Segmental morphometrics of the olive baboon (Papio anubis): a longitudinal study from birth to adulthood

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Abstract

The linear dimensions and inertial characteristics of the body are important in locomotion and they change considerably during the ontogeny of animals, including humans. This longitudinal and ontogenetic study has produced the largest dataset to date of segmental morphometrics in a Catarrhini species, the olive baboon. The objectives of the study were to quantify the changes in body linear and inertial dimensions and to explore their (theoretical) mechanical significance for locomotion. We took full-body measurements of captive individuals at regular intervals. Altogether, 14 females and 16 males were followed over a 7-year period, i.e. from infancy to adulthood. Our results show that individual patterns of growth are very consistent and follow the general growth pattern previously described in olive baboons. Furthermore, we obtained similar growth curve structures for segment lengths and masses, although the respective time scales were slightly different. The most significant changes in body morphometrics occurred during the first 2 years of life and concerned the distal parts of the body. Females and males were similar in size and shape at birth. The rate and duration of growth produced substantial size-related differences throughout ontogeny, while body shapes remained very similar between the sexes. We also observed significant age-related variations in limb composition, with a proximal shift of the centre of mass within the limbs, mainly due to changes in mass distribution and in the length of distal segments. Finally, we observed what we hypothesize to be ‘early biomechanical optimization’ of the limbs for quadrupedal walking. This is due to a high degree of convergence between the limbs’ natural pendular periods in infants, which may facilitate the onset of quadrupedal walking. Furthermore, the mechanical significance of the morphological changes observed in growing baboons may be related to changing functional demands with the onset of autonomous (quadrupedal) locomotion. From a wider perspective, these data provide unique insights into questions surrounding both the processes of locomotor development in primates and how these processes might evolve.

Key words: development; inertial properties; locomotion; Papio anubis; primate model.

Introduction

The locomotor apparatus of animals can be considered best adapted morphologically when its size and shape are well suited to the principal mode of locomotion of the species (see Witte et al. 1991; Preuschoft & Günther, 1994; Preuschoft et al. 1996). Primates are of special interest because, unlike many mammals with more specialized modes of locomotion, they adopt a variety of positional modes every day (Hunt et al. 1996) that involve a diverse range of substrate reaction forces (in terms of magnitude and direction) and moments. This is easily understood in view of the complex three-dimensional environment in which they evolved. Their locomotor apparatus should thus reflect, in some respects, an adaptive trade-off between efficiency, stability (static and dynamic), manoeuvrability and flexibility of movement.

It has been hypothesized that adaptations of the body to specific locomotor modes are likely to become more pronounced with development (Isler et al. 2006). This would make differential growth an important process for the locomotor transitions observed during primate ontogeny.
(e.g. Doran, 1997; Dunbar & Badam, 1998; Wells & Turnquist, 2001; Sarringhaus et al. 2014). Newborn primates are unable to walk: it commonly takes several weeks to months (depending on the species) to achieve this with a semblance of interlimb coordination, and from months to years to become fully efficient and autonomous. Their locomotor development is affected by complex processes involving different factors such as neuro-muscular maturation, experience and changes in body dimensions (e.g. Nakano, 1996; Adolph & Avolio, 2000; Garwicz et al. 2009; Kimura & Yaguramaki, 2009; Adolph et al. 2012). Adult catarrhine primates, including humans and their closest relatives, exhibit an impressive diversity in their morphometrics (i.e. linear dimensions, segmental masses, moment of inertia) in parallel with a wide diversity of locomotor modes. Yet these species appear to follow a common pattern of differential growth (Leigh et al. 2009). Consequently, studying the development of the locomotor apparatus in full is crucial to a better understanding of the ontogeny of locomotion potentially leading to specialized modes or generalized repertoires. Furthermore, such studies could improve our knowledge about the evolvability of locomotor profiles, i.e. ‘the ability of random variations to sometimes produce improvement’ (Wagner & Altenberg, 1996).

This paper focuses on changes in morphometrics and their (theoretical) mechanical significance during the ontogeny of the catarrhine olive baboon, *Papio anubis*, based on a longitudinal study of a large sample of captive individuals. At the adult stage, baboons are specialized for quadrupedal locomotion, yet they are also capable of adopting a variety of positional modes, including bipedalism (e.g. Rose, 1977). Just after birth, baboons only possess grasping abilities, but it has been shown that growing infants rapidly include quadrupedal walking and suspensory, climbing and leaping behaviour in their repertoire. As a result, infants exhibit a more diverse repertoire than adults (Druelle & Berillon, 2013). Furthermore, according to previous behavioural and biomechanical observations (Altmann & Samuels, 1992; Raichlen, 2005a; Druelle et al. in press), an autonomous locomotor profile is established in the first year of life, during which the functional demands on the limbs change in parallel with significant morphological changes. Such modifications in size and shape seem to adapt the body optimally for quadrupedal walking (Raichlen, 2005a). However, potential predisposing traits in the intrinsic morphodynamic properties of the limbs (i.e. those morphometrics that determine the intrinsic dynamics of the limb segments) may also favour early quadrupedal locomotion (Druelle et al. in press). In adult baboons, it has been demonstrated that there is a divergence in mass distribution between the forelimbs and hindlimbs that creates a natural pendular period (NPP) convergence (Raichlen, 2004). It has been previously argued that exhibiting good NPP convergence between the four limbs is likely to minimize muscle activity, and therefore energy expenditure, during quadrupedal locomotion (Myers & Steudel, 1997; Raichlen, 2004). It has further been hypothesized that the divergence in limb mass distribution might reflect the need to maintain precise grasping abilities in the hands in order to forage successfully after infants become less reliant on gripping the mother with the feet for transport (Raichlen, 2005b). In this context, we have produced the largest dataset of segmental morphometrics for a Catarrhini species to date and made it available for further exploration, e.g. in biomechanical studies such as (inverse) dynamic analyses of movement or musculoskeletal modelling. Based on this dataset and given that mass in newborn baboons is distally concentrated into the fore- and hindlimbs and that the body centre of mass has a more cranial position, we first hypothesized that, during early life, the morphology must adjust rapidly to meet the mechanical demands of developing the locomotor repertoire. It should be noted that in baboons, this repertoire tends to be quadrupedal, but that other locomotor abilities are also present. Nevertheless, according to our recent study on the early development of quadrupedal walking in baboons (Druelle et al. in press), the NPP of the four limbs should be strongly convergent initially. Adult baboons are sexually dimorphic (Leigh et al. 2009) and sex-related differences have been observed in the timing and intensity of their growth patterns (Glassman et al. 1983; Coelho, 1985; Glassman & Coelho, 1987; Leigh et al. 2009) and in their biomechanics (see hindlimb compliance in Patel et al. 2013). With regard to morphometrics, we hypothesized that females would have a lower rate of mass gain and that this pattern would end earlier than in males.

**Materials and methods**

**Study site and subjects**

This longitudinal study was conducted at the CNRS Primatology Station (UPS 846) at Rousset-sur-Arc, France, where approximately 300 baboons are bred and housed. From this population, we regularly measured 14 females and 16 males from infancy (minimum age 2 months) to adulthood; this experiment was conducted over a period of 7 years (Letter of approval MP/01/15/02/08 from the Midi-Pyrénées Region Ethics Committee, February 2008). The individuals were measured every 3 months until they were 2 years old and then every 6 months. All the individuals studied are part of the same troop and live together in a large park with various enrichments (Anvari et al. 2014; Berillon et al. 2010; Druelle & Berillon, 2013; see Supporting Information Table S1 for information on the individuals).

**Collecting morphometrics**

Our measurement protocol allows the use of the geometric model developed by Crompton et al. (1996), as previously applied to baboons (Raichlen, 2004, 2005b; Druelle et al. 2016a). External linear measurements of the nine body segments were taken (head, including the neck, trunk, upper arm, forearm, hand, thigh, shank, foot and tail). The landmarks used are the same as in the previous
studies (see Appendix in Schoonaert et al. 2007). The animals were anaesthetized to take the measurements (maximum duration, 20 min), and by convention, the limbs were measured on the left side only in order to ensure consistency between successive sessions and individuals, as well as to limit the duration of the anaesthesia. We obtained individual segment dimensions (lengths and anteroposterior and medio-lateral diameters) and estimated the inertial properties of the body segments from the model:

- segment mass, using segment average densities obtained from cadavers (Berillon et al. pers. obs.);
- segment centre of mass (CoM), calculated from the proximal joint of the segment (from the hip joint for the trunk segment);
- segment mass moment of inertia in the sagittal (I_x) and coronal (I_y) planes; the third moment of inertia in the horizontal plane was not calculated for this study because of its low impact on locomotor mechanics (cf. Schoonaert et al. 2007). Note that the moment of inertia about the segment CoM can be find using the parallel axis theorem (see Crompton et al. 1996; Raichlen, 2004).

We calculated inertial parameters for the whole body and the whole forelimb and hindlimb. Using the mathematical and geometric software GeoGebra 5.0, we designed free body diagrams of baboons at two extreme stages of their development, separately for males and females. We used a typical body posture (quadrupedal standing) observed in the wild with an average elbow joint angle of 155° and knee joint angle of 138° for all morphotypes (see Patel et al. 2013). We calculated the position of the whole body centre of mass (BCoM) in this posture, as well as the fore- and hindlimb centre of mass (CoM1 and CoM2, respectively). The CoM position was estimated as follows (Miller et al. 1973):

\[
\text{CoM} = \frac{\sum_{i=1}^{n} m_i r_i}{\sum_{i=1}^{n} m_i}
\]

where \( m_i \) is the mass of one segment and \( r_i \) is the distance of the segment CoM from the joint of origin (e.g. hip or shoulder). We then calculated the radius of gyration of the whole limb (RG) in the straight position. RG is a measure of overall limb mass distribution; it represents the distance of a point mass from the proximal joint that would give an equivalent inertia to the original limb. This is expressed as the square root of the whole limb’s mass moment of inertia about the proximal joint (which may be the shoulder or the hip), divided by the mass of the limb considered:

\[
\text{RG} = \sqrt{I/M}
\]

where \( I \) is the whole limb’s mass moment of inertia about the proximal joint (calculated using the parallel axis theorem; see Raichlen, 2004), \( M \) is the mass of the whole limb. RG can be normalized via a percentage of limb length. The natural pendular period (NPP) is a good way to represent the limb’s intrinsic morpho-dynamics (Myers & Steudel, 1997; Raichlen, 2004).

To model individual growth patterns, we regressed each individual’s absolute segmental morphometrics (length, mass, \( l_x, l_y \)) against the age at the time of measurement. Because each variable is age-dependent, we used type I regression models and their reliability was tested using the Fisher test. The slope and intercept values obtained for the significant models were used to test whether individual growth patterns followed the general pattern of growth, i.e. when all longitudinal data are grouped together. After controlling for the normality of the sample of individual growth patterns (Shapiro-Wilks test), we applied the one sample t-test to assess whether the mean of the sample of individual growth patterns was equal to the mean of the general growth pattern. This analysis was done using R 2.15 software (The R Foundation, Vienna, Austria, https://www.rproject.org/). To test differences between sexes, we used the slope values and the intercept values of the individual models. Comparisons were performed with permutation tests for independent samples (StatXact 3.1 software, Cytel, Inc., Cambridge, MA, USA).

Major age-related changes in segment lengths and masses were assessed using principal component analyses (PCA) following Druelle et al. (2016a). The set of variables was made dimensionless before being entered into the PCA: the length variables were divided by the cube root of the total body mass measured and the segment masses were divided by the total estimated body mass (Hof, 1996; Raichlen, 2005b). First, the PCA allowed us to reduce the number of variables. Secondly, it allowed us to determine the most significant changes occurring in the body morphometrics during growth. Thirdly, we plotted the principal component (the interpretability of the first eigenvalue was assessed using the broken-stick method (Frontier, 1976]) against age and we used a polynomial of degree \( n \) with a least-squares fit in order accurately to describe the period in which the most significant changes occur; linear models were not reliable here because of the autocorrelation of the distribution of residuals. The best fitting model is obtained when the coefficient of determination is considered high, i.e. \( R^2 > 0.5 \), and higher degrees do not further improve the fit (note that the end of the curve is likely to be biased because of the end of the dataset, which is why we have indicated this part with a dashed line). To estimate the developmental period when the amount of variation changes, i.e. decrease or increase in the growth rate, we calculated the second derivative of each polynomial model. A change in the sign of the second derivative curve indicates a point of inflection. The period of growth rate decrease or increase was determined from the median of the previous period, or the median between two points of inflection (see Fig. 1 for an example). We also performed a PCA, including general body morphometrics, i.e.

\[
\text{NPP} = 2\pi \sqrt{\frac{l}{M \times g \times \text{CoM}}}
\]

where \( g \) is the gravitational constant (9.8 m s\(^{-2}\)). We calculated the NPP of the fore- and hindlimbs for extended limbs with the hand and foot in a straight line with other segments. This procedure enables comparisons with previous studies (Myers & Steudel, 1997; Raichlen, 2004).

Statistics

In accordance with Crompton et al. (1996) and Raichlen (2004), the reliability of the external measurements procedure was assessed by calculating an estimated total body mass via the sum of the segment masses per individual. This estimated mass \( (\hat{m}) \) was compared with that directly measured with an electronic scale \( (x) \) yielding an absolute error calculation:

\[
\frac{|x - \hat{m}|}{x} \times 100
\]
segment lengths, segment masses and the total body mass, on 10 age–sex classes (0–6 months, 1½–2 years, 3–3½ years, 4½–5 years, 6–6½ years). This analysis was added for two reasons: first, to produce general age-related baboon morphotypes (coupled body size and body shape) and show how these change; secondly, to describe general differences between males and females during ontogeny for a confirmation of our previous tests.

Furthermore, according to previous contributions showing the importance of the shape and size of limbs in locomotion (see Kilbourne & Hoffman, 2013), we specifically analyzed the relationships between whole fore- and hindlimbs in straight positions. For each variable, i.e. limb length, limb mass, limb CoM, limb moment of inertia about the proximal joint, and limb NPP, we plotted the hindlimb against the forelimb. Because the two pairs of limbs vary together with age, we modelled these relationships using the reduced major axis (RMA), which is an orthogonal regression (type 2 model). The general growth pattern was thus assessed using the identity line: if the slope of the latter falls inside the confidence interval (95%) of the slope of the RMA model, this indicates a similar (isometric) growth pattern between fore- and hindlimbs. As we did for the full dataset, we validated the general growth pattern using comparisons with the mean of the sample of individual patterns (one sample t-test after controlling for normality of the sample). In addition, we plotted the dimensionless CoM and the dimensionless RG (CoM and RG are normalized via a percentage of limb length) against age and used the polynomial method previously described in order to assess the transitions. The significance level is 0.05 for all tests.

Results

Validity of the geometric model

The total body mass estimated from the direct external measurements is very close to that measured on a scale. We obtained, for females, a mean absolute error of 4.98 ± 1.69% (mean ± standard deviation) per individual measured and 5.16 ± 1.3% for males. These values strongly validate the reliability of the procedure and confirm the accuracy of the geometric model when applied to baboons; it was also validated previously on cadavers of different primate species using double-pendulum experiments (Crompton et al. 1996; Raichlen, 2004). This geometric model makes it possible to perform in vivo analyses of morphological parameters, thus offering new opportunities such as longitudinal follow up and observation, at the individual level, of performance–morphology interactions (Raichlen, 2005a, 2006; Young et al. 2007; Raichlen et al. 2009; Van Dam et al. 2010, 2011) and behaviour–morphology interactions (Druelle et al. 2016a,b).

Total body mass

Figure 2 shows that the polynomial model gives a good estimation of the increase in total body mass for female and male baboons ($R^2 = 0.93$ and 0.97, respectively). We observed two points of inflection for males, indicating an increase in the rate of mass gain at around 3.3 years of age and growth ending at around 6.55 years of age. In females, the general pattern is more linear than in males. We observed one point of inflection indicating a decrease in mass gain at around 4.9 years of age. This indicates the end of growth for females (Table 1).

Individuality within the general growth pattern

Table 2 shows the absolute segment linear and inertial parameters via the slopes (a) and intercepts (b) of the linear regression models. All the models are strongly significant ($P < 0.0001$) and illustrate the general growth pattern of
the olive baboon. Individual growth patterns closely match this general pattern. In females, no significant difference was found after testing for comparisons between individual and general patterns (Supporting Information Table S2). In males, we found only one significant difference for the foot mass (Supporting Information Table S3).

Regarding sex-related differences during ontogeny, it is clear that males exhibit a higher growth rate in length and mass for all the segments. The intercept values were not tested for differences because the lack of measurements at birth makes the origin of the calculated models redundant (see the negative values sometimes obtained: Table 2). However, by taking the data from the first measurement session, i.e. when the individuals were youngest [between 2 and 5 months for males (n = 7) and females (n = 7)], we were able to test early sex-related differences. Only one significant difference was found: the length of the hand is longer in males than in females (79 and 74 mm, respectively, permutation test: $P = 0.0061$).

### Changes in morphotypes (dimensionless segment lengths and masses) with age

The PCA produces a new principal component that explains a large fraction of the variance in segment lengths (36.12% in females and 34.7% in males) and in segment masses

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**Table 1** Polynomial regression models for total body mass.

| Sex       | Polynomial equation | $R^2$ | Point of inflection (year) | Variation in rate of change (year) |
|-----------|---------------------|-------|----------------------------|-----------------------------------|
| Female    | $-0.04x^3+0.3x^2+1.57x+1.39$ | 0.93  | 2.75                       | 4.9                               |
| Male      | $-0.04x^4+0.52x^3+1.83x^2+4.61x+0.61$ | 0.97  | 1.55-5.05                 | 3.3-6.55                          |

**Table 2** General growth pattern of segment inertial parameters represented by the linear regression model ($y = a \times \text{age} + b$).

| Segment | Length (mm) | Mass (kg) | $I_x$ (kg mm$^2$) | $I_y$ (kg mm$^2$) |
|---------|-------------|-----------|-------------------|-------------------|
|         | $a$ | $b$ | $a$ | $b$ | $a$ | $b$ | $a$ | $b$ |
| Females | | | | | | | | |
| Head    | 10.75 | 114.72 | 0.18 | 0.29 | 1830.37 | 5.08 | 1849.83 | -8.40 |
| Trunk   | 33.58 | 233.79 | 1.10 | 0.46 | 74465.39 | 45258.43 | 74056.72 | -44859.45 |
| Upper arm | 16.02 | 102.67 | 0.08 | 0.01 | 829.34 | -588.09 | 843.05 | -600.66 |
| Forearm | 17.11 | 111.14 | 0.04 | 0.04 | 551.64 | -266.72 | 550.84 | -266.53 |
| Hand    | 7.35  | 81.19  | 0.02 | 0.03 | 73.45  | -6.74  | 70.58  | 6.71   |
| Thigh   | 15.82 | 108.61 | 0.17 | 0.05 | 1734.90 | -1155.79 | 1793.15 | -1200.92 |
| Shank   | 15.62 | 111.39 | 0.05 | 0.03 | 641.62 | -301.36 | 645.83 | -305.03 |
| Foot    | 11.46 | 118.61 | 0.03 | 0.05 | 222.34 | 29.11  | 217.17 | 27.98  |
| Tail    | 20.92 | 291.90 | 0.03 | 0.05 | 957.71 | 365.70 | 957.71 | 365.70 |

$P$-value sex-related

| Segment | Head | Trunk | Upper arm | Forearm | Hand | Thigh | Shank | Foot | Tail |
|---------|------|-------|-----------|---------|------|-------|-------|------|------|
| Females | 10.75 | 33.58 | 16.02     | 17.11   | 7.35 | 15.82 | 15.62 | 11.46| 20.92|
| Males   | 16.98 | 42.38 | 20.92     | 21.24   | 10.25| 20.15 | 20.46 | 15.79| 34.13|

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(46.85% in females and 47.62% in males); the first observed eigenvalues, 3.25, 3.12, 4.22 and 4.29, respectively, always exceed the first eigenvalue provided by the broken-stick method, 2.83. Regarding segment lengths, the first dimension is compiled from the same segments for both sexes: the head [factor score (FS) female: 0.9; male: 0.86], the hand (FS female: 0.89; male: 0.84), the foot (FS female: 0.89; male: 0.84) and the tail (FS female: 0.77; male: 0.78) are positively correlated. This dimension is strongly correlated with age for females: \( r = -0.89 \) (\( P < 0.0001 \)) and males: \( r = -0.83 \) (\( P < 0.0001 \)); Fig. 3 illustrates these relationships. The 4th order polynomial gives a good estimate of growth-related changes: females: \( R^2 = 0.9 \); males: \( R^2 = 0.86 \).

Regarding segment masses, the first dimension is compiled from the same segments for both sexes: the head (FS female: 0.9; male: 0.86), the hand (FS female: 0.88; male: 0.87), the foot (FS female: 0.84; male: 0.8) and the tail (FS female: 0.75; male: 0.77) are negatively correlated with trunk mass (FS female: −0.72; male: −0.78). This dimension is strongly correlated with age for females: \( r = -0.88 \) (\( P < 0.0001 \)) and males: \( r = -0.86 \) (\( P < 0.0001 \)); Fig. 4 illustrates these relationships. The polynomial is a good estimate of these changes: females: \( R^2 = 0.86 \) (5th order polynomial), males: \( R^2 = 0.85 \) (5th order polynomial).

Using the second derivative method (see Statistics), we were able to detect the significant transitions in the growth patterns of segment length and segment mass (Table 3).

**Segment length**

In females, the greatest changes occur between birth and 1.65 years (Fig. 3). The growth rate is slower between 1.65 and 4.05 years and accelerates between 4.05 and 5.9 years of age, when growth stops. In males, the greatest changes occur between birth and 2 years of age. The rate of change is slower between 2 and 4.8 years and accelerates between 4.8 and 6.3 years of age, when growth stops.

**Segment mass**

A similar pattern appears for segment masses, but the changes occur earlier (Fig. 4). In females, the greatest changes occur between birth and 1.1 years. The rate of change is slower between 1.1 and 2.9 years and accelerates between 2.9 and 4.75 years of age, when growth stops (at 4.9 years for total mass gain). In males, the greatest changes occur between birth and 1.25 years. The rate of change is

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**Fig. 3** Relationship between the projection of the individuals on PC1 (relative segment lengths) and age. (a) All females and sessions plotted together; (b) all males and sessions plotted together.

**Fig. 4** Relationship between the projection of the individuals on PC1 (relative segment masses) and age. (a) All females and sessions plotted together; (b) all males and sessions plotted together.
slower between 1.25 and 3.55 years and accelerates between 3.55 and 6.2 years of age, when growth stops (at 6.55 years for total mass gain).

Based on these changes in body morphometrics, we were able to design the two extreme morphotypes of the baboon at birth and at the beginning of adulthood. Figure 5 illustrates these changes and their impact on the position of the BCoM as well as CoM1 (forelimb) and CoM2 (hindlimb). During development, we clearly observed a caudal shift of the BCoM into the trunk and a proximal shift of the CoM into the fore- and hindlimbs. The PCA applied to the 10 age-related morphotypes (see Statistics) produced two newly composed principal components that explain a large fraction of the variance (91.72%; Table 4). The two eigenvalues observed exceed the eigenvalues provided by the broken-stick method (PC1: $13.50^2 > 2.83$, PC2: $3.92 > 1.83$). Therefore, the first composed plan can be interpreted and allows general comparisons between females and males (Fig. 6). PC1 represents age-related variations in relative segment masses (except for the shank) and shows that the most significant differences occur during the first year of life. Female morphotypes, at all stages of their

| Sex      | Polynomial equation | $R^2$ | Point of inflection (year) | Variation in rate of change (year) |
|----------|---------------------|-------|----------------------------|------------------------------------|
| Segment lengths |
| Female  | $0.0187x^4 - 0.3035x^3 + 1.7885x^2 - 5.16x + 5.5505$ | 0.9   | 3.35-4.75                 | 1.65-4.05-5.9                      |
| Male    | $0.0114x^4 - 0.2191x^3 + 1.5418x^2 - 5.1015x + 5.8506$ | 0.86  | 4.05-5.55                 | 2-4.8-6.3                          |

| Segment masses |
| Female  | $-0.0106x^3 + 0.2079x^2 - 1.5057x + 5.0409x^2 - 8.7328x + 6.8916$ | 0.86  | 2.25-3.55-5.95          | 1.1-2.9-4.75                       |
| Male    | $-0.0053x^3 + 0.1313x^2 - 1.176x + 4.7429x^2 - 9.2897x + 7.5563$ | 0.85  | 2.55-4.55-7.85          | 1.25-3.55-6.2                      |

Fig. 5 Morphotypes of female (age-class 1, $n = 12$ and age-class 2, $n = 10$) and male (age-class 1, $n = 13$ and age-class 2, $n = 10$) baboons at two extreme stages of their ontogeny. The crosses represent the segment CoM, the small dots represent the joints between segments. All segment lengths are made dimensionless via the cube root of the total body mass.
Relative length

| Dimensions | Factor 1 | Factor 2 |
|------------|----------|----------|
| Hand       | 0.9491   | 0.0935   |
| Forearm    | 0.9747   | 0.0607   |
| Upper arm  | -0.9337  | -0.2665  |
| Thigh      | -0.9599  | -0.0151  |
| Shank      | 0.5338   | 0.2562   |
| Trunk      | -0.979   | 0.0258   |
| Head       | 0.9816   | -0.0816  |
| Tail       | 0.9626   | 0.1875   |
| Foot       | 0.9337   | 0.3263   |
| Total body mass | -0.788 | -0.5466 |

Numbers in bold indicate the main contributing factors.

Fig. 6 Projection of the 10 age–sex classes on the first composed plan of the PCA applied to dimensionless values (segment lengths and masses) and total body mass. Female baboons are represented with white squares and males are represented with black triangles. Numbers indicate the stage in development: 1 = 0–6 months; 2 = 1½–2 years; 3 = 3–3½ years, etc.

development, are closer to the adult morphotype than in males. Furthermore, in females, there is no difference between morphotypes 4 and 5. According to this axis, the last two female morphotypes and the last male morphotype are very similar. PC2 represents relative limb length and total body mass. At all stages in their development, males are relatively heavier and have shorter relative limb lengths than females. In males, a significant increase in total body mass occurs in the period between morphotypes 4 and 5, which corresponds to the acceleration in mass gain described previously (Table 1); this was not found in the female growth pattern.

Whole limb inertial properties: forelimb vs. hindlimb

Judging by the high values of the coefficients of determination, the limb growth pattern is well estimated by RMA models (Fig. 7). There is no difference between the general pattern of limb growth from the full dataset and the individual pattern (Supporting Information Tables S4 and S5). The general patterns of growth are similar for male and female baboons (see Figs 3 and 4). Therefore, it is possible to describe a general pattern of limb growth in baboons. During ontogeny, forelimb length, from the full dataset, is always and consistently smaller than hindlimb length [slope (95% CI): females 1.08 (1.05–1.1); males 1.09 (1.07–1.1)]. From the full dataset, the proximal shift of the CoM is more pronounced in hindlimbs than in forelimbs [slopes (95% CI): females 0.81 (0.77–0.86); males 0.79 (0.76–0.82)]. From the full dataset, the mass and moment of inertia of the hindlimbs about the proximal joint gradually increase relatively to the forelimbs [slopes (95% CI): 1.81 (1.76–1.87) and 1.78 (1.74–1.81), 1.50 (1.46–1.55) and 1.41 (1.38–1.44) for females and males, respectively].

Regarding NPP convergence, although the coupling between forelimb and hindlimb is always close during development (Fig. 8), the linear models show a trend towards a gradual increase in the forelimb NPP relatively to the hindlimb NPP [from the full dataset, slope (95% CI): females 0.82]. From the full dataset, the mass and moment of inertia of the hindlimbs about the proximal joint gradually increase relatively to the forelimbs [slopes (95% CI): 1.81 (1.76–1.87) and 1.78 (1.74–1.81), 1.50 (1.46–1.55) and 1.41 (1.38–1.44) for females and males, respectively].

Polynomial models yield good estimates for age-related variations in CoM and RG into fore- and hindlimbs (i.e. normalized via a percentage of limb length), as seen from the high values of the coefficients of determination (Fig. 9). Using the second derivative method (see Statistics), we determined the periods in which the CoM shifts proximally and RG changes in magnitude, towards a shorter radius, within the limbs (Table 6).

In females, three periods were plotted for the hindlimbs, and two periods for the forelimbs. In the hindlimbs, the
greatest proximal shift of the CoM occurs between birth and 0.95 years. The shift continues at a lower rate from 0.95 to 2.7 years of age, when the shift ends. In the forelimbs, the proximal shift of the CoM is close to the linear model. With the polynomial model, there is a more significant shift from birth to 1.85 years. This continues at a slightly slower rate from 1.85 to 5.4 years of age, when the shift comes to an end.

In males, three periods were plotted for hindlimbs and forelimbs. The greatest changes occur between birth and 1.35 years for both pairs of limbs. The shift continues at a slower rate from 1.35 to 4 years for hindlimbs and forelimbs, respectively, then accelerates in the hindlimbs from 4 to 6.65 years of age, when the shift comes to an end. In the forelimbs, it accelerates from 4.2 to 6.85 years of age, when it ends.

As seen in Fig. 9, the mass is more proximally concentrated in the hindlimbs than in the forelimbs at all stages of development in both sexes. In both pairs of limbs, baboons experience a proximal migration of the CoM. The results for the RG follow those for the CoM.
Discussion

Growth pattern

The shapes of the growth curves for skeletal length and segment mass are very similar. Although the same number of phases (variation in rates of change) is observed for segment length and mass, the timescales vary. The greatest changes occur in the first phase, corresponding to infancy; the second involves a significant decrease in the rates of change, which speed up again in the third phase up to adulthood. Our first hypothesis was that newborn baboons would have distally concentrated limb masses and that a new morphological pattern would develop rapidly to meet the mechanical demands of the onset of a quadrupedally oriented locomotor profile. Our results support this hypothesis: the greatest changes in body morphometrics occurred during the first year of life for segment mass and during the first 2 years for segment length. This involves a decrease in relative lengths and masses in the distal segments, i.e. the hand, foot, head and tail. From the general morphotypes, we also observed a relative increase in proximal segment masses, i.e. the thigh and upper arm, and in the trunk, as well as a relative decrease in forearm mass; no changes occurred in relative lengths for these segments. There is therefore a proximal shift of the CoM within the limbs that appears to depend mainly on the changes in relative segment masses and lengths (for autopodia); note that the CoM into each segment does not change with age. This growth pattern may be related to changes in limb functional demands because it matches the period in which baboons gradually become more active and autonomous, mainly through play and exploratory activities (Rose, 1977). According to our results, the next developmental period involves changes at a slower growth rate. There is a third and final period during which the growth rate increases slightly up to the end of development. The observation of these different phases is consistent with the work of Leigh et al. (2009), who observed the occurrence of growth spurts in baboons.

Although there are very few ontogenetic data of this kind for Catarrhini species in general, our results can nevertheless be compared with those of Turnquist & Wells (1994) for another cercopithecoid species, the rhesus macaque, and of Schoonaert et al. (2007) for a hominoid species, the chimpanzee. As previously proposed by Leigh et al. (2009), the olive baboon’s pattern of growth is similar, in some respects, to that of these two species. In macaques, the relative mass of the proximal limb segments increases while the relative masses of the distal limb segments and the head decrease. There is also a slight shift in the relative lengths of the fore- and hindlimbs in macaques, with the hindlimbs becoming relatively longer, whereas the slight difference in the limb lengths of baboons remains unchanged with age. Furthermore, the forearm and shank seem to remain constant throughout ontogeny in macaques, which is not the case for the relative mass of the forearm in baboons (cf. results reported here on general morphotypes). Because

Table 5 Comparisons of fore- and hindlimb NPPs during baboon ontogeny.

| Sex   | Age-class | FL mean | HL mean | FL–HL abs%diff | FL–HL Paired test |
|-------|-----------|---------|---------|----------------|------------------|
| Female | 0–1       | 1.11    | 1.11    | 2.56           | 0.7693           |
|       | 1–2       | 1.21    | 1.21    | 1.82           | 0.7250           |
|       | 2–3       | 1.28    | 1.28    | 1.66           | 0.3553           |
|       | 3–4       | 1.35    | 1.33    | 1.88           | 0.0008           |
|       | 4–5       | 1.38    | 1.36    | 1.99           | <0.0001          |
|       | 5–6       | 1.40    | 1.37    | 2.48           | <0.0001          |
| Male  | 0–1       | 1.12    | 1.11    | 2.18           | 0.0578           |
|       | 1–2       | 1.22    | 1.21    | 2.09           | 0.0099           |
|       | 2–3       | 1.30    | 1.28    | 1.91           | 0.0022           |
|       | 3–4       | 1.37    | 1.35    | 2.31           | 0.0002           |
|       | 4–5       | 1.45    | 1.42    | 2.63           | <0.0001          |
|       | 5–6       | 1.50    | 1.45    | 3.65           | <0.0001          |
|       | 6–7       | 1.50    | 1.46    | 3.25           | 0.0001           |

Fig. 8 Relationship between the NPP of the hindlimb and the NPP of the forelimb in female (a) and male (b) baboons. The dashed line is the line of identity. The linear equations represent the RMA regressions calculated with the full dataset for each variable and for each sex. The coefficients of determination indicate high reliability of the linear models in describing this relationship with age.
keeping more intrinsic muscle mass in forearms increases manual grasping abilities, this difference may be related to the fact that macaques spend more time in trees than baboons do. Similarly, the growth pattern of chimpanzees (Schoonaert et al. 2007) points to a shift from distal to proximal limb mass distribution, mainly through a decrease in the relative mass of the hand and foot and an increase in the relative mass of the thigh. There is also a significant decrease in the relative head mass but no increase in the relative trunk mass. Unlike macaques and baboons, chimpanzees have longer forelimbs than hindlimbs throughout their ontogeny. Nevertheless, the relative mass of the forearm and shank remains constant throughout ontogeny, as in macaques, and the upper arm does not increase in relative mass. This points to a need to maintain good grasping abilities in the hands in order to move efficiently in trees.

Schoonaert et al. (2007) reported that the decrease in the relative mass of the foot and hand may be due to a decrease in dimensionless segment lengths with age. Our results also suggest this relationship because the reduction in the relative mass of autopodia (hands and feet) parallels the reduction in the relative length of these segments. Preuschoft (2004) suggested that the independence of the grip is ensured by short intrinsic muscles of the hands and feet: longer digits would therefore require stronger and heavier muscles. Although having relatively longer and/or heavier distal segments has been correlated with grasping ability (Jungers & Fleagle, 1980; Raichlen, 2005b; Lawler, 2006; Druelle et al. 2016b), it has also been suggested that intrinsic morphological proportions, i.e. the length of digits

Table 6 Age-related migration of the CoM and RG into the limbs.

| Sex | Limbs | Polynomial equation | $R^2$ | Point of inflection (year) | Variation in rate of change (year) |
|-----|-------|---------------------|-------|---------------------------|---------------------------------|
|     | Centre of mass | |       |                           |                                 |
| F   | Fore  | $-0.0315x^3 + 0.3586x^2 - 2.0254x + 43.82$ | 0.50  | 3.75                       | 1.85-5.4                        |
|     | Hind  | $-0.0183x^3 + 0.3393x^2 - 2.294x^2 + 7.014x - 10.483x + 39.504$ | 0.62  | 1.95-3.45-5.75             | 0.95-2.7-4.6                    |
| M   | Fore  | $0.0239x^4 - 0.3997x^3 + 2.2135x^2 - 5.3761x + 45.38$ | 0.53  | 2.75-5.65                  | 1.35-4.2-6.85                   |
|     | Hind  | $0.0277x^5 - 0.4423x^4 + 2.3898x^3 - 5.8444x^2 - 37.679$ | 0.65  | 2.75-5.25                  | 1.35-4-6.65                     |
|     | Radius of gyration | |       |                           |                                 |
| F   | Fore  | $-0.0311x^3 + 0.356x^2 - 2.3023x + 67.763$ | 0.50  | 3.85                       | 1.9-5.45                        |
|     | Hind  | $-0.0232x^3 + 0.4283x^2 - 2.88x^3 + 8.7323x^2 - 13.111x + 62.431$ | 0.62  | 1.95-3.45-5.75             | 0.95-2.7-4.6                    |
| M   | Fore  | $0.0324x^4 - 0.5396x^3 + 2.9761x^2 - 7.1829x + 70.207$ | 0.55  | 2.75-5.55                  | 1.35-4.15-6.8                   |
|     | Hind  | $0.0381x^5 - 0.6054x^4 + 3.2282x^3 - 7.8193x^2 - 60.294$ | 0.65  | 2.65-5.25                  | 1.3-3.95-6.65                   |

Fig. 9 Relationship between the position of the CoM (1) and age, and the position of the RG (2) and age in female (a) and male (b) baboons. The white colour represents the forelimb, the black colour represents the hindlimb. The polynomial models are represented in the point clouds.
relative to metapodials, may be a substantial determinant of grasping performance (Lemelin & Schmitt, 2007; Young & Heard-Booth, 2016). In addition, having relatively longer distal segments may also facilitate some locomotor tuning (e.g. Lawler, 2006) by providing a larger area of support that potentially increases stability in quadrupedal locomotion at an early age.

Sex-related differences

Sex-related differences have been previously observed in the duration and rate of the growth pattern of baboons (Glassman et al. 1983; Glassman & Coelho, 1987; Leigh et al. 2009). Our large ontogenetic and longitudinal sample confirms this. All the linear models based on absolute values (segment length, mass and moment of inertia) differ between males and females, which means that most sex-related differences can be directly attributed to the higher absolute segment lengths and masses in male baboons, also resulting in higher values for the moment of inertia. Nevertheless, comparing infant male and female baboons in early infancy revealed only one significant difference, in hand length. Therefore, just after birth, male and female baboons are very similar in shape and size. Regarding total body mass, a rapid increase begins at around 3½ years of age in males and lasts until growth ceases (at 6½ years), whereas in females no changes were observed in the rate of mass gain, and growth stops earlier, at around 5 years of age. However, when corrected for total body mass effect, we observed a very similar pattern of growth for both sexes. According to the PCAs, males and females undergo the same transitional periods with similar changes in relative segment masses and lengths. Nevertheless, these transitional periods always occur later in males than in females, who reach adult body dimensions at an earlier age: before 6 years of age for relative segment length and before 5 years of age for relative segment mass, whereas males reach adult dimensions in relative length and mass a little after 6 years of age. Regarding the proximal shift of the limb CoM, the pattern appears to be similar between sexes in the hindlimbs, but subject to different processes in the forelimbs. In females, the shift is almost linear until around 5½ years of age, while the proximal shift observed in males follows the typical pattern found for other body dimensions, i.e. with considerable changes initially followed by a decrease in the growth rate, and a final acceleration just before growth ceases.

Development of the limbs

Our results show that the forelimbs are shorter than the hindlimbs in baboons and that this relationship is consistent throughout their development. It should be noted that the decrease in the intermembral index (without the hands and feet) observed at around 5 months of age by Shapiro & Raichlen (2006) in *Papio cynocephalus* is not marked here when observing the full duration of development, but is likely to become apparent with a shorter observation window. Three aspects have been considered: (i) the difference in shape between fore- and hindlimbs relatively to limb length, (ii) the impact of development on limb shape and (iii) the actual difference in shape between fore- and hindlimbs, i.e. relative to the proximal joint.

(i) Quadrupedal locomotion involves differential use of the limbs: hindlimbs are commonly used for propulsion while forelimbs are used more for braking (Demes et al. 1994). Furthermore, in primates, there is a need to maintain stronger grasping abilities in the hands than in the feet for foraging and manipulative functions (e.g. Pouydebat et al. 2014). Therefore, a disparity between mass distribution patterns in the fore- and hindlimb would reflect variations in the functional roles of the limbs during development (e.g. Rose, 1977; Raichlen, 2005b; Patel et al. 2013). Our results support this assumption in the sense that the CoM (linear effects) and RG (rotational effects) are always more distally positioned, relative to limb lengths, in the forelimbs than in the hindlimbs. For example, at 3½ years of age, we observed that the forelimb CoM is approximately in the same relative position as the hindlimb CoM was at birth (39.92 and 39.02%, respectively, in females; 40.24 and 38%, respectively, in males), whereas the latter is more proximally concentrated (7.09% more proximal in females and 6.29% in males).

(ii) Our results show that during early development, infant baboons experience a proximal shift of the CoM and RG in the fore- and hindlimbs. Regarding the kinematics of their quadrupedal walking, it has been shown that, at a very early age, infant baboons adopt longer stride lengths and lower stride frequencies because of their distally concentrated limb mass (Raichlen, 2005a). Furthermore, they exhibit total power outputs (i.e. the sum of internal and external power) similar to other quadrupedal mammals developing a more proximally concentrated limb mass (Raichlen, 2006). Therefore, it seems that infant baboons are using a trade-off mechanism that adapts their kinematics to reduce the energy expenditure induced by the movement of distally heavy limbs. Nevertheless, the CoM migration found in the present study is likely to reduce the rotational inertia of distal limb segments, and therefore the energy costs of swinging limbs in adults walking quadrupedally (Wickler et al. 2004). This supports the assumption that morphological adaptations become more pronounced during ontogeny.

(iii) After birth, actual fore- and hindlimb CoM and RG are very similar relative to their respective proximal
joints, i.e. the shoulder and hip. However, as a result of the proximal migration of the CoM described above, the actual forelimb CoM is more distally positioned than the actual hindlimb CoM at the end of development. On this point, it appears that, at an early age, differences in fore- and hindlimb mass distribution patterns compensate for differences in fore- and hindlimb lengths by creating fore- and hindlimb NPP convergence (see Myers & Steudel, 1997; Raichlen, 2004). With age, there is a loss of this convergence due to changes in limb composition. Although, during walking, specific mechanisms such as limb angular excursion may actively correct potential NPP divergence in adults (hindlimb excursion is relatively greater in baboons, Larson et al. 2001), it appears that swinging a limb according to its NPP reduces muscle activity, and therefore the metabolic energy demand. In this context, good convergence between fore- and hindlimbs should facilitate limb movement because the freely chosen frequency for the four limbs would not (theoretically) require any muscle activity, making locomotion more efficient. When we considered the limbs in a straight position (which is not a behavioural posture in baboons but allows interspecies comparisons), we found that forelimbs and hindlimbs in females converge to the same NPP up to 3 years of age, and in males up to 1 year of age. After 1 year, male forelimb and hindlimb NPPs diverge significantly, i.e. the NPP of the hindlimb is always smaller than the NPP of the forelimb, although their relationship remains very close (Table 5). This divergence appears only after 3 years of age in females. Our values for adult male [mean (95% CI): 3.25 (1.05)] and female [2.48 (0.55)] Papio anubis are consistent with those found by Raichlen (2004) in adult Papio cynocephalus [2.35 (1.36)]. Otherwise, the values we obtained for infant baboons [i.e. between 0 and 2 years of age, females: 2.11 (0.33), males: 2.13 (0.3)] are close to those obtained by Myers & Steudel (1997) for Canis familiaris [1.83 (1.26)]. Consequently, very young baboons, although immature in terms of neuromuscular control and with distally heavy limbs, exhibit limb mechanical properties that should facilitate the onset of quadrupedal walking (see Druelle et al. in press).

The findings of this study are of great interest because this biomechanical optimization (through NPP convergence) appears to predispose infant baboons for quadrupedal walking. According to Rose (1977), the first locomotor experience of infants occurs on the ground, just a few days after birth, when they squirm and tumble forwards as the hindlimbs start making propulsive movements while the forelimbs are unable to coordinate. After a few weeks, the fore- and hindlimbs seem sufficiently coordinated for quadrupedal walking. We also observed that this mode dominates their repertoire from the first locomotor stages (Druelle et al. 2016b) and is very soon performed efficiently (biomechanically speaking; Raichlen, 2006). Two important developmental processes are therefore described here: in parallel with major changes in limb shape that are related to changing functional demands, baboons exhibit ‘early biomechanical optimization’ for quadrupedal locomotion that certainly favours a rapid developmental transition toward this mode. This is in line with a growing body of literature showing that infant primates have a musculoskeletal system characterized by a capacity for a particular type of performance that could compensate for the immaturity of the system and thus promote survival during the early stages of life (e.g. Ruff, 2003; Young, 2005; Young et al. 2010a; Young & Heard-Booth, 2016). In an evolutionary perspective, early morphological adaptation of the limbs as observed in olive baboons may point to a high degree of integration between the fore- and hindlimbs (Young et al. 2010b). Such a developmental pattern (at the genotype level) is likely to be highly constrained by few possibilities for character variability, which would result in low degree of limb evolvability in baboons (Wagner & Altenberg, 1996).

To summarize, the changes in limb shapes and axial body related to differential growth (allometry) are significant during the ontogeny of olive baboons and appear clearly related to changes in functional demands. Besides this growth pattern, an early morphological predisposition for quadrupedal locomotion appears to be present. Similarities with the pattern of growth of Catarrhini species can be easily described, such as the proximal shift in limb mass distribution, as well as the caudal migration of the body centre of mass into the trunk; nevertheless, more ontogenetic data are necessary on Asian hominoids to confirm this Catarrhini-related growth pattern. Furthermore, the NPP convergence observed in infant baboons does not seem to be present in other Catarrhini species, at least in chimpanzees (Schoonaert et al. 2007). Such a morphological adaptation points to a high degree of locomotor specialization for quadrupedal locomotion in baboons, which does not, however, impede their other modes of locomotion. Consequently, studying variations in the morphological development of species that are phylogenetically close appears to be important to better understand the locomotor profiles exhibited in adult extant primates and how they might have evolved.

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Author contributions

G.B. designed the study and provided the financial and material resources. G.B., F.D., K.A. and V.M. conducted the experiments and acquired the data. F.D. and G.B. analyzed the experimental data. F.D., G.B. and P.A. interpreted the results and drafted the first version of the manuscript. All the authors revised the manuscript and approved the final draft.

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

Table S1. Information on the subjects.

Table S2. (a) Individual growth pattern of segment lengths represented by the linear regression model (y = a + b*age) in females. (b) Individual growth pattern of segment masses represented by the linear regression model (y = a + b*age) in females. (c) Individual growth pattern of segment inertia (x axis) represented by the linear regression model (y = a + b*age) in females. (d) Individual growth pattern of segment inertia (y axis) represented by the linear regression model (y = a + b*age) in females.

Table S3. (a) Individual growth pattern of segment lengths represented by the linear regression model (y = a + b*age) in males. (b) Individual growth pattern of segment masses represented by the linear regression model (y = a + b*age) in males. (c) Individual growth pattern of segment inertia (x axis) represented by the linear regression model (y = a + b*age) in males. (d) Individual growth pattern of segment inertia (y axis) represented by the linear regression model (y = a + b*age) in males.

Table S4. Individual linear regression models (RMA) in female baboons.

Table S5. Individual linear regression models (RMA) in male baboons.