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1	Systemic patterns of trabecular bone across the human and chimpanzee skeleton
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3	Short title: Human and chimpanzee systemic trabecular patterns
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Abstract

Aspects of trabecular bone architecture are thought to reflect regional loading of the skeleton, and thus differ between primate taxa with different locomotor and postural modes. However, there are several systemic factors that affect bone structure that could contribute to, or be the primary factor determining, interspecific differences in bone structure. These systemic factors include differences in genetic regulation, sensitivity to loading, hormone levels, diet, and/or activity levels. Improved understanding of inter/intraspecific variability, and variability across the skeleton of an individual, is required to properly interpret potential functional signals present within trabecular structure. Using a whole-region method of analysis, we investigated trabecular structure throughout the skeleton of humans and chimpanzees. Trabecular bone volume fraction (BV/TV), degree of anisotropy (DA) and trabecular thickness (Tb.Th) were quantified from high resolution micro-computed tomographic scans of the humeral and femoral head, third metacarpal and third metatarsal head, distal tibia, talus and first thoracic vertebra. We find that BV/TV is, in most anatomical sites, significantly higher in chimpanzees than in humans, suggesting a systemic difference in trabecular structure unrelated to local loading regime. Differences in BV/TV between the forelimb and hindlimb do not clearly reflect differences in locomotor loading in the study taxa. There are no clear systemic differences between the taxa in DA and, as such, this parameter may reflect function and relate to differences in joint loading. This systemic approach reveals both the pattern of variability across the skeleton and between taxa, and helps identify those features of trabecular structure that may relate to joint function.

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Keywords: Cancellous bone, Functional morphology, *Homo sapiens*, *Pan troglodytes*, Locomotion, Sedentism, Hominids

Introduction

The behaviour of extinct species can be reconstructed from plastic features of bony morphology
that reflect an individual's behaviour during life (Ruff et al., 2006). Experimental studies have
demonstrated the ability of bone to adapt to external loading (e.g. Lanyon, 1974; Robling et al.,
2002; Mori et al., 2003; Pontzer et al., 2006; Barak et al., 2011; Wallace et al., 2013), a process
often referred to as Wolff's Law (Wolff, 1986; Martin et al., 1998), or more generally as bone
functional adaptation (Cowin, 2001; Ruff et al., 2006). Trabecular bone has potential for
reconstructing the behaviour of fossil taxa (Kivell, 2016), as it remodels rapidly during life in
response to strain (Ehrlich and Lanyon, 2002), in comparison to the slower rate of remodelling of
cortical bone (Eriksen, 1986, 2010). Thus, the structure of trabecular bone could provide
information about the mechanical loading history of a joint, in terms of both the load magnitude
and direction. Studies among primates, including fossil specimens, have attempted to identify
behavioural signals in trabecular structure with varying degrees of success (e.g. Fajardo and
Müller, 2001; Ryan and Ketcham, 2002b; Griffin et al., 2010; Ryan and Shaw, 2012; Tsegai et
al., 2013; Skinner et al., 2015; Stephens et al., 2016; Zeininger et al., 2016). The ultimate goal
and framework within which these studies have been conducted is to first identify trabecular
differences in living species that are related to behaviour, for example locomotor or manipulatory
behaviours. Once this relationship between structure and behaviour has been established,
similarities between the trabecular structure of fossil specimens and living taxa could be used to
infer specific behaviours, or joint loading regimes, in fossil species.
However, the relationship between trabecular structure and behaviour in extant species is often
unclear. For example, many trabecular bone analyses have focused on the primate proximal
humerus (e.g. Fajardo and Müller, 2001; Fajardo et al., 2007; Ryan and Walker, 2010; Shaw and

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Ryan, 2012; Scherf et al., 2013; Scherf et al., 2015) and, for historical reasons (Skedros and
Baucom, 2007), the proximal femur (e.g. Fajardo and Müller, 2001; MacLatchy and Müller,
2002; Ryan and Ketcham, 2002a, b, 2005; Scherf, 2008; Ryan and Walker, 2010; Saparin et al.,
2011; Ryan and Shaw, 2012; Shaw and Ryan, 2012). However, few of these studies have found
clear differences in the trabecular structure of these joints that can be directly related to
locomotor mode and predicted joint function. Where structural differences in trabecular
architecture have been identified across locomotor groups, there is often no clear biomechanical
explanation, and trabecular architecture is not always consistent with predictions based on
biomechanical models. For example, studies of strepsirrhines have found that trabeculae within
the femoral head was more uniformly oriented in vertical clinging and leaping species compared
with slow climbing and/or quadrupedal taxa (MacLatchy and Müller, 2002; Ryan and Ketcham,
2002b, 2005). However, finite element analysis of the femoral head was unable to identify
differences in bone strain at a range of load orientations in vertical clinging and leaping Galago
compared to slow quadrupedal/climbing Loris (Ryan and van Rietbergen, 2005). This implies
that different trabecular structures may be able to mitigate stress in similar ways, and that joint
loading at the femoral (and potentially humeral) heads may actually be more similar than
predicted across divergent locomotor modes (Ryan and van Rietbergen, 2005; Fajardo et al.,
2007).
Since the first three-dimensional analysis of trabecular structure in primates (Fajardo and Müller,
2001), trabecular architecture has been described across a range of species and anatomical sites.
This body of work has revealed particular interspecific patterns in the variation of trabecular
structure, which suggests that any given species may have a similar trabecular structure across
several elements of their skeleton. As a notable example, recent humans have been shown to

91	have low trabecular bone volume throughout the postcranial skeleton, including highly-loaded
92	lower limb bones, such as the femur (e.g. Maga et al., 2006; Griffin et al., 2010; Tsegai et al.,
93	2013; Chirchir et al., 2015; Ryan and Shaw, 2015; Saers et al., 2016; Stephens et al., 2016;
94	Chirchir et al., 2017). In contrast, chimpanzees tend to have high bone volume across different
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96	skeletal elements in comparison to other hominoids (e.g. Maga et al., 2006; Griffin et al., 2010;
97	Tsegai et al., 2013). Although few trabecular studies include bonobos, their metacarpals and
98	metatarsals have the highest bone volume amongst the great apes (Griffin et al., 2010; Tsegai et
99	al., 2013), which is not readily explained by variation in body size, locomotor mode, or activity
	level (Susman et al., 1980; Doran, 1992, 1993a). Although bone volume fraction is the trabecular
100	parameter most strongly correlated with bone stiffness (Stauber et al., 2006; Maquer et al.,
101	2015), it does not seem to correspond directly to predictions of joint loading based on locomotor
102	mode.
103	There are several genetic and environmental factors, other than specific locomotor behaviours,
104	that could have a systemic effect on bone remodelling and trabecular structure (Bertram and
105	Swartz, 1991; Ruff et al., 2006; Kivell, 2016). Aspects of loading that are not evidently related to
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107	specific positional or locomotor behaviours include loading magnitude due to body mass (Doube
108	et al., 2011; Fajardo et al., 2013; Ryan and Shaw, 2013), differences in loading frequency
109	associated with overall activity levels (Lieberman, 1996), and other factors that may affect the
110	frequency, magnitude or orientation(s) of load and thus potentially impact remodelling of both
111	cortical and trabecular bone (Rubin and Lanyon, 1985; Frost, 1987; Skerry and Lanyon, 1995;
	Wallace et al., 2013). Genetic factors that might contribute to species-specific trabecular
112	structure include hormonal differences or differences in bone regulation, even between closely
113	related species (Lovejoy et al., 2003; Behringer et al., 2014a; Behringer et al., 2014b), between

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males and females (Riggs and Melton, 1995; Reginster and Burlet, 2006; Eckstein et al., 2007)
or at different life stages (Riggs and Melton, 1995; Tanck et al., 2001; Reginster and Burlet,
2006; Ryan and Krovitz, 2006; Gosman and Ketcham, 2009). These genetic differences may also
manifest as phylogenetic differences in bone structure, unrelated to locomotor mode (Fajardo et
al., 2013; Ryan and Shaw, 2013). Other aspects of the environment, such as diet and the
intestinal microbiome, could also have a systemic effect on bone structure (Prentice, 1997; Shea
et al., 2002; Cashman, 2007; Cao et al., 2009; Charles et al., 2015; McCabe et al., 2015). As the
rate of remodelling of bone is higher during growth, behaviours during development may be
more important for explaining trabecular morphology than those during adulthood (Bertram and
Swartz, 1991; Pettersson et al., 2010). This is of particular relevance for African apes, as the
percentage of knuckle-walking and suspension change significantly during development (Doran,
1992, 1997; Sarringhaus et al., 2014; Sarringhaus et al., 2016), although long bone cross-
sectional geometry in African apes continues to change into adulthood and reflect locomotor
behaviour at different life stages (Ruff et al., 2013; Sarringhaus et al., 2016; but see Demes et al.,
1998; Demes et al., 2001; Lieberman et al., 2004; Carlson, 2005). Trabecular morphology may
differ due to anatomical location (Morgan and Keaveny, 2001; Eckstein et al., 2007; Wallace et
al., 2015); for example, distal limb elements may be adapted to have a lower bone mass (bone
mineral density measured using pQCT and multiplied by joint size) and BV/TV than more
proximal limb elements (Chirchir, 2015; Saers et al., 2016).
The absence of detailed locomotor, positional and biomechanical data on particular primate
species may also contribute to limited identification of clear functional signals in trabecular
bone. For example, accurate information on locomotor frequencies is rare, in part because
several primate taxa are challenging to study in the wild due to lack of habituated populations,

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rarity of the species itself, and/or high density forest cover (Crompton et al., 2010). Many
species, especially hominoids, engage in multiple positional and locomotor behaviours (Hunt,
1991; Thorpe and Crompton, 2006; Myatt et al., 2011), beyond often over-simplified locomotor
categories. Furthermore, due to the difficulty -both ethically and practically- of studying the
biomechanics of locomotion in humans and especially non-human primates, there is little
accurate biomechanical data concerning loading orientations and joint reaction forces to inform
trabecular studies. Morphological differences related to locomotion have been investigated in
primate taxa through finite element analysis (e.g. Ryan and van Rietbergen, 2005; Richmond,
2007; Nguyen et al., 2014). Although finite element analyses enable more informed predictions,
they are often limited by a necessity to artificially reduce the complexity of the trabecular
structure (due to computational limitations) and a lack of validation (Richmond et al., 2005;
Ryan and van Rietbergen, 2005; Strait et al., 2005; Nguyen et al., 2014). Thus it is difficult to
determine which behaviour, or combinations of behaviours, are reflected in trabecular bone
structure.
To fully understand the functional significance of the trabecular bone structure of fossil
hominins, we need to further explore variation in trabecular bone across the skeleton of living
species. Previous studies have largely focused on one anatomical site (e.g. DeSilva and Devlin,
2012; Tsegai et al., 2013; Stephens et al., 2016) or region (Lazenby et al., 2011a; Schilling et al.,
2014; Tsegai et al., 2017), or have been limited to comparisons between the humerus and femur
(Fajardo and Müller, 2001; Ryan and Walker, 2010; Ryan and Shaw, 2012; Shaw and Ryan,
2012), and thus lack the context of how trabecular structure in any particular element or region
might reflect, at least in part, a broader systemic pattern. Several recent studies have addressed
the question of why previous comparative studies of trabecular bone have found notably gracile

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bone in modern humans. Chirchir et al. (2015) conducted an analysis of trabecular structure
across several skeletal elements in a sample of modern humans, fossil hominins and other extant
primates, showing that gracile trabecular structure in humans is a relatively recent (i.e. Holocene)
phenomenon. Ryan and Shaw (2015) further demonstrated, through a 3D volume of interest
analysis of trabecular structure in the proximal femur of modern humans varying in subsistence
strategies (foragers vs. agriculturalists), that gracile bone structure of recent humans is likely
linked to a reduction in overall activity level with the adoption of agriculture. This gracilisation
of the skeleton of agriculturalists is apparent across the lower limb, in the proximal and distal
epiphyses of the femur and tibia, although all populations share a proximo-distal reduction in
bone volume and increase in anisotropy (Saers et al., 2016). A similar pattern of gracilisation in
recent humans, compared to a Neolithic population, is also present in the proximal humerus
(Scherf et al., 2015). Chirchir et al. (2017) quantified trabecular bone fraction from pQCT data in
the forelimb and hindlimb of five groups of modern humans, with a range of lifestyles, from
foraging to industrial sedentary populations. This revealed a reduction in hindlimb robusticity
with increased sedentism, and more variable changes in forelimb robusticity. Variability in
trabecular architecture across the skeleton of recent humans has been documented, largely in the
clinical literature. There is high intra-individual variability in trabecular structure, with low
correlation between anatomical sites in several measures of trabecular architecture, quantified
using 2D and 3D stereological methods (Amling et al., 1996; Parkinson and Fazzalari, 2003),
pQCT (Groll et al., 1999; Chirchir, 2016), and microCT (Hildebrand et al., 1999; Ulrich et al.,
1999; Eckstein et al., 2007). However, as yet, no study has conducted a comprehensive
trabecular analysis, including parameters other than trabecular bone volume, across several
skeletal elements in humans in comparative context with other primates. Thus, it remains

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unknown how potential systemic patterns in trabecular bone might vary intraspecifically and

interspecifically. In this study we address this issue through quantification of trabecular bone volume fraction (BV/TV), degree of anisotropy (DA) and trabecular thickness (Tb.Th) in several anatomical sites within associated skeletons of recent humans and chimpanzees. Based on previous findings described above, we test three predictions: first, we predict that chimpanzees will have a higher BV/TV throughout the skeleton compared to humans (Maga et al., 2006; Griffin et al., 2010; Tsegai et al., 2013; Chirchir et al., 2015). Second, as humans and chimpanzees adopt locomotor behaviours that involve differential loading of the forelimb and hindlimb, we predict that BV/TV will be relatively similar across both limbs in chimpanzees, while BV/TV will be low across the forelimb compared to the hind limb in humans. Previous studies have demonstrated that humeral and femoral head trabecular structure does not reflect this difference in locomotor loading (Fajardo and Müller, 2001; Ryan and Walker, 2010; Shaw and Ryan, 2012), thus in this study we aim to test whether this pattern is consistent in other elements of the fore- and hindlimb. Third, as trabecular fabric has previously been associated with load direction and variability, we expect DA to differ between taxa in ways that reflect loading differences (Ryan and Ketcham, 2002b; Barak et al., 2013b; Su et al., 2013). Although Tb.Th is strongly correlated with body size (Doube et al., 2011; Barak et al., 2013a; Fajardo et al., 2013; Ryan and Shaw, 2013), it is also highly correlated with BV/TV (Barak et al., 2013a), and as such could parallel the systemic pattern of BV/TV. However, since the taxa in this study sample have a similar body mass, we predict that there will be no differences in trabecular thickness between these taxa, as has been found in general in previous studies (Cotter et al., 2009; Scherf et al., 2013; Ryan and Shaw, 2015; Zeininger et al., 2016; but see Barak et al., 2013b; Su and Carlson, 2017).

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Methods

207 Sample

Trabecular bone structure was analysed in the skeletons of *Pan troglodytes* (N = 7) and recent Homo sapiens (N = 7) individuals. Full details of the study sample are shown in Table 1. All chimpanzee specimens belong to a single subspecies, P. t. verus, and were wild-collected skeletons from the Taï National Park, Republic of Côte d'Ivoire. The human sample was collected from two skeletal collections: one from a 19th century cemetery in Inden. Germany and the other from 13-15th century medieval cemeteries in Canterbury, UK. All specimens were free from external signs of pathology. Trabecular architecture was quantified in two anatomical locations in the forelimb (humeral head and third metacarpal head [MC3]), four anatomical sites in the hindlimb (femoral head, distal tibia, talus, and third metatarsal head [MT3]) and one site in the axial skeleton (first thoracic vertebra [T1]) (Fig. 1). These anatomical sites were chosen to include elements from both limbs, and an element from the axial skeleton that is less affected by differential loading of the fore- and hindlimb. We aimed to sample all bones of the forelimb and hindlimb from the same side, but when elements were not adequately preserved, all elements from either the forelimb or hindlimb were taken from the contralateral side where possible. For example, if the right femur was absent, then the femur, tibia, talus and MT3 were taken from the left side where possible.

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Micro-CT scanning

All specimens were CT scanned using either a SkyScan 1173 or a BIR ACTIS 225/300 scanner housed at the Department of Human Evolution, Max Planck Institute for Evolutionary Anthropology (Leipzig, Germany). All scans were reconstructed as 16-bit tiff stacks with isotropic voxel sizes of 21-38 µm. All specimens were reoriented into standardised anatomical positions and were downsampled, due to computational constraints, using Avizo 6.3. Specimens

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were analysed at a range of resolutions (25-45µm), with adequate representation of trabeculae as demonstrated by the relative resolution (4.25-9.83), which indicates how many pixels represent the average trabecular strut (Sode et al., 2008). Following this, all specimens were segmented using the Ray Casting Algorithm of Scherf and Tilgner (2009).

Trabecular bone quantification

Analysis of trabecular bone structure was conducted using an in-house script in medtool v3.9 (www.dr-pahr.at), following Gross et al. (2014). Morphological filters were used to automatically segment the cortical and trabecular bone, resulting in definition of three materials: (1) cortical bone, (2) trabecular bone and (3) air inside the bone (Fig. 2A). In this way, the trabecular bone throughout an entire region (or the whole bone, in the case of the talus) could be analysed. Tb.Th was calculated using the BoneJ plugin (v1.3.12; Doube et al., 2010) for ImageJ (v1.46r; Schneider et al., 2012) from the segmented trabecular only region (Fig. 2B). To quantify the other trabecular parameters in medtool (following protocols outlined in Gross et al., 2014), a 2.5mm background grid was applied to each specimen, and a 5mm spherical volume of interest was used to measure BV/TV at each node of the background grid. A 3D tetrahedral mesh was created of the inner region of the bone (Fig. 2C), to which each node was assigned a BV/TV value (Fig. 2D) interpolated from the background grid. A mesh size of 1mm was used for the larger specimens (humeral head, femoral head, distal tibia, and talus) and a mesh size of 0.5mm for the smaller specimens (MC3, MT3, and T1). As the background grid size was constant for the sample, the results are independent of mesh size. The overall BV/TV was calculated as the mean of all elements in the 3D region of interest (ROI; see below). The mean intercept length method was used to calculate the local fabric tensor for each tetrahedron and these were normalised by the determinants (Luisier et al., 2014). Similar to BV/TV, an arithmetic mean of

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Il of the second order fabric tensors was computed within the ROI. The DA was calculated as ne DA = 1 - [smallest eigenvalue/largest eigenvalue], such that a DA of 1 represents "complete" nisotropy (i.e. all trabeculae are aligned, and there are no crossing trabeculae) and a DA of 0 eflects complete isotropy (i.e. there is no preferential alignment of trabeculae). Often the DA is ound between a DA of 1 representing isotropy and a DA > 1 representing increasing anisotropy, owever here we use an alternative, "normalised" DA. both humans and chimpanzees trabecular bone of the long bone epiphyses extends beyond the piphysis and into the shaft. As such, the ROIs for long bones were defined in order to sample as nuch of the trabecular bone-filled region as possible, which could potentially contribute to ystemic differences in trabecular structure. For each skeletal element the ROI was defined as ollows (Fig. 1). For the proximal humerus, this was defined as the point where curvature of the umeral head begins to expand from the shaft both medially and laterally (Fig. 1A). In the roximal femur, the femoral head was extracted with the inferior margin being at the most nferior point of the femoral head and the medial margin at the most medial point of the femoral ead (Fig. 1B). In the proximal femur, it was only possible to sample the femoral head, and small egion of the femoral neck, due to computational constraints in processing large data sets. The OI in the distal tibia was defined distally where curvature of the shaft begins in both medial and nterior views, which is at the proximal extent of the fibular notch (Fig. 1C). In the MC3 and IT3, the distal end (head) was defined as the point at which the shaft curves laterally in almar/plantar view (Fig. 1D & E). In the T1, only the trabeculae in the vertebral body were uantified (Fig. 1F). For the talus the trabecular bone in the entire element was quantified. Identification of homologous regions is complex due to the potential effect of differences in

location and size of the region being analysed. For example, sometimes dramatic differences in

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quantification of trabecular bone structure have been found with variation in position or size of small volumes of interest within a bone or epiphysis (Fajardo and Müller, 2001; Kivell et al., 2011; Lazenby et al., 2011b). Here, our 3D ROI includes a much larger region of trabecular structure (e.g. the entire epiphysis), but quantified values may also be affected by how the ROI is defined between taxa. Therefore, a test of intra-observer error was conducted for the humerus and tibia of one human and one chimpanzee, with the ROI defined five times on five consecutive days. The percentage difference in BV/TV compared to the original quantified value, ranged from -0.97% to 0.22% for the humerus and from -2.29% to 0.73% for the tibia.

Statistical analysis

Statistical analysis was conducted using R v3.3.2 (R Core Team, 2016) and ggplot2 (Wickham, 2009) for plot generation. Due to small sample sizes non-parametric tests were used. Taxonomic differences in trabecular structure at each anatomical site were tested for using Mann-Whitney U tests between taxa. To identify systemic patterns within species, Friedman tests were used to identify whether there were overall significant differences between the ranks of anatomical sites in humans and in chimpanzees. Following the results of the Friedman tests, Wilcoxon exact tests with p-values corrected with a post-hoc Bonferroni adjustment, were used to identify significant pairwise differences between anatomical sites within humans and within chimpanzees.

Differences in the systemic pattern between taxa were identified by comparing the results of within-species Wilcoxon exact tests. To identify correlations between trabecular parameters in different regions within humans and within chimpanzees. Spearman's correlation test was used with p-values corrected with a post-hoc Bonferroni adjustment. For all statistical tests a p-value < 0.05 was considered significant.

Results

Taxonomic differences

Means and standard deviations of trabecular parameters in each anatomical region and results of Mann-Whitney U tests for significant differences between species are shown in Table 2. Figure 3 shows box-and-whisker plots of the results for each taxon. There were no significant differences in Tb.Th between chimpanzees and humans in any anatomical region. Chimpanzees had significantly higher BV/TV than humans in the humeral, femoral, and MT3 heads as well as the talus. Chimpanzees also had significantly more anisotropic trabeculae in the humeral head and T1, and less anisotropic trabeculae in the talus and MT3.

Taxonomic differences in the patterning of BV/TV are further illustrated in Figure 4, where the BV/TV values are shown for each individual. In one human individual BV/TV values were much higher in every anatomical region, and this is the only individual that overlapped with chimpanzees in humeral, metatarsal, femoral, and talar BV/TV. Excluding this specimen from the statistical comparisons presented above led to significantly lower BV/TV in the human MC3 (p = 0.03), while the BV/TV values in the thoracic vertebra and tibia approached significance (p = 0.05).

Intraspecific and interspecific systemic patterns

Comparisons of trabecular structure within individuals are presented in Table 3, as the mean rank of each element for each trabecular parameter. This demonstrates the systemic pattern of trabecular bone structure within each taxon, with elements having a higher mean rank indicating generally higher values in that anatomical region across individuals. Across both chimpanzees and humans, all hindlimb elements, except for the MT3, had a higher mean rank for Tb.Th than

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the forelimb and axial elements, and the humerus had a higher mean rank for Tb.Th than the
metacarpal. In chimpanzees, the order of mean ranks of the different anatomical sites for BV/TV
was similar to that of Tb.Th. The only difference was a switch between the humerus and the T1.
In humans, the ranks of anatomical sites for BV/TV followed the pattern for Tb.Th less closely.
Notably, the humerus was the lowest ranking element for BV/TV in humans. The mean ranks of
DA differed between the taxa. Within the hindlimb of chimpanzees, the DA had the highest
mean rank in the tibia, MT3 and femur, with the talus having the most isotropic trabeculae. The
pattern in humans differed from that of chimpanzees in that the MT3 had a higher DA rank
compared to the other hindlimb anatomical sites. In the forelimb, the MC3 had a higher mean
rank for DA than the humerus in both taxa.
Results of Friedman tests (Table 3) indicated the presence of significant differences between
ranks of anatomical sites in all three trabecular parameters in both humans and chimpanzees.
Post-hoc Wilcoxon test comparisons with a Bonferroni adjustment are shown in Table 4. For
Tb.Th (Table 4), significant differences were largely due to thicker trabecular bone in the femur,
tibia and talus compared to other elements in both taxa. The humerus had significantly thicker
trabeculae than the MT3 in humans, and both the MC3 and MT3 in chimpanzees. Significant
differences in BV/TV between elements were largely due to low BV/TV in the human humerus
and to high BV/TV in the chimpanzee femur and talus (Table 4). Significant differences in DA
were largely due to high DA in the tibia and low DA in the talus in chimpanzees. In humans,
most significant differences were due to the high DA of the MT3

Trabecular correlations between anatomical sites

Spearman's correlation tests, to identify whether trabecular parameters were correlated between anatomical sites within each taxon, revealed only two significant correlations. In chimpanzees,

there was a significant correlation in Tb.Th between the humerus and femur (r = 0.96, p = 0.01) and between the talus and MT3 (r = 1.00, p < 0.01). There were no significant correlations between anatomical sites in humans.

348 Discussion

This study provides the first comprehensive 3D analysis of potential systemic patterns in trabecular architecture across the skeleton of humans and chimpanzees using a whole bone/region approach. We find both similarities and differences in regional patterning of trabecular structure across individuals and between taxa. Due to substantial variation in the morphology of the bones/epiphyses included in this study, direct comparison of trabecular bone architecture between anatomical sites is complex, as it may be influenced by factors such as articular surface area or the proximo-distal location of the element (Chirchir, 2015; Saers et al., 2016; for cortical bone see Lieberman et al., 2003). However, by identifying both shared and distinct systemic patterns of trabecular structure, relative (rather than absolute) comparisons can be made across anatomical sites and between taxa. In this comparative context, we find that the systemic pattern of BV/TV, Tb.Th and DA differs between chimpanzees and humans. However, this pattern is not always consistent across the skeleton, or clearly related to joint function based on predicted loading during locomotion.

Taxonomic differences in BV/TV

Recent modern humans have been found to have a lower BV/TV than non-human primates in various anatomical sites (e.g. Maga et al., 2006; Griffin et al., 2010; Shaw and Ryan, 2012; Scherf et al., 2013; Tsegai et al., 2013; Chichir et al., 2015; Ryan and Shaw, 2015), thus we predicted that chimpanzees would have higher BV/TV in all anatomical regions sampled in our study. We find general support for this hypothesis, with chimpanzees having significantly higher

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368	BV/TV than humans in the humeral, femoral and MT3 heads and the talus, and higher mean BV/
369	TV values, but not significantly so, in the distal tibia, MC3 and T1. Thus, using a whole-bone/
370	region approach across the skeletons of the same individuals, our results provide further support
371	of a general pattern of higher BV/TV in chimpanzees compared with humans documented in
372	previous studies.
373	Recent trabecular analyses have demonstrated the potential influence of activity levels on
374	trabecular architecture in modern humans, including BV/TV quantified from micro-CT scans or
375	converted from pQCT measures of volumetric mineral density (Chirchir et al., 2015; Ryan and
376	Shaw, 2015; Scherf et al., 2015; Saers et al., 2016; Chirchir et al., 2017). Recent modern humans
377	have lower BV/TV, calculated from pQCT scans, in both the upper and lower limb compared to
378	early modern humans and other fossil hominins, including <i>H. neanderthalensis</i> and members of
379	Australopithecus (Chirchir et al., 2015). The trabecular architecture in the centre of the proximal
380	humerus of recent modern humans is weaker (e.g. lower BV/TV and Tb.Th) than in Neolithic
381	modern humans (5,700-4,900BP) (Scherf et al., 2015). The timing of this reduction in BV/TV
382	may be related to changes in overall activity level, with recent mobile foragers having stronger
383	bone (higher BV/TV, higher Tb.Th, lower bone surface to volume ratio) in the proximal and
384	distal femur and tibia compared to recent sedentary agriculturalists (Ryan and Shaw, 2015; Saers
385	et al., 2016) and differences in trabecular BV/TV, quantified using pQCT, in particular of the
386	lower limb, can be related to subsistence strategy in recent populations (Chirchir et al., 2017).
387	In the sample included in this study, one human individual has higher BV/TV in every region of
388	the skeleton, which overlaps with chimpanzees in all anatomical locations. Unfortunately, no
389	historical information is available regarding the activity level or occupation of this individual.
390	However, it provides further support for a systemic pattern of trabecular BV/TV that could be

related to systemic factors, such as higher activity levels promoting bone remodelling throughout the skeleton (Lieberman, 1996). Across canids, felids and cercopithecines, species with longer travel distances have a higher relative trabecular bone mass, quantified from pQCT, than species with shorter travel distances, indicating the potential influence of overall activity on trabecular structure in a range of taxa (Chirchir et al., 2016a).

An explanation is not readily available for the high BV/TV in chimpanzees, in comparison to both active populations of humans and other primate taxa. In the femoral head, chimpanzees have higher BV/TV than closely related *Gorilla* and modern humans, having the highest BV/TV amongst 32 primate taxa (Ryan and Shaw, 2013), and when compared to humans with different subsistence strategies (Ryan and Shaw, 2015). In the humeral head, chimpanzees have higher BV/TV than Neolithic modern humans, recent modern populations and *Pongo* (Scherf et al., 2013; Scherf et al., 2015). Thus, activity levels alone may not explain the systemic difference in BV/TV between humans, chimpanzees, and other primate taxa. This is of particular importance for functional inferences drawn from trabecular structure in fossil hominins, where some anatomical regions or isolated specimens are also characterised by high trabecular BV/TV, similar to or higher than that of chimpanzees (Barak et al., 2013b; Chirchir et al., 2015; Skinner et al., 2015).

Functional signals in systemic patterns of BV/TV

We predicted that the patterns of trabecular BV/TV in the forelimb and hindlimb of chimpanzees and humans would reflect differential loading during locomotion, such that quadrupedal chimpanzees would have more similar BV/TV values in the forelimb and hindlimb, whereas bipedal humans would have higher BV/TV in the hindlimb elements. It is important to make comparisons between elements at a similar anatomical location due to the proximo-distal

414	decrease in trabecular bone mass (bone mineral density measured using pQCT and multiplied by
415	joint size) and BV/TV in hominoids and populations of humans with different subsistence
416	strategies (Chirchir, 2015; Saers et al., 2016). Thus, here we discuss differences between the
417	humeral and femoral head and between the MC3 and MT3 head.
418	We find that both chimpanzees and humans have significantly higher BV/TV in the femoral head
419	compared with the humeral head. This is consistent with previous comparisons of trabecular bone
420	in the humerus and femur in a range of anthropoid species, where all individuals (Fajardo and
421	Müller, 2001; Ryan and Walker, 2010), or the majority of individuals (Shaw and Ryan, 2012),
422	were found to have higher BV/TV in the femoral head compared to the humeral head. Mean
423	trabecular BV/TV, derived from micro-CT and pQCT, is higher in the femoral head compared to
424	the humeral head in extant chimpanzees, modern humans, early modern humans, and H .
425	neanderthalensis (but not in Australopithecus africanus) (Chirchir et al., 2015; Chirchir, 2016),
426	but this difference is not significant in modern humans (Chirchir, 2016). Previous analyses of
427	proximal femoral trabecular properties in humans, although not incorporating the humeral head,
428	or the same anatomical sites as the present study, have also found relatively high trabecular BV/
429	TV in the femoral neck (Amling et al., 1996; Eckstein et al., 2007 [in men but not women]) and
430	femoral head (Hildebrand et al., 1999; Ulrich et al., 1999; Parkinson and Fazzalari, 2003)
431	compared to other anatomical sites analysed (but see Chirchir, 2016).
432	However, the skeletal pattern is more complex when the BV/TV of other anatomical sites is
433	considered. We find that, compared to other anatomical regions, chimpanzees have very high
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435	femoral BV/TV, having the highest mean rank of all anatomical sites, whereas in humans femoral
436	BV/TV ranks lower than the talus. In contrast, humeral BV/TV in humans has the lowest mean
	rank, whereas in chimpanzees it ranks above the MT3 and MC3. Thus, chimpanzees have

relatively high femoral BV/TV and humans have very low humeral BV/TV, compared to other
anatomical sites. This finding supports our prediction that trabecular BV/TV would reflect
reduced loading of the human forelimb, but the pattern in chimpanzees does not support our
prediction of similar loading between the two limbs. This could be due to the 'hindlimb driven'
quadrupedal locomotion of chimpanzees, and other primate taxa, whereby the hindlimb
experiences greater vertical reaction forces than the forelimb, and propulsion is driven by the
hindlimb (Kimura et al., 1979; Demes et al., 1994). Thus, high BV/TV in femoral head of
chimpanzees and other primate taxa may reflect this difference in function of the hindlimb
during quadrupedal locomotion.
Comparisons between the MC3 and MT3 also do not support the hypothesis of higher BV/TV in
the hindlimb of humans and more similar BV/TV between the forelimb and hindlimb of
chimpanzees. On average, both humans and chimpanzees have higher BV/TV in the MC3
compared to the MT3, and, in contrast to our predictions, this pattern is more pronounced in
humans. In all human specimens in the study sample, and in 57% of the chimpanzees, the MC3
has higher BV/TV than the MT3, with this difference being significant in humans. This is
consistent with previous findings, where on average bone density in humans is higher in the
metacarpal head while in chimpanzees it is higher in the metatarsal head (Chirchir et al., 2015).
Thus, comparisons of BV/TV (derived both from micro-CT and pQCT scans) between the MT3
and MC3 does not reflect higher loading of the human hindlimb and more equal loading of the
forelimb and hindlimb in chimpanzees. These patterns identified between the femoral and
humeral heads, the MC3 and MT3, and throughout the skeleton may reflect the complex
relationship between mechanical load, activity level, and anatomical site (Judex et al., 2004;
Wallace et al., 2012; Wallace et al., 2013; Wallace et al., 2015).

)	Taxonomic differences and systemic patterning of DA and Tb.Th
L	Trabecular structure across the skeleton of humans and chimpanzees supports our prediction that
2	there would be no consistent taxonomic differences in DA. We found no consistent pattern in
3	DA values across the seven anatomical regions within each species. Humans had significantly
	more anisotropic trabeculae in the talus and MT3, and significantly more isotropic trabeculae in
	the humeral head and T1 compared to chimpanzees. This variability between taxa and
	anatomical sites may indicate that DA is primarily reflecting differences in joint loading (see
	below).
	Tb.Th has previously been found to scale with body size in a range of primate taxa and
	anatomical sites (Doube et al., 2011; Barak et al., 2013a; Fajardo et al., 2013; Ryan & Shaw,
	2013), but also to correlate with BV/TV (Barak et al., 2013a). Here, in support of our prediction,
	we found no significant differences in absolute Tb.Th between humans and chimpanzees.
	Considering the smaller body size of chimpanzees, this indicates that they have relatively thick
	trabeculae compared to humans, however due to the small difference in body size this is unlikely
	to lead to significant differences. We did, however, find that the systemic pattern of Tb.Th
	followed a similar pattern in both taxa, being generally higher in the hindlimb (femoral head,
	talus and distal tibia) and lower in the forelimb (humerus and MC3) in both taxa. This is
	supported by previous comparisons of Tb.Th between the humerus and femur, which found
	thicker femoral trabeculae in most taxa/individuals (Ryan and Walker, 2010; Shaw and Ryan,
	2012; Ryan and Shaw, 2013). However, the MT3 had thin trabecular bone compared to the rest
	of the hindlimb in both humans and chimpanzees, despite different loading regimes between
	these two taxa. Differences in BV/TV, but not Tb.Th, indicate potential differences in trabecular
	in the control of the

number (Tb.N) between these taxa. Previous studies have found differences in Tb.N between

humans and chimpanzees (e.g. distal tibia: Su, 2011; Barak et al., 2013b; vertebra: Cotter et al., 2009; femoral head: Ryan and Shaw, 2012; Shaw and Ryan, 2012; humeral head: Ryan and Shaw, 2012; Shaw and Ryan, 2012; Scherf et al., 2013) with chimpanzees having more numerous trabeculae, although this is not the case for the talus (Su, 2011; DeSilva and Devlin, 2012) or calcaneus (Kuo et al., 2013; Zeininger et al., 2016).

Functional signals in systemic patterns of DA

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The degree of anisotropy of trabeculae, and other related measures such as primary trabecular orientation and elongation index, are often able to distinguish between locomotor mode, especially when comparisons are made between different regions of an epiphysis (e.g. MacLatchy and Müller, 2002; Ryan and Ketcham, 2002b; Maga et al., 2006; Griffin et al., 2010; Hebert et al., 2012; Barak et al., 2013b; Su et al., 2013; Zeininger et al., 2016; Su and Carlson, 2017). However, not all trabecular analyses have identified differences in DA or orientationbased variables between locomotor groups (e.g. Fajardo et al., 2007; Kuo et al., 2013). In general, DA is thought to reflect the range of joint positions in which a joint experiences high loads, with more uniformly aligned trabeculae being associated with more stereotypical load orientations, and more isotropic trabeculae with a greater range of adopted joint positions (Fajardo and Müller, 2001; Ryan and Ketcham, 2002b). There is evidence of a systemic pattern in a proximo-distal increase in DA in the human femur and tibia (Saers et al., 2016), which is also found in the present study between the proximal femur and distal tibia. However, this could be a structural adaptation to the proximo-distal reduction in BV/TV, or could be related to other factors, such as differences in gross morphology, and thus loading stereotypy, between the femur and tibia (Saers et al., 2016).

505	We predicted that DA in the hindlimb and humeral head of chimpanzees would reflect
506	differences in loading between the study taxa. In general, humans experience more stereotypical
507	loading of the hindlimb than chimpanzees, whose locomotor repertoire includes knuckle-walking
508	quadrupedalism and several arboreal behaviours (e.g. climbing, clambering and suspension) that
509	require a greater range of joint positions (Hunt, 1991; Doran, 1992, 1993b, 1997; Sarringhaus et
510	al., 2014). We find some support for this prediction. The hindlimb of humans has significantly
511	higher DA in the MT3 head and talus compared to chimpanzees, perhaps reflecting the more
512	stereotypical loading during bipedalism, especially in the foot. Moreover, DA is significantly
513	higher in the MT3 than the MC3 of humans, but not in chimpanzees. However, this is not the
514	case for the distal tibia, where chimpanzees have higher DA (contrary to Barak et al., 2013b). In
515	the chimpanzee forelimb, we find significantly higher DA in the humeral head (contrary to
516	Scherf et al., 2013), and higher mean DA in the metacarpal head (supporting the findings of
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518	Tsegai et al., 2013; Chirchir et al., 2016b) compared to humans. In the T1 we find significantly
519	higher DA in chimpanzees compared to humans. A previous analysis of DA in eighth thoracic
520	vertebra found no significant difference in DA between chimpanzees and humans, but did
	identify a negative correlation between BV/TV and DA in humans, which was absent in non-
521	human apes (Cotter et al., 2009), indicating a complex interplay between these trabecular
522	parameters in the spinal column.
523	Although DA appears to correspond with the type of loading in some anatomical sites, other
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525	anatomical areas do not (e.g. the humeral head and distal tibia), nor do they always support the
	findings of previous studies. This may be related to the whole-region method applied in this
526	study, where trabecular bone from a larger region is quantified, in comparison to previous studies
527	where DA was measured in smaller sub-regions (e.g. volume of interest). Whether trabecular

alignment in a small subregion, or in an entire region, is a better indicator of overall loading is unclear. Another potential explanation, is that our predictions of joint loading are often oversimplified, and the impact of different behaviours on bone structure is unknown. For example, a lower DA might have been expected for the chimpanzee humeral head, based on their adoption of a range of arboreal behaviours and thus varied load orientations. However, knucklewalking is the most frequent locomotor behaviour used by adult chimpanzees (Doran, 1992; Sarringhaus et al., 2014), and as such, may contribute more to trabecular anisotropy than less frequent arboreal locomotor bouts.

Trabecular structure and articular morphology

Comparisons of trabecular bone structure between anatomical regions, or indeed of the same anatomical region between different taxa, are potentially influenced by differences in the gross morphology of the articular region, and by articular function. Primate taxa differ in relative articular surface area and absolute articular size, due to differences in both the magnitude of load and the range of joint excursion, which can be related to locomotor mode (Ruff, 1988; Godfrey et al., 1991; Ruff and Runestad, 1992; Godfrey et al., 1995; Ruff, 2002). Moreover, the relationship between articular surface area and joint mobility may differ between joint types; for example in a ball-and-socket joint, an increase in surface area may have more of an impact on joint mobility than in a hinge joint (Ruff, 2002). Although our discussion has focused largely on the comparative context, i.e. differences in the systemic pattern between humans and chimpanzees, it is important to recognise the potential impact of these aspects of external joint morphology on the findings of this study. It is beyond the scope of the present study to explore this further, however, it is an important and relatively unexplored area of trabecular research (but see Rafferty and Ruff, 1994). Future research into systemic patterns of trabecular structure

should further investigate the relationship between trabecular morphology and external articular morphology, both within and between taxa.

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Conclusion

Here we demonstrate that an understanding of the way in which trabecular bone varies across the skeleton can have important implications for inferring joint load, function, and ultimately behaviour, from trabecular structure. Chimpanzees and humans have systemically different trabecular BV/TV throughout their skeleton, such that humans (except for one individual within our sample) had lower BV/TV in all anatomical regions compared with chimpanzees. However, differences in BV/TV between the humeral and femoral head and the MC3 and MT3 do not directly reflect predicted differences in loading of the fore- and hindlimb in each taxon. Rather, overall BV/TV may be driven by other factors, such as overall activity level (Ryan and Shaw, 2015). Mean Tb.Th values across the skeleton do not differ significantly between chimpanzees and humans, and trabeculae are generally thicker in the hindlimb compared with the forelimb in both taxa. These systemic patterns must be considered when inferring the magnitude of joint load in any one skeletal area (e.g. high BV/TV may not necessarily reflect solely higher load/activity levels). This is particularly true, but also especially challenging, when inferring function in fossil taxa when only isolated elements are preserved, and thus potential systemic patterns are unknown. In contrast to BV/TV, the degree to which trabeculae are preferentially oriented (DA) did not differ consistently across the skeleton within chimpanzees or humans. Although the pattern of DA across different skeletal elements did not always fit our predictions, the pattern suggests that trabecular alignment may more directly reflect differences in the magnitude and direction of joint loading, and thus behaviour, than BV/TV (and Tb.Th).

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Author contributions

- Concept/design: ZJT, MMS, JJH, TLK; Acquisition of data: ZJT; Data analysis/interpretation: ZJT,
- 575 MMS, DHP, TLK; Drafting and revision of the manuscript: ZJT, MMS, DHP, JJH, TLK; Approval of the
- article: ZJT, MMS, DHP, JJH, TLK

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863 Tables

Table 1. Study sample

Taxon	Collection ¹	Specimen ID	Sex	Elements
H. sapiens	UG	INDEN_91	M	R Hum, R MC3 R Fem, R Tib, R Tal, L MT3 T1
H. sapiens	UG	INDEN_113	M?	R Hum, L MC3 R Fem, L Tib, L Tal, L MT3 T1
H. sapiens	UG	INDEN_118	F	R Hum, R MC3 R Fem, L Tib, L Tal, L MT3
H. sapiens	UG	INDEN_311	M	T1 R Hum, R MC3 R Fem, L Tib, L Tal, R MT3 T1
H. sapiens	UK	NGA_88_SK_766	U	L Hum, L MC3 R Fem, R Tib, R Tal, R MT3 T1
H. sapiens	UK	NGA_88_SK_825	U	R Hum, R MC3 L Fem, L Tib, R Tal, R MT3 T1
H. sapiens	UK	NGA_88_SK_880	U	R Hum, R MC3 L Fem, L Tib, L Tal, L MT3 T1
P. troglodytes verus	MPIEVA	MPITC_11781	M	L Hum, L MC3 R Fem, R Tib, R Tal, R MT3
P. troglodytes verus	MPIEVA	MPITC_11778	F	T1 L Hum, L MC3 R Fem, R Tib, R Tal, R MT3
P. troglodytes verus	MPIEVA	MPITC_14996	F	T1 L Hum, L MC3 R Fem, R Tib, R Tal, R MT3
P. troglodytes verus	MPIEVA	MPITC_15001	F	T1 L Hum, L MC3 R Fem, R Tib, R Tal, R MT3
P. troglodytes verus	MPIEVA	MPITC_15002	F	T1 L Hum, L MC3 R Fem, R Tib, R Tal, R MT3
P. troglodytes verus	MPIEVA	MPITC_15012	M	T1 R Hum, L MC3 R Fem, R Tib, R Tal, L MT3
P. troglodytes verus	MPIEVA	MPITC_15013	F	T1 L Hum, L MC3 R Fem, R Tib, L Tal, R MT3 T1

MPIEVA – Max Planck Institute for Evolutionary Anthropology, UK – University of Kent, UG – University of
 Göttingen

²M – Male, F – Female, U – Unknown, ? – indicates uncertainty concerning sex. Data taken from collection records.

Table 2. Trabecular structure in each taxon across anatomical sites. Mean values with standard deviation in parentheses, and p-values resulting from Mann-Whitney U tests between taxa. Significant differences are shown in bold.

Element	Taxon	Tb.Th (mm)	BV/TV (%)	DA
Humerus	Ното	0.21 (0.02)	12.72 (4.07)	0.11 (0.04)
	Pan	0.22 (0.02)	25.32 (3.82)	0.17 (0.02)
	p-value	0.90	< 0.01	< 0.01
MC3	Ното	0.19 (0.02)	21.25 (3.16)	0.20 (0.08)
IVICS	Pan	0.19 (0.02)	22.75 (1.58)	0.20 (0.08)
	- ****	0.18 (0.01)	0.16	0.23 (0.04)
	p-value	0.32	0.10	0.40
T1	Homo	0.22 (0.04)	21.29 (5.91)	0.12 (0.05)
	Pan	0.20 (0.02)	26.08 (3.78)	0.18 (0.05)
	p-value	0.38	0.16	0.03
Femur	Ното	0.26 (0.03)	22.72 (5.45)	0.16 (0.05)
Tomar	Pan	0.33 (0.07)	38.58 (6.85)	0.08 (0.09)
	p-value	0.07	<0.01	0.13
Tibia	Ното	0.26 (0.02)	21.66 (3.11)	0.29 (0.06)
1101a	Pan	0.24 (0.03)	25.98 (4.31)	0.29 (0.00)
	p-value	0.24 (0.03)	0.10	0.34 (0.03)
	P			
Talus	Ното	0.27 (0.03)	26.26 (3.43)	0.11 (0.06)
	Pan	0.31 (0.04)	35.94 (3.87)	0.02 (0.03)
	p-value	0.07	< 0.01	< 0.01
MT3	Ното	0.17 (0.02)	17.54 (3.47)	0.31 (0.03)
	Pan	0.18 (0.03)	22.89 (3.93)	0.22 (0.03)
	p-value	0.90	0.01	<0.01

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		Rank			
Taxon	Element	Tb.Th	BV/TV	DA	
Ното	Humerus	3.43	1.00	2.29	
	MC3	2.29	4.29	4.57	
	T1	3.57	4.43	2.43	
	Femur	5.71	5.14	3.71	
	Tibia	6.00	4.29	6.14	
	Talus	6.00	6.57	2.29	
	MT3	1.00	2.29	6.57	
	p-value	<0.01	<0.01	<0.01	
Pan	Humerus	3.57	3.57	3.29	
	MC3	2.14	2.00	5.29	
	T1	3.00	3.71	3.86	
	Femur	6.57	6.85	2.57	
	Tibia	4.86	4.00	6.86	
	Talus	6.43	6.14	1.00	
	MT3	1.43	1.71	5.14	
	p-value	<0.01	<0.01	<0.01	

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Table 4. Comparison between anatomical regions within each taxon. P-values from pairwise Wilcoxon tests with a post-hoc Bonferroni correction between all anatomical sites in *Homo* (shaded) and *Pan* (unshaded). Significant differences are shown in bold.

		Humerus	MC3	T1	Femur	Tibia	Talus	MT3
Tb.Th	Humerus		0.146	1.000	0.086	0.049	0.024	0.012
	MC3	0.024		1.000	0.012	0.012	0.012	1.000
	T1	1.000	0.795		1.000	0.233	0.367	0.795
	Femur	0.049	0.012	0.024		1.000	1.000	0.012
	Tibia	1.000	0.012	0.795	0.551		1.000	0.012
	Talus	0.012	0.012	0.012	1.000	0.147		0.012
	MT3	0.367	1.000	1.000	0.024	0.086	0.012	
BV/TV	Humerus		0.086	0.147	0.049	0.086	0.012	0.551
	MC3	1.000		1.000	1.000	1.000	0.367	0.367
	T1	1.000	0.795		1.000	1.000	1.000	1.000
	Femur	0.086	0.012	0.049		1.000	1.000	1.000
	Tibia	1.000	1.000	1.000	0.086		0.795	0.367
	Talus	0.012	0.012	0.024	1.000	0.024		0.086
	MT3	1.000	1.000	1.000	0.024	1.000	0.012	
DA	Humerus		0.551	1.000	1.000	0.012	1.000	0.012
	MC3	0.049		0.551	1.000	1.000	0.551	0.024
	T1	1.000	1.000		1.000	0.012	1.000	0.012
	Femur	1.000	0.147	1.000		0.233	1.000	0.012
	Tibia	0.012	0.086	0.024	0.012		0.012	1.000
	Talus	0.012	0.012	0.012	0.795	0.012		0.012
	MT3	0.086	1.000	1.000	0.551	0.049	0.012	

Figure legends

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Figure 1. Region of interest defined for each element. Grey boxes represent the definition of each region in specimens of *Pan* for (A) humeral head, (B) femoral head, (C) distal tibia, (D) third metacarpal head, (E) third metatarsal head, and (F) first thoracic vertebral body (shown in a midsagittal section, as transverse process obscures a clear view of the vertebral body). For the talus, not shown here, trabecular structure was quantified throughout the entire bone.

Figure 2. Quantification of trabecular bone. (A) Segmented voxel data where cortex, trabecular bone and air inside the bone are assigned different grey values. (B) Trabecular only region which was imported into BoneJ to measure Tb.Th. (C) 3D tetrahedral mesh of cortex and inner region of bone. (D) Each element in the tetrahedral mesh of the inner region was assigned a BV/TV value, as visualised here where regions of low BV/TV are in blue and high BV/TV in red.

Figure 3. Variation in trabecular bone structure across the skeleton of *Homo* and *Pan*. Boxplots showing (A) Tb.Th, (B) BV/TV and (C) DA in the humeral head (Hum), third metacarpal head (MC3), femoral head (Fem), distal tibia (Tib), talus (Tal), third metatarsal head (MT3), and first thoracic vertebra (T1) in *Homo* (red) and *Pan* (blue). Significant differences are indicated by brackets with * for p<0.05 and ** for p<0.01.

Figure 4. Systemic differences in BV/TV across the skeleton of *Homo* (red) and *Pan* (blue). BV/TV in each individual of *Homo* (red) and *Pan* (blue) in the humeral head (Hum), third metacarpal head (MC3), femoral head (Fem), distal tibia (Tib), talus (Tal), third metatarsal head (MT3), and first thoracic vertebra (T1)

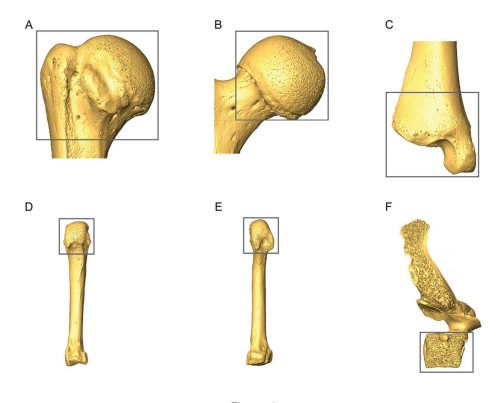


Figure 1 110x80mm (300 x 300 DPI)

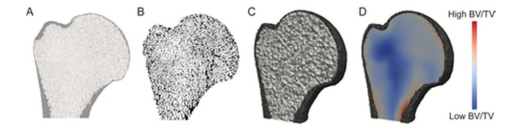


Figure 2 39x10mm (300 x 300 DPI)

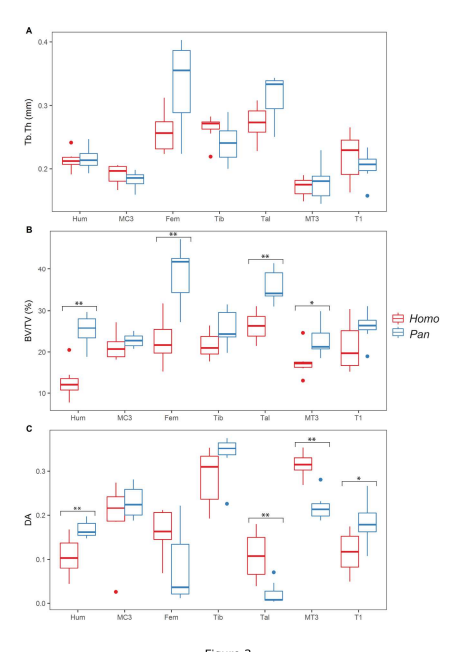
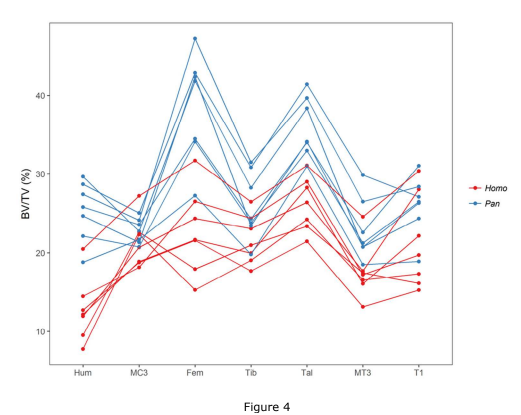


Figure 3 $208 \times 295 \text{mm} (300 \times 300 \text{ DPI})$



87x63mm (300 x 300 DPI)