Fossils reject climate change as the cause of extinction of Caribbean bats

J. Angel Soto-Centeno1,2 & David W. Steadman2

1Department of Mammalogy, American Museum of Natural History, New York, NY 10024 USA, 2Department of Ornithology, Florida Museum of Natural History, University of Florida, Gainesville, FL 32611, USA.

We combined novel radiocarbon dates of bat fossils with time-scaled ecological niche models (ENM) to study bat extinctions in the Caribbean. Radiocarbon-dated fossils show that late Quaternary losses of bat populations took place during the late Holocene (< 4 ka) rather than late Pleistocene (> 10 ka). All bat radiocarbon dates from Abaco (Bahamas) that represent extirpated populations are younger than 4 ka. We include data on six bat species, three of which are Caribbean endemics, and include nectarivores as well as insectivores. Climate-based ENMs from the Last Glacial Maximum to the present reflect overall stability in distributions, with suitable climatic habitat being present over time. In the absence of radiocarbon dates, bat extinctions had been presumed to take place during the last glacial-interglacial transition (ca. 10 ka). Now we see that extirpation of bats on these tropical islands is more complex than previously thought and primarily postdates the major climate changes that took place during the late Pleistocene-Holocene transition.

The Caribbean islands, also known as the West Indies, once were home to at least 73 species of non-volant mammals (insectivores, primates, rodents, and sloths), all of which were endemic to one or few islands1–3. Sometime after the Last Glacial Maximum (LGM, ca. 25 to 18 ka), however, 79%–84% of the Caribbean land mammals became extinct4,5. It is estimated that humans arrived in the Caribbean about 7–6 ka and the effects of human activity for millennia after are well documented6,7. While various hypotheses have been proposed to explain the late Quaternary extinction of non-volant land mammals in the Caribbean, the timing of these losses is compatible with the concept that direct (e.g., hunting) and indirect (e.g., habitat modification) anthropogenic activities often were involved4,8,9.

In contrast to non-volant land mammals, bats (order Chiroptera) show much less extinction during the late Quaternary in the Caribbean, with an estimate that only ca. 18% of species (12 of 66) became extinct during that time4,10. Extirpation (population-level loss) appears to have been more common than extinction (species-level loss) in Caribbean bats; 33% (22 of 66 species) of extant Caribbean bats are known to have lost a population from one or more islands5. The extirpations and extinctions of Caribbean bats have been attributed primarily to reductions in land area and changes in climate and habitat during the last glacial-interglacial transition (the Pleistocene-Holocene Transition, or PHT; centered on 11–9 ka), which included changes in island size and in the distribution of caves and cave environments associated with this climate change4,11.

The past 2.5 million years have been characterized by at least 22 glacial-interglacial cycles with rising and falling sea levels that caused major changes in island size, shape, and isolation12,13. Because the last glacial interval lies within the time that can be radiocarbon dated (up to ca. 50 ka), special attention has been given to the transition from the LGM (ca. 25–18 ka) cold climate to the warmer interglacial climate of the Holocene14 (HOL; ca. 10–0 ka).

Ecological niche models (ENMs; also called species distribution models, or SDMs) are used to estimate the relationship between species occurrence and the environmental and spatial characteristics where they occur15. ENMs are widely used to study biogeographic, conservation, ecological, and evolutionary questions16–18. ENM projections allow us to hindcast species distributions across time, which in combination with fossil data provide a strong framework to study extinctions and especially extirpations. In this paper, we examine previous hypotheses about the loss of insular bats using three ecologically distinct cave-dwelling species (Macrotus waterhousii, Monophyllus redmani, and Pteronotus pumilio) that overlap in distribution and have been subjected to extirpation in the Caribbean. Specifically, we combine radiocarbon dates on fossils with modern (present) and fossil-validated past climate ENMs to provide new evidence on extirpation of Caribbean bats. If late Quaternary losses of these bats were driven by climate change during the PHT, then we predict that: 1) radiocarbon dated fossils from extirpated populations of bats should date to the PHT or older (i.e., > 9 ka); 2) model projections to the past...
should approximate the distribution of fossils but not the present species distribution; and 3) availability of suitable climate as assessed by ENMs should be smaller in the present than in the LGM projections and have little overlap, resulting in no stability in distributions across time.

Results

Fossils and Last Appearance Radiocarbon Dates. Establishing the time of prehistoric extinctions requires accurate Last Appearance Radiocarbon Dates (LADs). Nonetheless, until now, direct LADs on individual bones have not been done for Caribbean bats; $^{14}$C dates on other materials from various bat-bearing Caribbean fossil sites range from ca. 21,500 to 600 Cal BP\(^ {19-22}\), although most sites with bat fossils remain undated, even indirectly. To address this situation, we obtained six new AMS $^{14}$C dates from purified collagen of individual fossil humeri from five extirpated species of bats from Ralph’s Cave on Abaco (Table 1; Fig. 1). For perspective, we also examined 2000 bat fossils (representing 20 species) from 20 fossil sites across the Greater Antilles and The Bahamas. Here we focus on the nine species from four families identified from Ralph’s Cave, a site described by Steadman and Franklin\(^ {21}\). Of the 97 fossil bat specimens from Ralph’s Cave, 51 represent species that still exist on Abaco, namely Eptesicus fuscus (Big Brown Bat; N = 18 specimens), Chironatalus tumidifrons (Bahamian Funnel-eared Bat; N = 1), Erophylla sezekorni (Buffy Flower Bat; N = 2), and Macrotrus waterhousii (Waterhouse’s Leaf-nosed Bat; N = 2), Pteronotus parnelli (Parnell’s Mustached Bat; N = 5), Lasiurus major (Greater Antillean Long-tongued Bat; N = 30). Five of these nine species are extirpated from Abaco, namely Monophyllus redmani (Greater Antillean Long-tongued Bat; N = 2), Pteronotus parnelli (Parnell’s Mustached Bat; N = 7), Natalus primus (Cuban Funnel-eared Bat; N = 18), Lasiusus minor (Minor Red Bat; N = 5), and Myotis auroriparius (Southeastern Myotis; N = 14).

The six AMS $^{14}$C dates that we determined for the five extirpated species were 1820–1630 Cal BP for Monophyllus redmani (Great Antillean Long-tongued Bat; N = 2), Pteronotus parnelli (Parnell’s Mustached Bat; N = 7), Natalus primus (Cuban Funnel-eared Bat; N = 18), Lasiusus minor (Minor Red Bat; N = 5), and Myotis auroriparius (Southeastern Myotis; N = 14).

Table 1 | Accelerator Mass Spectrometer (AMS) radiocarbon ($^{14}$C) dates for individual humeri representing extirpated populations of bats from Ralph’s Cave, Abaco, The Bahamas\(^ b \) and a single extant population from Trouing Jeremie, Département-du-Sud, Haiti\(^ c \). Specimens from Ralph’s Cave are listed in chronological order. All determinations were done at Beta Analytic, Inc., Miami, FL. For laboratory and calibration methods see www.radiocarbon.com.

| Species                  | Skeletal element | Sample number | $^{14}$C (%) | Conventional $^{14}$C age (yr BP) | Calibrated $^{14}$C age (Cal BP, 2 $\sigma$) |
|--------------------------|------------------|---------------|--------------|-----------------------------------|-----------------------------------------------|
| Monophyllus redmani\(^ b \) | Humerus          | Beta-358786   | –17.6        | 1810 ± 30                         | 1820–1630                                     |
| Pteronotus parnelli\(^ b \) | Humerus          | Beta-345516   | –18.0        | 3310 ± 30                         | 3630–3460                                     |
| Lasiusus minor\(^ c \)    | Humerus          | Beta-360273   | –19.3        | 3580 ± 30                         | 3690–3560                                     |
| Natalus primus\(^ c \)    | Humerus          | Beta-358785   | –18.4        | 3390 ± 30                         | 3700–3570                                     |
| Pteronotus parnelli\(^ b \) | Humerus          | Beta-345515   | –18.6        | 3490 ± 30                         | 3840–3690                                     |
| Macrotrus waterhousii\(^ h \) | Humerus          | Beta-345518   | –20.3        | 2030 ± 30                         | 2060–1900                                     |

Table 1: Accelerator Mass Spectrometer (AMS) radiocarbon ($^{14}$C) dates for individual humeri representing extirpated populations of bats from Ralph’s Cave, Abaco, The Bahamas and a single extant population from Trouing Jeremie, Département-du-Sud, Haiti. Specimens from Ralph’s Cave are listed in chronological order. All determinations were done at Beta Analytic, Inc., Miami, FL. For laboratory and calibration methods see www.radiocarbon.com.

Modern and Past Locality Records. The three focal species of bat vary in their modern and past distributions (Supplementary Fig. S1). We compiled 168 modern localities and 41 fossil localities for Ma. waterhousii (Supplementary Fig. S1), which occurs today throughout The Bahamas, Cuba, Hispaniola, and Jamaica, and has been documented as a fossil on Puerto Rico\(^ {16,18,24,25}\), we compiled 143 modern and 16 fossil localities for My. auroriparius (Supplementary Fig. S1), which now occurs in the Greater Antilles, Crooked and Acklins Islands (The Bahamas), and the Turks and Caicos Islands\(^ {26}\), with a fossil record from Abaco (herein). We compiled 115 modern and 13 fossil localities for P. parnelli (Supplementary Fig. S1), which currently occurs throughout the Greater Antilles\(^ {27}\) and also is widespread on the mainland from Mexico to Brazil\(^ {28}\), it formerly occurred on Abaco and on Antigua\(^ {10,21,29}\) (also see Results herein).

Figure 1 | Bat humeri. (a), (c), (e), (g), and (j) are late Holocene fossils from Ralph’s Cave, Abaco. (h) is a late Holocene fossil from Trouing Jeremie, Haiti. (b), (d), (f), (i), and (k) are modern specimens. Monophyllus redmani (a, b), Myotis auroriparius (c, d), Lasiusus borealis cf. minor (e, f), Natalus primus (g), Macrotrus waterhousii (h, i). Pteronotus parnelli (j, k). Mo. redmani UF 4826, Haiti (b). My. auroriparius UF 24049, Florida (d). L. borealis cf. minor UF 30161, Florida (f). Ma. waterhousii UF 20812, Haiti (i). P. parnelli UF 6947, Guatemala (k). Fossil photographs by JAS-C.
Table 2 | Overlap index (OI) and relative range size (RRS) estimates for Macrotus waterhousii, Monophasys redmani, and Pteronotus parnellii on the Caribbean. Time of the predicted distributions is indicated as Current (CUR), Holocene (HOL), and Last Glacial Maximum (LGM). Threshold values were determined using the least training presence threshold LPT95%. RRS for LGM distributions corrected for current island area are indicated with an asterisk (*). Negative RRS values indicate that the more recent range is bigger than the past (i.e. range expansion). Conversely, positive RRS values indicate that the more recent range is smaller than the past (i.e. range contraction).

| Species          | Threshold Value | CUR-HOL  | HOL-LGM | CUR-LGM  | CUR-HOL  | HOL-LGM | CUR-LGM |
|------------------|-----------------|----------|---------|----------|----------|---------|---------|
| Ma. waterhousii  | 0.2             | 0.893    | 0.592   | 0.531    | -0.116   | 0.806   |
| Mo. redmani      | 0.22            | 0.865    | 0.984   | 0.952    | -0.147   | 1.718   | 1.369   |
| P. parnellii     | 0.15            | 0.729    | 0.890   | 0.830    | -0.308   | 1.511   | 0.919   | 0.141*  |

Modern and Fossil-Validated Paleo-ENMs. ENMs inferring present distributions performed well for all three species of bat with average Area Under the Curve (AUC) values of 0.810 ± 0.015 for Ma. waterhousii, 0.829 ± 0.018 for Mo. redmani, and 0.791 ± 0.015 for P. parnellii. Model projections to the HOL correctly predicted 87.5% and 92% of fossil localities of Mo. redmani and P. parnellii, respectively (Supplementary Fig. S2). In the case of Ma. waterhousii, HOL model projections had a true positive fraction of 93%.

The Effect of Climate on Species Distributions. The earth’s climate underwent a dramatic shift from late Pleistocene glacial to Holocene interglacial conditions, with most of the change (especially in sea level) taking place during the PHT from 15 to 9 ka14,30–32. The ca. 120 m rise in sea level during this period had a large effect on island area in much of the Caribbean13,34, which would have reduced the amount of terrestrial habitats, perhaps (on some islands) beyond the level needed to sustain viable populations of organisms31. Based on paleo-ENM projections, we observed changes in the relative range size of predicted habitat that indicate a range contraction from LGM to HOL, followed by a small range expansion from HOL to the present for all three species of bat (Table 2; Supplementary Fig. S2). Despite the trend of range change from past to present, the amount of overlap in the distributions across time was at least 53% in Ma. waterhousii, 86% for Mo. redmani, and 72% for P. parnellii (Table 2). We represent these areas of overlap as stability maps, which include the intersection of distribution models across the three time periods for each species of bat (Fig. 2).

We also estimated Relative Range Size (RRS) and Overlap Index (OI) corrected for the current island area for each species to examine the distributional changes that have occurred on land areas that have not been affected by sea level change. For Ma. waterhousii, the corrected RRS suggest a very small contraction in range size from LGM to HOL, and also from HOL to the present. The high degree of overlap between HOL and present predicted distributions (i.e., 89%) suggests that the available suitable habitat has remained nearly unchanged throughout the Holocene (Table 2; Fig. 2). RRS values corrected for island size in Mo. redmani and P. parnellii show a range contraction from LGM to HOL and an expansion from HOL to the present (Table 2; Supplementary Fig. S2). Nonetheless, the magnitude of these changes is small because the RRS estimates do not deviate much from zero, which would indicate no change in relative range over time.

Discussion

Bats often dominate modern Neotropical mammalian faunas in numbers of species and individuals35. Certainly they are the dominant order of native mammal in the Caribbean today. The integration of ENMs with large fossil datasets and last appearance radiocarbon dates provides a validated ecological explanation to examine community turnover in bats. Previous studies proposed that late Pleistocene climate change (and associated changes in habitat and island area) was the primary cause of extinction of Caribbean bats10,11. Here we provide evidence that extirpated populations of bats were able to persist on Abaco, the northernmost edge of the distribution in each species, for at least five to seven millennia after the last glacial-interglacial climate transition. These results contradict the climate change hypothesis and thus require alternative explanations for the extirpation and extinction of Caribbean bats.

We hypothesized that if late Quaternary losses of these bats were driven by climate change during the PHT, then radiocarbon dated fossils from extirpated populations of bats should be older than 9 ka. In spite of suggestions to the contrary10,11,19,20, the direct and associated AMS 14C dates we present here show that extirpations took place into the late Holocene (<3.8 ka, in some cases <1.8 ka). The late Holocene existence of these bat populations also suggests that late Pleistocene changes in the size and distribution of Bahamian caves and cave environments are unlikely causes for the extirpation of these populations. Furthermore, associated AMS 14C dates presented by Steadman et al.21 and Pregill et al.29 show that two additional extirpated species (Pteronotus parnellii, Mormoops blainvilli) and an extinct one (Phyllonycteris major) were lost from Antiqua no earlier than ca. 4300 to 1800 Cal BP. If climatic changes during the PHT were the primary driver of the losses of Caribbean bats, it is difficult to understand why these species survived for at least 5000 years before becoming extinct.

ENMs have been successfully integrated with other approaches to estimate the magnitude of the effect that past climate change has had on various species17,36–38. The geographically explicit aspect of validated ENM projections over time provides an ideal method to examine biogeographic hypotheses about changes in the distribution of an organism in the face of major climatic changes. Our second goal was to develop rigorous ENMs that accurately represented the changes in distributions over time for three species of Caribbean bats. Our results of fossil-validated ENM projections to HOL and LGM climate for Ma. waterhousii, Mo. redmani, and P. parnellii were highly accurate with true positive fractions of 94%, 86%, and 83%, respectively. The paleo-ENMs confirmed the stability of suitable habitat in the past for all three species. Projections into present-day climate were also highly accurate with true positive fractions falling over 94%. Therefore, we believe that our ENM projections adequately represent the potential distribution based on suitable habitat for each bat species across the Caribbean. Areas of distributional overlap across all three time periods represent areas of stability, which have been considered as refugia for some continental organisms37. For insular bats, areas of stability suggest that a considerable portion of the land area that has remained above water over time has been part of the suitable habitat available for each species to sustain viable populations.
Despite this, overlap in suitable habitat for all three species of bat was at least 73% between the HOL and the present, and at least 60% between LGM and HOL on land that lies above sea level today.

It is important to note that the island groups where bat extirpations have been documented and radiocarbon dated (see Results, Steadman et al., 23, and Pregill et al., 29) had over 80% reduction in island area from LGM to the present (Supplementary Table S1). Our ENM results show that, even when a Caribbean island underwent a considerable reduction of land area during the PHT (considering the estimated low sea level of -135 m for the LGM42), the minimum amount of suitable habitat to sustain viable populations of *Ma. waterhousii*, *Mo. redmani*, and *P. parnelli* probably was unchanged. Although we recognize that reductions in the population sizes of *Ma. waterhousii* (Puerto Rico) and of *Mo. redmani* and *P. parnelli* (Bahamas) probably took place during the PHT, we find it unlikely that reduced island area alone would have eliminated these or any of the other bat populations that survived into the late Holocene but now are gone.

Prehistoric (and historic) human impacts on island species and ecosystems feature considerable habitat alteration, especially when the prehistoric peoples were horticulturalists who altered natural fire regimes on large scales43–45 as in the Bahamas, where Amerindians first colonized from 1.5 to 1.0 ka44,46. Bat extirpations documented for the northern Lesser Antilles21 occurred following a period of increased anthropogenic activity (ca. 5300 Cal BP) that is well documented on the nearby island of Puerto Rico47. Therefore, it seems plausible that human-induced habitat changes contributed to the extirpation of bats of the northern Lesser Antilles. While the precise mechanism leading to extirpation (e.g., direct predation, habitat loss, invasive plants and animals, etc.) is undetermined, the late Holocene demise of bat populations in the Bahamas coincides with similar land mammal, reptile, and bird losses on other Caribbean islands23,47,48.

The radiocarbon dates we present here, combined with climate ENMs, demonstrate that at least five species of bats withstood climate change and reduced land area during the Pleistocene-Holocene Transition on Abaco, only to become extirpated during the late Holocene, a time when climate conditions were largely similar to those of today48. Because Abaco (and other Bahamian islands) lost a much greater proportion of its land during the PHT sea-level rise than Greater Antillean islands23, we would expect that new research will discover that most extirpated populations or extinct species of bats in the Caribbean also survived well into the Holocene. We challenge other researchers (and ourselves) to develop a more comprehensive extinction chronology for Caribbean bats through additional AMS 14C dating. Only in this way will we be able to make archipelago-wide generalizations about when the numerous populations and species were lost.

**Methods**

**Modern and Fossil Localities.** Locality records with WGS84 latitude/longitude coordinates were obtained for modern specimens of *Macrotus waterhousii* (Waterhouse’s Leaf-nosed Bat), *Monophyllus redmani* (Greater Antillean Long-tongued Bat), and *Pteronotus parnelli* (Parnell’s Mustached Bat) with GPS in the field and from the MaNIS database (http://manisnet.org). Each of these species is ecologically distinct. For example, among cave dwelling species, *Ma. waterhousii* prefers the use of cool caves, whereas *Mo. redmani* and *P. parnelli* prefer hot caves. Similarly, these bats fit in different dietary guilds, with *Ma. waterhousii* as a gleaning insectivore, *Mo. redmani* primarily a nectarivore, and *P. parnelli* a hawking insectivore. Bat fossils (cataloged in the Vertebrate Paleontology Collection, Florida Museum of Natural History, University of Florida; UF) were identified to species by comparing them to modern bones (especially humeri, mandibles, and skulls) from the UF Mammalogy Collection. Fossil occurrences of all three species were georeferenced following the guidelines of MaNIS. Additional (non-UF) fossil localities for each species were obtained from the literature, and then georeferenced. Every locality representing an extirpated population was included in the fossil datasets to evaluate the performance of the paleo-ENMs. We plotted distributions for the three species of bat separately in ArcGIS v10.2 (ESRI, Redlands, CA) to visually assess the quality of spatial data and to correctly georeferenced localities to the nearest 0.001° following their locality description.
Table 3 | Model performance statistics based on AICc for the best and default Ecological Niche Models of three species of West Indian bats. Performance statistics were estimated using ENMtools (Warren et al. 2010). Selected model features include: quadratic (Q), product (P), hinge (H), and threshold (T). Model testing statistics are negative log likelihood (-lnL) and sample size corrected Akaike Information Criterion (AICc).

| Species            | Model Features | Parameters | -lnL  | AICc  |
|--------------------|----------------|------------|-------|-------|
| Macrotus waterhousii | QPH features   | 22         | 3974.89 | 7977.25   |
|                    | Auto features  | 32         | 4047.91  | 8169.82  |
| Monophyllus redmani | OPT features   | 33         | 3634.59  | 7335.37  |
|                    | Auto features  | 31         | 3680.78  | 7430.91  |
| Pteronotus parnellii | QPH features  | 18         | 3614.24  | 7266.91  |
|                    | Auto features  | 20         | 3614.42  | 7271.83  |

Last Appearance Radiocarbon Dates (LADs). Last Appearance Radiocarbon Dates (LADs) represent the latest dated records of a population or species, and as such can provide useful conservative estimates of the time of extirpation or extinction63. From two fossil localities (Ralph’s Cave, Abaco, Bahamas, 26.24961° N, -77.19021° W; and Trouen Jeremie, Département-du-Sud, Haiti, 18.20999° N, 74.01011° W), we established a chronology based on Accelerator Mass Spectrometer (AMS) radiocarbon (14C) dates from purified collagen extracted from different individual bones of the extinct sloth Neocuxus comes29. These dates ranged from 9490–9130 Cal BP to 7790–7610 Cal BP (early Holocene). We selected one humerus from Ma. waterhousii to the date the occurrence of this bat at Trouen Jeremie.

Ecological Niche Modeling (ENM). We used presence-only data analyzed in a maximum entropy approach to develop present and paleo-ecological niche models (paleo-ENMs) for Ma. waterhousii, Ma. redmani, and P. parnellii. We used climate data from 19 WorldClim variables for present climate, the Holocene (HOL: ca. 6 ka), and the Last Glacial Maximum51 (LGM; ca. 21 ka). The use of two time periods in the past allowed us to reconstruct distributions at a similar but possibly wetter and slightly warmer period (HOL) and at a cooler and drier period (LGM) compared to present conditions52,53. Both paleo-climate conditions were obtained from statistically downscaled estimates of past conditions from the ESCAM3 and MIROC models available from WorldClim (see www.worldclim.org for downsampling details). We used ESCAM3 and MIROC models because they showed good performance in previous paleo-climate reconstructions54. The three climate data sets were clipped to a regional extent of the Caribbean using ArcGIS v10.2. To correct for co-linearity among the 19 climate variables, we performed a Pearson Correlation in R v2.15.0 (correlation coefficient > 0.8; 24) and removed correlated climatic variables for each time period. The resulting datasets feature seven uncorrelated climatic variables that represent the climate of the Caribbean (Temperature — bio1, annual mean; bio2, mean diurnal range; bio3, isothermality; bio7, annual range; and Precipitation — bio12, annual; bio14, driest month; bio18, warmest quarter). We extracted present climate information for our species data and for a random background climate dataset of 1250 points to sample the accessible area of background climate for each species of bat. Random background points that were within 1 km of a species locality record were removed. This restriction of background is recommended to reduce error in model parameterization, validation, and comparison55.

We generated ENMs using maxent v3.3.3 because its predictive performance is consistently high compared to that of other methods56–60. We randomly partitioned species localities into 75% training and 25% testing datasets with 100 bootstrap replicates of 400 data points. Model performance is evaluated by model parameterization61–64. We developed multiple ENMs combining different features in maxent and evaluated the complexity of each model using AICc implemented in ENMTools v1.4.3.165. The parameters used in the final ENMs included a regularization multiplier of 1, and a combination of quadratic, product, threshold, and hinge features (Table 3). We projected all ENMs to present, HOL, and LGM background climate for the West Indies to obtain our final models.

Evaluation of model performance for each species of bat was assessed using the area under the receiving operating characteristic curve (AUC). AUC is a threshold independent measure that varies from 0 to 1, where a score of 1 represents perfect discrimination and a score of 0.5 represents a model no better than random66. We considered an AUC score ≥ 0.7 to represent good model predictions67–69. Some authors question the accuracy of AUC for evaluating model performance70 (e.g., when using presence only data). Therefore, we also quantified model overfitting directly by comparing threshold-dependent omission rates with theoretical omission levels71. We applied a lowest presence threshold rule of 95% (LPT95%) to the average models from maxent logistic output72. Under this rule, pixels with values equal or higher than the selected threshold were considered as suitable conditions for a species to sustain viable populations, which allowed us to generate binary (i.e., presence/absence) predictions. We chose LPT95% because it is a relaxed threshold for which predicted model training datasets resulted in at least 95% of locality points contained within suitable habitat73,74 (i.e., a 5% omission rate). The use of a relaxed threshold when generating past ENM projections allowed enough sensitivity to determine long-term suitable areas where species could maintain viable populations over time. The resulting presence-absence distribution maps for the three time periods of each species were summed and corrected for the present island area to develop stability maps (Fig. 2). Predicted areas of stability represent areas where each species is predicted to occur across the present, HOL, and LGM time periods. All range estimates were performed in ArcGIS v10.2.

Validation and Evaluation of Models Across Time. Paleo-ENMs were developed as model projections onto HOL and LGM climate using modern locality records. Therefore, paleo-ENMs cannot be evaluated based on AUC alone. We were interested in assessing how accurately our paleo-ENMs could predict the fossil locality data because inferences on how climate change affects an organism depend on how well the model can predict potential distributions in the past. For this assessment, we estimated the true positive fraction (i.e., correctly predicted fossil localities) to determine the fit of paleo-ENM projections over time for each species of bat. We used the LPT95%–transformed models to evaluate the effects of climate change on each species of bat from the present and paleo-ENMs by estimating two indices: relative range size (present ≥ past RRS = c/m – 1, or present ≤ past RRS = −1 × [m/c – 1]) and overlap index (OI = o/m; where m = present climate conditions, c = past climate conditions, and o = amount of overlap between present and paleo-ENMs)75. We used RRS to compare the size and magnitude of change of predicted present vs. paleo-ENMs. A negative RRS value implies that the paleo-distribution was smaller than the present distribution, a positive RRS value implies that the paleo-distribution was larger than at present, and a value of zero indicates no change76. OI was used to determine the magnitude of overlap between present and paleo-distributions. Overlap in predicted distributions relates to areas that remained climatically suitable and could have provided suitable habitat where bat populations persisted over time. Because eustatic sea level changes caused major changes in land distribution77,78, we estimated RRS and OI values corrected for the present-day area of the islands to isolate the effect of changes in island area on species distributions. These corrected estimates of the persistence of suitable climate and habitat are associated with areas that may have been occupied by each species throughout the late Quaternary.
13. Cuffey, K. & Marshall, S. Substantial contribution to sea-level rise during the last interglacial from the Greenland ice sheet. Nature 404, 591–594 (2000).
14. Denton, G. H. et al. The last glacial termination. Science 328, 1652–1658 (2010).
15. Franklin, J. Mapping species distributions: spatial inference and predictions. 317 (Cambridge University Press, 2009).
16. Waltari, E. & Hickerson, M. J. Late Pleistocene species distribution modelling of North Atlantic intertidal invertebrates. J. Biogeogr. 40, 249–260 (2012).
17. Rahmeh, a. C., Hickerson, M. J., Haddad, C. F. B., Rodrigues, M. T. & Moritz, C. Stability predicts genetic diversity in the Brazilian Atlantic forest hotspot. Science 323, 785–789 (2009).
18. Soto-Centeno, J. A., Barrow, L. N., Allen, J. M. & Reed, D. L. Reevaluation of a late-Holocene bird community from a Quaternary cave deposit in Cuba. Caribb. J. Sci. 39, 371–377 (2003).
19. Jiménez-Vázquez, O., Condón, M. M. & García-Cancio, E. Vertebrados post-glaciares en un residuario fosil de Tyto alba scopoli (Aves: Tytonidae) en el occidente de Cuba. Rev. Mex. Mastozoól. 9, 85–112 (2005).
20. Steadman, D. W., Pregill, G. K. & Olson, S. L. Fossil vertebrates from Antigua, Lesser Antilles: evidence for late Holocene human-caused extinctions in the West Indies. Proc. Natl. Acad. Sci. USA 81, 4448–4451 (1984).
21. Steadman, D. W. & Takano, O. M. A late-Holocene bird community from Hispaniola: refining the chronology of vertebrate extinction in the West Indies. The Holocene 23, 936–944 (2013).
22. Steadman, D. W. & Franklin, J. Changes in a West Indian bird community since the late Pleistocene. J. Biogeogr. DOI:10.1111/jbi.12148 (2014).
23. Choate, J. R. & Birney, E. C. Sub-recent Insectivora and Chiroptera from Puerto Rico, with the description of a new bat of the genus Stenodermus. J. Mammal. 49, 400–412 (1968).
24. Anderson, S. Macrotrus waterhousei. Mamm. Species 1, 1–4 (1969).
25. Homan, J. A. & Jones, K. J. Monophyly redisnan. Mamm. Species 57, 1–3 (1975).
26. Herd, R. M. Pteronotus parnellii. Mamm. Species 209, 1–5 (1983).
27. Clare, E. L. et al. Diversification and reproductive isolation: cryptic species in the only New World high-duty cycle bat, Pteronotus parnellii. BMC Evol. Biol. 13, 26 (2013).
28. Pregill, G. K., Steadman, D. W., Olson, S. L. & Grady, F. V. Late Holocene fossil vertebrates from Burma Quarry, Antigua, Lesser Antilles. Smithsonian. Contrib. Zool. 463, 1–27 (1988).
29. Clark, P. U. & Mix, A. C. Global change: ice sheets by volume. Nature 394, 610–613 (1998).
30. Clark, P. U. et al. The Last Glacial Maximum. Science 235, 710–714 (2009).
31. Hadly, E. A., Spaeth, P. A. & Li, C. Niche conservatism above the species level. Proc. Natl. Acad. Sci. USA 104, 18977–19002 (2007).
32. Berman, M. J. & Gnievcki, P. L. The colonization of the Bahamas archipelago: a reappraisal. World Archaeol. 26, 421–441 (1995).
Additional information

Supplementary information accompanies this paper at http://www.nature.com/scientificreports

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Soto-Centeno, J.A. & Steadman, D.W. Fossils reject climate change as the cause of extinction of Caribbean bats. Sci. Rep. 5, 7971; DOI:10.1038/srep07971 (2015).

This work is licensed under a Creative Commons Attribution-NonCommercial-NoDerivs 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder in order to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc-nd/4.0/