to 138 A. cristatellus, whereas A. carolinensis and A. equestris perch higher in the canopy 139 (Losos 2009). We can therefore test the hypothesis that more ecologically similar congeners influence the presence of A. cristatellus in Miami. Lastly, the thermal 140 141 biology of anoles in general, and A. cristatellus in particular, is well studied (Losos 142 2009). Previous studies detail the thermal preferences, thermal tolerances, and

143 field body temperatures of A. cristatellus from numerous sites in Puerto Rico and 144 Miami (e.g., Huey 1974; Huey and Webster 1976; Hertz 1992; Leal and Gunderson 145 2012; Kolbe et al. 2012; Gunderson and Leal 2012), allowing us to evaluate if the 146 effects of urban vegetation on thermal microclimates are relevant to A. cristatellus. 147 A primary goal of this study is to contrast how abiotic and biotic aspects of 148 the urban environment influence the current distribution and abundance of a 149 recently introduced species to better understand its potential for future spread. We 150 survey lizards and vegetation characteristics on a tree-by-tree basis using transects 151 across putative distribution boundaries, and test for relationships at the landscape 152 level between the presence of *A. cristatellus* and GIS-based data attributes of forest 153 cover and impervious surfaces. We predict that 1) urban vegetation structure and 154 arrangement will influence the occurrence and abundance of A. cristatellus. In 155 particular, we predict that *A. cristatellus* will be associated with denser vegetation 156 and forested areas, which produce relatively cooler microclimates. Based on 157 previous ecological studies (Losos 2009), we also predict 2) negative associations 158 between A. cristatellus and its more ecologically similar congeners in Miami. 159 Specifically, A. sagrei and A. distichus overlap most with A. cristatellus in their 160 structural microhabitat (i.e., the height, diameter, and type of perch), which should 161 lead to stronger interspecific competition.

162

163 Methods

164 Study area

165 We conducted this study in the Miami metropolitan area, where the initial 166 sites of introductions for *A. cristatellus* are documented. Genetic analyses confirmed 167 two independent introductions from geographically and genetically distinct native-168 range sources in Puerto Rico (Kolbe et al. 2007). The Key Biscayne population is 169 from San Juan, Puerto Rico and was first detected in a residential area on the island 170 in 1975 (Schwartz and Thomas 1975; Bartlett and Bartlett 1999). The South Miami 171 population is from northeast Puerto Rico and was found in a different residential 172 area in 1976 (Wilson and Porras 1983). The Key Biscayne population is \sim 5 km from 173 the mainland population separated by a bridge to Virginia Key and the Rickenbacker 174 Causeway to the mainland. The two introduction sites are ~ 12 km apart across 175 Biscayne Bay. The bulk of the study area is residential with detached single units, 176 considerable tree cover, and low-traffic, two-lane roads. Also present are 177 commercial districts, high-traffic roads up to six lanes, open parklands, urban 178 forests, and waterways such as canals, lakes, and coastal areas (Fig. 1, 179 Supplementary Fig. 1). 180

181 Study design and sampling

In summer 2009, we collected preliminary data on *A. cristatellus* presence in the Miami area by conducting block-by-block walking surveys radiating from the initial points of introduction in South Miami and Key Biscayne. Based on these data, we established five 610 m x 1100 m plots in South Miami, each crossing an observed transition from presence to absence of *A. cristatellus*. In each plot, we established three to six roadside transects running perpendicular to the transition zone

188 (Supplementary Fig. 2). On our initial visit to each transect in June 2010, we 189 measured tree characteristics (Table 1) and, using binoculars when needed, 190 observed Anolis lizards on trees with a trunk diameter >10 cm growing in the 191 roadside public right-of-way. Although lizards use smaller trees, the availability of 192 such trees was limited along roadsides and in yards. All species have multiple 193 diagnostic features, which aided accurate species identification. Following this 194 initial survey, we returned to each transect two more times to survey the same trees 195 for the presence and total number of lizards of the five *Anolis* species. One to three 196 trained observers were present for each survey, with at least two observers in most 197 cases. Data from these transects were used to evaluate whether the presence of A. 198 cristatellus was related to the presence of congeners and to the characteristics of the 199 trees and surrounding vegetation (Table 1).

200 Given patterns of species coexistence from these transects, we conducted 201 visual encounter surveys to determine if congener presence affects the relative 202 abundance and habitat use of A. cristatellus (Crump and Scott 1994; Kolbe et al. 203 2008). Surveys consisted of walking at a constant pace for 15 minutes and 204 recording the species, time, sex, and perch characteristics (i.e., height and diameter) 205 for all undisturbed lizards observed. We compared relative abundance at sites with 206 predominantly A. cristatellus (n=10 surveys) to sites with A. distichus and A. sagrei in 207 addition to A. cristatellus (n=6 surveys). Because A. distichus and A. sagrei co-occur throughout most of Miami, we were unable to find nearby sites with only one of 208 these species. We supplemented data on perch characteristics with opportunistic 209 210 observations of all three species at the same sites.

211 The presence-absence data from transects, visual encounter surveys, and 212 opportunistic surveys allowed us to map the distribution of *A. cristatellus* in Miami 213 (Figs. 1 and 2). In addition to the intensive sampling within the core areas of South 214 Miami and Key Biscayne, we also investigated potential localities throughout Miami-215 Dade County including municipal parks and regional waste transfer stations. 216 Preliminary surveys showed A. cristatellus was present in forest patches in some 217 parks associated with waste transfer stations, suggesting transport of vard waste 218 from houses to regional stations as a potential dispersal mechanism. 219 220 GIS analysis of forest cover and impervious surfaces 221 To complement analyses based on transect data, we conducted a geographic 222 information system (GIS) analysis of the study area using ArcGIS version 10.2 223 (Environmental Systems Research Institute, Redlands, CA) and publicly available 224 GIS layers of impervious surface (MRLC 2011) and forest fragmentation (NOAA 225 2010). The MRLC Percent Developed Imperviousness layer, a raster dataset with 226 30-meter pixels, provides the average percentage of human-made impervious 227 surface for each pixel. The NOAA forest fragmentation layer, a raster dataset with

228 30-meter pixels, distinguishes between four fragmentation types: 1) "core forest"

refers to forested pixels that are not adjacent to any non-forested pixels, 2) "patch

230 forest" refers to forested pixels in small patches that are not large enough to contain

a 90m x 90m block of forest, 3) "perforated forest" refers to forested pixels adjacent

to small non-forested patches that are not large enough to contain a 90m x 90m

233 block of non-forested area, and 4) "edge forest" refers to forested pixels adjacent to

234 larger non-forested patches that contain at least one 90m x 90m block of non-

forested area. We used GIS to generate 11 attributes describing forests andimpervious surfaces (Table 2).

237

238 Thermal microclimates and lizard body temperatures

239 To investigate the range of possible thermal microclimates available to A. 240 cristatellus in Miami during peak summer temperatures, we measured the 241 temperature under two trees—one with an open canopy (*Thrinax radiata*, DBH = 12 242 cm) and another with a heavily shaded canopy (*Chukrasia tabularis*, DBH = 145 cm). 243 We used painted, hollow, copper models the size of lizards with an iButton inside to 244 estimate temperatures a lizard would experience in each location without 245 behavioral or physiological thermoregulation (Hertz 1992; Gunderson and Leal 246 2012; Ackley et al. 2015a). Temperatures were recorded every 15 minutes on the 247 north, south, east, and west sides of each tree at a height of 1.5 m from 1700 h on 248 July 17 to 1000 h on July 21, 2014. To estimate the percentage of open canopy, we 249 took hemispherical canopy photographs in each cardinal direction using a 180° 250 fisheye lens and analyzed them using Gap Light Analyzer version 2.0 (Frazer et al. 251 1999).

To estimate the range of field body temperatures (T_b) for *A. cristatellus* in Miami, we sampled lizards and random locations at three sites that varied in species composition and vegetative structure. For comparative purposes, we also sampled *A. sagrei*, which has both higher field T_b and thermal tolerances than *A. cristatellus* (Corn 1971; Lee 1980; Gunderson and Leal 2012; Kolbe et al. 2012, 2014). The sites included a bike path along a canal where both species were sampled, a residential

258 area where only A. sagrei was sampled, and a forested area where only A. cristatellus 259 was sampled. For each undisturbed lizard captured, we recorded T_b, air 260 temperature 1 cm above the substrate where the lizard was perched, and copper 261 lizard model temperature at the same location as the lizard using a thermocouple 262 probe connected to a digital thermometer (Omega HH802U). We then took a 263 hemispherical canopy photo to estimate canopy openness as described above. For 264 comparison, we took copper lizard model temperature, air temperature, and a 265 canopy photo at randomly selected locations within each study site. Lizard T_b and 266 random location data were collected between 0800 to 1400 h, which is a high-267 activity time of day during the summer.

268

269 Data analysis

270 Occupancy and estimates of detection probabilities were calculated using single 271 season occupancy models (MacKenzie et al. 2002) in the program PRESENCE (Hines 272 2006). Likelihood models calculated in PRESENCE all assume that 1) any site where a 273 species is present remains occupied, 2) species may or may not be detected when present, 274 but are not detected when absent, and 3) the detection of a species at one sampling site is 275 independent of detection at all other sites (MacKenzie et al. 2002). A minimum of two 276 sample occasions is required for model estimation. We conducted three repeat surveys at 277 each sampling point. Occupancy models to calculate estimates of detection were 278 produced with all surveyors (Rick Stanley [RS], PV, and JJK) independently as 279 covariates as well as using full identity models including all surveyors.

280 We used generalized linear models (GLM; McCullagh and Nelder 1989; R 281 2013) with binomial (presence-absence) and continuous (abundance) response 282 variables. This allowed us to form linear and quadratic relationships between the 283 response and explanatory variables (Broennimann et. al. 2012), which were 284 standardized to normalize their distributions. Explanatory variables included the 285 tree characteristics of trunk diameter, canopy diameter, distance to nearest plant, 286 distance to nearest tree, and overstory canopy cover (Table 1). Model selection was 287 performed using a stepwise procedure based on the Akaike information criterion 288 (AIC; Akaike 1974). We conducted three separate analyses using presence-absence 289 as the response variable. First, we compared transect sections with A. cristatellus 290 present (but not necessarily occupying every tree) versus sections where A. 291 *cristatellus* was absent; second, we compared the presence versus absence of *A*. 292 *cristatellus* on all trees pooled; and third, we compared the presence versus absence 293 of A. cristatellus on trees within only the sections of transects with A. cristatellus 294 present. We then repeated the latter two analyses using *A. cristatellus* abundance as 295 the response variable.

When analyzing the GIS-based attributes, we conducted two separate
analyses. First, we divided street blocks from each transect into those with *A*. *cristatellus* present versus absent and compared attributes derived from GIS (Table
Second, we used presence and absence points for individual observations
throughout the Miami metropolitan area to test for relationships with GIS-derived
attributes, restricting the data set to no more than one observation per block.

302 For categorical explanatory variables (Table 1), we used likelihood ratio tests 303 to compare tree characteristics between sections of transects with A. cristatellus 304 present versus absent. When evaluating A. cristatellus abundance in relation to 305 categorical tree characteristics, we used t-tests or analyses of variance (ANOVA) as 306 appropriate. We tested for a relationship between the presence-absence of A. 307 cristatellus and the presence-absence of the four congeners using likelihood ratio 308 tests. We tested for effects using all trees sampled, as well as only those trees on the 309 sections of transects with A. cristatellus present. Analyses were conducted for trees 310 on each plot separately and with trees from all plots pooled.

Relative abundances from the visual encounter surveys were not normally distributed, so we used a Wilcoxon test to evaluate whether differences existed between sites. In particular, we predicted relative abundance of *A. cristatellus* would decrease when it is with other *Anolis* species compared to being alone. Logtransformation achieved normality for perch height and diameter, and we tested for a difference in these perch characteristics for *A. cristatellus* between sites with and without congeners using t-tests.

We compared lizard T_b and copper lizard model temperatures at the same locations using linear regression. Using this calibration, we adjusted model temperatures to make them directly comparable to lizard T_b for both species. We averaged model temperatures by hour and plotted them against time of day. We compared these model temperature estimates (i.e., non-thermoregulating lizards) to field T_b collected at the same time of year, and literature estimates of preferred T_b and high temperature tolerance (i.e., critical thermal maximum, or CTmax) for *A*.

325	cristatellus (Huey and Webster 1976). To investigate variation in field T_b of lizards,
326	we conducted an analysis of covariance (ANCOVA) testing for differences among
327	groups (i.e., A. cristatellus, A. sagrei, and random locations) with air temperature,
328	time of day, and canopy openness as covariates. We used the Johnson-Neyman
329	procedure (White 2003) to determine the range of covariate values in which
330	temperatures differed between groups when regression slopes were heterogeneous
331	(i.e., a significant interaction between the main effect and covariate).
332 333 334 335	Results
336 337	Anolis cristatellus distribution in Miami
337 338	The combination of opportunistic surveys, visual encounter surveys, and
339	transects resulted in fine-scale distribution data for A. cristatellus in key parts of the
340	Miami metropolitan area (n=362 presence points and n=483 absence points; Fig. 1,
341	Supplementary Fig. 1). This species has expanded its core range from the original
342	point of introduction no more than 2 km to the west, south, and east in South Miami,
343	and \sim 7 km to the northeast. A six-lane highway (i.e., the Dixie Highway/US 1) to the
344	northwest of the core South Miami distribution appears to limit unaided dispersal.
345	The introduction to Key Biscayne expanded across the majority of the island, but not
346	across the bridge to Virginia Key or causeway to mainland Miami.
347	We detected seven disjunct populations ranging from < 1 to 20 km from the
348	core distribution in South Miami. These sites included several Miami-Dade County
349	Parks (i.e., Chapman Fields, Kendall Indian Hammock, and Oak Grove) as well as the
350	University of Miami campus and three residential areas. We found A. cristatellus at

two of 13 waste transfer stations in Miami-Dade County (i.e., Chapman Fields and
Sunset Kendall), but only when adjacent to forested parks. Most waste transfer
stations had only a few widely spaced trees and were surrounded by residential or
commercial areas. Other species were present at all waste transfer stations with *A. distichus* and *A. sagrei* being the most common.

356

357 Tree characteristics

358 359 Transect surveys yielded observations on a total of 1035 trees. At least one 360 anole was present on 79% of the trees, and *A. cristatellus* occupied 25% of the trees 361 (Table 3). The best models to estimate detection probabilities for each species were 362 single season occupancy models including all three surveyors. Estimates of among-363 surveyor detection probability for the focal species, A. cristatellus, ranged from 0.50-0.96 364 among sites, and average detection probability for each surveyor for all sites was 365 estimated at 0.63-0.90 (Supplementary Table 1). Total detection probability for the full 366 model (all surveyors) was estimated at 0.88 (± 0.12). These estimates for detection were 367 high and therefore detection probabilities were not considered influential in subsequent 368 analyses. The most likely models of tree characteristics found *A. cristatellus* 369 associated with trees having larger trunks, larger canopies, greater percent of 370 overstory canopy cover, and closer to other plants and trees (Tables 1 and 4). These 371 results suggest that A. cristatellus occupies relatively shady and densely vegetated 372 areas.

373 Sections of transects with *A. cristatellus* present had a greater proportion of 374 native trees (X^2 =12.3, df=1, P<0.001, n=937) and trees with smooth bark (X^2 =14.4, 375 df=4, P<0.01, n=1035) as compared to transect sections with *A. cristatellus* absent.

In contrast, transect sections with and without *A. cristatellus* did not differ in the proportion of palm trees (X^2 =0.5, df=1, P=0.46, n=1028) or the number of trunks on trees (X^2 =4.3, df=2, P=0.12, n=1035).

379 Models for the abundance of A. cristatellus showed similar results with 380 increased abundance associated with trees having larger trunks and canopies, 381 greater percent of canopy cover, and closer to other plants (Table 4). Anolis 382 *cristatellus* abundance was twice as high on non-palm compared to palm trees 383 (t=2.7, df=1026, P<0.01) and highest on trees with multiple trunks $(F_{2,1032}=14.9, P<0.01)$ 384 P<0.0001), which were often large *Ficus* trees. Abundance did not differ between 385 native and non-native trees (t=-1.08, df=935, P=0.28) or among bark textures 386 (F_{4,1030}=1.57, P=0.18).

387 Analyses of *A. cristatellus* presence using GIS-based attributes were 388 consistent with transect surveys. Blocks with A. cristatellus present had more trees 389 per km, greater canopy cover, denser vegetation, and less impervious surface area 390 (Table 5A). Similarly, when analyzing the presence-absence points across Miami, A. 391 *cristatellus* was present at locations with less impervious surface and closer to 392 larger blocks of forest but not smaller forest patches (Fig. 2; Supplementary Fig. 3; 393 Table 5A). The percentage of forested area was three times greater in the core area 394 of A. cristatellus' distribution compared to the study area as a whole (Supplementary 395 Table 2). Moreover, the percentage of the core area with high impervious surface (> 40%) was about half as much as the study area as a whole (Supplementary Table 2). 396 397

399 Pooling all trees sampled, A. distichus and A. sagrei were both significantly 400 more likely to be absent when A. cristatellus was present than expected by chance, 401 with effects involving A. sagrei being much stronger (Table 6A). The presence of A. 402 *carolinensis* or *A. equestris* had no effect. When evaluating each plot separately, a 403 negative effect was observed with A. sagrei for most plots, and with A. carolinensis 404 and *A. distichus* in a few plots (Table 6A). This suggests congeneric interactions may 405 differ among plots. All comparisons for individual transects were non-significant 406 (results not shown).

407 Dividing each transect into sections based on A. cristatellus presence or 408 absence, only A. sagrei was more likely to be absent where A. cristatellus was 409 present (Table 6B). There was no interaction with the less abundant species A. 410 carolinensis and A. equestris. In contrast to the analyses of all trees pooled, this 411 analysis revealed no relationship between occurrence of *A. distichus* and *A.* 412 *cristatellus* (Table 6B). Potential interactions for *A. cristatellus* appear to be 413 strongest with A. sagrei, followed by A. distichus, but little evidence existed for 414 interactions with A. carolinensis or A. equestris.

415

416 Relative abundance and habitat use

417 Relative abundance estimates from visual encounter surveys were consistent 418 with the negative relationship between the presence of *A. cristatellus* and two of its 419 congeners in Miami. *Anolis cristatellus* was four times more abundant at sites with 420 no congeners than in sites occupied by *A. distichus* and *A. sagrei* (mean±SE: 45.3±2.5 421 versus 11.0±3.2 per survey; Wilcoxon: Z = 3.21, P<0.01). Furthermore, at sites with

422 congeners, *A. cristatellus* perched nearly twice as high (mean±SE: 79.0±4.2 v.

423 47.2±1.8; t=6.38, df=608, P<0.0001) and on trunk substrates twice as wide

424 (mean±SE: 18.6±1.5 v. 9.5±0.67; t=6.1 df=604, P<0.0001), suggesting a possible shift

425 in habitat use in the presence of congeners.

426

427 Thermal microclimates and lizard body temperatures

428 We investigated the thermal consequences of canopy cover by comparing 429 copper lizard model temperatures under trees with open versus shaded canopies. 430 The percentage of overstory canopy cover ranged from 31-46% for the open canopy 431 tree versus 89-91% for the shaded canopy tree (Supplementary Fig. 4). Model 432 temperatures for the two trees were similar through the night from ~1900 h to 433 \sim 0800 h (Fig. 3). After 0800 h, model temperatures on the open canopy tree 434 increased quickly, exceeding both shaded tree temperatures and preferred 435 temperatures of A. cristatellus from 1000-1800 h. While there was little variation in 436 model temperatures among the sides of the shaded tree, temperatures on the sides 437 of the open tree differed substantially from one another, with a maximum difference 438 of 5.7°C at 1000 h.

Lizard T_b and model temperatures showed a strong positive correlation (r=0.91; P<0.0001, n=52), suggesting that models accurately reflected lizard body temperatures. ANCOVA results showed all three covariates had significant positive effects on T_b/model temperatures (canopy openness: $F_{1,83}$ =46.42, P<0.0001; air temperature: $F_{1,83}$ =7.97, P=0.006; time of day: $F_{1,83}$ =23.51, P<0.0001; whole model R²=0.67). *Anolis sagrei* field body temperatures (mean±SE: 31.2°C±0.4) were

445 significantly higher than *A. cristatellus* temperatures (mean±SE: 28.8°C±0.4;

446	F _{2,83} =3.79, P=0.03; Tukey's HSD post hoc test P<0.05; Fig. 4). However, because the
447	species by canopy openness interaction was significant this main effect should not
448	be interpreted directly but only in conjunction with the covariate. The relationship
449	between temperature and canopy openness had a much steeper slope for A.
450	<i>cristatellus</i> compared to <i>A. sagrei</i> and random points (P<0.05 for comparison of
451	slopes; Fig. 4). The Johnson-Neyman procedure supported T_b differences between
452	A. cristatellus and both A. sagrei and random points for relatively closed canopies
453	(i.e., < 15% openness). In summary, all covariates had significant effects on lizard
454	T_b , but <i>A. cristatellus</i> had lower T_b compared to <i>A. sagrei</i> and the two species
455	appeared to thermoregulate differently in closed canopy areas.

456 457

458 **Discussion**

459

460 Since its introduction to South Miami over 40 years ago, A. cristatellus has 461 spread only modestly by diffusion ($\sim 0.2-0.25$ km/yr), much slower than some of the 462 other introduced Anolis species in Miami and invasive species in general (Lockwood 463 et al. 2007; Davis 2009). A recent analysis shows an order of magnitude faster spread rates on average for exotic lizards and snakes (~3 km/yr) and invaders to 464 465 the Nearctic (~5 km/yr; Liu et al. 2014). Results from our study suggest that both abiotic and biotic factors contribute to the limited spread of *A. cristatellus* in urban 466 Miami. The fragmentation of suitable habitat is an abiotic constraint. The presence 467 468 of *A. cristatellus* is strongly associated with forest habitats, which result in cooler 469 and more humid microclimates (e.g., Wong and Yu 2005; Georgi and Zafiriadis 2006;

Millward et al. 2014; Ackley et al. 2015a; Fig. 3). Because forests are patchily
distributed in Miami (Fig. 2a), dispersal by diffusion will be limited by
fragmentation caused by canals, non-forest habitats, and areas of impervious
surface, such as buildings, roads, and parking lots (Fig. 2). Therefore, humanmediated dispersal may be an important mechanism for moving *A. cristatellus* to
isolated patches of suitable habitat, which lizards are unable to reach via natural
diffusion.

477 Interactions with ecologically similar congeners may be a biotic constraint. 478 Anolis cristatellus is spreading into areas occupied by one or more additional Anolis 479 species. As expected, we found negative associations between A. cristatellus and 480 ecologically similar A. sagrei and A. distichus, but weak or no relationship between 481 the occurrence of *A. cristatellus* and *A. carolinensis* or *A. equestris*, which typically 482 perch higher in the canopy (Losos 2009). Ultimately, the relative abundance of each 483 species and the extent to which they overlap on niche axes, such as structural 484 habitat and thermal microclimate, will determine whether and how quickly A. 485 *cristatellus* spreads to new areas.

486

487 Effect of urban vegetation on the spread of A. cristatellus

The presence and abundance of introduced *A. cristatellus* in Miami are positively associated with larger trees, denser vegetation, greater canopy cover, proximity to forest, and less impervious surface. These features are indicative of forest patches in the urban environment including parks and certain residential areas. Previous studies show patterns of urban vegetation can be related to

493 numerous factors including socio-economics, remnant natural habitats, and 494 neighborhood age and history (e.g., Nowak et al. 1996; Martin et al. 2004; Grove et 495 al. 2006; Jenerette et al. 2007; Luck et al. 2009; Boone et al. 2010; Forman 2014). 496 Anolis cristatellus was present in several tropical hardwood hammocks, including 497 parks outside of its core distribution in South Miami. These disjunct populations 498 suggest dispersal limitation, not lack of suitable habitat, slows the spread of A. 499 *cristatellus* outside its core area in South Miami. The patchwork of scarce suitable 500 forested habitat in Miami will continue to limit the spread of A. cristatellus by 501 diffusion, making human transport an important dispersal mechanism. The 502 presence of A. cristatellus at forested parks located adjacent to spatially isolated 503 regional waste transfer stations suggests yard waste collection and transport may 504 be one such method of dispersal.

505 Vegetation in some residential areas within the core distribution of A. 506 cristatellus can change rapidly over short distances, likely affecting the ability of A. 507 *cristatellus* to spread to new areas. The transition from presence to absence of A. 508 *cristatellus* coincides with an abrupt increase in impervious surface and a loss of 509 forest habitat in some areas (see Fig. 2). The current distribution of *A. cristatellus* 510 includes mostly higher-income neighborhoods including parts of Coconut Grove, 511 Coral Gables, Pinecrest, and Key Biscayne (American Community Survey 2013; see 512 also Ackley et al. 2015b). Socio-economic factors influence surface temperatures 513 primarily through their impact on vegetation cover (Grove et al. 2006; Jenerette et al. 2007; Boone et al. 2010); such that areas with dense, mature tree canopies will 514 produce relatively cooler microclimates suitable for *A. cristatellus*. These underlying 515

516	effects of urban vegetation on available microclimates provide a mechanistic
517	understanding of the current distribution of A. cristatellus in Miami. Other studies of
518	urban and fragmented landscapes show species presence connected with other key
519	resources, such as prey availability (e.g., Sullivan et al. 2014), shelter availability
520	(e.g., Fischer et al. 2005), and structural habitat (e.g., Sarre et al. 1995; Garden et al.
521	2007; Santos et al. 2008) as well as urban development (e.g., Germaine and
522	Wakeling 2001). Future studies should test whether socio-economic factors are
523	correlated with vegetation and microclimates, and thus potentially useful for
524	predicting the spread of <i>A. cristatellus</i> in Miami.
525	
526	Thermal microclimates
527	We found substantial temperature differences between copper lizard models
528	on open versus shaded trees (Fig. 3). Non-thermoregulating lizards would
529	experience a temperature difference of up to a 7.6°C in the morning (1000 h) and a
530	5.7°C in the afternoon (1600 h). Open trees, but not shaded ones, experienced
531	temperatures exceeding observed field T_b for <i>A. cristatellus</i> in the summer in Miami
532	(Fig. 3). Denser overstory vegetation will produce relatively cooler microclimates
533	favorable for A. cristatellus in the city. Shade from vegetation cooled buildings up to
534	11.7°C during summer conditions in Toronto, Canada (Millward et al. 2014), and
535	shade from individual trees in city parks decreased average air temperatures by
536	10% and increased relative humidity by $18%$ in Thessaloniki, Greece (Georgi and
537	Zafiriadis 2006). Ackley et al. (2015a), using copper lizard models, found that
538	microclimates in areas with mesic landscaping were $5-10^{\circ}$ C cooler than those in

539 native xeric landscapes, even though the mean surface temperature of Phoenix, 540 Arizona, USA was 3°C warmer than the surrounding desert. Interestingly, surface 541 temperatures in Phoenix were related to vegetation during the daytime and the 542 proportion of paved area during the night (Buyantuyev and Wu 2010). Daytime 543 temperatures may limit activity or approach maximum thermal limits, whereas 544 nighttime temperatures likely influence metabolic costs during times of inactivity. 545 Whether the distribution of *A. cristatellus* in Miami is limited by daytime 546 temperatures driven by vegetation, nighttime temperatures related to impervious 547 surfaces, or both is a key question for future study.

548 Copper lizard model temperatures do not account for the ability of lizards to 549 thermoregulate. If suitably cool microhabitats were nearby, lizards in open areas 550 could behaviorally thermoregulate to preferred temperatures by shuttling between 551 warm and cool spots, at the cost of increased movement rates. The cost of 552 thermoregulation is predicted to be lower in more open sites because of the shorter 553 distance to sunny patches, which lowers the energetic cost of shuttling between sun 554 and shade (Huey 1974; Huey and Slatkin 1976; Huey and Webster 1976; Angilletta 555 2009). Accordingly, previous studies of *A. cristatellus* in Puerto Rico found that 556 lizards actively thermoregulate in open habitats, but thermoconform in less 557 variable, closed canopy habitats (Huey and Webster 1976). This versatility in 558 thermoregulatory behavior suggests that A. cristatellus might occupy both open and closed canopy sites in Miami; however, A. cristatellus is generally restricted to 559 560 relatively closed canopy locations (< 22% canopy openness; Fig. 4).

561 Our results suggest at least two possible explanations for this pattern. The 562 first is that A. cristatellus uses relatively cooler microclimates strictly due to its 563 thermal requirements: open canopy areas in Miami may be too warm relative to the 564 preferred temperature and upper thermal limit of *A. cristatellus*, and thus not 565 suitable for this species (Fig. 3). A second possibility is that A. cristatellus is 566 excluded from warmer areas by the presence of *A. distichus* and *A. sagrei*, which 567 both occupy warmer thermal niches than *A. cristatellus* (Huey and Webster 1976; 568 Lee 1980; Gunderson and Leal 2012; Leal and Gunderson 2012; Kolbe et al. 2012, 569 2014; this study). The relative importance of these two factors on limiting the 570 spread of *A. cristatellus* in the Miami area is an open question. These hypotheses 571 need to be comprehensively evaluated by including sites where each species is 572 present in the absence of the other as well as locations where they coexist. The 573 importance of microclimates to competitive interactions between the species, 574 allowing coexistence or contributing to competitive exclusion, warrants further 575 investigation.

576 During the summer in Miami, shade from urban vegetation is expected to 577 produce microclimates closer to the preferred body temperature of *A. cristatellus* as 578 compared to more open areas (Fig. 3). Higher activity rates are predicted when 579 lizards are closer to their preferred temperature (Gunderson and Leal 2015), 580 allowing lizards to better forage, mate, defend their territories, and escape from 581 predators. Mean body temperatures for A. cristatellus in Miami (28.8°C) and at low-582 elevation, mesic sites in Puerto Rico (~ 29°C from numerous sites; Huey and 583 Webster 1976; Hertz 1992a,b; Gunderson and Leal 2012) were similar to preferred

584 temperatures for lizards from three locations in Puerto Rico (range = $29.0-29.6^{\circ}$ C; 585 Huey and Webster 1976; Fig. 3). This suggests that some aspects of the thermal 586 niche of *A. cristatellus* are conserved between introduced populations in South 587 Miami and their low-elevation source population in northeast Puerto Rico (Kolbe et 588 al. 2007). This similarity in field body temperatures occurs despite shifts in other 589 aspects of their thermal niche, specifically the introduced population in South Miami 590 rapidly acquired the ability to tolerate lower temperatures relative to its source 591 population in Puerto Rico (see Kolbe et al. 2012; Leal and Gunderson 2012). 592

593 Effect of species interactions on the spread of A. cristatellus

594 Interspecific interactions, primarily competition, are thought to be important 595 factors structuring both native and introduced Anolis lizard communities (Losos et 596 al. 1993; Losos 2009). Previous experimental studies of anoles have found effects 597 on abundance and structural habitat use consistent with interspecific competition 598 when species coexist (e.g., Pacala and Roughgarden 1982; Rummel and 599 Roughgarden 1985; Leal et al. 1998; Stuart et al. 2014). In accordance with 600 predictions based on ecological similarity (primarily perch height), A. cristatellus 601 presence showed the strongest negative association with A. sagrei, followed by A. 602 distichus, and in a few cases with A. carolinensis and A. equestris. Thus, ecological 603 similarity of interacting species may provide important information for predicting 604 patterns of establishment and range expansion dynamics for introduced species. 605 The negative relationship between A. cristatellus and A. sagrei in Miami may 606 be explained by resource competition and agonistic interference (Salzburg 1984,

607 Losin 2012). When *A. cristatellus* was experimentally removed from plots five years 608 after its initial introduction in 1981, A. sagrei rapidly shifted back to the structural 609 habitat previously occupied by *A. cristatellus* — off the ground, on to trunks, and to 610 shadier sites (Salzburg 1984). We found consistent patterns, with A. cristatellus 611 occupying higher and broader perches as well as shadier microhabitats when 612 sympatric with A. sagrei. Additionally, A. cristatellus was far less abundant when 613 coexisting with congeners compared to when alone. However, competitive 614 interactions between A. cristatellus and A. sagrei may have changed over time with 615 their coexistence. Thirty years later, at the same site as Salzburg's experiment, 616 another removal experiment did not influence habitat use or body condition of these 617 two species (Losin 2012). Furthermore, A. sagrei lizards found sympatric with A. 618 *cristatellus* were less aggressive toward this species compared to *A. sagrei* from 619 allopartric populations (Losin 2012). Aggressive individuals facilitated the rapid 620 range expansion of western bluebirds in the northwestern U.S., but following 621 displacement of mountain bluebirds, their aggressive behavior decreased rapidly 622 (Duckworth & Badyaev 2007). Given that A. sagrei is ubiquitous in Miami and A. 623 *cristatellus* is still spreading, the opportunity exists to study resource use and 624 aggression of A. sagrei before and after the arrival of A. cristatellus.

625

626 Summary

The occurrence of introduced *A. cristatellus* in Miami is strongly associated
with forest habitat — dense vegetation, high canopy cover and low impervious
surface — and the lack of congeners, particularly *A. sagrei*. Given the correlative

630	nature of our analyses, it is difficult to tease apart the causal effects of urban
631	vegetation and species interactions for limiting the spread of <i>A. cristatellus</i> .
632	However, because A. sagrei already occupies nearly all habitats in Miami and forest
633	habitat is highly fragmented across the city, we predict that dispersal to forest
634	habitat will be the primary factor limiting future spread of <i>A. cristatellus</i> . Human-
635	mediated, sometimes long-distance, dispersal is likely to contribute to spread as
636	well as movement by diffusion through corridors of suitable habitats.
637	
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- Table 1. Variables recorded at each tree along transects designed to cross the
- 648 distributional boundary of *A. cristatellus* in the Miami metropolitan area. Tree
- 649 characteristics relate to the focal tree sampled and its surrounding vegetation. A
- total of 1035 trees were sampled on 19 transects in five plots. Congener presence
- 651 refers to the four other species of *Anolis* lizards with distributions that overlap with
- 652 *A. cristatellus* in Miami.
- 653

Variable	Description
Tree characteristics:	
Tree species	Species of tree
Native/non-native	Native or non-native tree species
Palm/non-palm	Palm or non-palm tree species
Trunk number	Number of trunks \geq 10 cm at 1.35 m height;
	Single, double, or multiple (> 2 trunks)
Bark texture	<i>Overlapping</i> (overlapping protrusions covering the trunk, such as palms covered with the bases of dead leaf pedicles); <i>shallow furrows</i> (bark with furrows, gaps, or cracks ≥ 0.5 cm and no overlapping or flaking); <i>deep furrows</i> (bark separating or flaking ≥ 0.75 cm over an area $\geq 2x4$ cm) or <i>smooth</i> (lacking any of the above characteristics)
Trunk diameter	Diameter (cm) of trunk at 1.35 m height
Canopy diameter	Mean canopy diameter (m) estimated from several
	measurements of the radius of the canopy
Distance to nearest plant	Distance (m) to the nearest stem at 1.35 m height
Distance to nearest tree	Distance (m) to the nearest tree with a diameter ≥ 30 cm at 1.35 m height
Overstory canopy cover	Mean percent overstory canopy cover both facing towards and away from the road as measured using a spherical densiometer
Congener presence:	
A. carolinensis	Presence/absence of A. carolinensis
A. distichus	Presence/absence of A. distichus
A. equestris	Presence/absence of A. equestris
A. sagrei	Presence/absence of A. sagrei

Table 2. GIS attributes derived from maps of forest fragmentation (NOAA 2010), percentage of human-made impervious surface (MRLC 2011), and variables recorded along transects.

GIS Attribute	Description
Block length	Length of each block in km
Trees per km	Number of trunks / Block length * 1000
Block canopy density	Number of trunks * Mean canopy diameter / Block length
Canopy cover GIS	Proportion of a 5-m buffer around the block transects including pixels classified as any type of forest
Distance to nearest forest	Distance (m) from each point to the center of the nearest pixel of any type of forest
Distance to nearest patch forest	Distance (m) from each point to the center of the nearest pixel of patch forest
Distance to nearest block forest	Distance (m) from each point to the center of the nearest pixel of non-patch forest (i.e., either perforated, edge or core forest)
Forest class	Indicates the type of forest for a point: 0 (no forest), 1 (patch), 2 (perforated/edge/core)
Impervious surface (1 pixel)	Value of the impervious raster pixel where each point is located (30m by 30m)
Impervious surface (9 pixels)	Average value of the 9 impervious raster pixels around each point (90m by 90m)
Impervious surface (25 pixels)	Average value of the 25 impervious raster pixels around each point (150m by 150m)

Table 3. Counts for the number of lizards present on surveyed trees (n=1035) and the percent of trees occupied by the five *Anolis* species encountered on transects in Miami. Results for each of three surveys are shown for A) all data, B) sections of transects where *A. cristatellus* is present, and C) sections of transects where *A. cristatellus* is absent. The combined surveys column indicates whether a species was present on a tree in at least one of the three surveys; these data were used in analyses.

A) 1st survey Number of lizards							2nd survey Number of lizards							3rd Numbe	l surv er of	•		Combined surveys Number of lizards			
Species	0	1	2	3	≥4	%	0	1	2	3	≥4	%	0	1	2	3	≥4	%	Absent	Present	%
A. carolinensis	986	45	4	0	0	4.7	967	64	3	1	0	6.6	940	87	6	1	0	9.1	856	179	17.3
A. cristatellus	895	107	23	7	3	13.5	873	119	29	12	2	15.7	895	115	17	5	3	13.5	781	254	24.5
A. distichus	865	133	28	8	1	16.4	829	142	50	10	4	19.9	788	170	54	16	7	23.9	636	399	38.6
A. equestris	1029	6	0	0	0	0.6	1029	6	0	0	0	0.6	1013	6	0	0	0	0.6	1018	17	1.6
A. sagrei	838	135	43	10	9	19.0	812	157	47	8	11	21.5	831	142	36	11	14	19.6	660	375	36.2

B) 1st survey Number of lizards					2nd survey Number of lizards							3rd umbe	surv r of l		ds		Combined surveys Number of lizards				
Species	0	1	2	3	≥4	%	0	1	2	3	≥4	%	0	1	2	3	≥4	%	Absent	Present	%
A. carolinensis	554	25	4	0	0	5.0	553	30	0	0	0	5.1	539	41	2	1	0	7.5	492	91	15.6
A. cristatellus	443	107	23	7	3	24.0	421	119	29	12	2	27.8	443	115	17	5	3	24.0	329	254	43.6
A. distichus	495	70	15	2	1	15.1	483	72	23	4	1	17.2	476	78	22	4	3	18.4	383	200	34.3
A. equestris	578	5	0	0	0	0.9	579	4	0	0	0	0.7	579	4	0	0	0	0.7	571	12	2.1
A. sagrei	492	66	17	3	5	15.6	496	61	16	4	6	14.9	512	51	11	4	5	12.2	428	155	26.6

C)	N		t surv er of		rds		N	2no umbo	d sur er of		rds		3rd survey Number of lizards					Combined surveys Number of lizards				
Species	0	1	2	3	≥4	%	0	1	2	3	≥4	%	0	1	2	3	≥4	%	Absent	Present	%	
A. carolinensis	432	20	0	0	0	4.4	414	34	3	1	0	8.4	401	46	4	0	0	11.1	364	88	19.5	
A. cristatellus																					0.0	
A. distichus	370	63	13	6	0	18.1	346	70	27	6	3	23.5	311	92	32	12	4	31.0	253	199	44.0	
A. equestris	451	1	0	0	0	0.2	450	2	0	0	0	0.4	434	2	0	0	0	0.5	447	5	1.1	
A. sagrei	346	69	26	7	4	23.5	316	96	31	4	5	30.1	319	91	25	7	9	29.3	232	220	48.7	

Table 4. Inferential statistics based on tree characteristics showing the most likely generalized linear models for presenceabsence of *A. cristatellus* when A) dividing sections of transects into areas with *A. cristatellus* present versus absent (two groups), B) comparing the presence versus absence of *A. cristatellus* on all trees pooled, and C) comparing the presence versus absence of *A. cristatellus* on trees within only the sections of transects with *A. cristatellus* present, and for abundance of *A. cristatellus* for D) all trees pooled and E) within only the sections of transects with *A. cristatellus* present.

A)

Effect df	Error df	Z	P
1	1031	0.09	0.06
1	1030	0.13	0.16
1	1029	-0.53	< 0.0001
1	1028	0.41	< 0.0001
	Effect df 1 1 1 1	1 1031 1 1030 1 1029	110310.09110300.1311029-0.53

Note: second best model \triangle AIC = 0.8; including the Distance to Nearest Tree effect; P=0.27.

B)

Effect	Effect df	Error df	Z	Р
Canopy diameter	1	1031	0.27	< 0.001
Distance to nearest plant	1	1030	-0.86	< 0.0001
Distance to nearest tree	1	1029	-0.20	0.05
Overstory canopy cover	1	1028	0.90	< 0.0001

Note: second best model \triangle AIC = 1.6; including the Trunk Diameter effect; P=0.53.

C)

Effect	Effect df	Error df	Z	Р
Trunk diameter	1	580	0.51	< 0.0001
Distance to nearest plant	1	579	-0.87	< 0.0001
Overstory canopy cover	1	578	0.72	< 0.0001

Note: second best model \triangle AIC = 1.37; including the Distance to Nearest Tree effect; P=0.43.

D)

Effect	Effect df	Error df	Z	Р
Trunk diameter	1	1031	4.46	< 0.0001
Canopy diameter	1	1030	2.24	0.03
Distance to nearest plant	1	1029	-5.71	< 0.0001
Overstory canopy cover	1	1028	3.65	< 0.0001

Note: second best model Δ AIC=1.23; including the Distance to Nearest Tree effect; P=0.38.

E)

Effect	Effect df	Error df	Z	Р
Trunk diameter	1	579	0.68	< 0.0001
Distance to nearest plant	1	580	-0.34	< 0.0001
Overstory canopy cover	1	578	0.32	< 0.0001

Note: second best model Δ AIC=1.59; including the Canopy Diameter effect; P=0.52.

Table 5. Inferential statistics for presence-absence of *A. cristatellus* showing the most likely generalized linear models when A) comparing street blocks with *A. cristatellus* present versus absent (n=93) and B) comparing the presence versus absence of *A. cristatellus* in 30 x 30 pixels (n=839). Selection of most favored models was supported by a likelihood ratio test against reduced models containing only the intercept term (A: $\chi^2 = 51.19$, *P* <0.0001; B: $\chi^2 = 203.24$, *P* <0.0001).

A)

Effect	Effect df	Error df	Z	Р
Distance to nearest plant	1	92	-1.799	0.07
Mean overstory canopy cover	1	92	2.007	0.05
Trees per km	1	92	1.865	0.06
% Impervious-raster	1	92	-2.386	0.02

Note: second best model \triangle AIC = 1.25; including the Distance to Nearest Tree effect; *P* = 0.39.

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Effect	Effect df	Error df	Ζ	Р
Distance to nearest patch forest	1	838	8.752	< 0.0001
Distance to nearest block forest	1	838	-6.446	< 0.0001
Impervious surface (1 pixel)	1	838	-2.947	< 0.01
Impervious surface (25 pixels)	1	838	-4.318	< 0.0001

Note: second best model \triangle AIC = 1.06; including the Forest_Class effect; *P* = 0.39.

Table 6. Results of likelihood ratio tests to determine whether the presence-absence of *A. cristatellus* on trees in Miami was related to the presence-absence of each of its four congeners. We combined trees on transects from each of the five plots analyzing A) all trees pooled and B) only trees from sections of transects with *A. cristatellus* present.

A)									
		A. carolinensis		A. distichus		A. sagrei		A. equestris	
Plot	Ν	X ²	Р	X ²	Р	X ²	Р	X ²	Р
Charles	163	5.71	0.02	17.92	< 0.0001	18.90	< 0.0001	0.26	0.61
Le Jeune	125	0.25	0.61	0.27	0.60	4.21	0.04	7.32	< 0.01
Ludlum	270	0.59	0.44	1.32	0.25	7.81	< 0.01	0.03	0.86
Maynada	137	6.29	0.01	2.23	0.14	3.68	0.06	0.74	0.39
SW 104	340	0.56	0.45	0.85	0.36	28.91	< 0.0001	0.76	0.38
All Plots Combined	1035	2.38	0.12	7.21	< 0.01	59.19	< 0.0001	2.31	0.13

B)

	A. carolinensis			A. distichus		A. sagrei		A. equestris	
Plot	Ν	X ²	Р	X ²	Р	X ²	Р	X ²	Р
Charles	87	1.12	0.29	2.24	0.13	0.68	0.41	0.10	0.75
Le Jeune	80	0.42	0.52	0.13	0.72	1.95	0.16	4.60	0.03
Ludlum	215	0.00	0.99	0.39	0.53	7.52	< 0.01	0.04	0.85
Maynada	48	5.55	0.02	5.35	0.02	1.70	0.19	1.32	0.25
SW 104	153	0.20	0.65	0.53	0.47	3.34	0.07	2.77	0.10
All Plots Combined	583	0.71	0.40	1.58	0.21	22.23	< 0.0001	1.08	0.30

Figures Captions

Fig. 1. Location of our study area in metropolitan Miami-Dade County of southeast Florida showing the core area in South Miami and sampling locations for *A. cristatellus*. Inset photo of a male *A. cristatellus* perched on a brick wall (J. Kolbe).

Fig. 2. Maps of a portion of the core area near Coconut Grove showing *A. cristatellus* presence and absence points and A) the four types of forest fragments and B) the percentage of impervious surface within 75 m.

Fig. 3. Mean temperatures for copper lizard models placed on the trunks of two trees, one with an open canopy and the other with a shaded canopy, in each cardinal direction. Points are hourly means collected over a 3.5-day period in July 2014 (error bars are omitted for clarity). Patterned shading (gray) shows the range of field T_b for *A. cristatellus* in Miami during each hour from 0800-1400 from this study as well as the preferred T_b (light gray) and the critical thermal maximum (dotted line) of *A. cristatellus* measured for populations from Puerto Rico (Huey and Webster 1976).

Fig. 4. Relationships between lizard field body temperature or copper lizard model temperature and significant covariates from the ANCOVA: a) canopy openness, b) air temperature, and c) time of day for *A. cristatellus* (black circles), *A. sagrei* (white circles), and copper lizard models at random locations (gray circles) in South Miami. Separate slopes are shown for the significant temperature by canopy openness interaction.

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Fig. 1.

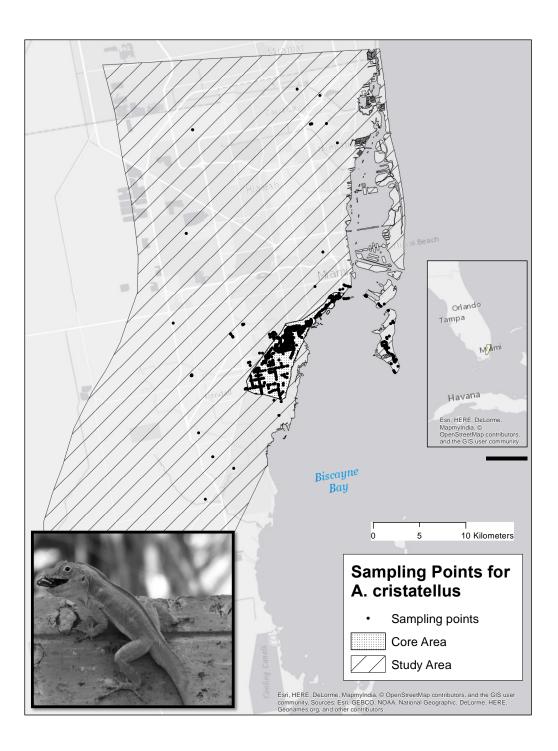
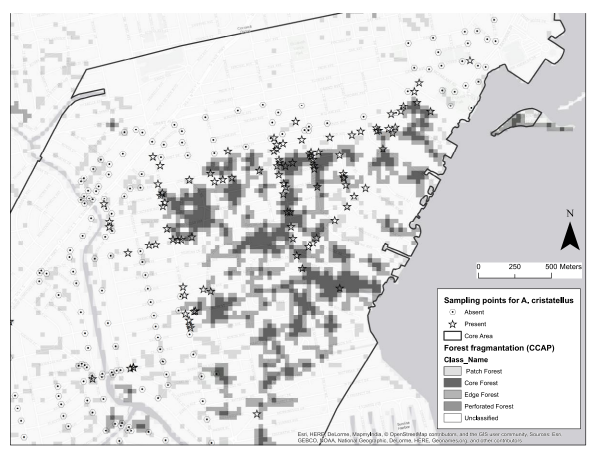
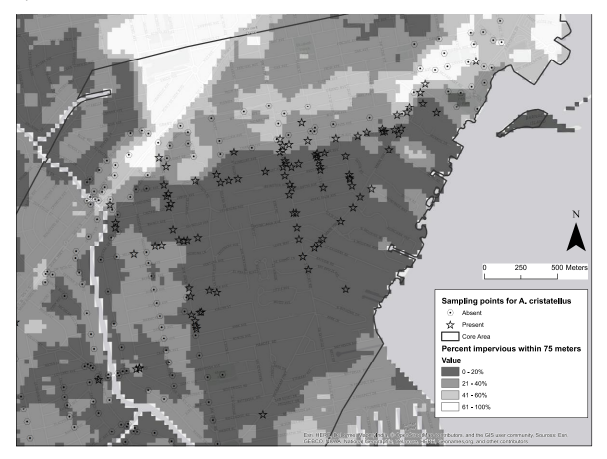


Fig. 2.









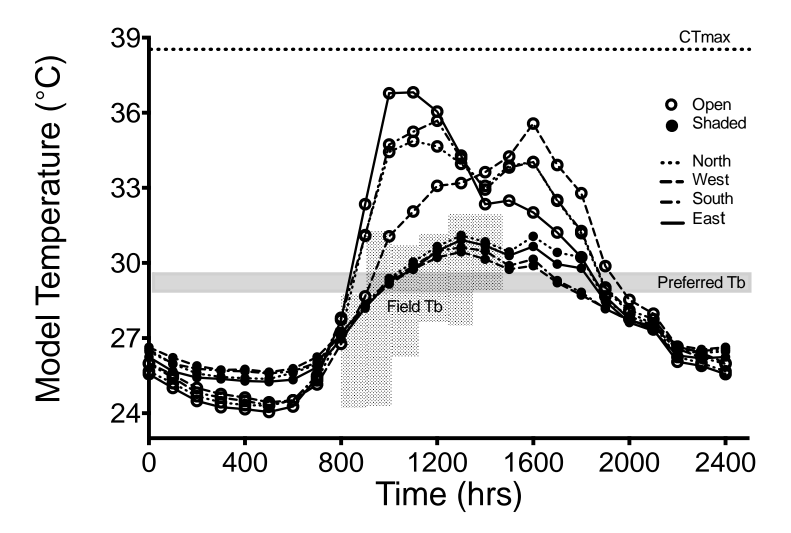
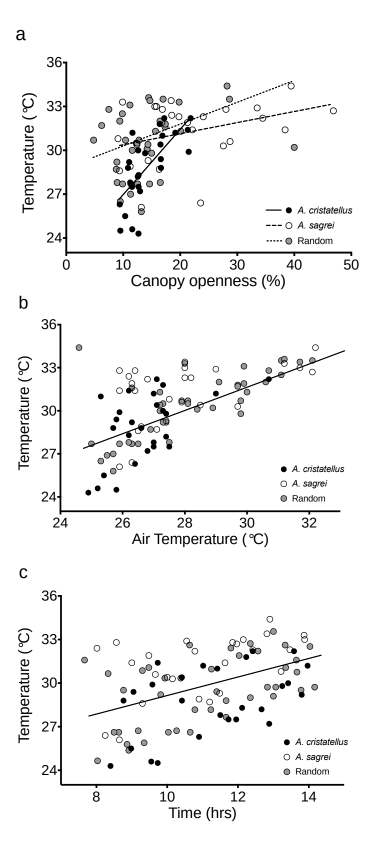


Fig. 4.



References

Ackley JW, Angilletta MJ Jr., DeNardo D, Sullivan B, Wu J (2015a) Urban heat island mitigation strategies and lizard thermal ecology: landscaping can quadruple potential activity time in an arid city. Urban Ecosyst DOI 10.1007/s11252-015-0460-x.

Ackley JW, Wu J, Angilletta MJ Jr., Myint SW, Sullivan B (2015b) Rich lizards: how affluence and land cover influence the diversity and abundance of desert reptiles persisting in an urban landscape. Biol Conserv 182:87-92.

Akaike H (1974) A new look at the statistical model identification. IEEE Transactions on Automatic Control 19:716–723.

Akbari H, Pomerantz M, Taha H (2001) Cool surfaces and shade trees to reduce energy use and improve air quality in urban areas. Solar Energy 70:295-310.

American Community Survey (2013) United States Census Bureau. Available from http://www.census.gov (accessed August 2015)

Anderson L, Burgin S (2008) Patterns of bird predation on reptiles in small woodland remnant edges in peri-urban north-western Sydney, Australia. Landscape Ecol 23:1039-1047.

Angilletta MJ Jr (2009) Thermal adaptation: a theoretical and empirical synthesis. Oxford University Press, Oxford

Arnfield AJ (2003) Two decades of urban climate research: a review of turbulence, exchanges of energy and water, and the urban heat island. Int J Climatol 23:1-26.

Bartlett RD, Bartlett PP (1999) A field guide to Florida reptiles and amphibians. Gulf Publishing, Houston.

Bonnington C, Gaston KJ, Evans KL (2014) Squirrels in suburbia: influence of urbanisation on the occurrence and distribution of a common exotic mammal. Urban Ecosyst 17:533-546.

Boone CG, Cadenasso ML, Grove JM, Schwarz K, Buckley GL (2010) Landscape, vegetation characteristics, and group identity in an urban and suburban watershed: why the 60s matter. Urban Ecosyst 13:255-271.

Broennimann O, Fitzpatrick MC, Pearman PB, Petitpierre B, Pellissier L, Yoccoz NG, Thuiller W, Fortin M-J, Randin C, Zimmermann NE, Graham CH, Guisan A (2012) Measuring ecological niche overlap from occurrence and spatial environmental data. Global Ecol Biogeo 21:481–497. Buyantuyev A, Wu J (2010) Urban heat islands and landscape heterogeneity: linking spatiotemporal variations in surface temperatures to land-cover and socioeconomic patterns. Landscape Ecol 25:17-33.

Cadenasso ML, Pickett STA, Schwarz K (2007) Spatial heterogeneity in urban ecosystems: reconceptualizing land cover and a framework for classification. Front Ecol Environ 5:80-88.

Case TJ, Bolger DT, Petren K (1994) Invasions and competitive displacement among house geckos in the Tropical Pacific. Ecology 75:464-477.

Chen WY, Jim CY (2008) Assessment and valuation of the ecosystem services provided by urban forests. In: Ecology, planning, and management of urban forests international perspective. Springer, New York, pp 53–83.

Corn MJ (1971) Upper thermal limits and thermal preferenda for three sympatric species of *Anolis*. J Herpetol 5:17-21.

Crump ML, Scott NJ Jr (1994) Visual encounter surveys, p. 84–92. In: Measuring and Monitoring Biological Diversity: Standard Methods for Amphibians. W. R. Heyer, M. A. Donnelly, R. W. McDiarmid, L. A. C. Hayek, and M. S. Foster (eds.). Smithsonian Institution Press, Washington, D.C.

Davis MA (2009) Invasion Biology. Oxford University Press.

Duckworth RA, Badyaev AV (2007) Coupling of dispersal and aggression facilities the rapid range expansion of a passerine bird. PNAS 104:15017-15022.

Forman RTT (2014) Urban Ecology. Cambridge University Press.

Fischer J, Lindenmayer DB, Barry S, Flowers E (2005) Lizard distribution patterns in the Tumut fragmentation "Natural Experiment" in south-eastern Australia. Biol Conserv 123:301-315.

Frazer GW, Canham CD, Lertzman KP (1999) Gap Light Analyzer (GLA), Version 2.0: Imaging software to extract canopy structure and gap light transmission indices from true-colour fisheye photographs, users manual and program documentation. Copyright © 1999: Simon Fraser University, Burnaby, British Columbia, and the Institute of Ecosystem Studies, Millbrook, New York.

Garden JG, McAlpine CA, Possingham HP, Jones DN (2007) Habitat structure is more important than vegetation composition for local-level management of native terrestrial reptile and small mammal species living in urban remnants: a case study from Brisbane, Australia. Austral Ecol 32:669-685. Georgi NJ, Zafiriadis K (2006) The impact of park trees on microclimate in urban areas. Urban Ecosyst 9:195-209.

Germaine SS, Wakeling BF (2001) Lizard species distributions and habitat occupation along an urban gradient in Tucson, Arizona, USA. Biol Conserv 97:229-237.

Grove JM, Troy AR, O'Neil-Dunne JPM, Burch WR, Cadenasso ML, Pickett STA (2006) Characterization of households and its implications for the vegetation of urban ecosystems. Ecosystems 9:578-597.

Gunderson AR, Leal M (2012) Geographic variation in vulnerability to climate warming in a tropical Caribbean lizard. Funct Ecol 26:783-793.

Gunderson AR, Leal M (2015) Patterns of thermal constraint on ectotherm activity. Am Nat 185: In press.

Hamdi R, Schayes G (2008) Sensitivity study of the urban heat island intensity to urban characteristics. Int J Climatol 28:973-982.

Hines JE (2006) PRESENCE - Software to estimate patch occupancy and related parameters. USGS-PWRC http://www.mbr-pwrc.usgs.gov/software/presence.html.

Huang L, Li J, Zhao D, Ahu J (2008) A fieldwork study on the diurnal changes of urban microclimate in four types of ground cover and urban heat island of Nanjing, China. Building and Environ 43:7-17.

Hertz PE (1992) Temperature regulation in Puerto Rican *Anolis* lizards: a field test using null hypotheses. Ecology 73:1405–1417.

Huey RB (1974) Behavioral thermoregulation in lizards: importance of associated costs. Science 184:1001–1003.

Huey RB, Webster TP (1976) Thermal biology of *Anolis* lizards in a complex fauna: the *cristatellus* group on Puerto Rico. Ecology 57:985–994.

Huey RB, Slatkin M (1976) Cost and benefits of lizard thermoregulation. Quart Rev Biol 51:363-384.

Irschick DJ, Carlisle E, Elstrott J, Ramos M, Buckley C, Vanhooydonck B, Meyers J, Herrell A (2005) A comparison of habitat use, morphology, clinging performance and escape behaviour among two divergent green anole lizard (*Anolis carolinensis*) populations. Biol J Linnean Soc 85:223-234. Jenerette GD, Harlan SL, Brazel A, Jones N, Larsen L, Stefanov WL (2007) Regional relationships between surface temperature, vegetation, and human settlement in a rapidly urbanizing ecosystem. Landscape Ecol 22:353-365.

Kolbe JJ, Glor RE, Schettino LR, Lara AC, Larson A, Losos JB (2007) Multiple sources, admixture, and genetic variation in introduced *Anolis* lizard populations. Conserv Biol 21:1612-1625.

Kolbe JJ, Colbert PL, Smith BE (2008) Niche relationships and interspecific interactions in Antiguan lizard communities. Copeia 2008:261-272.

Kolbe JJ, VanMiddlesworth PS, Losin N, Dappen N, Losos JB (2012) Climatic niche shift predicts thermal trait response in one but not both introductions of the Puerto Rican lizard *Anolis cristatellus* to Miami, Florida, USA. Ecol Evol 2: 1503-1516.

Kolbe JJ, Ehrenberger JC, Moniz HA, Angilletta MJ (2014) Physiological Variation among Invasive Populations of the Brown Anole (*Anolis sagrei*). Physiol Biochem Zool 87:92-104.

Kolbe JJ, Battles AC, Avilés-Rodríguez KJ (2015) City slickers: poor performance does not deter *Anolis* lizards from using artificial substrates in human-modified habitats. Funct Ecol doi: 10.1111/1365-2435.12607

Kraus F (2009) Alien reptiles and amphibians: a scientific compendium and analysis. Springer.

Leal M, Gunderson AR (2012) Rapid change in the thermal tolerance of a tropical lizard. Am Nat 180:815-822.

Leal M, Rodríguez-Robles JA, Losos JB (1998) An experimental study of interspecific interactions between two Puerto Rican *Anolis* lizards. Oecologia 117:273-278.

Lee JC (1980) Comparative thermal ecology of two lizards. Oecologia 44:171-176.

Lever C (2003) *Naturalized reptiles and amphibians of the world.* Oxford University Press.

Liu X, Li X, Liu Z, Tingley R, Kraus F, Guo Z, Li Y (2014) Congener diversity, topographic heterogeneity and human-assisted dispersal predict spread rates of alien herpetofauna at a global scale. Ecol Lett 17:821-829.

Lockwood JL, Hoopes MF, Marchetti MP (2007) *Invasion Ecology*. Blackwell Publishing.

Losin NJE (2012) The evolution and ecology of interspecific territoriality: studies of *Anolis* lizards and North American wood-warblers. Dissertation. University of California Los Angeles.

Losos JB (1994) Integrative approaches to evolutionary ecology: *Anolis* lizards as model systems. Ann Rev Ecol Syst 25:467-493.

Losos JB (2009) *Lizards in an Evolutionary Tree: Ecology and Adaptive Radiation of Anoles*. University of California Press, Berkeley.

Losos JB, Marks JC, Schoener TW (1993) Habitat use and ecological interactions of an introduced and a native species of *Anolis* lizard on Grand Cayman. Oecologia 95:525-532.

Luck GW, Smallbone LT, O'Brien R (2009) Socio-economics and vegetation change in urban ecosystems: patterns in space and time. Ecosystems 12:604-620.

MacKenzie DI, Nichols JD, Lachman GB, Droege S, Royle JA, Langtimm CA (2002) Estimating site occupancy rates when detection probabilities are less than one. Ecology 83:2248-2255.

Marnocha E, Pollinger J, Smith TB (2011) Human-induced morphological shifts in an island lizard. Evol Appl 4:388-396.

Martin CA, Warren PS, Kinzig AP (2004) Neighborhood socioeconomic status is a usefull predictor of perennial landscape vegetation in residential neighborhoods and embedded small parks of Phoenix, AZ. Landscape Urban Plan 69:355-368.

McCullagh P, Nelder J (1989) Generalized linear models, 2nd ed. Chapman and Hall/CRC, Boca Raton.

Millward AA, Torchia M, Laursen AE, Rothman LD (2014) Vegetation placement for summer built surface temperature moderation in an urban microclimate. Environ Man 53:1043-1057.

Multi-resolution land characteristics (MRLC) consortium (2011) The national land cover database: percent developed imperviousness. Available from http://www.mrlc.gov

National Oceanic and Atmospheric Administration (NOAA) Office for Coastal Management (2010) Coastal Change Analysis Program (C-CAP) Regional Land Cover Database. Available from http://www.coast.noaa.gov/dataregistry/search/dataset/info/forestfrag

Nowak DJ, Rowntree RA, McPherson EG, Sisinni SM, Kerkmann ER, Stevens JC (1996) Measuring and analyzing urban tree cover. Landscape Urban Plan 36:49-57.

Pacala S, Roughgarden J (1982) Resource partitioning and interspecific competition in two two-species insular *Anolis* lizard communities. Science 217:444-446.

Perry G, Buchanan BW, Fisher RN, Salmon M, Wise SE (2008) Effects of artificial night lighting on amphibians and reptiles in urban environments. In *Urban Herpetology* (Mitchell JC, Jung Brown RE, Bartholomew B, eds). Society for the Study of Amphibians and Reptiles, pp. 239-256.

Petren K, Case TJ (1996) An experimental demonstration of exploitation competition in an ongoing invasion. Ecology 77:118-132.

Pianka ER (1981) Competition and niche theory. In *Theoretical Ecology* (May RM, ed). Blackwell, pp 167-196.

Pickett STA, Cadenasso ML Grove JM, Nilon CH, Pouyat RV, Zipperer WC, Costanza R (2001) Urban ecological systems: linking terrestrial ecological, physical, and socioeconomic components of metropolitan areas. Annu Rev Ecol Syst 32:127-157.

R Core Team (2013) R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. Available from http://www.R-project.org/

Ramalho CE, Hobbs RJ (2011) Time for a change: dynamic urban ecology. Trends Ecol Evol 27:179-188.

Rizwan AM, Dennis YCL, Liu C (2008) A review of the generation, determination and mitigation of Urban Heat Island. J Environ Sci 20:120-128.

Royle JA, Nichols JD (2003) Estimating abundance from repeated presence-absence data or point counts. Ecology 84:777-790.

Rummel J, Roughgarden J (1985) Effects of reduced perch-height separation on competition between two *Anolis* lizards. Ecology 66:430-444.

Salzburg MA (1984) *Anolis sagrei* and *Anolis cristatellus* in Southern Florida: a case study in interspecific competition. Ecology 65:14-19.

Santos T, Diaz JA, Perez-Tris J, Carbonell R, Tellera JL (2008) Habitat quality predicts the distribution of a lizard in fragmented woodlands better than habitat fragmentation. Animal Conserv 11:46-56.

Sarre S, Smith GT, Meyers JA (1995) Persistence of two species of gecko (*Oedura reticulata* and *Gehyra variegata*) in remnant habitat. Biol Conserv 71:25-33.

Schwartz A, Thomas R (1975) A checklist of West Indian amphibians and reptiles. Special Publication. Carnegie Museum of Natural History 1:1-216.

Shochat E, Warren PS, Faeth SH, McIntyre NE, Hope D (2006). From patterns to emerging processes in mechanistic urban ecology. Trends Ecol Evol 21:186-191.

Stuart YE, Campbell TS, Hohenlohe PA, Reynolds RG, Revell LJ, Losos JB (2014) Rapid evolution of a native species following invasion by a congener. Science 346: 463-466.

Sullivan BK, Sullivan KO, Vardukyan D, Suminski T (2014) Persistence of Horned Lizards (*Phrynosoma* spp.) in urban preserves of Central Arizona. Urban Ecosyst 17:707-717.

White CR (2003) Allometric analysis beyond heterogeneous regression slopes: use of the Johnson-Neyman technique in comparative biology. Physiol Biochem Zool 76:135-140.

Wilson LD, Porras L (1983) The ecological impact of man on the south Florida herpetofauna. University of Kansas Museum of Natural History Special Publication No. 9:1-89.

Wong NH, Yu C (2005) Study of green areas and urban heat island in a tropical city. Habitat International 29:547-558.

Supplementary Table 1. Estimates of detection probability calculated from singleseason presence-absence occupancy models using sampling covariates of each surveyor and all surveyors combined. Models were built using three repeat surveys of 38 independent sampling points. Mean values are shown with 1 SE in parentheses. No variation exists for PV detection probabilities because this surveyor was present for each sampling event. Low detection probabilities for *A. equestris* likely reflect the low numbers observed for this species (Table 3). Estimates of detection probabilities for the three most common species were sufficiently high (> 0.87) to consider detection unlikely to influence subsequent analyses of presence-absence and abundance.

	Model			
Species	psi(.),p(JJK)	psi(.),p(PV)	psi(.),p(RS)	psi(.),p(All Surveyors)
A. cristatellus	0.9003	0.6340	0.7781	0.8807
	(0.039)	(0.000)	(0.035)	(0.020)
A. sagrei	0.7368	0.9298	0.5921	0.9298
	(0.033)	(0.000)	(0.024)	(0.012)
A. distichus	0.7018	0.8772	0.6097	0.8772
	(0.028)	(0.000)	(0.028)	(0.006)
A. carolinensis	0.6524	0.7240	0.5766	0.7302
	(0.021)	(0.000)	(0.020)	(0.010)
A. equestris	0.4021	0.2323	0.4479	0.2056
	(0.014)	(0.000)	(0.013)	(0.009)

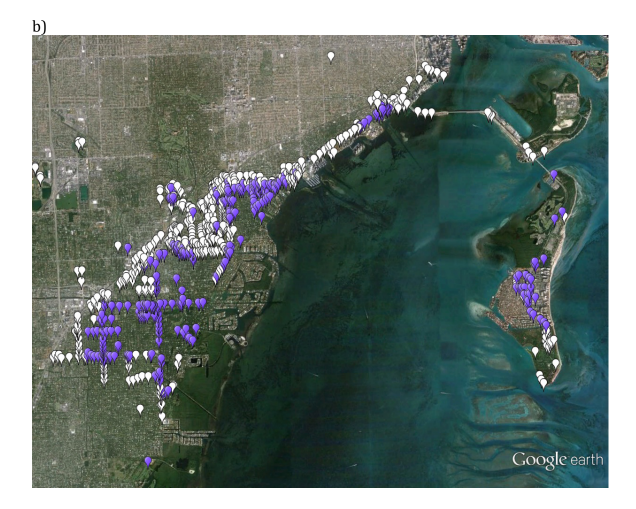
Supplementary Table 2. Proportion of forest fragmentation using previously defined forest categories and impervious surface cover using two spatial scales for the study area and core area in the Miami.

Attribute	Category	Proportion of	Proportion of	
		Study Area (1,511 km²)	Core Area (33.63 km ²)	
Forest fragmentation	Core Forest	0.02	0.06	
	Patch Forest	0.03	0.06	
	Perforated Forest	0.00	0.01	
	Edge Forest	0.03	0.13	
	Non-forest	0.92	0.73	
Impervious surface cover (within 15m)	0-20%	0.43	0.59	
	21-40%	0.27	0.28	
	41-60%	0.18	0.08	
	61-100%	0.13	0.06	
Impervious surface cover (within 75m)	0-20%	0.47	0.57	
	21-40%	0.20	0.26	
	41-60%	0.16	0.09	
	61-100%	0.17	0.08	

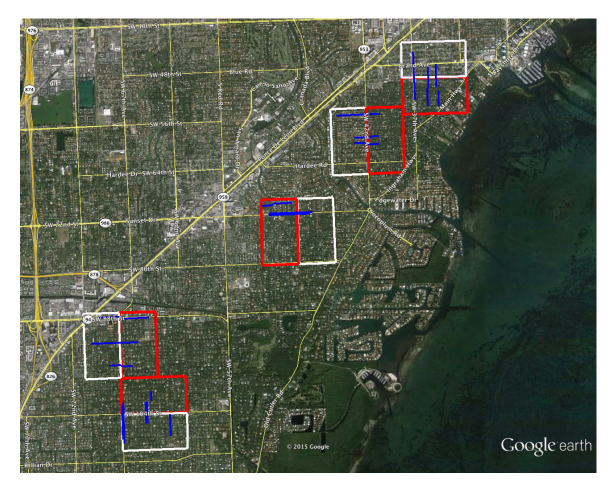
Supplementary Fig. 1. The distribution of *A. cristatellus* in the Miami metropolitan area showing presence (purple) and absence (white) points for a) the entire area surveyed including regional waste transfer stations and municipal parks and b) a view of the core distribution in South Miami and Key Biscayne (see attached .kmz file).



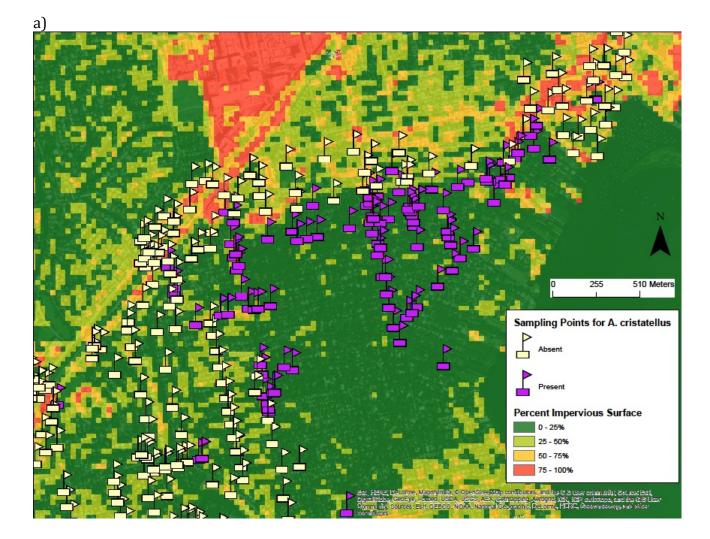
a)

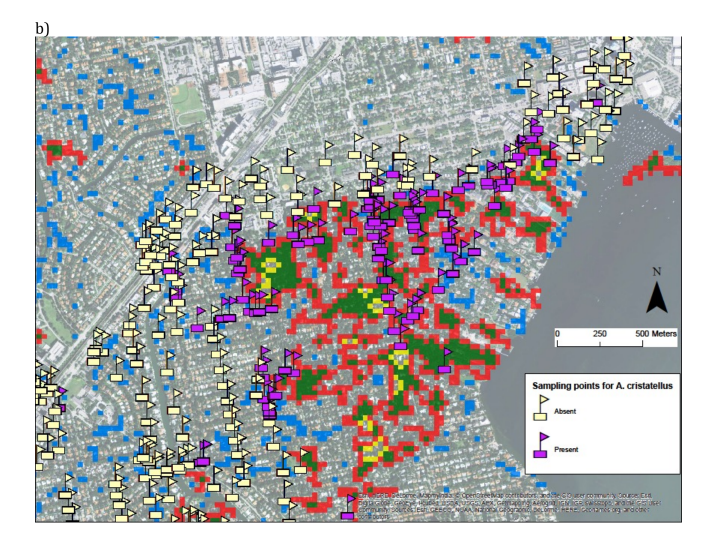


Supplementary Fig. 2. Map showing the location of the five plots and 19 transects in South Miami used to survey for the presence of *A. cristatellus*. We established plots crossing observed transitions from presence (red rectangles) to absence (white rectangles) of *A. cristatellus* based on preliminary surveys. In each plot, we established three to six roadside transects (blue lines) running perpendicular to the transition zone.



Supplementary Figure 3. GIS-based map for a portion of the distribution of *A. cristatellus* in Coconut Grove showing presence and absence points and a) percentage of impervious surface divided into four categories and b) the four forest fragmentation types. This is a color version of Fig. 2.





Supplementary Figure 4. Overstory canopy photos facing in the four cardinal directions for representative a) shaded and b) open canopy trees. Percentage of overstory cover is shown next to each photo.

