Effects of infant age and sex, and maternal parity on the interaction of lactation with infant feeding development in chimpanzees

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Abstract

The interaction between infant feeding and maternal lactational physiology influences female inter-birth intervals and mediates maternal reproductive trade-offs. We investigated variation in feeding development in 72 immature wild chimpanzees (*Pan troglodytes schweinfurthii*) at Ngogo, Kibale National Park, Uganda, and made inferences about maternal lactation over the course of infancy. We compared the percentage (%) of time that mothers nursed infants as a function of infant age and assessed how hourly rates and bout durations of nursing and foraging varied in association with differences in offspring age, sex, and maternal parity. Nursing % times, rates and durations were highest for infants ≤ 6 months old but did not change significantly from 6 months to 5 years old. Nursing continued at a decreasing rate for some 5- to 7-year-olds. Infants ≤ 6 months old foraged little. Foraging rates did not change after 1 year old, but foraging durations and the % time devoted to foraging increased with age. Independent foraging probably became a dietary requirement for infants at 1 year old, when their energy needs may have surpassed the available milk energy. Infants spent as much time foraging by the time they were 4 to 5 years old as adults did. No sex effect on infant nursing or foraging was apparent, but infants of primiparous females had higher foraging rates and spent more time foraging than the infants of multiparous females did. Although no data on milk composition were collected, these findings are consistent with a working hypothesis that like other hominoids, chimpanzee mothers maintained a fixed level of lactation effort over several years as infants increasingly supplemented their growing energy, micronutrient and hydration needs via independent foraging. Plateauing lactation may be a more widespread adaptation that allows hominoid infants time to attain the physiology and skills necessary for independent feeding, while also providing them with a steady dietary base on which they could rely consistently through infancy, and enabling mothers to maintain a fixed, predictable level of lactation effort.
Introduction

Infant feeding development (or infant nutritional development) in mammals is the transition from complete dependence on maternal milk to nutritional independence [1–3]. Stages of feeding development include exclusive suckling; transitional feeding, which starts when infants first consume non-milk food; and weaning, which ends after the last nursing bout with milk transfer [1,2]. Comfort nursing (without milk transfer), in which immature individuals continue to make non-nutritive nipple contacts after the completion of lactation [4,5], may also occur. When it does, it leads to different physiological and behavioral weaning ages unless nutritional and behavioral weaning occur at the same time, as is often the case [4,6]. A physiologically weaned individual has stopped ingesting maternal milk because milk transfer has ended but is not yet behaviorally weaned if they still make nipple contacts. A behaviorally weaned individual has stopped making nipple contacts but might already have been physiologically weaned.

Changes in suckling and foraging by infants as they age influence the scheduling and duration of feeding development stages and affect early-life rates of growth and survival [3,7]. Furthermore, since nursing and foraging behaviors reflect the distribution of infants’ dietary needs for milk and non-milk foods, changes in infant feeding behavior illuminate the interaction between infant nutritional requirements over the course of development and maternal lactational physiology. This is because milk synthesis is largely a maternal physiological response to mechanical nipple stimulation by the suckling infant [8–13]. The interaction between infant feeding behavior and maternal lactational physiology can influence inter-birth intervals and mediate important maternal trade-offs between current and future reproduction and number versus quality of offspring [3,14–17].

In primates, the time spent independently foraging by immature individuals generally increases with increasing age [18–25], which is a proxy for size, development of ecological competency, and total metabolic needs. Cross-species data show that, counter-intuitively, the length of time spent nursing can either decrease, increase, or remain stable through the majority of infancy until it finally decreases and stops at behavioral weaning. When nursing gradually decreases with increasing infant age (e.g. olive baboon, Papio anubis: [26]; rhesus macaque, Macaca mulatta: [9,10]; blue monkey, Cercopithecus mitis stuhlmanni: [24]; vervet monkey, Chlorocebus pygerythrus: [27]; mountain gorilla, Gorilla beringei: [28]; chimpanzee, Pan troglodytes schweinfurthii at Gombe: [29], and Mahale: [23]), maternal lactation effort should decline over time because females reduce lactational investment as their offspring grow and develop foraging competence [5,10,30,31].

Alternatively, when infants nurse progressively more frequently and/or for longer per day as they age (e.g. chimpanzee, Pan troglodytes schweinfurthii at Kanyawara: [22]), maternal lactation effort may increase with infant age to support the increasing energetic needs of larger, developing infants [18,32]. Increasing nursing behavior may thus reflect increased maternal milk synthesis over time, because while infants spend more time foraging as they gradually attain independent feeding abilities, they also continue to obtain much of the energy they need for growth and maintenance from milk.

Finally, nursing behaviors may remain relatively constant through most of infancy (e.g. Japanese macaque, Macaca fuscata: [33]; rhesus macaque, Macaca mulatta: [11]; yellow baboon, Papio cynocephalus: [18]; gelada, Theropithecus gelada: [34]; chimpanzee, Pan troglodytes schweinfurthii at Gombe: [21] and Ngogo: this study; orangutan, Pongo pygmaeus wurmbii: [35]), which may reflect a regular rate of maternal milk synthesis. Milk transfer rates in such cases presumably remain constant and females maintain steady lactational investment while offspring attain independent feeding abilities, with infants meeting their additional growth-,
developmental-, and size-related energy and nutrient needs through increased foraging during
the transition to nutritional independence [1,3,32,35–38]. Offspring could thus rely on a stable
and predictable milk supply through most of infancy. Anticipating the milk supply would
allow infants to compensate for gaps between their energy needs and the energy available from
milk by foraging independently.

The sucking efficiency of infants, however, can also increase with age as infants develop the
ability to coordinate suckling rate with swallowing and respiration. Consequently, while time
spent nursing can change or remain constant as infants age, the amount of milk transfer can
increase or remain the same [39]. Similarly, the composition and nutrient density of milk
secreted may change with infant age and development and is rarely investigated below gross
nutrient composition level [40].

The age at which infants first ingest non-milk foods, and thus the age at which transitional
feeding begins, does not necessarily overlap with the age at which infants must regularly sup-
plement their diet with non-milk foods because they have physiologically outgrown the energy
available to them from maternal milk [3,38]. For instance, non-human ape infants may not
physiologically need to feed on non-milk foods until around 1 year old, even though they often
sample non-milk foods several months before this age [3,4,38]. This point in feeding develop-
ment is also different from the age at which immature primates are developmentally ready to
rely more heavily on adult foods, which could help predict when survival is possible—though
potentially still unlikely—without the continuing nutritional support of maternal milk
[23,36,41,42]. Infant chimpanzees, for example, may start to rely more heavily on solid foods
at 3 years old [23,41], and they spend as much time as adults foraging independently by 4 to 5
years old, which is also the age range within which behavioral and physiological weaning usu-
ally occurs [4,21,22]. Indeed, orphan chimpanzees in the wild can only survive without nursing
after 3 years old, and their survival chances decline if their mother dies before they reach age 5
[43,44]. Thus, while the point at which primate infants quantitatively increase their nutritional
dependence on solid food and reduce their dependence on maternal milk may happen some-
time before physiological weaning, continued nursing and maternal milk transfer to infants
after this point is adaptive for females and their offspring [1,3,36,42].

Considerable intra-specific variation overlies species-typical patterns in infant feeding
behavior. Variation in infant feeding patterns can lead to differential rates of growth and devel-
opment and to variation in the length of infant dependency on mothers [3,16,36]. Several fac-
tors are associated with variation in nursing and milk intake rates [40], including maternal
health condition [45–49], parity and reproductive experience [13,49–51], quantity and quality
of milk produced [9,51–54]; maternal activities outside of infant care [18,34]; alloparental care
[55,56]; infant sex [13,51,57–59], hunger [39,60], age, and nursing efficiency [61]; and varia-
tion in food availability [25,62]. Solid food intake rates of immature individuals may vary with
age [18–25,63] and sex [24,64]; the extent to which others share food with them [65–67]; and
variation in the ease of processing different types of food and temporal variation in food avail-
ability [25,62,67].

Data on infant diets from additional sites provide insights into how life history theory
explains intraspecific variation in lactational effort and feeding ontogeny and help reveal
which associated traits are more plastic or more constrained [1,38,68]. Wild chimpanzee
infant feeding behavior has been described at several sites, but detailed descriptions of
nursing and the development of independent foraging have not previously been available for
wild chimpanzees (Pan troglodytes schweinfurthii) at Ngogo, Uganda. Previous fecal
stable nitrogen isotope analyses indicated a physiological weaning age at Ngogo of 4 to 4.5
years old [4]. While many infants were also behaviorally weaned by this age, several
showed prolonged comfort nursing and were not behaviorally weaned until 5.5 to 6 years
old, or even 7 to 8 years old, based on observations of nipple contacts [4,6]. Males were behaviorally, and probably physiologically, weaned later than females [6]. Stable carbon and nitrogen isotope data showed that the proportions of milk to non-milk food in the diets of infants gradually decreased through most of infancy, consistent with the occurrence of a gradual physiological weaning process over several years [4]. However, data on how nursing and independent foraging behaviors are expressed through infancy are needed to understand whether this gradual physiological weaning process results from reduced maternal lactational investment over the course of infancy or from infants relying progressively more on independent foraging while maintaining their reliance on milk [6]. Here we report new results of analyses on feeding development in the Ngogo chimpanzee community using behavioral data on nursing and foraging by 72 immature individuals ranging from 0 to 9 years old. To delineate variation in infant feeding and to infer how maternal lactation changed over the course of infancy, we assessed how infant age, sex and maternal parity affected the percentage (%) of time that infants spent nursing and independently foraging and the rates and durations of nursing and foraging bouts.

Methods

Study site and species

The Ngogo study site, in the center of Kibale National Park, Uganda, comprises about 35 km² of mostly old growth forest and also includes colonizing forest, grasslands, and swamp areas of Acanthus pubescens and Cyperus papyrus [69,70]. Dry seasons are separated by rainy seasons that occur from March to June and September to November [70,71].

The Ngogo chimpanzee community numbered between 199 and 207 individuals, including 54 to 57 adult females, 26 to 33 adult males, 30 to 34 juvenile and adolescent females, 33 to 42 juvenile and adolescent males, and 40 to 53 infants during the study periods. The community was in the process of undergoing a permanent fission by the 2018 study period [72]. Chimpanzees at Ngogo eat primarily fruit, especially a fig species, Ficus mucuso, as well as relying heavily on leaves, and to a lesser extent on other plant parts, such as pith and stems [70,73]. They also hunt and eat a variety of mammals, particularly red colobus monkeys (Procolobus badius) [74–76]. Females at Ngogo do not form dominance hierarchies [77]. Female conflicts are generally rare, and when they do occur, clear or consistent outcomes, with decided winners and losers, are uncommon [77–79].

Study subjects

We studied 72 immature chimpanzees aged 0 to 9 years old. None had younger maternal siblings—that is, their mothers did not have new infants during the data collection period. These 72 individuals were offspring of 56 adult females, as 16 adult females contributed two offspring to our dataset. We refer to individuals aged 0 to ≤ 5 years old as “infants” [80]. Data on the first appearance of infants and on the last appearance of their mothers prior to births were used to estimate ages of study subjects. Estimated ages varied mostly from one or several days to within a month. For two individuals, birth date estimates were within several months and were partly based on comparisons of their nutritional, physical and social independence relative to other infants with precise age estimates. Data collectors could individually recognize the chimpanzees. Researchers learned to recognize individual chimpanzees based on distinguishing facial features and color patterns, scars, ear knicks, missing fingers, limbs or eyes, and hair color and distribution on the head or body.
Data collection

Data were collected on directly observed daytime feeding of immature chimpanzees by IB from January to March 2013 and September 2013 to June 2014, and by IB, CC and KJD from January to April 2018 using focal animal sampling [81]. IB trained CC and KJD in the field to help ensure consistency and reliability in data collection between the three observers. During 1-hour samples, observers continuously recorded the frequency and duration focal subjects spent nursing (i.e., making nipple contact) and foraging (i.e., ingesting non-milk or solid foods). We excluded times when study subjects were on their mothers' ventrums with their faces not visible. Focal subjects were foraging when they independently looked for, picked, bit, chewed and ingested food that they acquired themselves. We excluded exploratory mouthing or handling of vegetation that did not lead to ingestion. Focal samples were terminated if the subject was out of view for more than 10 minutes. When feeding bouts lasted beyond the end of the 1-hour focal sample, observers tried to record the end time of the bout. Initial selection of focal subjects on a daily basis was either random or aimed at those individuals for whom data were particularly needed. After completing a sample, observers chose another individual from among those present and subsequently aimed to cycle through all individuals present in the same order for the rest of the day. Observers attempted to collect behavioral data that were evenly distributed between mornings (7am-12pm) and afternoons (12pm-5:30pm) to minimize the potential effects of diurnal variation in activities. We obtained a total of 1245.2 focal sampling hours and a mean of 12.4 hours per focal subject (Table 1), not including time that subjects were out of view. Focal follows were done over a mean of 8 ± 4 (SD) different days per study subject by age category.

All aspects of this research were approved by the Uganda Wildlife Authority (UWA), Uganda National Council for Science and Technology (UNCST), Makerere University in Uganda, the University of Toronto’s Office of Research Ethics and Environmental Health and Safety, and Université de Montréal’s Comité de déontologie de l’expérimentation sur les animaux (CDEA). Additional information regarding the ethical, cultural, and scientific considerations specific to inclusivity in global research is included in the Supporting Information (S1 Checklist).

Data analyses

We separated the behavioral data from birth to year 1 into two 6-month increments (0 to ≤ 6 months old, > 6 months to ≤ 1 year old), as the end of exclusive suckling and beginning of

| Infant age (years) | Number of focal individuals (female, male) | Mean number of focal hours (SD) |
|--------------------|-------------------------------------------|--------------------------------|
| 0–0.5              | 14 (9, 4, 1 sex unknown)                  | 12.05 (6.97)                    |
| > 0.5–1            | 10 (6, 4)                                 | 13.37 (6.59)                    |
| > 1–2              | 19 (11, 8)                                | 13.14 (3.60)                    |
| > 2–3              | 22 (9, 13)                                | 11.86 (5.52)                    |
| > 3–4              | 17 (9, 8)                                 | 11.58 (5.42)                    |
| > 4–5              | 7 (4, 3)                                  | 14.38 (5.95)                    |
| > 5–6              | 4 (1, 3)                                  | 9.50 (3.19)                     |
| > 6–7              | 4 (2, 2)                                  | 16.64 (13.95)                   |
| > 7–8              | 2 (0, 2)                                  | 11.88 (6.61)                    |
| > 8–9              | 1 (0, 1)                                  | 6.27 (na)                       |
| Total              | 100 (51, 48, 1 sex unknown)               | 12.39 (6.51)                    |

https://doi.org/10.1371/journal.pone.0272139.t001
transitional feeding in apes can occur before infants are 1 year old [1,3,4,63]. We pooled behavioral data on older individuals into 1-year increments (> 1 to 2 years old, > 2 to 3 years old, etc.; Table 1). To determine the amount of time that immature chimpanzees spent nursing or foraging, we calculated behavioral % times for each study subject by dividing the total number of hours they spent nursing or foraging by the total number of focal sampling hours at each age category, multiplied by 100. Nursing and foraging behaviors were considered distinct bouts when separated by at least 1 minute [9,27,28]. To obtain hourly behavioral rates, we divided bout frequencies of nursing or foraging by the total focal sampling hours at each age category for each study subject. We used nursing rates because they have been shown to correlate positively with milk synthesis and negatively with the resumption of ovulation [10,27,36]. To determine mean nursing bout durations, we included only complete bouts for which we saw both beginnings and endings. We obtained mean nursing bout durations for each age category by subtracting the onset of nursing from the end time for a given bout, adding all bout durations, and dividing the sum by the number of nursing bouts for each infant. Foraging bout durations varied much more than those of nursing bouts and we had fewer complete foraging bouts (observed from beginning to end), than incomplete bouts (either beginning, end, or both not seen). Complete foraging bouts were skewed towards those with the shortest durations. To determine mean foraging bout durations for each age category, we included both complete and incomplete bouts and subtracted the onset of foraging (or time when observation of foraging started) from the end time (or time when observation ended) for a given bout, adding all bout durations, and dividing by the number of foraging bouts for each infant.

Statistical analyses. We conducted Generalized Estimating Equations (GEE) analyses [82–84] to evaluate how the % time spent nursing and foraging and the rate and duration of nursing and foraging bouts varied in relation age category, sex (male or female), and maternal parity (primiparous or multiparous). We conducted a first set of GEEs on all immature chimpanzees, 0 to ≤ 9 years old. Of the 72 individuals, 23 contributed data to multiple age categories. Our total sample size used in the first set of GEEs was thus 100 individuals by age category (Table 1).

As a further evaluation of how feeding varied during infancy, we conducted a second set of GEEs on infants > 6 months to ≤ 5 years old. We excluded subjects of the youngest age category (0 to ≤ 6 months old), because adjustment in lactation, nursing, and infant metabolic physiology shortly after birth [12,85–87] could lead to differences in nursing between the youngest infants and those infants in older age categories. We also excluded subjects from the age categories > 5 years old because most chimpanzees are physiologically and behaviorally weaned at Ngogo by 4.5 years old, and we expected that most nipple contacts after age 5 would be non-nutritive [4]. We included 55 infants aged > 6 months to ≤ 5 years old; 16 of these contributed data to multiple age categories, so the total sample size for the second set of GEEs was 75 infants by age category.

We included infant and mother identities as the grouping structure in the analyses to control for repeated measurements of the same subjects. We ran statistical analyses using SPSS version 27, with alpha set at 0.05, and applied a Bonferroni correction to the resulting p values to account for multiple testing of the same sample.

Results

Nursing

Infants ≤ 6 months old spent 5.85 (SD: ± 3.4) % of their time suckling, and nursed on average for 1.63 (± 0.51) bouts per hour and for 2.03 (SD: ± 0.73) minutes per nursing bout. After > 6 months and until ≤ 4 years old, mean time spent nursing and mean nursing bout rates stayed
around 3 (± 0.51) % and 1.00 (± 0.44) bout per hour, and nursing durations remained close to 2.00 (± 0.50) minutes per bout. Nursing started to decrease after 4 years old, although it sometimes continued until offspring were 7 years old (Figs 1 to 3).

Percent time spent nursing and nursing hourly rates were higher for younger infants and decreased with age (GEE: P < 0.001 and < 0.001, respectively; Table 2), but average nursing bout durations did not vary with age (GEE: P = 0.84). Repeating the analyses only among infants > 6 months to ≤ 5 years old showed no significant changes in nursing % times, rates, or durations with age (GEE: P = 0.64, P = 0.25, P = 0.95, respectively; Table 3). Neither the sex of the infants nor the parity of their mothers predicted variation in nursing parameters (Tables 2 and 3).

### Foraging

Infants ≤ 6 months old spent 0.84 (SD: ± 1.81) % of their time foraging and feeding on non-milk foods and foraged on average for 0.11 (± 0.20) bouts per hour and for 1.48 (± 2.12) minutes per foraging bout. Infants > 6 months to ≤ 1 year old spent 17.18 (± 15.97) % of their time foraging and showed on average 1.64 (± 0.49) foraging bouts per hour and 5.87 (± 4.02) minutes per bout. From > 1 year old and onward, offspring foraging rates varied little, with a mean value of 1.99 (± 0.61) bouts per hour. However, the daily % time spent foraging and foraging bout durations were higher in older infants and ranged from 24.85 (± 8.13) % and 8.02 (± 2.52) minutes per foraging bout for infants > 1 to ≤ 2 years old, to a mean of 43.25 (± 0.01) % and 10.26 (± 3.45) minutes per foraging bout for infants > 2 to ≤ 3 years old (Table 3).
### Table 2. Generalized Estimating Equations (GEEs) for correlations between feeding parameters of chimpanzees 0 to 9 years old and infant age, sex, and maternal parity.

| Infant feeding parameter | Infant or Maternal Characteristic | β     | SE  | 95% Wald confidence interval | Hypothesis Test |
|--------------------------|----------------------------------|-------|-----|-----------------------------|-----------------|
|                          |                                  |       |     |                             | Lower          |
|                          |                                  |       |     |                             | Upper          |
|                          |                                  |       |     |                             | Wald χ² (df = 1) | P       |
| Percent time spent nursing | Infant age                      | -0.54 | 0.11| -0.75 -0.33                 | 25.03           | 0.000   |
|                          | Infant sex                      | 0.02  | 0.36| -0.69 0.73                  | 0.004           | 1.00    |
|                          | Maternal parity                 | -0.70 | 0.44| -1.56 0.16                  | 2.56            | 0.22    |
| Nursing hourly rate      | Infant age                      | -0.15 | 0.02| -0.20 -0.11                 | 53.01           | 0.000   |
|                          | Infant sex                      | 0.01  | 0.08| -0.15 0.18                  | 0.03            | 1.00    |
|                          | Maternal parity                 | -0.09 | 0.09| -0.27 0.08                  | 1.06            | 0.61    |
| Nursing bout duration    | Infant age                      | -0.001| 0.001| -0.002 0.001               | 0.66            | 0.84    |
|                          | Infant sex                      | 0.000 | 0.002| -0.005 0.004               | 0.04            | 1.00    |
|                          | Maternal parity                 | -0.004| 0.002| -0.008 0.001               | 2.29            | 0.26    |
| Percent time spent foraging | Infant age                     | 9.30  | 1.41| 6.54 12.06                  | 43.63           | 0.000   |
|                          | Infant sex                      | 4.31  | 3.66| -2.86 11.48                 | 1.39            | 0.48    |
|                          | Maternal parity                 | -4.66 | 3.36| -11.24 1.93                 | 1.92            | 0.33    |
| Foraging hourly rate     | Infant age                      | 0.08  | 0.06| -0.04 0.20                  | 1.78            | 0.36    |
|                          | Infant sex                      | 0.35  | 0.20| -0.04 0.73                  | 3.13            | 0.15    |
|                          | Maternal parity                 | 0.40  | 0.21| -0.01 0.80                  | 3.59            | 0.12    |
| Foraging bout duration   | Infant age                      | 0.09  | 0.12| 0.05 0.12                   | 20.33           | 0.000   |
|                          | Infant sex                      | -0.01 | 0.03| -0.06 0.04                  | 0.13            | 1.00    |
|                          | Maternal parity                 | -0.01 | 0.04| -0.08 0.06                  | 0.05            | 1.00    |

https://doi.org/10.1371/journal.pone.0272139.t002

### Table 3. Generalized Estimating Equations (GEEs) for correlations between feeding parameters of chimpanzees 6 months to 5 years old and infant age, sex, and maternal parity.

| Infant feeding parameter | Infant or Maternal Characteristic | β     | SE  | 95% Wald confidence interval | Hypothesis Test |
|--------------------------|----------------------------------|-------|-----|-----------------------------|-----------------|
|                          |                                  |       |     |                             | Lower          |
|                          |                                  |       |     |                             | Upper          |
|                          |                                  |       |     |                             | Wald χ² (df = 1) | P value |
| Percent time spent nursing | Infant age                      | -0.19 | 0.19| -0.56 0.18                 | 0.98            | 0.64    |
|                          | Infant sex                      | 0.51  | 0.31| -0.10 1.12                 | 2.70            | 0.20    |
|                          | Maternal parity                 | -0.41 | 0.35| -1.10 0.28                 | 1.36            | 0.49    |
| Nursing hourly rate      | Infant age                      | -0.09 | 0.06| -0.20 0.02                 | 2.38            | 0.25    |
|                          | Infant sex                      | 0.01  | 0.09| -0.17 0.18                 | 0.01            | 1.00    |
|                          | Maternal parity                 | -0.01 | 0.11| -0.22 0.21                 | 0.003           | 1.00    |
| Nursing bout duration    | Infant age                      | 0.001 | 0.001| -0.001 0.003              | 0.52            | 0.95    |
|                          | Infant sex                      | 0.001 | 0.002| -0.004 0.005              | 0.10            | 1.00    |
|                          | Maternal parity                 | -0.001| 0.002| -0.006 0.003              | 0.24            | 1.00    |
| Percent time spent foraging | Infant age                     | 6.57  | 1.21| 4.19 8.95                  | 29.30           | 0.000   |
|                          | Infant sex                      | 1.07  | 2.42| -3.67 5.81                 | 0.20            | 1.00    |
|                          | Maternal parity                 | -5.51 | 2.14| -9.71 -1.32                | 6.64            | 0.02    |
| Foraging hourly rate     | Infant age                      | 0.002 | 0.03| -0.05 0.05                 | 0.01            | 1.00    |
|                          | Infant sex                      | 0.16  | 0.11| -0.05 0.38                 | 2.17            | 0.28    |
|                          | Maternal parity                 | -0.48 | 0.12| -0.71 -0.24                | 16.05           | 0.000   |
| Foraging bout duration   | Infant age                      | 0.03  | 0.01| 0.02 0.04                  | 33.91           | 0.000   |
|                          | Infant sex                      | -0.003| 0.01| -0.03 0.02                 | 0.07            | 1.00    |
|                          | Maternal parity                 | 0.000 | 0.01| -0.02 0.02                 | 0.001           | 1.00    |

https://doi.org/10.1371/journal.pone.0272139.t003
21.78) % and 22.18 (± 18.61) minutes per foraging bout for individuals in age categories > 6 years old (Figs 4 to 6).

Foraging rates did not vary significantly across age categories (GEE: 0 to < 9 years old: $P = 0.36$, Table 2; > 6 months to ≤ 5 years old: $P = 1.00$, Table 3), but the daily % times spent foraging and average bout durations increased with age (0 to ≤ 9 years old: $P < 0.001$ and < 0.001, respectively; > 6 months to ≤ 5 years old: $P < 0.001$ and < 0.001, respectively). Among all individuals ≤ 9 years old, foraging parameters did not vary significantly with sex or maternal parity (Table 2). However, while foraging bout durations did not vary with maternal parity ($P = 1.00$) for infants 6 months to 5 years old, infants of primiparous females spent more time foraging ($P < 0.05$; Fig 7) and showed higher foraging hourly rates ($P < 0.001$) than infants of multiparous females (Table 3). Foraging parameters of infants 6 months to 5 years old did not vary significantly with infant sex.

**Discussion**

**Infant feeding development and maternal lactation**

The amount of time spent nursing and the rates and durations of nursing bouts were highest for newborns, 0 to 6 months old, which reflects the fact that milk is the only major source of energy during this time. This could reflect high nutritional needs to support peak postnatal infant growth rates directly after birth, like those of human babies [88]. It may also indicate that like other primates, chimpanzee infants take time to acquire the ability to nurse efficiently.
Newborns might need longer and more frequent nipple contacts than older infants to obtain enough milk. Indeed, newborns, who showed a head-bobbing reflex in search of their mothers’ nipples like human neonates do [89], often took time to latch and sometimes had difficulty remaining latched. Mothers might also have needed time to adjust, as newborns usually required extensive physical support to stay higher on the chest, which aided in latching.

Nursing parameters decreased with age but did not change significantly between 6 months and 5 years old. This could mean that the rate of milk synthesis did not vary greatly for mothers during this time, consistent with the fact that lactational performance (volume, quality, and rate of milk secretion) is largely controlled by the rate of mechanical stimulation of the nipple through suckling [8–13]. Age-invariant nursing behavior at Ngogo is consistent with data from chimpanzees at Gombe [21], but not with reports that nursing changed through infancy among chimpanzees at Kanyawara [22] and Mahale [23]. These inter-site differences suggest that despite the group-level clustering in nursing behavior that can be shown within chimpanzee communities, this behavior is susceptible to a degree of plasticity similar to that characterizing other developmental parameters, such as post-natal growth [90].

However, infant suckling efficiency might improve with age, even though the % time that infants sucked, and the rates and durations of nursing bouts varied little. If this was the case, older infants would obtain more milk during each nursing bout than younger infants [39]. Thus, maternal lactation effort would increase as infants aged, instead of remaining stable.
Another limitation to interpretation is our lack of data on night-time nursing, which may be common, as it is in humans [91]. If so, the nursing rates, durations and % times presented here underestimate the daily amount of nursing.

Ape infants take longer than one year to attain nutritional independence and are behaviorally weaned several years after birth (e.g. white-handed gibbons, *Hylobates lar*: 2.0 to 2.5 years; orangutan, *Pongo pygmaeus*: 7 years; gorilla, *Gorilla*: 2.8 to 4.6 years; *P. troglodytes*: 4 to 5 years; reviewed in [1] and [92]). Regular and prolonged nursing until behavioral weaning completion may be a pattern typical for hominoids, including humans [1,3,21,35,38], and cercopithecoids that can also take longer than a year to wean offspring behaviorally (e.g. Japanese macaques, *Macaca fuscata*: [33]; rhesus macaques, *Macaca mulatta*: [11]; yellow baboons, *Papio cynocephalus*: [18]; geladas, *Theropithecus gelada*: [34]). We previously used fecal stable nitrogen isotopes to show a gradual decrease in the relative contribution of milk to solid food in the diet of chimpanzee infants 1 to 4.5 years of age [4]. Given that nursing % times, bout rates and durations varied little through most of infancy, but independent foraging increased with age, we argue that infants were effectively leading their own gradual physiological weaning process. The decreasing relative contribution of milk to solid food in the diet seems to result from infants increasing their reliance on foods they procure independently rather than mothers gradually taking away lactational investment over the years.

While some infants sampled non-milk foods as early as 2 to 3 months after birth [4], these comprised a negligible proportion of the diet for at least the first 6 months, in line with expectations of infant dietary needs for apes and humans [1,3,38]. A notable difference between

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**Fig 4. Percentage of focal observation time that immature chimpanzees spent foraging at different ages.** Box plots show median values (solid horizontal lines), 25th and 75th percentile values (box boundaries), highest and lowest values (whiskers) and outliers (circles).

https://doi.org/10.1371/journal.pone.0272139.g004
chimpanzees and humans is that by the time they are 6 months old, human infants need regular provisioning with complementary foods to sustain growth, while chimpanzees and other apes may regularly sample solid foods at this age, but do not require dietary supplementation with solid foods until months later [1,3,38,41,93]. Moreover, ape infants acquire most of their supplementary foods via their own foraging efforts, rather than via provisioning by others. Supplementation of maternal milk with solid foods has been predicted to take a primary dietary role at around 1 year of age in chimpanzees; this may be when infant energy needs surpass their mothers’ lactational capacities, and when infants thus must start meeting some of their own nutritional needs [1,3,38]. At Ngogo, this prediction is supported by a few lines of evidence. First, fecal stable isotope data indicated that the relative contribution of maternal milk to solid food in the diet began to decrease when infants were one year old [4], at which age infants start regularly incorporating solid foods in their diets. Second, foraging rates increased until chimpanzees were 1 year old, after which they remained stable, but foraging bout durations and % times continued to increase with age. This implies that by 1 year of age, infants were foraging as often as adult chimpanzees—which makes sense because chimpanzee infants foraged whenever their mothers did by this age [4,55,94]—but that they needed to forage longer per bout as they aged because they needed to supplement maternal milk with progressively more solid foods. Findings at other sites similarly showed that chimpanzee infants did not start feeding regularly on non-milk foods until close to 1 year of age, and that the overall time they spent foraging progressively increased through development [19,21–23].

Fig 5. Average foraging rates of immature chimpanzees at different ages. Box plots show median values (solid horizontal lines), 25th and 75th percentile values (box boundaries), highest and lowest values (whiskers) and extremes (asterisks).

https://doi.org/10.1371/journal.pone.0272139.g005
Adult chimpanzees at Ngogo feed on average for 47% of their day [95]. Immature chimpanzees reached the adult-level 40 to 50% daily foraging times at around 5 years old at Kanyawara [22] and at 4.5 years old at Gombe [21]. Likewise, immature chimpanzees at Ngogo reached adult-level daily % foraging times by 4 to 5 years old, when they fed on non-milk foods for an average of 46.7 (SD ± 6.0) % of their day. After 5 years old, daily % foraging times varied greatly from one age category to another and between individuals, which could be due to the small number of chimpanzees in our sample after this age.

Most chimpanzees at Ngogo attained nutritional independence and were physiologically weaned by 4.5 years old, but a few continued comfort nursing for years after physiological weaning [4,6]. In the present study, three of four 5- to 6-year-olds and two of four 6- to 7-year-olds made nipple contacts during focal observations. Fecal stable isotope data established the presence of suckling with milk transfer for one of these individuals, while nipple contacts were for comfort, without milk transfer, for the others [4,6]. Our findings here showed that comfort nursing did not just involve occasional brief nipple contacts when infants checked mammary glands, but instead involved nipple contacts with measurable durations that occurred commonly, albeit at decreasing rates. The notion that some immature chimpanzees at Ngogo attained nutritional independence years before they stopped behaving as infants through continued nipple contacts accords with assessments of chimpanzee feeding development at other sites (Kanyawara: [41]; Mahale: [23]). Comfort nursing after physiological weaning may allow infants time to adjust to the changes that occur in the mother-infant social relationship with the infant-juvenile transition, and may result in behavioral weaning that is distinct from physiological weaning [6,96].

Fig 6. Average foraging bout durations of immature chimpanzees at different ages. Box plots show median values (solid horizontal lines), 25th and 75th percentile values (box boundaries), highest and lowest values (whiskers), outliers (circles), and extremes (asterisks).

https://doi.org/10.1371/journal.pone.0272139.g006

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Infant sex

We previously found that infant males were weaned later than females behaviorally, and likely physiologically, and that they had greater proportions of milk in their diets than same-aged females [6]. Two explanations for the greater proportions of milk in male diets could be that males generally nursed more often or longer per bout than females, or that they foraged less than females. However, our results were not consistent with either of these possibilities. Alternatively, males might have obtained more milk during each nursing bout than females [39]. Detailed data on infant cheek and jaw movements while suckling would help us estimate the amount of milk that infants obtained per bout [13,52]. Alternatively, immature males could be less efficient foragers than females; data on bite rates or the rate of food items ingested during foraging bouts could help discern this. Our results, however, are in line with findings from chimpanzees at Gombe that similarly showed no significant developmental differences between male and female infants in nursing and most foraging behaviors [21].

Maternal parity

Infants of primiparous females foraged at higher rates and spent more time foraging than the infants of multiparous females. Primiparous females may produce lower quality milk and/or less milk than multiparous females [49,51,97–99] and may physiologically wean their infants later than their multiparous counterparts [100]. These differences could occur because...
primiparous females are less experienced mothers and new to lactation, while also often being
physiologically immature and still developing while caring for their first infants [101–105].
Infants of primiparous mothers may need to compensate for lower milk quantities or quality
by foraging more than infants of multipares do. However, the proportions of milk to solid
food (mother-infant differences in fecal stable nitrogen isotopes) in the age specific diets of
infants of primiparous and multiparous mothers were similar at Ngogo, as are inter-birth
intervals of primiparous and multiparous females [6]. Thus, the higher foraging parameters
found here for infants of primiparous mothers did not seem to lead to lower proportions of
milk to solid food in their diets or to earlier attainment of nutritional independence compared
to infants of multiparous mothers.

Females at Ngogo occasionally share substantial amounts of premasticated food with their
infants, and the food most often shared is also the most commonly eaten by the chimpanzees
(a fig, Ficus mucuso) [70,94]. We previously found that primiparous females shared premasti-
cated food with their infants less often than multiparous females [94]. It may thus be that the
infants of primiparous females needed to forage independently more to meet their dietary
needs because their mothers shared less food with them.

**Future directions**

Although we had a large dataset for a wild primate infant study, we had only enough data to
make inferences about lactation and infant feeding development using mixed cross-sectional
analytic methods. Collection of longitudinal data for individual mother-offspring pairs would
make it possible to establish more precise estimates of population-level averages. Longitudinal,
age-adjusted cohort data would also allow us to assess effects of temporal variation in the avail-
ability of foods other than milk. For example, analysis of dental barium levels of four orangu-
tans (P. abelii and P. pygmaeus) throughout the entire infant period showed that orangutans
can implement a cyclical pattern of lactation to accommodate changes in infant nursing dur-
ing seasonal fluctuations in fruit availability [62]. We could not evaluate the effects of temporal
variation in food availability on nursing. Ngogo is characterized by relatively high food abun-
dance and relatively low variance in fruit availability [106–108] compared to other chimpanzee
sites, and slight fluctuations in yearly food availability may have relatively little influence on
lactation and infant feeding.

Comfort nursing deserves more investigation. It occurs after physiological weaning [4] but
some suckling by infants who are not yet physiologically weaned might also have been non-
nutritive. However, the consistency of nursing behavior during infancy supports the argument
that nipple contacts were driven more by dietary than socio-emotional needs, given that spe-
cific quantities of milk would have been available at certain times due to the interaction
between suckling stimuli and maternal lactational physiology [8–10,13].

**Implications for hominin evolution**

Our aggregated findings are in line with the argument that chimpanzee mothers maintained a
stable pattern of lactation effort, which may be evolutionarily more predictable for infants as
they increasingly supplement their growing energy needs with non-milk foods and gradually
attain the ability to feed independently [35,38]. As in humans [1,3,109–113], lactation effort in
many apes might not change progressively over time, but instead might maintain a plateau
through most of infancy [38]. Among the different patterns of lactation that could have charac-
terized our early hominin ancestors, plateauing lactation is a good candidate for the common
ancestral strategy [1,38]. Multi-year plateauing of lactation might have allowed hominoid
infants time to attain the physiology and behavioral skills necessary for independent feeding,
while also providing a steady and consistent nutritional base and buffering them from the negative effects of poor nutritional intake in early life [1,3,68]. Plateauing lactation would also have enabled females to maintain fixed, predictable levels of lactation effort through the course of infancy.

Compared to extant apes, humans have evolved a suite of adaptive characteristics associated with cooperative breeding that enabled females to have relatively short lactation periods for each offspring, and to invest in new infants well before previous offspring reached feeding independence, thus allowing for increased female reproductive rates [1,68,109,114–120]. Continued investigations of inter-population and inter-individual variation in infant feeding, weaning and lactation in chimpanzees and other apes will shed light on the evolutionary underpinnings that would have allowed early hominins to go from an ape-like model to the contemporary human form of infant feeding and maternal lactational investment.

Supporting information

S1 Checklist. Inclusivity in global research.
(PDF)

S1 File. Data used in analyses.
(XLSX)

Acknowledgments

We are grateful to our field assistants in Uganda, Godfrey Mbabazi, Alfred Tumusiime, Lawrence Ndangizi, Ambrose Twineonujuni and Denis Kyalikunda, and our dry lab assistants, Deandra Chipilliquen and Mathieu Gaudreault, in Canada. Thank you to Samuel Angedakin and the late Jeremiah Lwanga for logistical support at Ngogo, and to the Uganda Wildlife Authority, the Uganda National Council for Science and Technology, and Makerere University for permissions to do research at Ngogo. We thank Andrew Robbins and an anonymous reviewer who provided valuable constructive feedback to improve this paper.

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