Positional Behavior of Introduced Monk Parakeets (*Myiopsitta monachus*) in an Urban Landscape

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Simple Summary: Positional behaviors comprise the entirety of animals’ locomotion and posture. Often, these positional behaviors are paired with information about substrate characteristics (e.g., orientation, diameter, texture, height) and frequency to gain an ecological perspective of when and why an animal utilizes a particular behavior. Thus far, quantitative studies of positional behavior have been limited to mammals, leaving a major gap in our understanding of how animals utilize their environment. In this study, we present the first quantitative report of positional behavior within Aves, presenting scan sampling data from an established colony of Monk Parakeets (*Myiopsitta monachus*) from Brooklyn, New York City. Parrots exhibited a strong preference for small and terminal branches, a selection which may reflect targeted foraging of new fruit growth and leaf-buds. We further observed that the gait transition from walking to sidling appears primarily driven by substrate size, with the former preferred on the ground and on large, broad substrates and the latter used to navigate smaller branches. Finally, we observed an increase in locomotor diversity on artificial versus naturally occurring substrates. This demonstrates the importance of a flexible behavioral repertoire in facilitating a successful transition towards an urban landscape in introduced species and underscores the need for further studies exploring positional behaviors among urban wildlife.

Abstract: Positional behaviors have been broadly quantified across the Order Primates, and in several other mammalian lineages, to contextualize adaptations to, and evolution within, an arboreal environment. Outside of Mammalia, however, such data are yet to be reported. In this study, we present the first quantitative report of positional behavior within Aves, presenting scan sampling data from a colony of Monk Parakeets (*Myiopsitta monachus*) from Brooklyn, New York City. Each scan recorded locomotor and postural behavior and information about weather condition, temperature, and substrate properties (e.g., type, size, orientation). A distinction was also recorded between natural and artificial substrates. Parrots exhibited a strong preference for small and terminal branches, a selection which may reflect targeted foraging of new fruit growth and leaf-buds. We further observed that the gait transition from walking to sidling appears primarily driven by substrate size, with the former preferred on the ground and on large, broad substrates and the latter used to navigate smaller branches. Finally, we observed an increase in locomotor diversity on artificial versus naturally occurring substrates. This demonstrates the importance of a flexible behavioral repertoire in facilitating a successful transition towards an urban landscape in introduced species.

Keywords: arboreal; locomotion; posture; Psittaciformes; Psittacidae; urban; exotic species; introduced species
1. Introduction

The use of arboreal substrates presents many challenges, reflecting the difficulties of navigating a complex three-dimensional environment using substrates of varying orientation, size, texture, compliance, and gap-distance [1–4]. Understanding how arboreal animals successfully negotiate such conditions has been the focus of numerous field- [5–8] and laboratory-based [9–13] studies for two primary reasons. First, it is well-established that humans derive from arboreal primates, and unravelling the selective pressures of our ancestors contextualizes our own evolutionary history [1,14,15]. Second, anatomically, modern humans are not well-adapted to arboreal life. Even individuals that regularly ascend trees for sustenance [16] perform poorly when compared to non-human primates. While the fundamental biomechanics of arboreal locomotion are well-established, there remain critical shortcomings to our analyses of arboreal behaviors, most notably with regards to taxonomic sampling of which, with limited exceptions [7,8,17–19], has been dominated by observational studies of primates (see [20] for a review). This sampling bias raises the question of whether observed behaviors associated with arboreal locomotion are indicative of tetrapods in general, mammals, or just primates.

Arboreal locomotion has evolved numerous times across tetrapods, and some lineages have become highly specialized for life in the trees [3,21–25]. Among birds, the parrots (Order: Psittaciformes) are a primarily arboreal lineage [26–28] that have evolved numerous anatomical features well-known to be associated with arboreal locomotion [29–31]. Briefly, these include long zygodactylous digits for grasping around supports [30,32], a distal elongation of the penultimate phalanx associated with a shortening of the proximal phalanges [32–35] that has been posited to increase grasping force [32,36], high mobility at the hip joint, and digital and tarsometatarsal pads that are textured and highly sensitive [37]. Further, Psittaciformes tend to have relatively short tarsometatarsi compared to other avian species [31]. This morphology serves to reduce limb length, allowing the animal to maintain closer contact to the substrate, thereby decreasing the gravitational pitching moment during climbing and rolling torques on thin arboreal substrates [38]. Parrots have also co-opted the feeding system and neck musculature to function as a third limb during climbing, an exaptation completely unique to the order [37,39]. Despite the well-known arboreal tendencies of parrots, there remains limited information about the way these animals exploit their environment in terms of locomotion and postures. Previous studies have been largely anecdotal in nature [40,41], and classifications of their locomotion and posture have been grossly categorical (e.g., “arboreal” and “perching”) [30,34].

Positional behaviors comprise the entirety of animals’ locomotion (i.e., behaviors that involve movement of the center of mass) and posture (i.e., behaviors that involve no movement of the center of mass) [42]. Often, these positional behaviors are paired with information about substrate characteristics (e.g., orientation, diameter, texture, height) and frequency to gain an ecological perspective of when and why an animal utilizes a particular behavior [5,6,42]. Traditionally, positional behaviors serve a corollary role in studies of evolutionary morphology. Studies of this sort usually proceed by noting salient differences in behavior and morphology in two or more species, establishing form/function relationships between the two, and attempting to explain the relationships using arguments derived from biomechanical principles [8,43–45]. Success in such an endeavor depends on the quality of both morphological and behavioral datasets. As such, the lack of detailed information hampers efforts to assess the evolutionary importance of behaviors and to construct realistic locomotor groupings or classifications [46]. More recently, gaining an understanding of positional behavior has been critical for conservation efforts, as such information is essential for focusing on the protection of critical habitats and for establishing naturalistic environments in zoological institutions and rehabilitation facilities [47]. Taken together, positional behavior is therefore an important component of the behavioral repertoire of a species.

This study represents the first analysis of positional behavior in wild parrots. Our study population comprises an introduced population of Monk Parakeets (Myiopsitta...
Animals Parakeets (Psittacidae: *Myiopsitta monachus*), are small (body length of 30 cm, wingspan of 48 cm, and body mass of 100 g), brightly colored green parrots with a greyish breast, greenish-yellow abdomen, long tail feathers, and zygodactylous feet [30,60]. Like most Psittaciformes, they are long-lived, with estimates ranging anywhere from 20 to 30 years [48,60]. Monk Parakeets usually feed on a variety of seeds, fruits, blossoms, leaf buds, tree parts, grasses, and insects [48,61,62]. Native to temperate and subtropical regions of Argentina and surrounding countries, the Monk Parakeets have become one of the most successful introduced species [55]. Self-sustaining introduced populations have become established throughout Europe and the United States [56,63,64]. The species adapts well to urban areas and has established populations in large cities, such as Miami, Chicago, and New York [48,55,61]. Part of the species’ ability to withstand harsh temperate conditions has been attributed to their nest-building behavior. Unlike most parrots that nest in tree hollows, Monk Parakeets build large stick-nests that house large communal colonies. The external heat radiating off each individual paired with the insulating properties of the nest allows the animals to maintain a safe core body temperature, even in freezing conditions [55,56,61,63,65].

2.2. Study Site

All data were collected from Green-Wood Cemetery (25th Street, Brooklyn, NY, 11232; N 40.65811, W 73.99460), which is a 193.4 ha cemetery in the western portion of Brooklyn, New York City. The Monk Parakeets have established a large nest situated among the gothic architecture of the main gate (Figure 1). Current surveys estimate approximately 36–50 individuals at Green-Wood Cemetery [48].

2.3. Data Collection

Behavioral observations were conducted whenever possible over 101 days between 17 January and 31 June 2021. The decision to end the study was based on two factors. First, by sampling from January through the end of June, we were able to capture the entire range of temperature extremes usually reported in New York State [66]. Second, the last novel behavior was observed on 21 March 2021, meaning there was an additional three months
of sampling where no new behaviors were recorded. As such, we are confident our study captured the entire range of positional diversity of Monk Parakeets at this study site.

Figure 1. A large population of introduced Monk Parakeets (*Myiopsitta monachus*) have established a nest in Green-Wood Cemetery, Brooklyn, NY.

We developed an ethogram (Table 1) for this study based on Hunt and colleagues [5], Dilger [40], Brockway [41], and a short (1–15 January 2021) ad libitum sampling session by the principal investigator. We only observed one behavior (i.e., uprighting) not on the original ethogram during the formal sampling period. Prior to the initiation of the study, all observers conducted training sessions (3 in total, consisting of 5–10 investigators each) with the principal investigator to gain familiarity and confidence in behavioral classification and sampling protocols. These training sessions were also used as an opportunity to assess the potential of interobserver error in the sampling protocol (see the Section 2.4 below).

Table 1. Ethogram for this study based on Hunt and colleagues [5], Dilger [40], Brockway [41], and a short (1–15 January 2021) ad libitum sampling session by the principal investigator.

| Positional Behavior | Description |
|---------------------|-------------|
| **Locomotion**      |             |
| Flight              | Locomotion in which no body part touches a substrate. The individual’s wings are slightly angled which allows them to deflect the air downward to generate lift. The trunk is held in a pronograde position. |
| Hover               | The wings are extended and flexed rapidly such that flight is achieved without significant movement accomplished in any direction. |
| Flight_short_interbranch | With a starting position of resting upright on a branch, this is a fast but short locomotion propelled by flapping the wings, which fully supports the body weight as the bird settles on another nearby branch. This behavior is differentiated from Leap_wing-assisted based on the relatively long distances between subsequent branches. |
| Bridge              | A torso pronograde gap-closing movement by anchoring the feet on a substrate while using the beak to reach across a gap to take hold of another substrate. Grip is retained until a secure position is established on the other side, then pulls the body across the open space to the substrate on the other side. |
Table 1. Cont.

| Positional Behavior       | Description                                                                                                                                                                                                                                                                                                                                 |
|---------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Climb                     | With an orthograde trunk orientation, forward motion is achieved as each hindlimb asynchronously protracts then retracts, allowing each foot to make alternating contact with the substrate. The weight-bearing is entirely on the hindlimbs without the support of the beak or wings. The feet may be flexed or extended depending on substrate size. The tail may be used as a prop. |
| Climb_beak-assisted       | With an orthograde trunk orientation, upward motion is achieved as each hindlimb asynchronously protracts then retracts, allowing each foot to make alternating contact with the substrate. Simultaneously, the neck is stretched, allowing the beak to grasp a substrate such that the weight-bearing is on the hindlimbs and the beak. After upward movement is achieved, the neck retracts and the beak releases its grasp of the substrate. The feet may be flexed or extended depending on the substrate size. The tail may be used as a prop. |
| Climb_wing-assisted       | With an orthograde trunk orientation, upward motion is achieved as each hindlimb asynchronously protracts then retracts, allowing each foot to make alternating contact with the substrate. Simultaneously, the wings are flapped. The torso is held pronograde to the substrate. The feet may be flexed or extended depending on the substrate size. The tail may be used as a prop. |
| Climb_wingbeak-assisted   | With an orthograde trunk orientation, upward motion is achieved as each hindlimb asynchronously protracts then retracts, allowing each foot to make alternating contact with the substrate. Simultaneously, the neck is stretched, allowing the beak to grasp a substrate, and at least one wing is flapped. After upward movement is achieved, the neck retracts and the beak releases its grasp of the substrate. The feet may be flexed or extended depending on the substrate size. The tail may be used as a prop. |
| Descent                   | With an orthograde trunk orientation, a head-first downward movement is achieved as each hindlimb asynchronously protracts then retracts, allowing each foot to make alternating contact with the substrate.                                                                                                                                 |
| Descent_beak-assisted     | With an orthograde trunk orientation, a head-first downward movement on the substrate is achieved as each hindlimb asynchronously protracts then retracts, allowing each foot to make alternating contact with the substrate. Simultaneously, the beak grasps the substrate such that the weight-bearing is on the beak and the hindlimbs. After downward movement is achieved, the neck retracts and the beak releases its grasp of the substrate. |
| Descent_wing-assisted     | With an orthograde trunk orientation, a head-first downward movement is achieved as each hindlimb asynchronously protracts then retracts, allowing each foot to make alternating contact with the substrate. Simultaneously, the wings are flapped. |
| Descent_wingbeak-assisted | With an orthograde trunk orientation, downward motion is achieved as each hindlimb asynchronously protracts then retracts, allowing each foot to make alternating contact with the substrate. Simultaneously, the neck is stretched, allowing the beak to grasp a substrate such that the weight-bearing is on the hindlimbs and the beak. During this time, at least one wing is flapping. After downward movement is achieved, the neck retracts and the beak releases its grasp of the substrate. |
| Hop                       | Bipedal locomotion, wherein the feet push off and land almost simultaneously on the substrate. The majority of the weight-bearing is on the hindlimbs, with no substantial support from any other body part. There is a period of free flight. As the individual contacts the substrate, the feet are in an extended position. The trunk is held horizontal. |
| Leap                      | A gap-crossing locomotion with a pronograde trunk orientation that primarily uses the hindlimbs to thrust forward. The hindlimbs and back are initially in flexed position and then are vigorously extended. There is a period of free flight until the hindlimbs land on the substrate. The wings are not involved. |
| Leap_wing-assisted        | A gap-crossing locomotion that primarily uses the hindlimbs to thrust forward. The flexed hindlimbs and back are forcefully extended with the assistance of the wings. There is an extended period of free flight until the individual grasps and lands on the substrate. The trunk is held in a pronograde position throughout the locomotion. This behavior is differentiated from Flight_short_interbranch based on the relatively short distances between subsequent branches. |
Table 1. Cont.

| Positional Behavior | Description |
|---------------------|-------------|
| Run                 | With a pronograde trunk orientation, forward motion is achieved as each hindlimb asynchronously protracts then retracts, allowing each foot to make alternating contact with the substrate. The weight-bearing is entirely on the hindlimbs without the support of another body part. The feet may be flexed or extended depending on the substrate size. |
| Sidling             | Involves sidewise progression along the perch, where one foot is moved before the other in a shuffling manner. |
| Suspensory          | Locomotion on a substrate using the hindlimbs, which are anchored around a substrate to support the full body weight, which is in an inverted position and moving along the substrate. |
| Uprighting          | From a hanging position, where the hindlimbs are anchored on a substrate with the body inverted, the bird uses its beak to propel its body weight over the branch and into an upright position. |
| Walk                | With a pronograde trunk orientation, forward motion is achieved as each hindlimb asynchronously protracts then retracts, allowing each foot to make alternating contact with the substrate. The weight-bearing is entirely on the hindlimbs without the support of another body part. The feet may be flexed or extended depending on the substrate size. |
| Posture             | |
| Cantilevering       | A postural position on a stable substrate with a near-vertical plane. Both feet are anchored and grasped firmly on the substrate as the trunk is held rigid and horizontal, and then the subject extends and reaches out. This posture is maintained for several seconds. |
| Cling               | A postural position on a substrate with a near-vertical plane. The feet are extended and claws are responsible for anchoring to the substrate and bearing most of the weight, with no significant support from other body parts as its trunk is held in a near-vertical orientation. |
| Hang                | A postural position on a substrate with a near-horizontal plane. The animal is inverted. Most of the body weight is supported by the feet grasping the substrate above the subject’s center of mass. The trunk may be pronograde or orthograde as the feet firmly grasp the substrate. |
| Perch               | Sitting upright and resting on a substrate with the hindlimbs flexed and phalanges grasping around the substrate. |
| Stand               | A postural position on a horizontal substrate. The hindlimbs support most of the weight, with no significant support from other body parts. The feet are splayed in an extended position and a pronograde trunk orientation. |

As the Monk Parakeets often travelled in large groups and distinguishing individuals was difficult, we used a five-minute interval scan sampling to collect behavioral data. Each scan lasted less than one minute. We scanned the Monk Parakeets from left to right or in a clockwise sweep to avoid potential bias toward given individuals. We recorded the behavior of all visible individuals during scanning, with no individual sampled twice. Number of individuals (mean ± standard deviation: 5.76 ± 4.32) observed per scan varied considerably. During each scan, we identified the predominant behavior of the sampled individual after observing it for five seconds. We also cataloged weather condition (i.e., sunny, overcast, snow flurries, rain), temperature, latitude and longitude, substrate type, and orientation (Table 2). For short flights or leaps between supports, the substrate was coded based on the landing site. No substrate or orientation code was scored if the animals’ feet were not in contact with the support (e.g., flight, hovering) during the five second within-scan observational period. All observers were equipped with 8 × 21 compact binoculars (BriGenius, Shenzhen, China) to aid in behavioral scoring.
Table 2. Definition and description of substrate conditions.

| Substrate Variable | Definition                                                                 |
|--------------------|---------------------------------------------------------------------------|
| Ground             | Ground and related substrates (e.g., rocks, roots, logs)                 |
| Very large         | Arboreal substrates with a diameter > 20 cm (e.g., tree trunks)          |
| Large              | Arboreal substrates with a diameter larger than the dorsoventral height of the animal but < 20 cm (e.g., tree boughs) |
| Medium             | Arboreal substrates with a diameter approximately equal to the dorsoventral height of the animal |
| Small              | Arboreal substrates with a diameter approximately equal to foot span (i.e., complete coverage by the hallux and foredigits) |
| Terminal           | Arboreal substrates with a diameter less than foot span (i.e., overlapping coverage by the hallux and foredigits) |
| Nest               | Communal stick nest                                                      |
| Artificial         | Man-made substrates (e.g., telephone wires and poles, building facades, roofs) |

Orientation

| Orientation | Definition |
|-------------|------------|
| Horizontal  | Angle between 0° and 22.5° |
| Oblique     | Angle between 22.5° and 67.5° |
| Vertical    | Angle between 67.5° and 90° |

2.4. Data Analysis

Following established methods [67, 68], we utilized pilot data collected during the training sessions (see above) to assess the possibility of interobserver error in data scoring. After the initial walkthrough of the study goals and an introduction to the ethogram and scan sampling protocols, each observer watched the nest for one hour (i.e., twelve scans) and was asked to score positional behavior, substrate type, and orientation. We used the crosstab function in MATLAB (MathWorks, Inc., Natick, MA) to calculate a $\chi^2$-statistic to assess whether the likelihood of scoring a particular positional behavior, substrate type, or orientation was influenced by the observer. We detected no significant interobserver variation in scoring with regards to locomotion (all $\chi^2$-values across three training groups ≤ 45.76; all $p$-values > 0.838), posture (all $\chi^2$-values across three training groups ≤ 6.95; all $p$-values ≥ 0.642), substrate type (all $\chi^2$-values across three training groups ≤ 42.88; all $p$-values ≥ 0.537), or orientation (all $\chi^2$-values across three training groups ≤ 4.78; all $p$-values > 0.936). Based on these results, we were satisfied that interobserver error in the main dataset (i.e., data collected after the training sessions, 101 days between 17 January and 31 June 2021) should be minimal.

To address the main dataset (i.e., data collected after the training sessions, 101 days between 17 January and 31 June 2021), we first divided the observational sample into locomotor and postural datasets. From these, we calculated proportions of different positional behaviors, substrate types, and orientation used by the animals during the sampling period. We used the crosstab function in MATLAB to conduct a series of $\chi^2$-tests to assess whether observing a particular positional behavior was influenced by substrate type, orientation, month, or weather condition. As weather condition had a low number of observations, we used Yate’s $\chi^2$-tests for these analyses.

We conducted a second analysis comparing locomotor diversity [20] between arboreal, terrestrial, and artificial substrates. Briefly, this study co-opted the Shannon–Wiener diversity index to calculate a singular measure of locomotor diversity. The Shannon–Wiener diversity index is traditionally used in the ecology literature to provide a statistically comparable metric to quantify the diversity of species composition in a specific area of interest (e.g., forest, transect, etc.) [69–72]. This study employs this same logic using the proportion of each locomotor behavior observed on a particular substrate to calculate
a substrate-specific locomotor diversity index. The locomotor diversity index (LDI) is calculated as:

\[
\text{Locomotor diversity index} = - \sum p_i \ln p_i
\]

where \(p_i\) is the proportion of a particular locomotor behavior (e.g., walking, leaping, climbing) out of the total combination of locomotor behaviors (e.g., proportion of walking + proportion of leaping + proportion of climbing). A low LDI indicates that Monk Parakeets frequently use only a few different locomotor behaviors on a particular substrate. High LDI values indicate that Monk Parakeets frequently use many different locomotor behaviors on a particular substrate.

3. Results

Across the six-month sampling period, we collected 11,246 observations comprising of 7337 postural (65.24%) and 3909 locomotor scans (34.76%) (Table S1). Throughout the study period, the temperature ranged from \(-3.89\) to \(36.11\) °C. Weather conditions varied between snow flurries (\(n = 11\) observations, 0.10%), overcast (\(n = 2954\) observations, 26.27%), rain (\(n = 539\) observations, 4.79%), and sun (\(n = 7742\) observations, 68.84%) (Table S2). In total, twenty observers contributed to the overall sample.

During non-aerial activities (i.e., excluding flight and hovering), Monk Parakeets spent a majority of their time on their large elaborate nest (\(n = 3777\) observations, 44.76%) (Figure 2 and Table S3). During non-aerial activities outside the nest, Monk Parakeets were mainly engaged in arboreal locomotion (\(n = 3424\) observations, 40.59%). Animals favored both terminal (\(n = 1670\) observations, 19.80%) and small-diameter branches (\(n = 1240\) observations, 14.70%), while the use of medium (\(n = 458\) observations, 5.43%), large (\(n = 38\) observations, 0.45%), and very large branches (\(n = 18\) observations, 0.21%) was less common. Terrestrial (\(n = 342\) observations, 4.05%) locomotion and posture and movement on artificial structures (\(n = 894\) observations, 10.60%) were also observed. During non-aerial locomotion, Monk Parakeets were primarily observed on horizontally (\(n = 5841\) observations, 69.24%) and obliquely oriented (\(n = 2163\) observations, 25.64%) supports, while the use of vertical supports (\(n = 432\) observations, 5.12%) was more limited (Table S5).

Locomotor behaviors were dominated by flight (\(n = 2639\) observations, 67.51%). However, during non-aerial activities (\(n = 1098\) observations), short flights within trees (\(n = 432\) observations, 39.34%), walking (\(n = 223\) observations, 20.31%), and sidling (\(n = 125\) observations, 11.38%) were the most common forms of locomotion, while anti-pronograde behaviors, sensu stricto Stern [14], were uncommon (bridging: \(n = 3\) observations, 0.27%; suspensory: \(n = 4\) observations, 0.36%; uprighting: \(n = 5\) observations, 0.45%). Climbing behaviors (pooled Climb, Climb_beak-assisted, and Climb_wing-assisted) accounted for 7.92% of non-aerial behaviors (\(n = 87\) observations) and were observed in generally equal proportions (Figure 3 and Table S1).

The time of year had a significant influence on locomotor behavior (\(\chi^2 = 504.62; p < 0.001\)), where walking became more common in March, April, May, and June, and leaping with the assistance of the wings, which was a fairly common locomotor behavior early in the study, was not observed after March (Figure 3). As a greater proportion of time was spent on the ground (see above), this was associated with a general increase in walking (Table S4). Weather also had a significant influence on locomotor behavior (\(\chi^2 = 252.58; p < 0.001\)), where non-aerial movements were less common during snow flurries and rain (Table S2).
Figure 2. Pie charts demonstrating the relative proportions of substrate orientation (a, b) and substrate type (c, d) collected during postural (a, c) and locomotor behaviors (b, d) from an established colony of Monk Parakeets (Myiopsitta monachus) from Brooklyn, New York City, during the study period.

Substrate had a significant influence on locomotor behavior (Figure 3; $\chi^2 = 1517; p < 0.001$). When on the nest, Monk Parakeets spent a majority of the time making short flights and then quickly landing nearby ($n = 233$ observations, 64.54%). Climbing the nest using only the hindlimbs was also common ($n = 37$ observations, 10.25%). Terrestrial locomotion was dominated by walking ($n = 169$ observations, 91.35%), but hopping ($n = 11$ observations, 5.95%) and running ($n = 5$ observations, 2.70%) were also observed. On all arboreal substrates, excluding those of large diameter, short flights to branches within the same tree were most common. On terminal branches, leaping using the wings was also quite common ($n = 67$ observations, 38.07%). On small and medium branches, sidling is the dominant locomotor mode (small: $n = 74$ observations, 35.24%; medium: $n = 29$ observations, 40.28%). Walking was also a major component of the locomotor repertoire on medium branches ($n = 12$ observations, 16.67%), and the primary form of locomotion on large branches ($n = 3$ observations, 42.86%). Climbing with the assistance of the beak was observed on all arboreal substrates (terminal: $n = 1$ observation, 0.57%; small: $n = 9$ observations, 4.29%; large: $n = 1$ observation, 14.29%; very large: $n = 4$ observations, 50.00%), excluding medium diameter branches. Movement on artificial substrates was variable, with walking being most common ($n = 25$ observations, 31.65%), followed by short flights to nearby substrates ($n = 15$ observations, 18.99%), leaping with the assistance of the wings ($n = 7$ observations, 8.86%), and equal contributions of climbing, climbing with assistance of the beak, climbing with assistance of the wings, and descent with assistance of the beak ($n = 5$ observations, 6.33%). Accordingly, locomotor diversity was greater on artificial supports (LDI = 2.14), compared to arboreal substrates (LDI = 1.78), the nest (LDI 1.35), or the ground (LDI = 0.35).

Substrate orientation significantly influenced the observed locomotor modes (Figure 3; $\chi^2 = 597.97; p < 0.001$). Across all orientations, short flights to nearby supports were common (Table S5). On horizontal substrates, walking ($n = 203$ observations, 35.74%) and sidling ($n = 76$ observations, 13.38%) comprised the greatest proportion of observed locomotor behaviors. On oblique supports, walking ($n = 20$ observations, 6.29%) and sidling ($n = 49$ observations, 15.41%) were also quite common but leaping with the assistance of the
wings ($n = 75$ observations, 23.58%) was the most frequently observed behavior. Excluding short flights to nearby supports, locomotion on vertical supports consisted primarily of climbing ($n = 37$ observations, 17.45%) and climbing with assistance of the beak ($n = 26$ observations, 12.36%).

Figure 3. Illustrations and proportions of locomotor behaviors (a) observed based on differing substrate types, orientations, weather conditions, and time of year (b), collected from an established colony of Monk Parakeets (*Myiopsitta monachus*) from Brooklyn, New York City, during the study period. The pie chart (c) illustrates the overall proportions of locomotor behaviors. The expanded pie chart illustrates proportions of non-aerial behaviors.

Postural behaviors were dominated by perching ($n = 7112$ observations, 96.93%; Table S1). The time of year had a significant influence on postural behavior ($\chi^2 = 167.25; p < 0.001$; Table S4), where standing increased significantly in May and June. As a greater proportion of time was spent on the ground (see above), this was associated with a general increase in standing. Weather also had a significant influence on postural behavior ($\chi^2 = 115.15; p < 0.001$), where standing and clinging were relatively more common in sunny and rainy conditions, respectively (Figure 4 and Table S2).
Based on our definitions of support size, we anticipated that Monk Parakeets would primarily utilize small-diameter substrates, as these would allow the foot to completely wrap around the substrate with no overlap between the toes and theoretically maximize grip force potential [53]. While the use of small-diameter supports was common, terminal branches were the most commonly used substrates by Monk Parakeets. Thus, Prediction 3 is partially supported. As discussed thoroughly in the primate literature, terminal branches allow animals new foraging opportunities as these are the primary site for new fruit growth.
and leaf-buds [1,74,75], which have been shown to be a favorite food item of the Monk Parakeets in Brooklyn [48]. However, movement on terminal branches is dangerous as such slight substrates may oscillate under the weight of the animal, increasing the risk of becoming unbalanced and falling [1,22,76]. While some arboreal species mitigate these concerns by adopting suspensory positional behavior [19,22], such a strategy was rarely observed among Monk Parakeets (Table S1). Instead, Monk Parakeets, most likely due to their small body mass, like many similarly sized primates [43,77], use their grasping feet to counteract toppling forces and maintain balance on the branch [54,78]. However, the diameter of terminal branches means the toes demonstrate substantial overlap between each other and arguably indicates that the digital flexors are primarily active on the early, ascending portion of a Hill-type length–tension curve [79]. While it is possible that the passive tendon-locking mechanism described above may supersede these considerations, there are insufficient data available to interpret how substrate diameter influences grasping ability in birds. Future work following Sustaita and Hertel [53] and Ward and colleagues [80] would help to address this gap.

Flight was the most observed form of locomotion among Monk Parakeets. Even when moving within a tree or around the nest, flight and hovering remained the primary form of movement. Thus, this aspect of Prediction 2 is supported. This finding is not surprising considering the lower cost of transport associated with flight compared to terrestrial locomotion [49–51]. While the energetic cost associated with hindlimb-based locomotion in parrots is unknown, it is likely that these costs are quite high. Compared to most other birds, parrots have relatively short tarsometatarsi that reduce the overall length of the limb [31], thereby requiring increased stride frequencies during locomotion [81,82]. Further, like primates, parrots have large feet with considerable mass positioned distally along the hindlimb [83,84]. Higher stride frequency and distal weight distribution are both associated with increased locomotor costs [82–85]. As such, it seems probable that the locomotor behavior of Monk Parakeets may be explained through considerations of the overall energetic budget. Future work should explore how differences in body size, a factor known to influence locomotor costs, influence the positional behaviors between parrot species and in other birds.

Walking and sidling were the most observed forms of hindlimb-driven locomotion. Both walking and sidling involved the alternating left/right patterning of the limbs. This finding is in accordance with Provini and Höfling [31], who note that parrots violate general patterns in avian locomotion by being a relatively small-bodied arboreal lineage that does not hop. While we did observe Monk Parakeets hop occasionally, this movement was rare (n = 28 observations) and most often observed on the ground. Thus, this aspect of Prediction 2 was rejected. The presence of hopping in Monk Parakeets rejects the idea of a possible neuromuscular constraint in the lineage [31] and highlights a gap in our understanding of why certain avian lineages hop while others do not. Further, it is unclear what factors trigger parrots to adopt hopping gaits.

The occurrence of walking and sidling was largely driven by substrate diameter. Generally, walking was observed on terrestrial and large-diameter substrates, while sidling became more common as substrate size decreased. To our knowledge, sidling was first described by Dilger [40], and involves sidewise progression along the perch, where one foot is moved before the other in a shuffling manner. Originally, Dilger [40] proposed that the use of sidling was reserved for slow speed progression, but Brockway [41] noted that sidling was adopted even at high speeds in budgerigars (Melopsittacus undulatus). Based on the marked shift between walking and sidling dependent on substrate diameter, a finding in accordance with Young et al. [86] in Rosy-faced Lovebirds (Agapornis roseicollis), we propose that the use of sidling represents a solution to an anatomical constraint. Compared to most tetrapod lineages, birds have greatly reduced the number of tarsal and metatarsal bones into a singular tarsometatarsus [87]. Further, articular surfaces with the knee proximally and phalanges distally are characterized with noticeable deep trochleae [28,87]. As such, the ability to generate parasagittal movements (e.g., inversion and eversion) at the distal
joints of the hindlimb is limited. By adopting sidling, birds can position their grasping feet perpendicular to the long axis of the substrate, thus ensuring grasping abilities on thin arboreal supports where inversion/eversion is not possible [86].

Contrary to Prediction 2, we did not observe a large proportion of beak-assisted climbing. Parrots are unique among tetrapods as they have co-opted the feeding system and neck musculature into an effective third limb to both propel and power the body during vertical ascent. Climbing with the assistance of the beak comprised 3.28% of substrate-based locomotion (Table S1) and was largely restricted to movement on vertical supports (Table S5). This is consistent with recent laboratory experiments that the use of the head as a third limb is only present at very steep inclines (e.g., 67.5°), and does not become ubiquitous until vertical ascent [39]. Monk Parakeets infrequently utilized vertical supports and were rarely observed on very large substrates (e.g., tree trunks). This is in stark contrast to the movements of woodpeckers, nuthatches, and treecreepers [21,29,38,88]. Webster and colleagues [4] suggested that the use of the beak as a third limb is more common among larger-bodied Psittaciformes, but this suggestion is anecdotal in nature and further work on the positional behavior of additional parrot species is required to verify this claim.

4.1. Positional Behavior in an Urban Environment

Urban habitats are dramatically modified from their natural state, creating unique challenges and selection pressures for organisms that reside in them. Winchell and colleagues [58] also noted that arboreal species are particularly impacted because the anthropogenic structures with which they interact differ from trees in structural, material, and surface properties. While multiple studies have demonstrated that locomotor behaviors and performance change in response to urban landscapes [59], to our knowledge this is the first dataset reporting how overall positional behavior is influenced. Ideally, we would directly compare the positional behavior of Monk Parakeets in their natural range in juxtaposition to the current dataset. In the absence of such data, we compared the locomotor behavior of Monk Parakeets on artificial substrates compared to natural substrates (i.e., terrestrial and arboreal substrates and the nest). Monk Parakeets were quite adept at movement on artificial substrates, and were often observed moving on telephone wires, concrete (e.g., building facades), marble (e.g., gravestones), and metal (e.g., vehicles). On these artificial structures, Monk Parakeets adopted a more diverse locomotor repertoire with less reliance on a single locomotor modality. Accordingly, locomotor diversity was higher when moving on artificial structures compared to natural substrates. Thus, Prediction 4 was supported. The data within this study are insufficient to determine why these changes occur between natural and artificial substrates but highlight the importance of behavioral flexibility in the locomotor behavior of urban species. It is unclear whether behavioral flexibility is the response to movement in urban environments, or whether it is an important trait of species able to colonize and persist in urban areas.

4.2. Limitations

A study of this nature is faced with a number of limitations that must be addressed. Most notable was the multiple observers that may introduce interobserver error in scoring. While training sessions and subsequent interobserver statistical analyses (see Section 3) suggest these effects to be minimal, such error cannot be removed entirely. This manuscript represents the culmination of student-led research aimed at introducing animal behavior research to a broader range of individuals and increasing inclusivity in the research process. As such, it was not possible to reduce the number of observers without compromising inclusivity and broadening participation in science. Another limitation is in the use of scan sampling, which was required because of the flighty nature of the animals and the inability to differentiate individuals. Scan sampling provides an unbiased assessment of the activity budget of an animal but suffers from missing rare or uncommon behaviors. This is evidenced by the almost exclusive use of perching and flight. However, these limitations do not detract from the overall findings and interpretations of the manuscript.
as we identified no new behaviors after the first three months of sampling (see above). Further, as the goal was to capture the range and frequency of certain behaviors, scan sampling was most appropriate [89]. Next, the study started in January. During this month, the observers were just becoming acquainted with the study site and animals. Accordingly, there are fewer observations for January than other months. This means we may have missed information about how Monk Parakeets behave in cold weather conditions. As the ability for these parrots to survive in these conditions is critical for understanding their success as introduced species, we hope future studies will focus more sampling effort on this critical time period. Finally, the most rigorous assessments of positional behavior are able to account for a relative abundance of substrates [19,90]. However, these types of analyses are usually limited to captive settings where substrate conditions can be manipulated or appropriately counted. Such relative abundance analyses were not possible within this study.

5. Conclusions

This study revealed several novel and potentially valuable insights into the locomotor repertoire of Monk Parakeets. Firstly, we determined that these birds exhibit a marked preference for small and terminal branches when perching arboreally, a selection which may reflect the ability of the digital flexor musculature to most effectively develop high grip forces and engage a passive tendon-locking mechanism in these tightly curled pedal postures. The preference for small and terminal branches is similar to what is observed in primates. In this way, Monk Parakeets likely demonstrate a myriad of anatomical characteristics similar to primates that allow for movement on such precarious substrates. Further work is required to assess if and how larger parrot species navigate the challenges of terminal branch movement. Secondly, we report that the gait transition from walking to sidling appears primarily driven by substrate size, with the former preferred on the ground and on large, broad substrates and the latter used to navigate smaller branches. Such a gait pattern has been poorly investigated in the literature and likely represents a novel behavioral solution to an anatomical constraint [86]. If so, the presence of sidling is likely present in many avian species, or at least those that spend significant time in the trees [31]. Thirdly, contrary to our initial prediction, climbing with the use of the beak was relatively rare and was primarily influenced by the orientation of the substrate. Perhaps this is not surprising as the use of the beak, at least in Rosy-faced Lovebirds [39], does not become common until orientations >67.5°. Thus far, beak use has only been investigated in small-bodied parrot species (this study, and [40–42]) and additional sampling effort across Psittaciformes is required to understand the conditions that elicit tri-pedal gaits in parrots. Finally, we observed an increase in locomotor diversity on artificial versus naturally occurring substrates, a shift which underscores the necessity of behavioral flexibility within introduced urban species to successfully navigate a novel, variable, and often challenging environment.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/ani12182372/s1, Table S1: Overall percentage (%) of positional behaviors collected from an established colony of Monk Parakeets (*Myiopsitta monachus*) from Brooklyn, New York City during the study period. Percentages of locomotor repertoire calculated for all behaviors and for non-aerial (i.e., excluding flight and hovering) behaviors; Table S2: Overall percentage (%) of weather conditions and the associated positional behaviors collected from an established colony of Monk Parakeets (*Myiopsitta monachus*) from Brooklyn, New York City during the study period. No substrate or orientation code was scored if the animals’ feet were not in contact with the support (e.g., flight, hovering); Table S3: Overall percentage (%) of substrate type and the associated positional behaviors observed on these substrates collected from an established colony of Monk Parakeets (*Myiopsitta monachus*) from Brooklyn, New York City during the study period. No substrate or orientation code was scored if the animals’ feet were not in contact with the support (e.g., flight, hovering); Table S4: Overall percentage (%) of sampling effort during the study period and the associated positional behaviors observed during each month collected from an established
A colony of Monk Parakeets (*Myiopsitta monachus*) from Brooklyn, New York City during the study period. No substrate or orientation code was scored if the animals’ feet were not in contact with the support (e.g., flight, hovering); Table S5: Overall percentage (%) of substrate-orientation and the associated positional behaviors observed on these orientations collected from an established colony of Monk Parakeets (*Myiopsitta monachus*) from Brooklyn, New York City during the study period. No substrate or orientation code was scored if the animals’ feet were not in contact with the support (e.g., flight, hovering).

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**References**

1. Cartmill, M. Arboreal Adaptations and the Origin of the Order Primates. In *The Functional and Evolutionary Biology of Primates*; Routledge: Oxford, UK, 1972; ISBN 978-1-315-13212-9.
2. Cartmill, M. Pads and Claws in Arboreal Locomotion. *Primate Locomot.* 1974, 1, 45–83. [CrossRef]
3. Preuschoft, H. What Does “Arboreal Locomotion” Mean Exactly and What Are the Relationships between “Climbing”, Environment and Morphology? *Z. Morphol. Anthrop.* 2002, 2, 171–188. [CrossRef]
4. Webster, C.; Jusufi, A.; Liu, D. A Comparative Survey of Climbing Robots and Arboreal Animals in Scaling Complex Environments. In Proceedings of the International Symposium on Robotics and Mechatronic, Rome, Italy, 16–19 February 2019; Yang, R., Takeda, Y., Zhang, C., Fang, G., Eds.; Springer International Publishing: Cham, Switzerland, 2019; pp. 31–45.
5. Hunt, K.D.; Cant, J.G.H.; Gebo, D.L.; Rose, M.D.; Walker, S.E.; Youlatos, D. Standardized descriptions of primate locomotor and postural modes. *Primates* 1996, 37, 363–387. [CrossRef]
6. Youlatos, D. Comparative locomotion of six sympatric primates in Ecuador. *Ann. Sci. Nat. Zool. Biol. Anim.* 1999, 20, 161–168. [CrossRef]
7. Youlatos, D. Locomotor and Postural Behavior of *Sciurus igniventris* and *Microsciurus flaviventer* (Rodentia, Sciuridae) in Eastern Ecuador. *Mammalia* 1999, 63, 405–416. [CrossRef]
8. Ercoli, M.D.; Youlatos, D. Integrating locomotion, postures and morphology: The case of the tayra, *Eira barbara* (Carnivora, Mustelidae). *Mamm. Biol.* 2016, 81, 464–476. [CrossRef]
9. Young, J.W. Substrate determines asymmetrical gait dynamics in marmosets (*Callithrix jacchus*) and squirrel monkeys (*Saimiri boliviensis*). *Am. J. Phys. Anthr.* 2009, 138, 403–420. [CrossRef]
10. Young, J.W. Gait selection and the ontogeny of quadrupedal walking in squirrel monkeys (*Saimiri boliviensis*). *Am. J. Phys. Anthr.* 2012, 147, 580–592. [CrossRef]
11. Young, J.W. Ontogeny of limb force distribution in squirrel monkeys (Saimiri boliviensis): Insights into the mechanical bases of primate hind limb dominance. *J. Hum. Evol.* 2012, 62, 473–485. [CrossRef]

12. Granatosky, M.C.; Tripp, C.H.; Schmitt, D. Gait kinetics of above- and below-branch quadrupedal locomotion in lemurid primates. *J. Exp. Biol.* 2016, 219, 53–63. [CrossRef]

13. Granatosky, M.C. Forelimb and hindlimb loading patterns during quadrupedal locomotion in the large flying fox (Pteropus vampyrus) and common vampire bat (Desmodus rotundus). *J. Zool.* 2018, 305, 63–72. [CrossRef]

14. Stern, J. Before Bipedality. *Yearb. Phys. Anthropol.* 1975, 19, 59–68.

15. Chester, S.G.B.; Williamson, T.E.; Bloch, J.I.; Silcox, M.T.; Sargis, E.J. Oldest skeleton of a plesiadapiform provides additional evidence for an exclusively arboreal radiation of stem primates in the Paleocene. *R. Soc. Open Sci.* 2017, 4, 170329. [CrossRef] [PubMed]

16. Venkataraman, V.V.; Kraft, T.S.; Dominy, N.J. Tree climbing and human evolution. *Proc. Natl. Acad. Sci. USA* 2012, 110, 1237–1242. [CrossRef]

17. Essner, R.L. Three-dimensional launch kinematics in leaping, parachuting and gliding squirrels. *J. Exp. Biol.* 2002, 205, 2469–2477. [CrossRef]

18. Essner, R.L. Morphology, locomotor behaviour and microhabitat use in North American squirrels. *J. Zool.* 2007, 272, 101–109. [CrossRef]

19. Granatosky, M.C.; Karantanis, N.E.; Rychlik, L.; Youlatos, D. A suspensory way of life: Integrating locomotion, postures, limb movements, and forces in two-toed sloths Choloepus didactylus (Megalonychidae, Folivora, Pilosa). *J. Exp. Zool. Part A: Ecol. Integr. Physiol.* 2018, 329, 570–588. [CrossRef]

20. Granatosky, M.C. A Review of Locomotor Diversity in Mammals with Analyses Exploring the Influence of Substrate-Use, Body Mass, and Intermembral Index in Primates. *J. Zool.* 2018, 306, 207–216. [CrossRef]

21. Bock, W.J.; Miller, W.D. The Scansorial Foot of the Woodpeckers, with Comments on the Evolution of Perching and Climbing Feet in Birds. In *American Museum Novitates. No. 1931*; Evolution of perching and climbing feet in birds; American Museum of Natural History: New York, NY, USA, 1959.

22. Cartmill, M. Climbing. In *Functional Vertebrate Morphology*; Hildebrand, M., Bramble, D.M., Liem, K.F., Wake, D.B., Eds.; Belknap: Cambridge, MA, USA, 1985; pp. 73–88.

23. Fischer, M.S.; Krause, C.; Lilje, K.E. Evolution of chameleon locomotion, or how to become arboreal as a reptile. *Zoology* 2010, 113, 67–74. [CrossRef]

24. Granatosky, M.C. Primate Locomotion. In *Encyclopedia of Animal Cognition and Behavior*; Vonk, J., Shackelford, T., Eds.; Springer International Publishing: Cham, Switzerland, 2020; pp. 1–7. ISBN 978-3-319-47829-6.

25. Aretz, J.M.; Brown, C.E.; Deban, S.M. Vertical locomotion in the arboreal salamander *Aneides vagrans*. *J. Zool.* 2021, 316, 72–79. [CrossRef]

26. Ksepka, D.T.; Clarke, J.A.; Grande, L. Stem Parrots (Aves, Halcyornithidae) from the Green River Formation and a Combined Phylogeny of Pan-Psittaciformes. *J. Paleontol.* 2011, 85, 835–852. [CrossRef]

27. Ksepka, D.T.; Clarke, J.A. A new stem parrot from the Green River Formation and the complex evolution of the grasping foot in Pan-Psittaciformes. *J. Verteb. Paleontol.* 2012, 32, 395–406. [CrossRef]

28. Zelenkov, N.V. The first fossil parrot (Aves, Psittaciformes) from Siberia and its implications for the historical biogeography of Psittaciiformes. *Biol. Lett.* 2016, 12, 20160717. [CrossRef] [PubMed]

29. Richardson, F. Adaptive Modifications for Tree-Trunk Foraging in Birds; University of California Press: Berkeley, CA, USA, 1942; Volume 46.

30. Abourachid, A.; Fabre, A.-C.; Cornette, R.; Höfling, E. Foot shape in arboreal birds: Two morphological patterns for the same pincer-like tool. *J. Anat.* 2017, 231, 1–11. [CrossRef]

31. Provisi, P.; Höfling, E. To Hop or Not to Hop? The Answer Is in the Bird Trees. *Syst. Biol.* 2020, 69, 962–972. [CrossRef]

32. Backus, S.B.; Sustaita, D.; Odhner, L.U.; Dollar, A.M. Mechanical analysis of avian feet: Multiarticular muscles in grasping and perching. *R. Soc. Open Sci.* 2015, 2, 140350. [CrossRef]

33. Hamrick, M.W.; Rosenman, B.A.; Brush, J.A. Phalangeal morphology of the Paromomyidae (??Primates, Plesiadapiformes): The evidence for gliding behavior reconsidered. *Am. J. Phys. Anthropol.* 1999, 109, 397–413. [CrossRef]

34. Hopson, J. Ecomorphology of Avian and Nonavian Theropod Phalangeal Proportions: Implications for the Arboreal versus Terrestrial Origin of Bird Flight. In *New Perspectives on the Origin and Early Evolution of Birds: Proceedings of the International Symposium in Honor of John H. Ostrom*; Gauthier, J., Gall, L.F., Eds.; Peabody Museum of Natural History, Yale University: New Haven, CT, USA, 2007; pp. 207–216. [CrossRef]

35. Kavanagh, K.D.; Shoval, O.; Winslow, B.B.; Alon, U.; Leary, B.P.; Kan, A.; Tabin, C.J. Developmental bias in the evolution of phalanges. *Proc. Natl. Acad. Sci. USA* 2013, 110, 18190–18195. [CrossRef]

36. Struble, M.K.; Gardner, J.; Gibb, A.C. Grasping Behavior in Birds Drives Pedal Adaptations. *Integr. Comp. Biol.* 2020, S60, E226.

37. Reader, L.L.; Carrier, D.R.; Goller, F.; Isaacs, M.R.; Crisp, A.M.; Barnes, C.J.; Lee, D.V. Climbing parrots achieve pitch stability using forces and free moments produced by axial-appendicular couples. *J. Exp. Biol.* 2022, 225. [CrossRef]

38. Norberg, U.M. Functional osteology and myology of the wing of the dog-faced bat Rousettus aegyptiacus (È.O. Geoffroy) (Mammalia, Chiroptera). *Zomorphology* 1972, 73, 1–44. [CrossRef]
39. Young, M.W.; Dickinson, E.; Flaim, N.D.; Granatoksky, M.C. Overcoming a 'forbidden phenotype': The parrot’s head supports, propels and powers tripod locomotion. *Proc. R. Soc. B Biol. Sci.* **2022**, *289*, 2020245. [CrossRef] [PubMed]

40. Dilger, W.C. The Comparative Ethology of the African Parrot Genus Agapornis. *Z. Tierpsychol.* **1960**, *17*, 649–685. [CrossRef]

41. Brockway, B.F. Ethological Studies of the Badgerigar (Melopsittacus Undulatus): Non-Reproductive Behavior. *Behaviour* **1964**, *22*, 193–222. [CrossRef]

42. Prost, J.H. A Definitional System for the Classification of Primate Locomotion. *Am. Anthr.* **1965**, *67*, 1198–1214. [CrossRef]

43. Goodenberger, K.E.; Boyer, D.M.; Orr, C.M.; Jacobs, R.L.; Femiani, J.C.; Patel, B.A. Functional morphology of the hallucal metatarsal with implications for inferring grasping ability in extinct primates. *Am. J. Phys. Anthr.* **2014**, *156*, 327–348. [CrossRef]

44. Fabre, A.-C.; Marigó, J.; Granatoksky, M.C.; Schmitt, D. Functional associations between support use and forelimb shape in strepsirrhines and their relevance to inferring locomotor behavior in early primates. *J. Hum. Evol.* **2017**, *108*, 11–30. [CrossRef]

45. Youlatos, D.; He, G.; Guo, S.; Li, B. Positional behavior, habitat use, and forelimb morphology of P.*

46. del Hoyo, J.; Elliott, A.; Sargatal, J.; Christie, D.A. (Eds.) *Handbook of the Birds of the World* Volume 1–16. Lynx Edicions: Barcelona, Spain, 2010; ISBN 978-84-9874-237-X.

47. South, J.M.; Pruett-Jones, S. Patterns of Flock Size, Diet, and Vigilance of Naturalized Monk Parakeets in Hyde Park, Chicago. *Am. J. Phys. Anthr.* **2005**, *131*, 261–271. [CrossRef] [PubMed]

48. Baldwin, S.C. Greater Efficiency Is Achieved by Birds, Fish—and Bicyclists. *Science* **1972**, *180*, 222–228. [CrossRef]

49. Alexander, R.M. The Merits and Implications of Travel by Swimming, Flying, and Running. *Science* **1977**, *198*, 52–58. [CrossRef]

50. Proctor, B.F. Ethological Studies of the Budgerigar (Melopsittacus Undulatus): Non-Reproductive Behavior. *Z. Tierpsychol.* **1964**, *22*, 193–222. [CrossRef]

51. Alexander, R.M. The Merits and Implications of Travel by Swimming, Flying, and Running. *Science* **1972**, *180*, 52–58. [CrossRef]

52. Olkowicz, S.; Kocourek, M.; Luˇ can, R.K.; Porteš, M.; Fitch, W.T.; Herculano-Houzel, S.; Nˇ emec, P. Birds have primate-like numbers of neurons in the forebrain. *Proc. Natl. Acad. Sci. USA* **2016**, *113*, 7255–7260. [CrossRef]

53. Sustaita, D.; Hertel, F. In vivo bite and grip forces, morphology and prey-killing behavior of North American accipiters (Accipitr- dae) and falcons (Falconidae). *J. Exp. Biol.* **2010**, *213*, 2617–2628. [CrossRef] [PubMed]

54. Sustaita, D.; Poudyebat, E.; Manzano, A.; Abdala, V.; Hertel, F.; Herrel, A. Getting a grip on tetrapod grasping: Form, function, and evolution. *Biol. Rev.* **2013**, *88*, 380–405. [CrossRef] [PubMed]

55. Preston, C.E.C.; Pruett-Jones, S.; Eberhard, J.R. Monk Parakeets as a Globally Naturalized Specie. In *Primate Locomotion: Recent Advances*; Strasser, E., Fleagle, J.G., Rosenberger, A.L., McHenry, H.M., Eds.; Springer US: Boston, MA, USA, 1998; pp. 5–29. ISBN 978-1-4899-0092-0.

56. Pruett-Jones, S.; Tarvin, K.A. Monk Parakeets in the United States: Population Growth and Regional Patterns of Distribution. In *Primate Locomotion: Recent Advances*; Strasser, E., Fleagle, J.G., Rosenberger, A.L., McHenry, H.M., Eds.; Springer US: Boston, MA, USA, 1998; pp. 5–29. ISBN 978-1-4899-0092-0.

57. Clemente, C.J.; Dick, T.; Wheatley, R.; Gaschik, J.; Nasir, A.F.A.A.; Cameron, S.F.; Wilson, R. Moving in complex environments: A biomechanical analysis of locomotion on inclined and narrow substrates. *J. Exp. Biol.* **2019**, *222*, 189654. [CrossRef]

58. Baldwin, S.C. *The Brooklyn Parrots FAQ: All about the Wild Monk Parakeets of Brooklyn, NY*, 1st ed.; Stephen C. Baldwin: New York, NY, USA, 2015.

59. Schmidt-Nielsen, K. Locomotion: Energy Cost of Swimming, Flying, and Running. *Science* **1972**, *177*, 222–228. [CrossRef]

60. Tucker, V.A. The Energetic Cost of Moving About: Walking and Running Are Extremely Efficient Forms of Locomotion. Much Greater Efficiency Is Achieved by Birds, Fish—and Bicyclists. *Am. Sci.* **1975**, *63*, 413–419.

61. Alexander, R.M. The Merits and Implications of Travel by Swimming, Flight, Running for Animals of Different Sizes. *Integr. Comp. Biol.* **2002**, *42*, 1060–1064. [CrossRef] [PubMed]

62. Olkowicz, S.; Kocourek, M.; Luˇ can, R.K.; Porteš, M.; Fitch, W.T.; Herculano-Houzel, S.; Nˇ emec, P. Birds have primate-like numbers of neurons in the forebrain. *Proc. Natl. Acad. Sci. USA* **2016**, *113*, 7255–7260. [CrossRef]

63. Sustaita, D.; Hertel, F. In vivo bite and grip forces, morphology and prey-killing behavior of North American accipiters (Accipitr- dae) and falcons (Falconidae). *J. Exp. Biol.* **2010**, *213*, 2617–2628. [CrossRef] [PubMed]

64. Sustaita, D.; Poudyebat, E.; Manzano, A.; Abdala, V.; Hertel, F.; Herrel, A. Getting a grip on tetrapod grasping: Form, function, and evolution. *Biol. Rev.* **2013**, *88*, 380–405. [CrossRef] [PubMed]

65. Hobson, E.A.; Avery, M.L.; Wright, T.F. The socioecology of Monk Parakeets: Insights into parrot social complexity. *J. Exp. Biol.* **2010**, *213*, 2617–2628. [CrossRef] [PubMed]

66. Lawler, R.R. Sifaka positional behavior: Ontogenetic and quantitative genetic approaches. *Am. J. Phys. Anthr.* **2006**, *131*, 261–271. [CrossRef] [PubMed]
68. Gebo, D.L.; Chapman, C.A. Habitat, annual, and seasonal effects on positional behavior in red colobus monkeys. *Am. J. Phys. Anthr.* 1995, 96, 73–82. [CrossRef]

69. Pielou, E.C. The measurement of diversity in different types of biological collections. *J. Theor. Biol.* 1966, 13, 131–144. [CrossRef]

70. Risser, P.G.; Rice, E.L. Diversity in Tree Species in Oklahoma Upland Forests. *Ecology* 1971, 52, 876–880. [CrossRef]

71. Spellerberg, I.F.; Fedor, P.J. A Tribute to Claude Shannon (1916–2001) and a Plea for More Rigorous Use of Species Richness, Species Diversity and the ‘Shannon–Wiener’ Index. *Glob. Ecol. Biogeogr.* 2003, 12, 177–179. [CrossRef]

72. Keylock, C. Simpson diversity and the Shannon-Wiener index as special cases of a generalized entropy. *Oikos* 2005, 109, 203–207. [CrossRef]

73. Quinn, T.H.; Baumel, J.J. The digital tendon locking mechanism of the avian foot (Aves). *Zoomorphology* 1990, 109, 281–293. [CrossRef]

74. Grand, T.I. A Mechanical Interpretation of Terminal Branch Feeding. *J. Mammal.* 1972, 53, 198–201. [CrossRef]

75. Dunbar, D.C.; Badam, G.L. Locomotion and Posture During Terminal Branch Feeding. *Int. J. Primatol.* 2000, 21, 649–669. [CrossRef]

76. Young, J.W.; Stricklen, B.M.; Chadwell, B.A. Effects of support diameter and compliance on common marmoset (*Callithrix jacchus*) gait kinematics. *J. Exp. Biol.* 2016, 219, 2659–2672. [CrossRef] [PubMed]

77. Schmitt, D.; Zeininger, A.; Granatosky, M.C. Patterns, Variability, and Flexibility of Hand Posture During Locomotion in Primates. In *The Evolution of the Primate Hand*; Kivell, T.L., Lemelin, P., Richmond, B.G., Schmitt, D., Eds.; Developments in Primatology: Progress and Prospects; Springer: New York, NY, USA, 2016; pp. 345–369. ISBN 978-1-4939-3644-1.

78. Lammers, A.R.; Gauntner, T. Mechanics of torque generation during quadrupedal arboreal locomotion. *J. Biomech.* 2008, 41, 2388–2395. [CrossRef] [PubMed]

79. Dickinson, E.; Young, M.W.; Kim, C.J.; Hadjiargyrou, M.; Granatosky, M.C. Force Generation Capabilities in the Parrot Hindlimb Reflect Preferential Perching Behaviors. In Proceedings of the Integrative and Comparative Biology, Phoenix, AZ, USA, 3–7 January 2022; Volume 561.

80. Ward, A.B.; Weigl, P.D.; Conroy, R.M. Functional Morphology of Raptor Hindlimbs: Implications for Resource Partitioning. *Ornithology* 2002, 119, 1052–1063. [CrossRef]

81. Reilly, S.M.; McElroy, E.J.; Biknevicius, A.R. Posture, gait and the ecological relevance of locomotor costs and energy-saving mechanisms in tetrapods. *Zoology* 2007, 110, 271–289. [CrossRef]

82. Pontzer, H. Effective limb length and the scaling of locomotor cost in terrestrial animals. *J. Exp. Biol.* 2007, 210, 1752–1761. [CrossRef]

83. Raichlen, D.A.; Pontzer, H.; Shapiro, L.J. A new look at the Dynamic Similarity Hypothesis: The importance of swing phase. *Biol. Open* 2013, 2, 1032–1036. [CrossRef]

84. Kilbourne, B.M.; Andrada, E.; Fischer, M.S.; Nyakatura, J.A. Morphology and motion: Hindlimb proportions and swing phase kinematics in terrestrially locomoting charadriiform birds. *J. Exp. Biol.* 2016, 219, 1405–1416. [CrossRef]

85. Heglund, N.C.; Taylor, C.R. Speed, Stride Frequency and Energy Cost per Stride: How Do They Change with Body Size and Gait? *J. Exp. Biol.* 1988, 138, 301–318. [CrossRef]

86. Young, M.W.; Lynch, S.K.; Dickinson, E.; Currier, A.A.; Davoli, E.C.; Hanna, C.S.; Fischer, H.M.; DiUbaldi, G.A.; Granatosky, M.C. Patterns of single limb forces during terrestrial and arboreal locomotion in rosy-faced lovebirds (*Psittaciformes: Agapornis roseicollis*). *J. Exp. Biol.* 2022, 225, jeb244571. [CrossRef]

87. Carril, J.; Barbeito, C.G.; Tambussi, C.P. Making a parrot zygodactyl foot: Osteology and morphogenesis of the tarsometatarsus in the monk parakeet (*Myiopsitta monachus*). *Zoology* 2020, 144, 125877. [CrossRef]

88. Fujita, M.; Kawakami, K.; Higuchi, H. Hopping and climbing gait of Japanese Pygmy Woodpeckers (*Picoides kizuki*). *Comp. Biochem. Physiol. Part A Mol. Integr. Physiol.* 2007, 148, 802–810. [CrossRef] [PubMed]

89. Cohen, J. *Statistical Power Analysis for the Behavioral Sciences*, 2nd ed.; Lawrence Erlbaum Associates: Hillsdale, NJ, USA, 1988; ISBN 978-0-8058-0283-2.

90. Stevens, N.J. Stability, limb coordination and substrate type: The ecorelevance of gait sequence pattern in primates. *J. Exp. Zool. Part A Comp. Exp. Biol.* 2006, 305, 953–963. [CrossRef] [PubMed]