



Predictors of Mammalian Diversity in the New York Metropolitan Area

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Urbanization can have profound consequences for mammalian biodiversity and is thought to contribute to patterns of species richness and community composition. Large cities can be particularly challenging environments for mammals because these habitats are often impacted by anthropogenic perturbations, including high human population density, fragmented habitats, and extensive human development. In this study, we investigated mammalian species richness, Shannon–Wiener diversity, and evenness in the most densely populated region in the United States: the New York metropolitan area. Specifically, we deployed camera traps from 2015 to 2019 to investigate six drivers of mammalian diversity across 31 greenspaces: (1) human population density, (2) patch size, (3) habitat type, (4) surrounding land cover, (5) geographical barriers to dispersal, and (6) habitat heterogeneity. We found that mammal community composition is largely influenced by a multitude of anthropogenic factors. Specifically, mammal species richness was higher in greenspaces with larger patch sizes and lower in greenspaces surrounded by more development. Moreover, Shannon–Wiener diversity and evenness were higher in urban natural landscapes than human-altered landscapes. In a subset of data that only included carnivores, we found that carnivore Shannon–Wiener diversity was higher in urban natural habitats and in sites with lower human population densities. Finally, we found that geographical barriers to dispersal contributed to both patterns of mammalian diversity and patterns of carnivore diversity: mammal taxa richness, Shannon–Wiener diversity, and evenness were all significantly higher on the continent (Bronx/Westchester) than on Long Island. These results suggest that preserving urban greenspaces is important for maintaining both mammalian and carnivore biodiversity and that management of mammals in cities should concentrate on maintaining large, connected, natural greenspaces.

Keywords: urban ecology, mammals, habitat heterogeneity, patch size, biodiversity, human population density, New York metropolitan area, land cover

INTRODUCTION

As urbanization continues to spread throughout the world, its consequences, including increased human population growth, habitat fragmentation, and deforestation, are becoming increasingly important to examine and understand, especially in terms of biodiversity (McKinney, 2002; McDonald et al., 2008). Urbanization has had profound impacts on ecosystem function and is

thought to be a major driver of biodiversity loss (Czech et al., 2000; Ray, 2000; Sala et al., 2000; Dobson et al., 2006; McCleery, 2010; Seto et al., 2012; Lewis et al., 2015). Mammals are sensitive to urbanization because their habitats within cities are often fragmented or replaced by human developments (Fernández and Simonetti, 2013; Łopucki and Kitowski, 2017). Carnivores are especially impacted by increased urbanization due to their large home ranges, human intolerance, limited diets, and low densities (Ordeñana et al., 2010; Stark et al., 2020). However, while urbanization is often associated with a lack of wildlife, recent studies have shown that species diversity, richness, and abundance are actually high in some urban greenspaces (Gehrt et al., 2009; Gallo et al., 2017; Matthies et al., 2017; Parsons et al., 2018). Indeed, some mammals, including coyotes (*Canis latrans*), raccoons (*Procyon lotor*), opossums (*Didelphis virginiana*), and foxes (e.g., *Vulpes vulpes*), are known as “urban adapters”, and have benefited from being in close proximity to developed areas reaping the benefits of increased food sources, decreased competition, and increased protection from large predators (McKinney, 2002; Adams and Lindsey, 2010; Gehrt et al., 2013; Lombardi et al., 2017). Understanding what factors influence patterns of mammalian community composition are important considerations in maintaining and protecting biodiversity in urban habitats. In this study, we used camera traps to investigate the impact of six drivers of mammalian diversity across greenspaces in the New York metropolitan area: (1) human population density (Magura et al., 2021), (2) patch size (Arrhenius, 1921), (3) habitat type (McDonnell and Pickett, 1990), (4) surrounding land cover (Cavia et al., 2009), (5) geographical barriers to dispersal (MacArthur and Wilson, 1967), and (6) habitat heterogeneity (Cornelis and Hermy, 2004).

At first glance it would seem human population density would negatively affect wildlife diversity, and indeed this is the case for many species and circumstances (Olifiers et al., 2005; Schipper et al., 2008; Stark et al., 2020). However, the impact of human population density on biodiversity is species-specific and scale-dependent (e.g., Prange et al., 2004; McCleery, 2010; Gehrt et al., 2013; Lombardi et al., 2017). On a global scale, as human population increases, more natural habitats are destroyed for residential, commercial, and agricultural use; mammals are forced from their habitats, and many do not survive (Lopes and Ferrari, 2000; Kasso and Bekele, 2014). Accordingly, several studies have reported a negative correlation between human population density and species richness (e.g., Gehrt and McGraw, 2007; Parsons et al., 2018; Stark et al., 2020). Another consequence of increased human population density is the occurrence of more hunting and culling in habitats once dominated by animals (Cullen et al., 2000). This is especially the case for large mammals that humans perceive as threats, such as coyotes (Nagy et al., 2017). Finally, although not as well-studied, vehicles have a major impact on mammalian diversity, in some years even greater than that of hunting (Forman and Alexander, 1998). Due to the abundance of roads located near forested areas, vehicular killings have contributed to overall mammal diversity loss in recent decades (Bashore et al., 1985; Forman and Alexander, 1998; Seiler, 2001; Shilling et al., 2021). While an inverse relationship between biodiversity and human population

density has been documented in many circumstances, on a local scale, species richness and abundance of certain taxa, such as free-ranging cats (*Felis catus*), raccoons (*P. lotor*), and certain rodents, are actually higher in areas of high human population density (Luck, 2007; Gehrt et al., 2013; Lombardi et al., 2017). This may be largely attributed to the high availability of anthropogenic food sources in areas with high human population densities (Contesse et al., 2004; Prange et al., 2004; McCleery, 2010; Magura et al., 2021). Furthermore, close proximity to humans in urban landscapes has been shown to reduce predation pressures for small and medium size mammal species (Ditchkoff et al., 2006; McCleery, 2010). This is predominantly due to the displacement of large predators as a result of high human presence and activity in centrally populated areas (Muhly et al., 2011). When taking all these factors into consideration, densely populated cities typically have a net negative effect on mammalian species richness and especially pose a threat for larger mammals (Gehrt and McGraw, 2007; Parsons et al., 2018; Stark et al., 2020).

Patch size, defined as the arrangement and size of usable land area, have a major influence on patterns of mammalian diversity (Lombardi et al., 2017). In highly developed areas, increased urbanization often results in habitat fragmentation or isolation effects, which reduces the amount of available living space (Fahrig, 2003; Cavia et al., 2009; Nielsen et al., 2014) and separates larger populations into smaller subpopulations that, individually, are less viable. Some species of mammals are less likely to tolerate isolated patches of habitats that result from increased urbanization because they require more space to acquire vital resources (Etter et al., 2002; Prange et al., 2004; Moll et al., 2020). Accordingly, several studies have reported a positive association between patch size and species diversity (Saunders et al., 1991; Iida and Nakashizuka, 1995; Fahrig, 2003). Nielsen et al. (2014) found that a minimum of 10 hectares of patch size is necessary to ensure high species richness in urban greenspaces. However, some mammals, such as large carnivores, require an even greater amount of greenspace sometimes extending five square kilometers and beyond (Stark et al., 2020). Thus, mammalian diversity is largely defined by species-area effects whereby species richness is a function of patch size (MacArthur and Wilson, 1967; Hodgkison et al., 2007; Matthies et al., 2017).

The type of habitat might also affect patterns of mammalian diversity (Gallo et al., 2017). In urban areas, some greenspaces are dominated by *human-altered habitats* while other greenspaces are mainly comprised of *urban natural habitats*. Human-altered habitats include parks, which are characterized by manicured vegetation and high human activity (Nielsen et al., 2014); golf courses, which are comprised of large, semi-artificial greenspaces where human activity varies temporally (Boone et al., 2008); and cemeteries (Gallo et al., 2017). Some species can flourish in human-altered habitats (urban adapters), while other species cannot (urban avoiders) (McKinney, 2002, 2006). In contrast to human-altered habitats, urban natural habitats are typically composed of undeveloped woodlands with secondary growth forest (Vernon et al., 2014) situated within an urban or human-altered matrix. Natural areas within urban environments remain crucial in supporting mammal populations (Baker and Harris, 2007). Several studies have found that the presence of natural

habitats within urban greenspaces allows for greater movement of multiple species (Radeloff et al., 2005; Gehrt et al., 2011; Bateman and Fleming, 2012; Scott et al., 2014).

The amount of developed land cover surrounding a habitat is another factor thought to be associated with mammalian diversity (Bierwagen, 2006; Cavia et al., 2009). Urban habitats often have decreased structural and functional connectivity between the highly developed cityscape and the less developed natural habitats within (LaPoint et al., 2015; Beninde et al., 2016). This reduced connectivity resulting from urbanization often leads to decreased distribution of smaller species (Zollner, 2000). Highly developed areas also have a high percent of impervious surface cover (i.e., roads, buildings, landscaping), and often have lower biodiversity than less developed areas (McKinney, 2008; Gallo et al., 2017). High impervious surface cover surrounding natural habitats like roads, buildings, and parking lots restrict home ranges and can lead to increased death by vehicle collision (Trombulak and Frissell, 2000; Seiler, 2001; Gehrt et al., 2013). These barriers to dispersal reduce gene flow and population viability relative to a connected metapopulation and can lead to decreased species richness (Bateman and Fleming, 2012). Mammals situated in habitats surrounded by highly developed land cover also experience high rates of habitat and resource disturbance (Blair and Launer, 1997) and the loss of natural vegetation to development (McKinney, 2008; McCleery, 2010). Together, these changes to habitats that encompass greenspaces place limitations on the species within the greenspace itself. Therefore, in order to maintain mammalian diversity, it is important to have patches of natural habitat and vegetation as well as increased connectivity between greenspaces and highly developed surrounding areas (Beninde et al., 2016; Gallo et al., 2017; Fidino et al., 2020).

Another factor that contributes to patterns of mammalian diversity is habitat heterogeneity. The habitat heterogeneity hypothesis proposes that the species diversity of an area will increase as the number of habitats increases (MacArthur and MacArthur, 1961). Hence, it is predicted that a greater number of habitats can offer more resources to a wider range of animals (Garden et al., 2007; Hodgkison et al., 2007; Matthies et al., 2017). In more heterogeneous habitats, there tend to be more habitat edges, creating more resources for supporting a greater number of mammals (Racey and Euler, 1982; Cramer and Willig, 2005). Accordingly, several studies of mammals have reported a positive association between habitat heterogeneity and species richness (Jepsen and Topping, 2004; Arques et al., 2014; Presley et al., 2019). Thus, urban greenspaces that have a variety of habitats are thought to be vital to mammalian diversity (Hodgkison et al., 2007).

Lastly, geographical barriers can also influence patterns of mammalian diversity. Specifically, islands that are bounded by water create barriers to dispersal for many species (Weckel et al., 2015). Hice and Schmidly (2002) found that continental regions exhibit higher species diversity than island regions mainly because they do not require the crossing of harsh waterways or busy bridges for colonization. Some mammals, like coyotes (Harrison, 1992) and raccoons (Prange et al., 2004), can cross these barriers and move between the mainland and the island,

which has been observed in major cities like New York (Weckel et al., 2015; Henger et al., 2020) and San Francisco (Sacks et al., 2006). Nonetheless, the intersection of urbanization and geographical barriers to dispersal are thought to lead to reduced mammal richness and evenness among urban habitats (Ekernas and Mertes, 2006; Weckel et al., 2015).

Our goal was to determine predictors of mammalian diversity in various greenspaces across the New York metropolitan area. To do so, we deployed camera traps across 31 study sites (**Figure 1** and **Table 1**). We hypothesized that mammal community composition is influenced by a multitude of anthropogenic factors including human population density (Magura et al., 2021), patch size (Arrhenius, 1921), habitat type (McDonnell and Pickett, 1990), surrounding land cover (Cavia et al., 2009), geographical barriers to dispersal (MacArthur and Wilson, 1967), and habitat heterogeneity (Cornelis and Hermy, 2004). Specifically, we predicted that mammalian diversity would be higher in study sites where there were lower human population densities, larger patch sizes, urban natural habitats, less developed land cover surrounding the core habitat, and more heterogeneous habitats. Because bodies of water are important geographical barriers to dispersal, we also predicted that mammal diversity would be greater in the mainland (Bronx/Westchester) than in Long Island, Manhattan, and Randall's Island. Identifying and understanding what factors influence patterns of mammalian diversity is critical to maintaining both mammalian populations and overall biodiversity.

MATERIALS AND METHODS

Field Sites

This research focused on mammalian diversity across various greenspaces in the New York metropolitan Area. Our study sites included 31 greenspaces throughout four geographical regions: (1) Long Island; (2) mainland (Bronx and Westchester); (3) Manhattan; and (4) Randall's Island (**Supplementary Table 1**). Of the 31 sites, 16 were located in Long Island, nine in the mainland, five in Manhattan, and one in Randall's Island. Within each of these regions, there were various types of greenspaces. Of the 16 sites on Long Island, one was a baseball field, one was a golf course, one was a ranch/farm, two were cemeteries, and the remaining eleven sites were urban parks. Of the nine mainland sites, all contained natural vegetation with one site also containing multiple recreational and tourist facilities. In Manhattan, four of the five sites were parks consisting mainly of natural vegetation, while the fifth site was a more developed urban park containing a mixture of natural vegetation and human-altered habitats. The Randall's Island site was comprised of a combination of open areas, natural vegetation, and human made recreational facilities. Each of the greenspaces contained one or more cameras to monitor wildlife and predict patterns of mammalian diversity.

Camera Surveys

Camera traps were deployed across 31 sites in the New York metropolitan area from December 2015 to June 2019 following

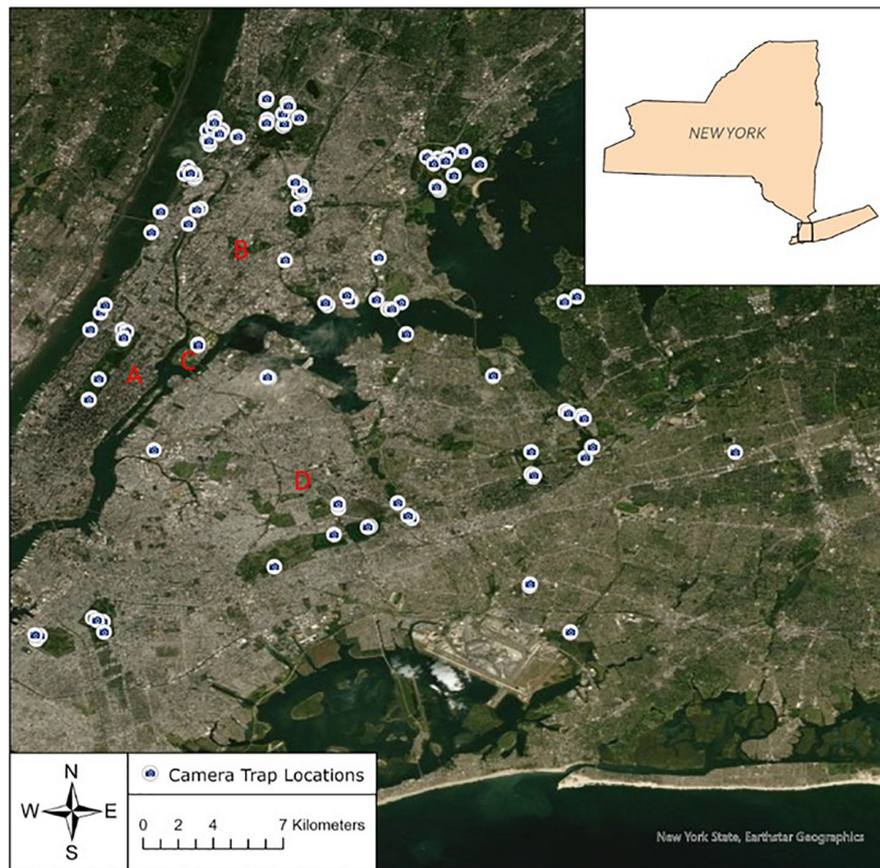


FIGURE 1 | Map of camera locations across 31 greenspaces in which mammals were surveyed in the New York metropolitan area, 2015–2019. A = Manhattan; B = Mainland (Bronx/Westchester); C = Randall's Island; D = Long Island.

the guidelines described in Nagy et al. (2016). Except for small rodents, which we classified at the family level (family Muridae), we identified all mammals to the species level using photographs captured by each camera trap. Throughout the five-year course of this study, three types of Reconyx cameras were used: RC55, PC800, and HC500 motion- and heat-activated camera traps (Reconyx, Inc., Holmen, WI, United States). Each of these cameras had 0.2 s trigger speed, 1080p resolution, and comparable infrared flash ranges (RC55 and HC500: 15 m, PC800: 21 m). There were two models of the RC55 used, with the older one having a 5.0×7.6 cm red/infrared flash primarily for night pictures, and the newer one with a single LED bulb for a “semi-covert” infrared flash. Using ArcGIS 9 and ArcGIS10 (ESRI, Redlands, CA, United States), random locations were selected for camera deployment in the different greenspaces with at least 500 m distance between each camera. Exact survey dates varied year-to-year but generally cameras were deployed either year-round or more commonly twice a year, first in early January through mid-April and again June through August. These twice-annual deployments allowed for the detection of seasonally-active, hibernating mammals (e.g., chipmunk: *Tamias striatus*; groundhogs: *Marmota monax*) and the detection of new coyote breeding sites in the winter—i.e., when occupancy across the

landscape is highest as dispersers establish new territories and pairs form (Nagy et al., 2016). At each location, cameras were deployed in a substantial patch of forest or undeveloped area. Because we had a limited number of camera traps, in cases in which a study area consistently surveyed the same mammals, we shifted camera traps to new locations within a greenspace to increase coverage. Many of the cameras required relocation due to theft, legal ramifications, or necessity for a better-suited placement. Each of these relocations were at least within ~ 50 m of the originally planned locations. The actual number of cameras per greenspace varied, however, the density of cameras in most cases was at least one camera per 0.5 km^2 in each of the greenspaces (Table 1). Across the five years of the study, the cameras were typically placed in the same general areas unless there were natural disasters or theft that previously influenced the camera placement.

Predictors of Mammalian Diversity

To gain better insight into what predicts mammalian diversity across an urban landscape, we modeled three response variables: (1) richness, defined as the number of different mammal taxa present in a given greenspace or region (Dorji et al., 2014); (2) evenness, defined as the abundance equality of mammal

TABLE 1 | Dates, location, and characteristics of study sites surveyed 2015–2019.

| Study site | Survey dates | Latitude, longitude ^A | Human population density people/km ^{2B} | Patch size (km ²) | Habitat type | Percent developed land cover (500 m) | Percent developed land cover (1,000 m) | Habitat heterogeneity (500 m) | Habitat heterogeneity (1,000 m) | Region | Number of cameras/km ² | Number of cameras deployed |
|------------------------|------------------|----------------------------------|--|-------------------------------|-----------------------|--------------------------------------|--|-------------------------------|---------------------------------|------------------|-----------------------------------|----------------------------|
| Alley Pond | 2/1/16–5/7/19 | 40.75742, –73.74266 | 28,552.03 | 2.82 | Urban Natural Habitat | 71.58 | 82.86 | 7 | 10 | Long Island | 2.48 | 7 |
| Bronx Park | 1/30/16–6/13/19 | 40.86273, –73.87398 | 139,345.89 | 2.95 | Urban Natural Habitat | 44.33 | 85.71 | 10 | 13 | Mainland | 3.39 | 10 |
| Central Park | 2/17/16–7/24/17 | 40.76635, –73.97463 | 271,920.26 | 3.43 | Human-Altered Habitat | 30.89 | 30.56 | 8 | 12 | Manhattan | 2.62 | 9 |
| Clearview Golf Course | 2/1/16–3/29/19 | 40.77767, –73.78519 | 34,319.93 | 0.48 | Human-Altered Habitat | 53.32 | 0 | 4 | 7 | Long Island | 6.25 | 3 |
| Cunningham | 2/3/16–5/7/19 | 40.74176, –73.76742 | 37,726.40 | 2.25 | Urban Natural Habitat | 41.83 | 33.33 | 7 | 8 | Long Island | 2.67 | 6 |
| Elmjack | 5/27/16–5/10/19 | 40.77693, –73.89096 | 65,241.80 | 0.058 | Human-Altered Habitat | 69.89 | 51.43 | 7 | 7 | Long Island | 51.72 | 3 |
| Ingrams Field | | | | | | | | | | | | |
| Ferry Point | 2/1/16–5/16/19 | 40.81174, –73.82832 | 31,890.52 | 1.32 | Human-Altered Habitat | 44.46 | 50 | 13 | 13 | Mainland | 3.03 | 4 |
| Forest Park | 6/13/16–5/10/19 | 40.70644, –73.84295 | 68,951.44 | 1.30 | Urban Natural Habitat | 16.38 | 0 | 6 | 6 | Long Island | 3.08 | 4 |
| Fort Washington | 2/18/16–9/1/16 | 40.84486, –73.94538 | 160,249.04 | 0.51 | Urban Natural Habitat | 48.8 | 28.13 | 6 | 10 | Manhattan | 5.88 | 3 |
| Francis Lewis | 2/3/16–2/8/19 | 40.79685, –73.82587 | 36,189.90 | 0.039 | Human-Altered Habitat | 59.56 | 43.75 | 7 | 7 | Long Island | 25.64 | 1 |
| Green-wood Cemetery | 1/26/17–5/13/19 | 40.65319, –73.99909 | 75,492.97 | 1.90 | Human-Altered Habitat | 62.04 | 54.29 | 3 | 4 | Long Island | 1.58 | 3 |
| Highbridge Park | 7/9/17–7/28/17 | 40.84889, –73.92803 | 184,104.12 | 0.262 | Urban Natural Habitat | 66.71 | 19.44 | 9 | 12 | Manhattan | 7.63 | 2 |
| Hutchinson | 3/7/16–5/8/19 | 40.833, –73.83879 | 31,890.52 | 0.031 | Urban Natural Habitat | 85.01 | 83.33 | 4 | 5 | Mainland | 32.26 | 1 |
| Idlewild Park | 2/10/16–3/23/19 | 40.6564, –73.74908 | 32,512.12 | 1.65 | Urban Natural Habitat | 22.67 | 0 | 9 | 11 | Long Island | 1.21 | 2 |
| Inwood Hill Park | 12/26/15–5/10/19 | 40.87247, –73.92971 | 89,468.55 | 0.789 | Urban Natural Habitat | 19.11 | 0 | 10 | 11 | Manhattan | 10.14 | 8 |
| Kings Point Park | 6/29/16–9/1/16 | 40.81459, –73.74606 | 5,881.86 | 0.70 | Urban Natural Habitat | 9.3 | 0 | 7 | 10 | Long Island | 2.86 | 2 |
| Maple Grove Cemetery | 3/7/16–5/10/19 | 40.71028, –73.82334 | 88,168.38 | 0.272 | Human-Altered Habitat | 88.89 | 80 | 4 | 4 | Long Island | 7.35 | 2 |
| Pelham Bay Park | 2/2/16–5/11/19 | 40.86636, –73.81177 | 31,890.52 | 2.23 | Human-Altered Habitat | 7.91 | 2.86 | 7 | 13 | Mainland | 6.73 | 15 |
| Prospect Park | 7/7/17–5/13/19 | 40.65661, –73.96745 | 139,567.73 | 2.46 | Urban Natural Habitat | 8.29 | 5.41 | 12 | 14 | Long Island | 1.63 | 4 |
| Pugsley Creek | 2/18/16–5/7/19 | 40.81244, –73.85221 | 69,530.82 | 2.40 | Urban Natural Habitat | 72.66 | 20 | 9 | 12 | Mainland | 1.67 | 4 |
| Queensline | 2/10/16–5/10/19 | 40.71709, –73.85794 | 95,769.99 | 0.198 | Urban Natural Habitat | 94.85 | 89.19 | 4 | 4 | Long Island | 15.15 | 3 |
| Railroad | 2/10/16–5/10/19 | 40.67924, –73.76794 | 47,282.82 | 0.152 | Urban Natural Habitat | 90.16 | 16.67 | 4 | 4 | Long Island | 13.16 | 2 |
| Randall's Island | 2/3/16–5/4/19 | 40.79188, –73.9234 | 63,011.82 | 0.092 | Human-Altered Habitat | 86.74 | 87.88 | 9 | 11 | Randall's Island | 21.74 | 2 |
| Ridgewood Highland | 1/26/18–5/16/18 | 40.68739, –73.8877 | 70,792.15 | 1.71 | Human-Altered Habitat | 40.96 | 8.33 | 11 | 11 | Long Island | 0.58 | 1 |
| Riverdale | 1/31/16–5/12/19 | 40.89881, –73.91566 | 72,364.27 | 0.51 | Urban Natural Habitat | 11.71 | 8.82 | 12 | 14 | Mainland | 21.57 | 11 |
| Riverside Park | 2/17/16–4/16/19 | 40.81059, –73.96693 | 208,386.12 | 0.35 | Human-Altered Habitat | 47.36 | 22.86 | 8 | 8 | Manhattan | 11.43 | 4 |
| Smiling Hogshead Ranch | 1/27/17–4/29/18 | 40.74252, –73.94411 | 25,236.84 | 0.008 | Human-Altered Habitat | 99.66 | 100 | 5 | 5 | Long Island | 125 | 1 |
| Soundview | 7/3/16–5/7/19 | 40.81164, –73.86387 | 69,530.82 | 0.719 | Urban Natural Habitat | 57.88 | 0 | 9 | 9 | Mainland | 4.17 | 3 |
| Starlight Park | 2/18/16–5/12/16 | 40.83183, –73.88263 | 131,408.23 | 0.011 | Urban Natural Habitat | 97.59 | 75.76 | 4 | 4 | Mainland | 90.91 | 1 |
| Van Cortlandt | 1/27/16–5/16/19 | 40.90697, –73.89154 | 51,011.11 | 4.65 | Urban Natural Habitat | 12.39 | 0 | 7 | 7 | Mainland | 3.44 | 16 |
| Willow Lake | 2/10/16–1/9/17 | 40.71763, –73.82987 | 67,286.60 | 0.665 | Urban Natural Habitat | 53.54 | 0 | 9 | 9 | Long Island | 1.5 | 1 |

^AFor sites containing multiple camera sites, the latitude and longitude was recorded based on the location at which there were the most photo captures.

^BThe human population density was calculated by averaging the population density of all of the zip codes surrounding each site.

taxa in a specific greenspace or region (Dorji et al., 2014); and (3) Shannon–Wiener diversity, defined as the proportional abundance of each mammal taxon from the total sample of individuals within the given surveyed community (Ekernas and Mertes, 2006). We conducted two analyses, one that included all mammal taxa identified in our camera trap surveys, and another that excluded humans and domestic dogs. In cases in which both analyses yielded the same best supported model, we report the results including all mammal taxa. In cases in which the best supported model differed between analyses, we report two sets of results, one including all mammal taxa and another excluding humans and domestic dogs. We included free-ranging cats in both analyses because these can include feral cats as well as free-roaming cats possibly subsidized by human foods (Elizondo and Loss, 2016).

For our analysis of interest—predictors of mammalian diversity in the greenspaces of the New York metropolitan area—we modeled six predictor variables (Table 2): (1) human population density; (2) patch area; (3) habitat heterogeneity; (4) type of habitat; (5) percent developed land cover; and (6) region. We also included cameras per square kilometer in our models to control for variation in camera deployment. The number of cameras deployed at each site ranged from one to sixteen with a mean of 4.45 cameras per site ($SD \pm 4.01$). Sampling effort ranged from 50 camera trap days to 2,279 camera trap days ($SD \pm 478$).

- Human population density.** To measure human population density, we used United States Zip Code data¹ to ascertain the number of individuals per square mile in the zip code in which the study site was located and then we converted these data to the number of individuals per square kilometer. In cases in which study sites were situated in more than one zip code, we calculated the mean human population density of the zip codes surrounding the study site.
- Patch size.** We calculated patch size using the “measure distance” tool on Google Maps.²
- Habitat type.** We classified study sites into two main categories based on habitat type. If a greenspace was comprised of greater than 50% secondary growth forest, we classified this habitat as *urban natural* (Gallo et al., 2017). On the other hand, if a greenspace was comprised of greater than 50% manicured lawns, athletic fields, playgrounds, and golf courses, we classified this habitat as *human-altered* (Gallo et al., 2017).
- Percent developed land cover.** ArcGIS Pro 2.6 and the National Land Cover Database³ were used to calculate percent developed land cover at each of the 31 greenspaces. We constructed buffers of 500 m to assess whether developed land cover in the immediate habitat impacted mammalian diversity (e.g., Gallo et al., 2017) and buffers of 1,000 m to assess whether land use surrounding a greenspace impacted mammalian diversity (e.g., Gomes et al., 2011). In cases in which the best supported model yielded the same results regardless of spatial scale, we report and discuss the model results based on land cover within 500 m (Gallo et al., 2017). Following Callaghan et al. (2019), Stark et al. (2020), and Goldstein et al. (2022), we combined the percent land cover of three different NLCD classes (“developed, low intensity”; “developed, medium intensity”; and “developed, high intensity”) into a superclass called “developed land cover.” Developed land cover included areas with a mixture of both constructed materials and vegetation, and where impervious surface cover accounted for at least 20 percent of the land cover.
- Habitat heterogeneity.** To calculate habitat heterogeneity, we counted the number of different habitat types based on 15 categories provided by the National Land Cover Database (Supplementary Table 2). We calculated habitat heterogeneity at two spatial scales: buffers of 500 and 1,000 m surrounding each of the 31 greenspaces. As with percent developed land cover, in cases in

¹<https://www.unitedstateszipcodes.org/>

²<https://www.google.com/maps>

³<https://www.mrlc.gov/data/legends/national-land-cover-database-2019-nlcd2019-legend>

TABLE 2 | Response and predictor variables for mammalian diversity among 31 greenspaces in the New York metropolitan area.

Response variables

| | |
|--------------------------|---|
| Richness | the number of different taxa present per study site |
| Shannon–Wiener diversity | the proportional abundance of each taxon from the total sample of individuals within the given surveyed community |
| Evenness | the abundance equality of a given species in a specific study site |

Predictor variables

| | |
|------------------------------|--|
| Human population density | the number of individuals per square kilometer in the zip code in which the study site was located |
| Patch size | the area in kilometers of each of the study sites |
| Habitat type | the type of greenspace that makes up greater than 50% of the patch size of the study site (coded as either <i>urban natural habitat</i> or <i>human-altered habitat</i>) |
| Percent developed land cover | the percentage of land cover type within a 500 or 1,000 m buffer comprised of a mixture of constructed materials and vegetation and where impervious surface cover accounts for at least 20% of the land cover |
| Habitat heterogeneity | the number of different greenspace types based on 15 classifications from the National Land Cover Database (2019) constituting each study site (Supplementary Table 2) |
| Region | geographical location of camera traps defined as: (1) mainland (Bronx/Westchester); (2) Manhattan; (3) Long Island; (4) Randall's Island |
| Cameras per km ² | number of cameras sites per study site divided by patch size of study site |

which the best supported model yielded the same results regardless of spatial scale, we report and discuss the model results based on land cover within 500 m (Gallo et al., 2017).

6. *Region*. We classified region into four categories: (1) Long Island; (2) Manhattan; (3) mainland (Bronx and Westchester); and (4) Randall's Island.

Statistical Analyses

All statistical analyses were completed using the R Project for Statistical Computing.⁴ To calculate measures of biodiversity, we used the R package *vegan*.⁵ First, we created a species accumulation curve of the 31 sampling sites to ensure adequate sampling effort (**Supplementary Figure 1**). Second, we calculated alpha diversity (mammal taxa richness) of each study site and gamma diversity (total mammal taxa richness) across all study sites. Third, we used capture frequencies of each mammalian taxon as a proxy for relative abundance (Pardo et al., 2018; Knowlton et al., 2019). To calculate relative abundance, we used the following calculation: number of independent photographs ÷ sampling effort × 100. Sampling effort was calculated as the sum of the number of active camera days. Independent photographs were calculated by counting the maximum number of individuals from the same taxon in the same photograph during hourly intervals. Only detections of the same taxon taken at durations greater than 60 minutes were considered as independent (Cusack et al., 2015; Lewis et al., 2015; Stark et al., 2020). Lastly, we calculated Shannon–Wiener diversity and evenness using the diversity function in *vegan*.

We performed multiple linear regressions to test for predictors of mammal taxa richness, Shannon–Wiener diversity, and evenness. Because carnivores might be more sensitive to the impacts of urbanization (Woodroffe, 2000; Crooks, 2002; Šálek et al., 2014), we also performed multiple linear regressions to detect patterns of species richness, Shannon–Wiener diversity, and evenness with a subset of the data that only included carnivores. For these analyses, we classified carnivores as a functional feeding group (Stark et al., 2020). Therefore, carnivores included all species from the order Carnivora as well as the Virginia opossum (*D. virginiana*), a marsupial that behaves functionally as a carnivore (Stark et al., 2020).

To perform all multiple linear regression models, we used the *stats* package in R. All predictor variables and response variables are described in **Table 2**. Because measures of richness incorporated count data, we modeled these response variables using the *glm* function with Poisson error distribution. Because Shannon diversity indices are strictly positive and continuous, we modeled Shannon diversity using the *glm* function with Gamma error distribution. Finally, we modeled evenness using the *lm* function and tested for model assumptions using the *gvlma* package (Peña and Slate, 2006). We found no violations of any of the model assumptions of linear models.

For all analyses, we performed multiple comparison tests of all possible parameter combinations using the *MuMIn* package,⁶ and we used Akaike information criterion (AIC) to select the best fitting models (Burnham and Anderson, 2004). In cases in which more than one model yielded an AICc difference of <2, we performed model averaging using the summed weight method (Burnham and Anderson, 2004; Grueber et al., 2011). The model-averaged coefficients were calculated using conditional R² (Nakagawa and Schielzeth, 2013).

Lastly, to detect for multicollinearity, we calculated generalized variance inflation factors (GVIFs) for model predictor variables using the package *car* (Fox and Weisberg, 2011). We found no evidence of problematic multicollinearity as all GVIFs were <2.5 (Fox, 2015).

RESULTS

Mammalian Taxa Richness, Diversity, and Evenness

Across the 31 study sites that were sampled, we observed a total of 15 mammal taxa including 8 carnivore species (**Table 3** and **Supplementary Table 3**). A species accumulation curve indicated that there was adequate sampling effort (Soberón and Llorente, 1993; **Supplementary Figure 1**). Measures of diversity (mean ± SD) varied across study sites. Across all mammals, taxa richness ranged from five taxa at both Starlight Park and Willow Lake to a maximum of 13 taxa at Riverdale Park (8.23 ± 2.43; **Table 4**). Shannon–Wiener diversity ranged from 0.703 at Ridgewood Highland to 1.788 at Van Cortlandt Park (1.26 ± 0.289; **Table 4**). Evenness ranged from 0.392 at Ridgewood Highland to 0.856 at Railroad Park (0.61 ± 0.12; **Table 4**). The three most common mammal taxa were free-ranging cats (*F. catus*), gray squirrels (*Sciurus carolinensis*), and raccoons (*P. lotor*); these were found at all 31 study sites. The most common carnivores were free-ranging cats (100% of study sites), raccoons (100% of study sites), and opossums (*D. virginiana*; 87% of study sites) (**Supplementary Table 3**). The three rarest taxa were mink (*Mustela vison*; 3% of study sites; n = 1 study site), groundhogs (*M. monax*; 19% of study sites; n = 6 study sites), and white-tailed deer (*Odocoileus virginianus*; 19% of study sites; n = 6 study sites). Carnivore species richness ranged from three species at Highbridge Park, Ridgewood Park, and Starlight Park to seven species at Inwood Hill Park, Pelham Bay Park, and Riverdale Park (4.58 ± 2.12). Carnivore Shannon–Wiener diversity ranged from 0.14 at Central Park to 1.32 at Ferry Point Park (0.71 ± 0.26). Carnivore evenness ranged from 0.13 at Central Park to 0.81 at Railroad Park (0.47 ± 0.02). The parks with the lowest diversity were mainly comprised of free-ranging cats, gray squirrels, humans, raccoons, small rodents (family Muridae), and opossums (**Supplementary Table 3**).

Predictors of Mammalian Taxa Richness

Based on model averages, we found that patch size and region were significant predictors of mammalian taxa richness at a scale

⁴<https://www.R-project.org/>

⁵<http://vegan.r-forge.r-project.org/>

⁶<https://r-forge.r-project.org/projects/mumin/>

TABLE 3 | Mammal taxa identified across the 31 study sites by region and number of sites.

| Taxa | Mainland | Manhattan | Long Island | Randall's Island | Number of sites |
|---|----------|-----------|-------------|------------------|-----------------|
| Chipmunk (<i>Tamias striatus</i>) | X | X | X | | 7 |
| Cottontail (<i>Sylvilagus floridanus</i>) | X | X | X | | 15 |
| Coyote (<i>Canis latrans</i>)* | X | X | X | | 11 |
| Domestic Cat (<i>Felis catus</i>)* | X | X | X | X | 31 |
| Domestic Dog (<i>Canis familiaris</i>)* | X | X | X | X | 23 |
| Gray Squirrel (<i>Sciurus carolinensis</i>) | X | X | X | X | 31 |
| Groundhog (<i>Marmota monax</i>) | X | X | X | | 6 |
| Human (<i>Homo sapiens</i>) | X | X | X | X | 27 |
| Mink (<i>Mustela vison</i>)* | | X | | | 1 |
| Raccoon (<i>Procyon lotor</i>)* | X | X | X | X | 31 |
| Red Fox (<i>Vulpes vulpes</i>)* | X | X | X | | 7 |
| Small Rodent (family Muridae) | X | X | X | X | 21 |
| Striped Skunk (<i>Mephitis mephitis</i>)* | X | X | X | | 11 |
| Virginia Opossum (<i>Didelphis virginiana</i>)* | X | X | X | X | 27 |
| White-Tailed Deer (<i>Odocoileus virginianus</i>) | X | | X | | 6 |

*Denotes species included in carnivore analysis.

of 500 m. Specifically, study sites with larger patch sizes harbored significantly more mammal taxa than study sites with smaller patch sizes ($\beta = 0.109$; 95% CI: 1.594–2.578; $p = 0.046$; **Figure 2A** and **Table 5**). Moreover, mammal taxa richness was lower in Long Island than on the mainland (Bronx and Westchester) ($\beta = -0.408$; 95% CI: -6.938 to -1.213; $p = 0.005$; **Figure 2B** and **Table 5**). However, at a scale of 1000 m, the effect of patch size and region on mammalian species richness was masked by the percent developed land cover surrounding urban greenspaces. Specifically, greenspaces surrounded by more developed land cover harbored significantly lower mammalian taxa richness than greenspaces surrounded by less developed land cover ($\beta = -0.006$; 95% CI: -0.012 to -0.0003; $p = 0.038$; **Figure 2C** and **Table 5**).

Predictors of Mammalian Shannon–Wiener Diversity

Across all mammal taxa, there were two model parameters that best predicted Shannon–Wiener Diversity at the scale of 500 m: habitat type and region. Specifically, Shannon–Wiener diversity was significantly higher in greenspaces where the dominant habitat type was an urban natural habitat rather than a human altered habitat ($\beta = -0.154$; 95% CI: -2.860 to -2.138; $p = 0.023$; **Figure 3A** and **Table 5**) and on the mainland (Bronx and Westchester) than Long Island ($\beta = 0.192$; 95% CI: 6.546 to 3.193; $p = 0.003$; **Figure 3B** and **Table 5**). When we excluded humans and domestic dogs from our analysis, at a scale of 500 m, one additional model parameter best predicted Shannon–Wiener diversity: human population density. Specifically, Shannon–Wiener diversity was significantly higher in greenspaces with lower human population densities than in greenspaces with higher human population densities ($\beta = 0.000002$; 95% CI: 0.0000002–0.000005; $p = 0.035$). At a scale of 1000 m, greenspaces surrounded by less developed land cover exhibited higher Shannon–Wiener diversity indices than greenspaces surrounded by more developed land cover ($\beta = 0.003$; 95% CI: 0.0003–0.005; $p = 0.030$; **Figure 3C** and **Table 5**).

Predictors of Mammalian Taxa Evenness

Across all mammal taxa, the model parameter that best predicted evenness across the 31 study sites was habitat type. Specifically, evenness was significantly higher in study sites where the dominant habitat type was an urban natural habitat than study sites where the dominant habitat type was a human-altered habitat ($\beta = 0.091$; 95% CI: 0.008–0.175; $p = 0.032$; **Figure 4** and **Table 5**). However, habitat type was not a significant predictor of evenness when we excluded domestic dogs and humans from the analysis.

Predictors of Carnivore Diversity

In analyses that specifically focused on carnivores, there were no significant predictors of carnivore species richness. Carnivore Shannon–Wiener diversity was higher in greenspaces with lower human population densities than in greenspaces with higher human population densities ($\beta = 0.00001$; 95% CI: 3.3 e-06–1.6 e-05; $p = 0.007$; **Figure 5A** and **Table 6**). Carnivore Shannon–Wiener diversity was also significantly higher on the mainland (Bronx and Westchester) than Long Island ($\beta = 0.777$; 95% CI: 3.438–1.225; $p = 0.002$; **Figure 5B** and **Table 6**). Further, Shannon–Wiener diversity of carnivores was higher in greenspaces where the dominant habitat type was urban natural rather than human altered ($\beta = -0.528$; 95% CI: -1.065 to -5.737; $p = 0.049$; **Figure 5C** and **Table 6**). Finally, the best model parameter for predicting carnivore evenness was region. Specifically, evenness was significantly lower in Long Island than on the mainland ($\beta = -0.203$; 95% CI: -3.411 to -6.5678; $p = 0.004$; **Figure 6** and **Table 7**).

DISCUSSION

In this investigation of predictors of mammalian diversity in the New York metropolitan area, we found ample support for the hypothesis that mammalian diversity is associated with anthropogenic factors and geographical barriers. Consistent

TABLE 4 | Average mammal taxa richness, Shannon Wiener diversity, and evenness for each of the 31 study sites.

| Study site | Richness | Shannon–Wiener | Evenness |
|------------------------|-------------|----------------|-------------|
| Alley Pond Park | 11 | 1.484 | 0.619 |
| Bronx Park | 11 | 1.425 | 0.594 |
| Central Park | 8 | 1.292 | 0.621 |
| Clearview Golf Course | 7 | 0.888 | 0.456 |
| Cunningham Park | 8 | 1.098 | 0.528 |
| Elmjack Ingrams Field | 7 | 0.768 | 0.395 |
| Ferry Point Park | 12 | 1.605 | 0.646 |
| Forest Park | 7 | 1.217 | 0.625 |
| Fort Washington Park | 10 | 1.599 | 0.695 |
| Francis Lewis Park | 5 | 1.331 | 0.827 |
| Green-wood Cemetery | 8 | 0.835 | 0.401 |
| Highbridge Park | 6 | 0.944 | 0.527 |
| Hutchinson | 8 | 1.339 | 0.644 |
| Idlewild Park | 7 | 1.241 | 0.638 |
| Inwood Hill Park | 10 | 1.632 | 0.709 |
| Kings Point Park | 8 | 1.212 | 0.583 |
| Maple Grove Cemetery | 6 | 1.151 | 0.642 |
| Pelham Bay Park | 13 | 1.437 | 0.560 |
| Prospect Park | 7 | 1.035 | 0.532 |
| Pugsley Creek Park | 11 | 1.194 | 0.498 |
| Queensline | 9 | 1.353 | 0.616 |
| Railroad Park | 6 | 1.534 | 0.856 |
| Randall's Island | 7 | 1.272 | 0.654 |
| Ridgewood Highland | 6 | 0.703 | 0.392 |
| Riverdale Park | 13 | 1.617 | 0.631 |
| Riverside Park | 6 | 0.940 | 0.525 |
| Smiling Hogshead Ranch | 6 | 1.016 | 0.567 |
| Soundview Park | 10 | 1.787 | 0.766 |
| Starlight Park | 5 | 1.181 | 0.734 |
| Van Cortlandt Park | 12 | 1.788 | 0.720 |
| Willow Lake | 5 | 1.188 | 0.738 |
| Mean (SD) | 8.23 (2.43) | 1.26 (0.289) | 0.61 (0.12) |

with our predictions, we found that mammal taxa richness was higher in greenspaces with larger patch sizes and less developed land cover suggesting that species richness is a function of habitat area (Arrhenius, 1921). Moreover, mammal taxa richness and Shannon–Wiener diversity were greater on the continent (Bronx/Westchester) than Long Island suggesting that there are barriers to dispersal that limit the movement of species between regions (Weckel et al., 2015). Further, both Shannon–Wiener diversity and evenness were higher in urban natural habitats than human-altered habitats. Measures of carnivore diversity were also associated with anthropogenic factors and geographical barriers. Among carnivores, Shannon–Wiener diversity was significantly lower in greenspaces with higher human population densities and in human-altered habitats. Further, both Shannon–Wiener diversity and evenness were significantly higher among carnivores on the mainland (Bronx/Westchester) than in Long Island. Collectively, these results provide insights into the drivers of mammalian community composition and are helpful for drawing

inferences on what factors contribute to mammalian diversity in urban environments.

Throughout the greenspaces of the New York metropolitan area, some mammalian taxa were more common than others. Three species were identified in all 31 study sites: free-ranging cats, gray squirrels, and raccoons. In addition to these three species, the greenspaces that exhibited low taxa diversity also tended to harbor small rodents (family Muridae) and opossums. The five taxa found at these low diversity sites—free-ranging cats, gray squirrels, raccoons, small rodents (family Muridae), and opossums—are known as urban adapters (McKinney, 2002; Adams and Lindsey, 2010; Gehrt et al., 2013; Lombardi et al., 2017). This is largely because these mammals have the capacity to exploit both urban greenspaces while also supplementing their diets with anthropogenic food sources (Cove et al., 2018; Guiry and Buckley, 2018; Nicholson and Cove, 2022). One of these urban adapters, free-ranging cats, is associated with several negative impacts, including the local extinction of native wildlife and the spread of certain pathogens (Gehrt et al., 2013).

In contrast to these animals, three mammalian species were rare among the greenspaces of the New York metropolitan area: minks, groundhogs, and white-tailed deer. Minks were identified in only one of the 31 greenspaces. This is probably because minks require wetland habitats and tend to reside in freshwater streams and rivers (Allen, 1984). Most of our study sites were not situated in wetland habitats, which might explain why the only mink identified was located in a greenspace adjacent to the Hudson River. Previous studies indicate that minks are typically found in sparsely populated rural areas and only occasionally in urban settings (Silva-Rodríguez et al., 2020). White-tailed deer were identified in six of the 31 greenspaces surveyed in this study. Five of these greenspaces were located on the mainland; one was located on Long Island. These findings suggest that the roads and waterways that separate the mainland from the islands make it particularly challenging for large-bodied mammals such as deer to disperse to highly urbanized islands with high human population densities (Michael, 1965; Long et al., 2010). Like deer, groundhogs were also identified in six of the 31 greenspaces surveyed in this study. Surveys indicate that both white-tailed deer and groundhogs are more likely to reside in rural and suburban habitats; however, both of these species have increasingly become more abundant in urban greenspaces in recent years (Gaughan and Destefano, 2005; Lehrer and Schooley, 2010). These findings suggest that minks, groundhogs, and white-tailed deer are urban avoiders (McKinney, 2002), but still have the capacity to exploit some urban habitats. Our findings suggest that urban greenspaces can be refuges for many mammalian taxa, especially urban adapters; however, in some cases, even urban avoiders can make use of certain urban greenspaces.

We found support for the idea that mammalian diversity is characterized by species-area effects. In support of this idea, patch size was positively correlated with species richness. This result is consistent with other studies that have also reported higher mammal taxa richness in greenspaces with relatively larger patch sizes (Yates et al., 1997; Hodgkison et al., 2007; Nielsen et al., 2014). These results also lend support to the species-area hypothesis, the idea that species richness is a function of

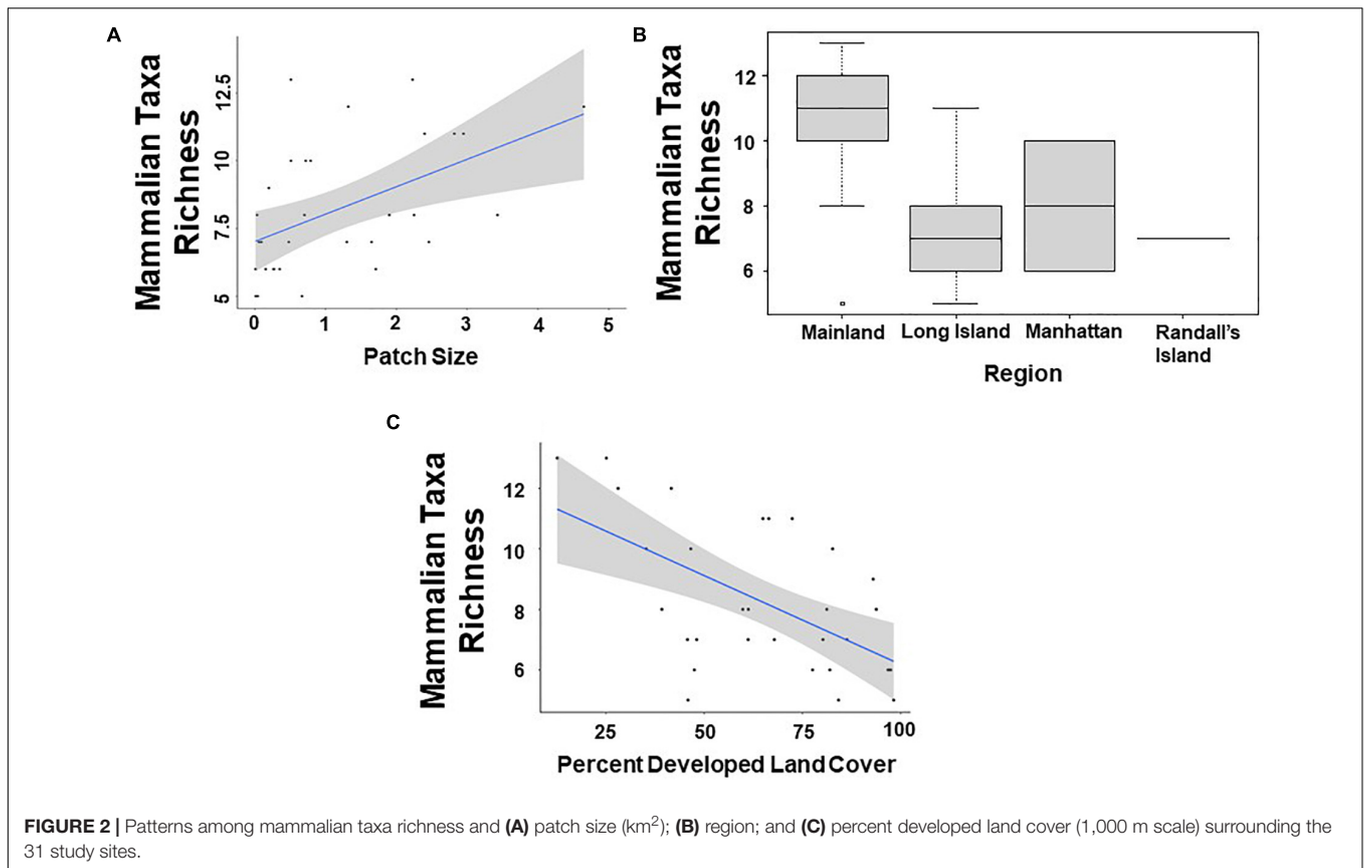


TABLE 5 | Best supported models for each response variable for mammalian diversity based on averaging of parameter estimates.

| Response Variables | Fixed effects | Estimate | Std. error | Adjusted SE | z-value | P-value |
|--|-----------------------------|----------|------------|-------------|---------|---------|
| Mammal taxa richness (500 m scale) | Patch size | 0.109 | 0.052 | 0.054 | 1.995 | 0.046* |
| | Heterogeneity (fine scale) | 0.025 | 0.025 | 0.026 | 0.979 | 0.327 |
| | Percent developed | -0.003 | 0.003 | 0.003 | 1.232 | 0.218 |
| | Human population density | 1E-06 | 1E-06 | 1E-06 | 0.886 | 0.376 |
| | Cameras per km ² | -0.004 | 0.003 | 0.003 | 1.534 | 0.125 |
| | Region—Long Island | -0.408 | 0.139 | 0.146 | 2.79 | 0.005** |
| | Region—Manhattan | -0.299 | 0.19 | 0.199 | 1.498 | 0.134 |
| | Region—Randall's Island | -0.4 | 0.392 | 0.411 | 0.974 | 0.33 |
| | Habitat Type (natural) | 0.091 | 0.132 | 0.138 | 0.659 | 0.51 |
| Mammal taxa richness (1000 m scale) | Percent developed | -0.006 | 0.003 | 0.003 | 2.074 | 0.038* |
| | Patch size | 0.07 | 0.054 | 0.057 | 1.243 | 0.214 |
| | Habitat type (natural) | 0.119 | 0.131 | 0.137 | 0.871 | 0.384 |
| | Heterogeneity (fine scale) | 0.036 | 0.028 | 0.029 | 1.248 | 0.212 |
| Shannon–Wiener diversity (500 m scale) | Human population density | 1E-06 | 1E-06 | 1E-06 | 1.746 | 0.081 |
| | Habitat type (natural) | -0.154 | 0.065 | 0.068 | 2.277 | 0.023* |
| | Region—Long Island | 0.192 | 0.062 | 0.065 | 2.971 | 0.003** |
| | Region—Manhattan | 0.014 | 0.11 | 0.114 | 0.121 | 0.904 |
| | Region—Randall's Island | -0.001 | 0.163 | 0.172 | 0.007 | 0.995 |
| Shannon–Wiener diversity (1,000 m scale) | Habitat type (natural) | -0.164 | 0.061 | 0.064 | 2.555 | 0.011* |
| | Percent developed | 0.003 | 0.001 | 0.001 | 2.172 | 0.030* |
| | Region—Long Island | 0.152 | 0.062 | 0.065 | 2.349 | 0.019* |
| | Region—Manhattan | 0.09 | 0.078 | 0.082 | 1.104 | 0.27 |
| | Region—Randall's Island | -0.038 | 0.162 | 0.17 | 0.224 | 0.823 |
| Mammalian taxa evenness (500 m scale) | Habitat type (natural) | 0.091 | 0.041 | 0.043 | 2.142 | 0.032* |
| | Patch size | -0.021 | 0.017 | 0.017 | 1.205 | 0.228 |
| | Percent developed | 0.001 | 0.001 | 0.001 | 0.881 | 0.378 |

Model-average coefficients (conditional average), standard error, adjusted SE, z-value, and P value of the averaged models are shown. *Denotes significance of <0.05. **Denotes significance of <0.01.

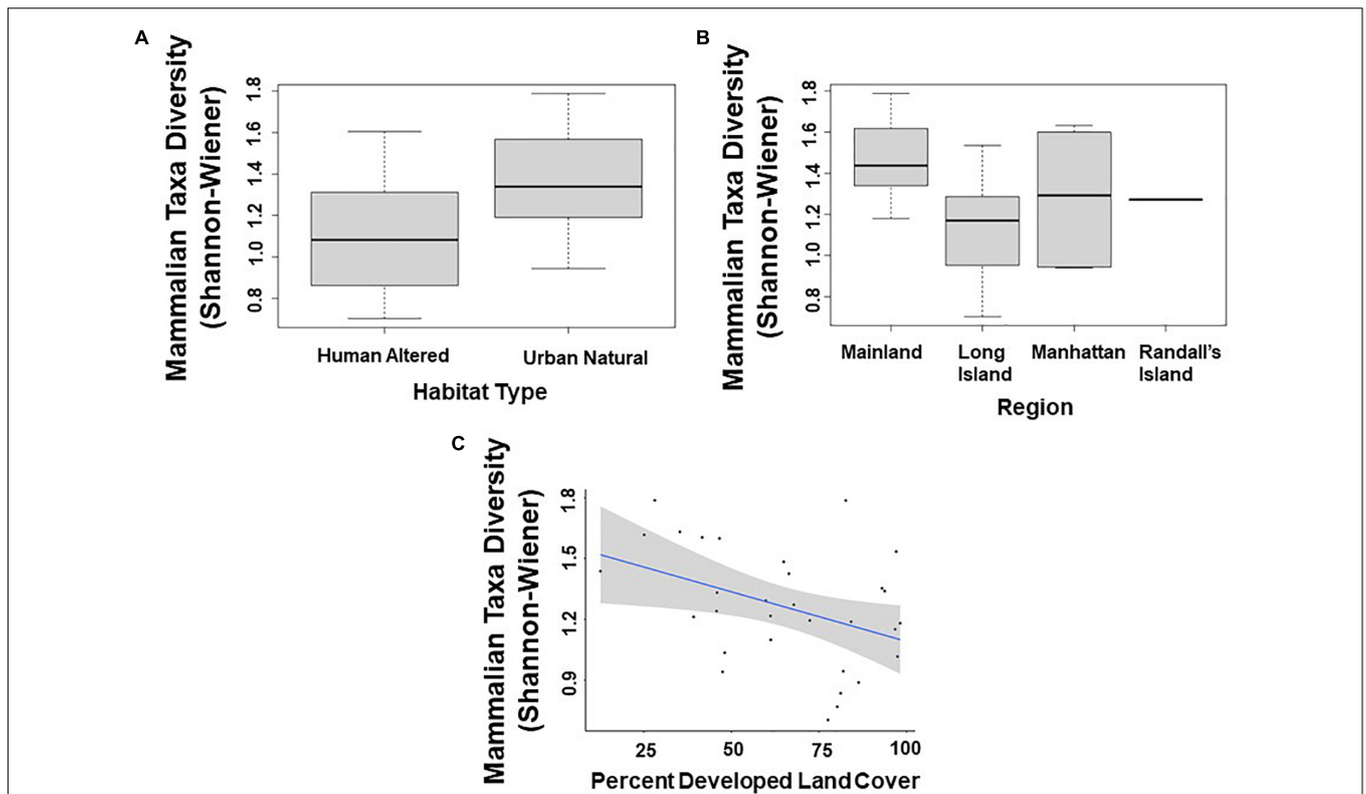


FIGURE 3 | Patterns among mammalian taxa diversity (Shannon–Wiener index) and **(A)** type of habitat; **(B)** region; and **(C)** percent developed land cover (1,000 m scale) across the 31 study sites. Points and whiskers on the plot represent the mean and standard error.

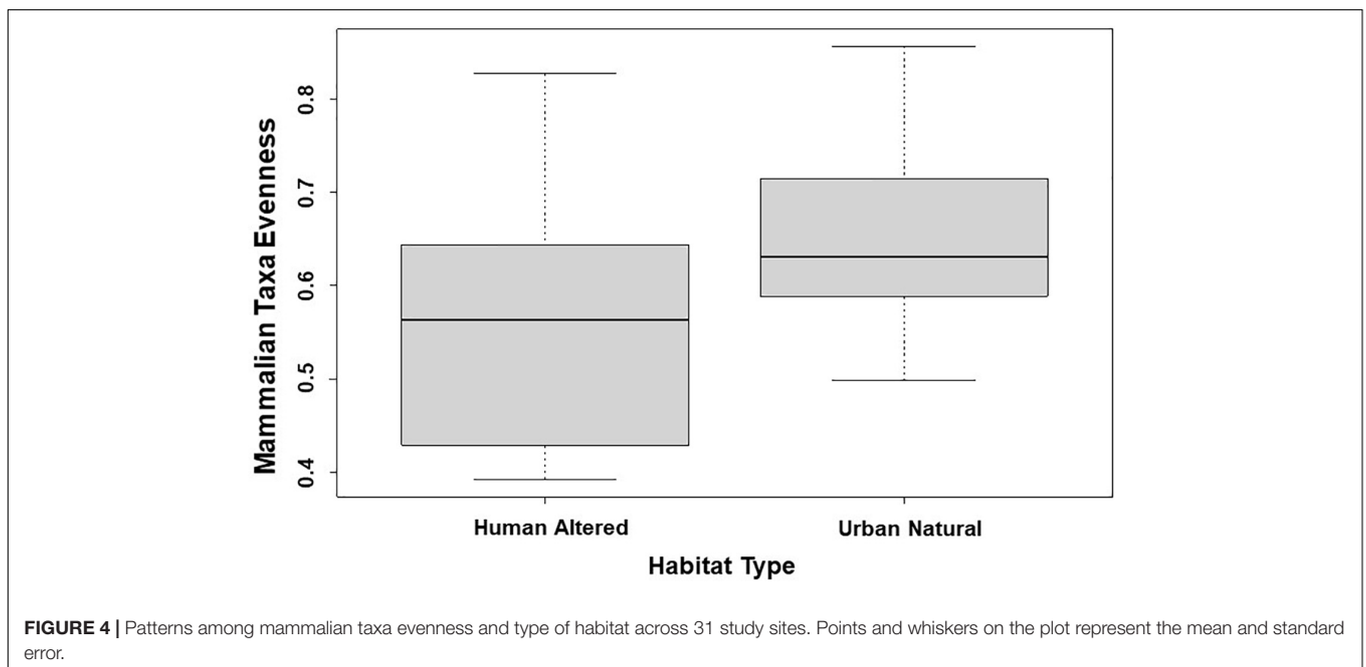


FIGURE 4 | Patterns among mammalian taxa evenness and type of habitat across 31 study sites. Points and whiskers on the plot represent the mean and standard error.

the amount of available habitat (Arrhenius, 1921; Turner et al., 2005). In the case of the current study, we find evidence of the species-area effect in an urban context.

Habitat type was a significant predictor of two measures of biodiversity: Shannon–Wiener diversity and evenness. First, we found that Shannon–Wiener diversity was higher in study sites

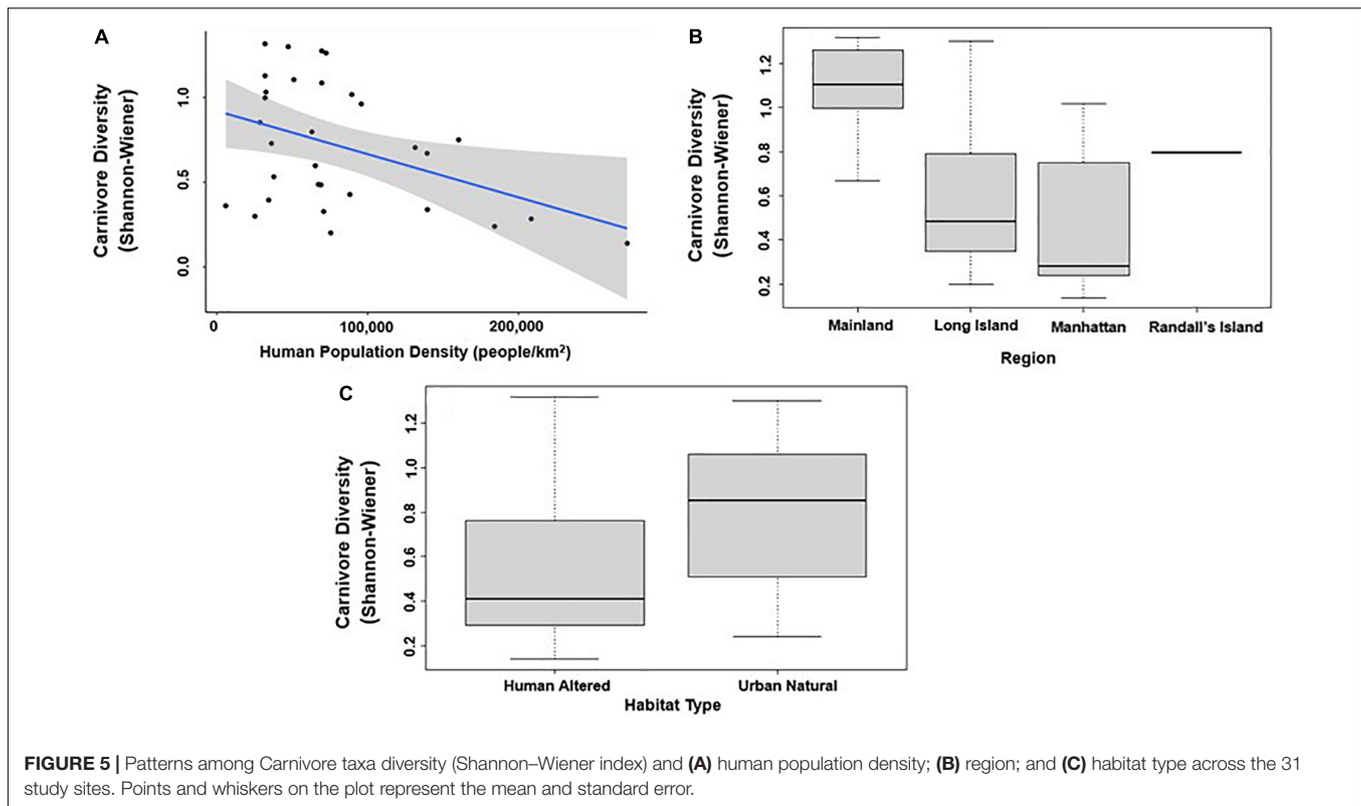


TABLE 6 | Best supported model for Carnivore Shannon–Wiener diversity (500 m scale) based on averaging of parameter estimates.

| Predictor variable | Fixed effects | Estimate | Std. error | t-value | P-value |
|--|--------------------------|----------|------------|---------|---------|
| Carnivore Shannon–Wiener diversity (500 m scale) | Human population density | 0.00001 | 3E-06 | 2.942 | 0.007** |
| | Region—Long Island | 0.777 | 0.223 | 3.484 | 0.002** |
| | Region—Manhattan | 0.157 | 0.474 | 0.331 | 0.743 |
| | Region—Randall's Island | −0.124 | 0.591 | −0.21 | 0.836 |
| | Habitat type (natural) | −0.528 | 0.256 | −2.065 | 0.049* |

Model-average coefficients (conditional average), standard error, t-value, and P value of the averaged models are shown. *Denotes significance of <0.05. **Denotes significance of <0.01.

where the dominant habitat was an urban natural habitat rather than a human-altered habitat. Although some mammals may thrive in human-altered habitats, like golf courses and cemeteries (Gallo et al., 2017; Wurth et al., 2020), most mammals tend to have higher persistence in urban natural areas, or habitats that consist of large patches of secondary growth forest and shrubs (Atwood et al., 2004; Baker and Harris, 2007). It has also been found that manicured lawns and artificial nocturnal illumination, as seen in parks, golf courses, and cemeteries, may limit dispersal (Mahan and O'Connell, 2005; Fitzgibbon et al., 2007). Second, we also found a positive correlation between habitat type and evenness. Specifically, we found that evenness was higher in study sites where more than fifty percent of the patch area was secondary growth forest. These findings are consistent with previous research that suggest that diversity in urban natural habitats is higher than in human-altered habitats (Parsons et al., 2018). Taken together, these two findings—higher Shannon–Wiener diversity and higher taxa evenness among

mammals in urban natural habitats—suggest that it is important for park managers to maintain at least 50% natural area within urban parks to support mammalian diversity.

Consistent with our predictions, we found that greenspaces surrounded by more developed land cover exhibited lower mammal taxa richness than greenspaces surrounded by less developed land cover. This result was only significant at a scale of 1000 meters, but at this scale, it masked the effects of any of the other predictor variables. This finding appears to contradict the urban refugia hypothesis (Stark et al., 2020), which suggests that urban greenspaces in highly developed areas are oases for biodiversity. More specifically, our results differ from the findings of a recent study in the New York metropolitan area that found support for the urban refugia hypothesis: Stark et al. (2020) found that the relative abundance of mammalian carnivores was higher in greenspaces with higher surrounding human development. In our analysis of mammalian carnivores, we found no significant relationship between human development and carnivore species

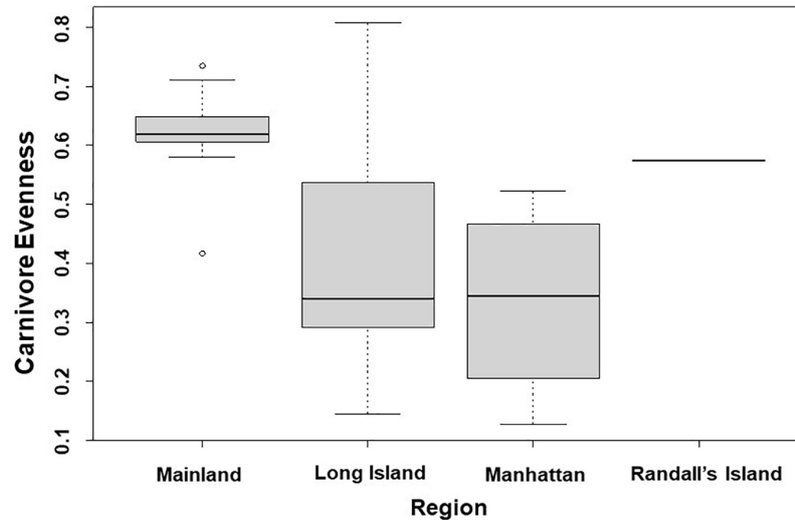


FIGURE 6 | Patterns among Carnivore evenness and region across the 31 study sites. Points and whiskers on the plot represent the mean and standard error.

TABLE 7 | Best supported model for Carnivore evenness (500 m scale).

| Predictor variable | Fixed effects | Estimate | Std. error | Adjusted SE | z-value | P-value |
|----------------------------------|--------------------------|-----------|------------|-------------|---------|---------|
| Carnivore evenness (500 m scale) | Patch size | -0.040 | 0.024 | 0.026 | 1.581 | 0.114 |
| | Percent developed | 0.001 | 0.001 | 0.001 | 1.193 | 0.233 |
| | Human population density | -0.000001 | 1E-06 | 1E-06 | 1.813 | 0.070 |
| | Region—Long Island | -0.203 | 0.067 | 0.07 | 2.895 | 0.004** |
| | Region—Manhattan | -0.177 | 0.132 | 0.136 | 1.3 | 0.194 |
| | Region—Randall's Island | 0.003 | 0.172 | 0.18 | 0.016 | 0.988 |
| | Habitat type (natural) | 0.122 | 0.06 | 0.063 | 1.956 | 0.050 |

Standard error, adjusted SE, z-value, and P value of the best model is shown. · Denotes $P < 0.1$. **Denotes significance of <0.01 .

richness. However, the Stark et al. (2020) study was conducted in nature preserves outside of New York City and in less populous areas than the current study. Specifically, the mean (\pm SD) human population density for the Stark et al. (2020) study was 7,305 (\pm 11,222) humans per square kilometer while the mean human population density in the current study was 81,451 (\pm 60,515) humans per square kilometer. Perhaps in highly urbanized, densely populated areas, a minimum amount of greenspace is required for urban greenspaces to function as refugia for mammals, especially carnivores. Another factor that might explain our result is that in highly developed areas, buildings, roads, and other anthropogenic structures reduce connectivity between habitats (LaPoint et al., 2015; Beninde et al., 2016). Obstruction in connectivity can also lead to resource depletion, habitat disturbance, and ultimately reduced species richness (Blair and Launer, 1997; Fidino et al., 2020). Additionally, our result might also be attributed to higher mortality rates caused by humans, including vehicle collisions (Trombulak and Frissell, 2000; Seiler, 2001; Collins and Kays, 2011) and poisoning (Brooks et al., 2020). Overall, our results suggest that highly developed areas pose a particularly daunting challenge for maintaining high levels of mammalian diversity.

Urban greenspaces located on the mainland (Bronx and Westchester) harbored greater mammal taxa richness and Shannon–Wiener diversity than urban greenspaces located on

urban islands (Long Island, Manhattan, Randall's Island). These findings suggest that geographical barriers to dispersal might impact patterns of mammalian diversity. In support of this idea, previous studies have documented multiple barriers to dispersal between the mainland (Bronx and Westchester) and the other three regions (Long Island, Manhattan, and Randall's Island) (Weckel et al., 2015; Nagy et al., 2016). Additionally, roads and waterways, as well as other barriers, play a large role in limiting dispersal (Oxley et al., 1974; Merriam et al., 1989; Baker and Harris, 2007). For example, coyotes have not established strong breeding populations yet in Long Island mainly because of bridges and rivers that separate the island from the Bronx and Westchester (Nagy et al., 2017). Our results lend support to the hypothesis that islands, even those in close proximity to the continent, can serve as barriers to dispersal.

The New York metropolitan area harbors several carnivore species that vary in their abundance and distribution. We found that the greenspaces surveyed in this study supported one or more of the following carnivores: coyotes (*C. latrans*), free-ranging cats (*F. catus*), domestic dogs (*Canis familiaris*), minks (*M. vison*), opossums (*D. virginiana*), raccoons (*P. lotor*), red foxes (*V. vulpes*), and striped skunks (*Mephitis mephitis*). Because of their low densities, need for large habitat area, and conflict with humans, many large carnivores once native to this region, including black bears (*Ursus americanus*), bobcats

(*Lynx rufus*), fishers (*Pekania pennanti*), and long-tailed weasels (*Neogale franata*) were not observed in any of the greenspaces surveyed in this study, although these mammals persist in nature preserves outside of New York City (Spinola et al., 2008; Stark et al., 2020; Nagy, unpublished data). In the present study, we found that Shannon–Wiener diversity among carnivores was higher in urban natural habitats and in greenspaces with lower human population densities. In support of these findings, previous studies have found that many carnivores tend to avoid human interaction, preferring less developed urban habitats with lower human population densities (Grinder and Krausman, 2001; George and Crooks, 2006; Gehrt et al., 2009). We also found that carnivore evenness was significantly higher in the mainland than in sites located in Long Island and Manhattan. This finding suggests that there might be differences in the habitat characteristics between the mainland sites (Bronx and Westchester) and the island sites (Manhattan and Long Island) including differences in floral diversity (Ekernas and Mertes, 2006), resource availability (Matthies et al., 2017), the historical range of various species (e.g., Rooney, 2001; Hody and Kays, 2018), and management strategies (McPhearson et al., 2016). Furthermore, the asymmetric distribution of carnivore communities on the islands might also be explained by barriers to dispersal between the mainland and island sites. Although large mammals, like carnivores, are more adept at crossing bridges and waterways than smaller mammals, these barriers still hinder and reduce the likelihood of movement between regions (Weckel et al., 2015; Henger et al., 2020). Collectively, these findings suggest that carnivores are particularly sensitive to urbanization and barriers to dispersal such as bridges and large bodies of water.

Because we did not trap or tag species and relied solely on camera data, one limitation of this study was that we were unable to identify individuals. Therefore, the abundance indices that we calculated are rough estimates and may not be representative of actual mammal abundances. In future studies, the use of mark-recapture or identification techniques to assess individual sightings would ensure a more accurate representation of abundance (Ekernas and Mertes, 2006; Lewis et al., 2015; Lombardi et al., 2017). The current study surveyed sixteen sites on Long Island, thirteen of which were located in Queens, two in Brooklyn, and one in Nassau County. Since the majority of the Long Island study sites were located in Queens, the results may not be representative of mammal diversity for all of the island. For future studies, it would be beneficial to have a more even distribution of study sites across the island to allow for more accurate comparison of regions.

CONCLUSION AND FUTURE DIRECTIONS

Our findings suggest that anthropogenic variables, such as percent developed land cover, patch size, and habitat type, influence patterns of mammalian diversity, and that it is crucial to understand and study the consequences of increasing urbanization on mammalian communities. There were some mammals in the current study that appeared to flourish across all urban greenspaces (i.e., urban exploiters) and were less

sensitive to urbanization, apparently because they can take better advantage of anthropogenic food resources. On a more nuanced level, our findings suggest that there are some habitat features that are more suitable for mammals that are sensitive to urbanization (i.e., urban avoiders), including large patch sizes and high proportions of secondary growth forest. Hence, while urbanization is generally detrimental to biodiversity, we found that there were greenspaces in the New York metropolitan area where mammalian diversity was high. This suggests that there is a potential to make cities more suitable habitats for many mammalian species and that a barren ecology is not inevitable. The current study provides correlative evidence that some features of urbanization are associated with patterns of mammalian biodiversity. Our findings might be helpful in terms of management and conservation efforts within similarly large urban centers. Based on our results, we recommend that planners and managers of cities focus their efforts on maintaining large, connected, natural greenspaces, especially when surrounded by highly developed areas with high human population densities. Urban ecological research would benefit from longitudinal studies conducted across multiple large cities that can document changes in diversity, abundance, and evenness of mammals.

DATA AVAILABILITY STATEMENT

The original contributions presented in this study are included in the article/**Supplementary Material**, further inquiries can be directed to the corresponding author.

ETHICS STATEMENT

Ethical review and approval was not required for the animal study because this was an observational study using photographs.

AUTHOR CONTRIBUTIONS

AB, CN, MW, DL, and BH designed the study. CN and MW collected the data. AB and BH analyzed the data. AB, DL, and BH wrote the manuscript. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fevo.2022.903211/full#supplementary-material>

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Table S1. Field Site Descriptions and Features

| Study Site | Site Description and Features |
|------------------------------|--|
| Alley Pond Park | greenspace characterized by woodlands and hilly terrain with sandy ridges, meadows, ponds, and surrounding wetlands ¹ . |
| Bronx Park | an urban park characterized by a natural streamside terrain with a mixture of red maple hardwood and floodplain forests along the banks of the Bronx River ² . |
| Central Park | an urban park dominated by large American elms and containing a multitude of athletic and recreational facilities ³ (Atha et al., 2020). |
| Clearview Golf Course | human-altered habitat consisting of a hilly terrain with a combination of open and tree lined fairways; in addition to natural forests, this location also contains artificial ponds ⁴ . |
| Cunningham Park | greenspace characterized by hilly terrain with many hiking trails, sports facilities, and developed greenways; this park contains a section of relatively untouched woodland containing mostly of oak-hickory trees ⁵ . |
| Elmjack Ingrams Field | athletic field lined by small, forested area on three sides and an empty lot on the fourth side; due to its recreational purposes, it has a manicured field with multiple patches of dirt. |
| Ferry Point Park | an urban park containing golf courses as well as sports fields and recreational pathways; due to its location and proximity to multiple parkways, it serves as a connection to multiple parks ⁶ . |
| Forest Park | greenspace made up of a combination of natural areas and manicured habitats; on the hilly terrain of the eastern side, it contains natural forests of great oaks, while on the western side it contains athletic fields and golf courses ⁷ (Glaeser, 2006). |
| Fort Washington Park | an urban park serving as a connection between parklands along a large portion of the Hudson River; home to many native trees and shrubs; filled with meadow grasses ⁸ . |
| Francis Lewis Park | greenspace where the majority of its area is above water while the remaining area is submerged under water; this park also allows for beach access unlike many other parks in New York City ⁹ . |
| Green-wood Cemetery | cemetery known as one of the largest urban grasslands in New York City; comprised of a rolling landscape with diverse flora, this cemetery provides an “urban oasis” when compared to its dense, urban surroundings (Charap et al., 2020). |
| Highbridge Park | an urban park that is set atop a rocky cliff and contains a variety of trees creating woodlands and glens ¹⁰ . |
| Hutchinson | patch of greenspace comprised of patchy forested areas, manicured lawns, and is flanked by a nearby parking lot. |
| Idlewild Park | urban greenspace characterized by a diverse array of habitats including woodlands, meadowlands, tidal wetlands, freshwater wetlands, and grasslands ¹¹ . |
| Inwood Hill Park | greenspace characterized by four plant communities: 1) salt marsh 2) forest 3) lawn 4) successional field, as well as seven types of vegetation; this park |

| | |
|-------------------------------|---|
| | also contains wooded forests as well as the last salt marsh in Manhattan ¹² (Loeb, 1986). |
| Kings Point Park | urban park that is located near the Long Island Sound and contains swamps consisting of red maples, sweetgum, sour gum, sassafras, as well as black birch; within the park, there are multiple manicured athletic fields as well as numerous hiking trails ¹³ (Greller et al., 2006). |
| Maple Grove Cemetery | cemetery situated at one of the highest elevations in Queens atop rocky hills and surrounded by railroads; this site contains a natural dense forest with manmade lakes ¹⁴ . |
| Pelham Bay Park | urban park situated between two bodies of water and comprised of a mixture of meadows, rocky shores, dense woodlands, and saltwater marshes; transitioning into a secondary forest from a wooded savannah, this park is home to oaks and low grassy vegetation, as well as golf courses, athletic fields, and hiking trails ¹⁵ (Ekernas and Mertes, 2006). |
| Prospect Park | urban park known for containing one of Brooklyn's last remaining and richest forests filled with the pin oaks, Norway maples, and sycamore maples; this site contains woodlands, wetlands, meadows, and the only lake in Brooklyn ^{16,17,18} . |
| Pugsley Creek Park | greenspace bordered by both two creeks from which the East River flows into; Pugsley Creek Park is one of the few parks that have remained mostly undeveloped with pristine freshwater wetlands ¹⁹ . |
| Queensline | greenspace characterized by a unique thin line of forested area lying on the border between Forest Hills and Rego Park, Queens; this patch of forest is perpendicular to the Long Island Railroad and is in close proximity to other greenspaces including parks, cemeteries, and athletic fields. |
| Railroad Park | greenspace located next to a railroad and known as a woodland preserve; this site is mostly fenced off and is mainly comprised of natural flora with some landscaping ²⁰ . |
| Randall's Island | large greenspace home to several different natural habitats including an urban forest, freshwater wetlands, a tidal marsh, meadows, coastal areas, as well as an urban farm; the park is dominated by wild grasses and herbaceous plants and is also home to white ash, common hackberry, and eastern cottonwood trees ^{21,22} . |
| Ridgewood Highland | urban park located within the Ridgewood Reservoir and home to a lush, dense forest as well as a freshwater pond; there are also playgrounds and athletic fields located within the park ²³ . |
| Riverdale Park | this urban park runs along a railroad beside the Hudson River in the Bronx and is home to a large oak hickory forest as well as a successional forest as a result of glacial activity ²⁴ . |
| Riverside Park | situated above the Hudson River, this park is characterized by hilly terrain and a combination of natural areas and fields, courts, and slopes for athletic purposes ²⁵ . |
| Smiling Hogshead Ranch | urban farm formed atop abandoned railroad tracks and containing both agricultural farm area and gardens ²⁶ . |
| Soundview Park | greenspace characterized by vast marshlands located in close proximity to the point in which the Bronx River changes from saltwater to the only freshwater |

| | |
|---------------------------|--|
| | river in New York; initially built atop a landfill, this park now contains athletic fields including courts, fields, and tracks ^{27,28} . |
| Starlight Park | urban park containing numerous athletic fields, playgrounds, picnic areas, and a dock for kayaking; our study site location was set in the Bronx River greenway, a small, forested area across from the Bronx River and parallel to Starlight Park ²⁹ . |
| Van Cortlandt Park | urban park known as the third-largest park in New York City with a secondary forest composed of a mixed hardwood forest of white ash, oaks, and tulip poplars; along with natural wetlands and woodlands, this park also contains multiple athletic spaces including a stadium, golf courses, and hiking trails (McDonnell et al.,1990). |
| Willow Lake | greenspace containing one of the largest bodies of water in Queens, surrounded by the parkland of Flushing Meadows Corona Park; this site contains thick reeds across the freshwater wetlands as well as a multitude of willows and other flower species ³⁰ . |

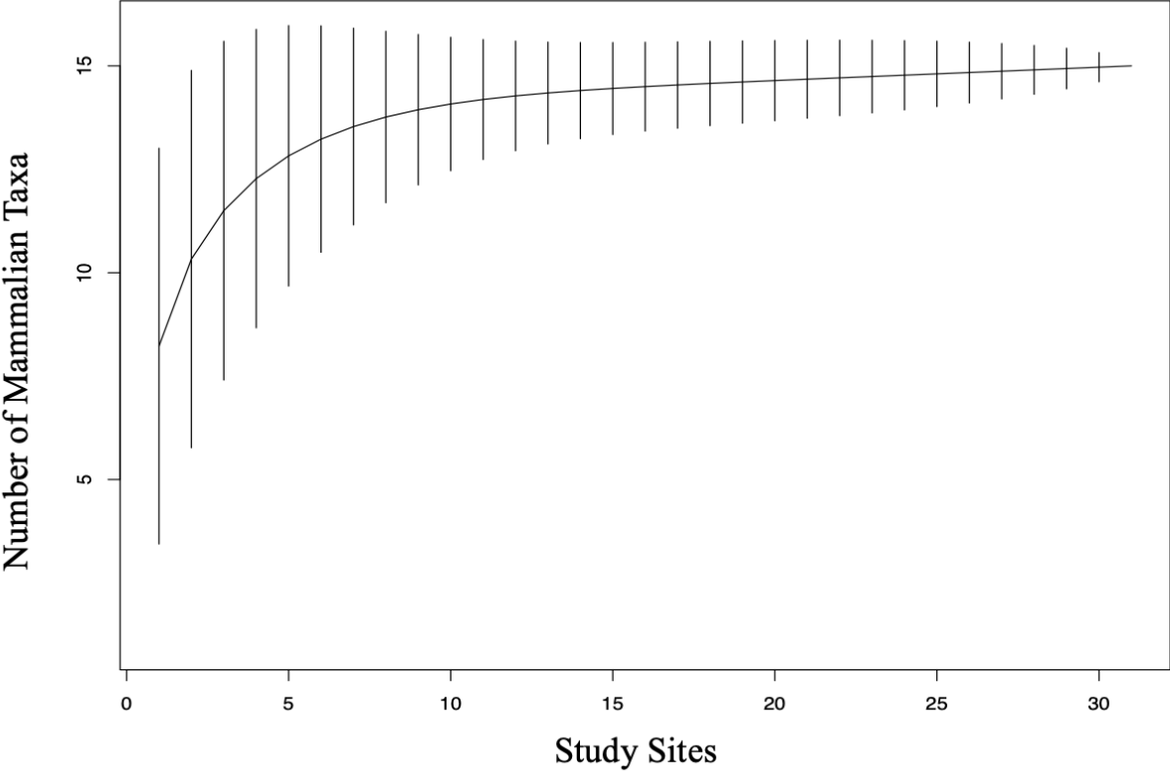
Table S2. National Land cover Database classes

| Class | Classification Description |
|-------------------------------------|--|
| Open water | sites consisting of open water with less than 25% soil or vegetation |
| Developed: Open space | sites with a combination of lawn grasses and built materials with less than 20% impervious surfaces (i.e., parks, manicured lawns, golf courses, housing developments) |
| Developed: Low intensity | sites containing a combination of natural vegetation and built materials with 20-49% impervious surfaces |
| Developed: Medium intensity | sites with both natural vegetation and built materials with impervious surfaces accounting for 50-79% of the land cover |
| Developed: High intensity | sites where impervious surfaces make up 80-100% of the land cover due to high development resulting from commercial and industrial activities |
| Barren land | sites consisting mainly of natural earthy material with less than 15% of the land cover being vegetation |
| Deciduous forest | sites that consist of more than 20% vegetation cover; predominantly comprised of trees over five meters tall with over 75% of the trees shedding leaves seasonally |
| Evergreen forest | sites with vegetation encompassing more than 20% of total land cover and trees that maintain their green color and do not shed leaves seasonally |
| Mixed forest | sites with more than 20% of their land cover consisting of vegetation with neither deciduous nor evergreens comprising more than 75% of the tree cover |
| Shrub/scrub | sites that have more than 20% of their land cover consisting of shrubs less than five meters tall |
| Grassland/herbaceous | sites that have 80% or more of total land cover as grass-like vegetation |
| Pasture/hay | sites that have more than 20% of their land cover as some form of grasses or legumes for animals to graze on or to produce hay |
| Crops | sites where crop vegetation accounts for over 20% of the total land cover and the area is continuously tilled and used for producing annual crops |
| Woody/wetlands | sites in which shrubland or forests comprise more than 20% of the total land cover, and the soil is intermittently soaked with water |
| Emergent herbaceous wetlands | sites that consist of more than 80% perennial herbaceous vegetation and the soil is intermittently soaked with water |

Table S3. Mammal taxa detection across the 31 study sites

| Study Site | Chipmunk | Cotton-tail Rabbit | Coyote | Free-ranging Cat ^A | Domestic Dog ^A | Gray Squirrel | Ground-hog | Human | Mink | Raccoon | Red Fox | Small Rodent | Striped Skunk | Virginia Opossum | White-tailed Deer | Total Richness | Camera Trap Days | Sampling effort by area |
|---|----------|--------------------|--------|-------------------------------|---------------------------|---------------|------------|-------|------|---------|---------|--------------|---------------|------------------|-------------------|----------------|------------------|-------------------------|
| Alley Pond Park | X | X | | X | X | X | | X | | X | X | X | X | X | | 11 | 1089 | 386 |
| Bronx Park | X | X | X | X | X | X | | X | | X | | X | | X | X | 11 | 835 | 283 |
| Central Park | X | | | X | X | X | X | X | | X | | X | | | | 8 | 362 | 106 |
| Clearview Golf Course | | X | | X | X | X | | X | | X | | | | X | | 7 | 201 | 419 |
| Cunningham Park | | | | X | X | X | | X | | X | X | X | | X | | 8 | 387 | 172 |
| Elmjack Ingrams Field | | | X | X | | X | | X | | X | | X | | X | | 7 | 132 | 2276 |
| Ferry Point Park | X | X | X | X | X | X | | X | | X | | X | X | X | X | 12 | 680 | 515 |
| Forest Park | | | | X | X | X | | X | | X | | X | | X | | 7 | 141 | 108 |
| Fort Washington Park | | X | | X | | X | X | X | X | X | | X | X | X | | 10 | 159 | 312 |
| Francis Lewis Park | | | | X | X | X | | X | | X | | | | | | 5 | 224 | 5744 |
| Green-wood Cemetery | | X | | X | | X | X | X | | X | | | X | X | | 8 | 165 | 87 |
| Highbridge Park | | | | X | | X | X | X | | X | | X | | | | 6 | 29 | 111 |
| Hutchinson | | | | X | | X | X | X | | X | X | | X | X | | 8 | 177 | 5710 |
| Idlewild Park | | | | X | X | X | | X | | X | | X | | X | | 7 | 246 | 149 |
| Inwood Hill Park | | X | X | X | X | X | | X | | X | X | | X | X | | 10 | 564 | 715 |
| Kings Point Park | | | | X | X | X | | | | X | X | X | | | X | 8 | 91 | 130 |
| Maple Grove Cemetery | | | | X | X | X | | X | | X | | | | X | | 6 | 404 | 1485 |
| Pelham Bay Park | X | X | X | X | X | X | | X | | X | X | X | X | X | X | 13 | 1280 | 574 |
| Prospect Park | | X | | X | X | X | | X | | X | | X | | | | 7 | 223 | 91 |
| Pugsley Creek Park | | X | X | X | X | X | | X | | X | | X | X | X | X | 11 | 350 | 146 |
| Queensline | | X | X | X | X | X | | X | | X | | X | | X | | 9 | 176 | 889 |
| Railroad Park | | | X | X | X | X | | | | X | | | | X | | 6 | 196 | 1289 |
| Randall's Island | | | | X | X | X | | X | | X | | X | | X | | 7 | 283 | 3076 |
| Ridgewood Highland | | X | | X | | X | | | | X | | X | | X | | 6 | 54 | 32 |
| Riverdale Park | X | X | X | X | X | X | X | X | | X | X | X | X | X | | 13 | 2279 | 4469 |
| Riverside Park | | | | X | X | X | | X | | X | | | | X | | 6 | 223 | 637 |
| Smiling Hogshead Ranch | | | | X | X | X | | X | | X | | | | X | | 6 | 195 | 24375 |
| Soundview Park | | X | X | X | X | X | | X | | X | | X | X | X | | 10 | 393 | 547 |
| Starlight Park | | | | X | | X | | X | | X | | | | X | | 5 | 65 | 5909 |
| Van Cortlandt Park | X | X | X | X | X | X | | X | | X | | X | X | X | X | 12 | 1176 | 253 |
| Willow Lake | | | | X | | X | | | | X | | X | | X | | 5 | 50 | 75 |
| Proportion of sites occupied | 0.23 | 0.48 | 0.35 | 1.00 | 0.74 | 1.00 | 0.19 | 0.87 | 0.03 | 1.00 | 0.23 | 0.68 | 0.35 | 0.87 | 0.19 | | | |
| "X" denotes detection of taxa on at least one camera trap at the given study site; ^A = domestic animal | | | | | | | | | | | | | | | | | | |

Figure S1: Species accumulation curve representing the number of mammalian taxa present across the 31 study sites



Footnotes

1. <https://www.nycgovparks.org/parks/Q001/>
2. <https://www.nycgovparks.org/parks/bronx-park/highlights/11414>
3. <https://www.centralparknyc.org/about>
4. <https://www.nycgovparks.org/parks/clearview-park-golf-course/history>
5. <https://www.nycgovparks.org/parks/cunningham-park/history>
6. <https://www.nycgovparks.org/parks/ferry-point-park/history>
7. <https://www.nycgovparks.org/parks/Q015/>
8. <https://www.nycgovparks.org/parks/fort-washington-park/history>
9. <https://www.nycgovparks.org/parks/francis-lewis-park>
10. <https://www.nycgovparks.org/parks/highbridge-park>
11. <https://www.nycgovparks.org/greening/nature-preserves/site?FWID=32>
12. <https://www.nycgovparks.org/parks/inwood-hill-park>
13. <https://www.gnparks.org/499/Major-Parks>
14. <http://www.maplegrove.biz/about/rich-history/>
15. http://pelhambaypark.org/?page_id=71
16. <https://www.dec.ny.gov/outdoor/57289.html>
17. <https://www.prospectpark.org/learn-more/archives/park-tree-history/>
18. <https://www.nycgovparks.org/parks/B073/>
19. <https://www.nycgovparks.org/parks/pugsley-creek-park/highlights/11647>
20. <https://www.nycgovparks.org/parks/railroad-park-q412>
21. <https://www.inaturalist.org/guides/1430>
22. <https://randallsisland.org/visit/waterfront-natural-areas/>
23. <https://queensmuseum.org/2020/02/ridgewood-reservoir-for-the-21st-century>
24. <https://www.nycgovparks.org/parks/riverdale-park>
25. <https://riversideparknyc.org/overview/>
26. <https://www.idealists.org/en/nonprofit/0e0735788f8e45a68144cf7449493f91-smiling-hogshead-ranch-queens>
27. <https://www.nycgovparks.org/parks/soundview-park>
28. <https://www.seagrant.sunysb.edu/hriver/pdfs/coastalchange/SoundviewPark/ProjectSummary-Narrative.pdf>
29. <https://www.nycgovparks.org/parks/starlight-park>
30. <https://www.nycgovparks.org/parks/flushing-meadows-corona-park/highlights/12907>

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