BRIEF COMMUNICATION

The Neanderthal patellae from Krapina (Croatia):
A comparative investigation of their endostructural
conformation and distinctive features compared to the
extant human condition

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Abstract

Objectives: The Neanderthal patella differs from that of extant humans by being
thicker anteroposteriorly and by having more symmetric medial and lateral articular
facets. However, it is still unclear to what extent these differences affect knee kinesiology.
We aim at assessing the endostructural conformation of Neanderthal patellae
to reveal functionally related mechanical information comparatively to the extant
human condition. In principle, we expect that the Neanderthal patella (i) shows a
higher amount of cortical bone and (ii) a trabecular network organization distinct
from the extant human condition.

Materials and Methods: By using micro-focus X-ray tomography, we characterized
the endostructure of six adult patellae from the OIS 5e Neanderthal site of
Krapina, Croatia, the largest assemblage of human fossil patellae assessed so far,
and compared their pattern to the configuration displayed by a sample of 22 recent
humans.

Results and Discussion: The first expectation is rejected, indicating that the patellar
bone might have not followed the trend of generalized gracilization of the human
postcranial skeleton occurred through the Upper Pleistocene. The second prediction
is at least partially supported. In Krapina the trabecular network differs from the com-
parative sample by showing a higher medial density and by lacking a proximal rein-
forcement. Such conformation indicates similar load patterns exerted in
Neanderthals and extant humans by the vastus lateralis, but not by the vastus media-
lis, with implications on the mediolateral stabilization of the knee joint. However, the
patterns of structural variation of the patellar network remain to be assessed in other
Neanderthal samples.

KEYWORDS
internal structure, Neanderthal, patella

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1 | INTRODUCTION

In European and Near Eastern Neanderthals, the morphology of the patella differs from the typical condition of extant humans (EHs) by being absolutely and relatively thicker anteroposteriorly and by having more symmetric medial and lateral articular facets associated with a more centrally positioned articular ridge and higher angles between the medial and lateral margins of the articular facets and the ridge (e.g., Boule, 1911–1913; Heim, 1982; Rosas et al., 2020; Trinkaus, 1983a, 2000; Trinkaus & Rhoads, 1999; Trinkaus et al., 2017; Vandermeersch, 1981). A recent geometric morphometric analysis of the assemblage from El Sidrón, Spain, has shown that the typically thick Neanderthal patella with symmetrical contact surfaces tends to be accompanied by a distally positioned lateral mass compared to the more central position observed in EH (Rosas et al., 2020). Conversely, overlapping proportions between the two taxa have been recorded for the kneecap maximum height and breadth (Rosas et al., 2020). In Neanderthals, along with a posterior displacement relative to the diaphyseal axes of the tibial tuberosity and condyles of the knee joint capsule and of the ligament attachments (e.g., Boule, 1911–1913; Fraipont, 1888; Heim, 1982; Lustig, 1915; Trinkaus, 1983a; Vandermeersch, 1981), a thicker patella has been suggested to increase the quadriceps femoris moment arm in the knee joint, thus making the Neanderthal legs more powerful in extension (Chapman et al., 2010; Miller & Gross, 1998; Trinkaus, 1983b). However, a close relationship between the degree of patellar facet asymmetry and the configuration of the distal femur has been questioned (Trinkaus & Rhoads, 1999). Indeed, while a more expanded lateral facet of the EH patella has been related to the medial tibiofemoral angle (Hartigan et al., 2011; Rosas et al., 2020), bicondylar angles in Neanderthal femora associated to more symmetric patellar facets are not distinguishable from those measured in EH (Tardieu & Trinkaus, 1994; Trinkaus, 2006). Accordingly, it does not appear to be a tight relationship between degree of patellar facet asymmetry and articular configuration of the distal femur and it is still unclear to what extent variation in patellar articular proportions affects knee kinesiology (Trinkaus, 2000).

More recently, it has been proposed that, compared with the Homo sapiens derived morphology, the condition of the Neanderthal patellar articular surfaces is related to a rotation of the tibia with respect to the distal femur, implying a mediolateral displacement of the patellar ligament which would increase contact force of the patella with the femoral condyles and affect the functional space for the expansion of the facets (Rosas et al., 2020).

Given the ability of the mechanosensitive cortical and trabecular bony tissues to adjust structurally during life to the site-specific loading environment (e.g., Hoechel et al., 2015; Katoh et al., 1996; Mazurier, 2006; Mazurier et al., 2010; Raux et al., 1975; Toumi et al., 2006, 2012; Townsend et al., 1975; Van Kampen & Huiskes, 1990), the assessment of patellar endostructural features such as cortical bone thickness variation and trabecular network arrangement have the potential to provide subtle functionally related information about the knee joint mechanical environment (Cazenave et al., 2019; Cazenave et al., 2022). A preliminary comparative study noninvasively investigating the inner organizational pattern of the adult patella in two Neanderthals (from Krapina, Croatia, and Regourdou, France), one anatomically modern fossil human (the Magdalénien from Chancelade, France), and in a sample of African- and European-derived Holocene individuals revealed marked differences not only between the Neanderthal and the fossil and Holocene humans, but also between the two Neanderthal specimens (Cazenave et al., 2020). Interestingly, despite a general reduction trend in postcrani al skeletal robusticity recorded from the Upper Pleistocene through the Holocene (Chirichiri et al., 2015, 2017; Ruff et al., 2015; Ryan & Shaw, 2015; Scherf et al., 2016), the analyses did not reveal any tendency toward gracilization for al l quantitative parameters reflecting bone robusticity considered in the study (Cazenave et al., 2020). Nonetheless, this evidence needs confirmation from a larger sample. Additionally, the existence of a set of distinctive inner features uniquely characterizing the Neanderthal patella needs to be verified and the extent of the differences in endostructural conformation (patterns) between Neanderthals and EH remains to be assessed.

Here we apply 2D and 3D techniques of virtual imaging to an X-ray microtomographic record to quantitatively characterize the internal structure of six patellae from the Neanderthal site of Krapina, Croatia, and compare the results to the EH condition (Cazenave et al., 2019, 2020; Hoechel et al., 2015; Toumi et al., 2006, 2012). In principle, we expect that the Neanderthal assemblage (i) shows, on average, a higher amount of cortical bone than measured in the comparative human sample used in this study and, based on the outer characteristics distinguishing the Neanderthal from the modern human patella (Rosas et al., 2020; Trinkaus, 2000). We also expect that the fossil patellae (ii) possess some features in both cortical bone topographic distribution and site-specific organization of the trabecular network clearly distinct from the EH condition, thus allowing to identify a unique Neanderthal endostructural pattern.

2 | MATERIALS AND METHODS

2.1 | Materials

The fossil sample consists of three right and three left patellae representing a minimum of four (see below) Neanderthal adult individuals of unknown sex selected because of their relative preservation conditions from the assemblage of 16 patellae from the Eemian (OIS 5e) Croatian site of Krapina (Kricun et al., 1999; Radovčić et al., 1988; Rink et al., 1995).

The specimen Krapina 215.1-Pa. 1 (hereafter referred as Pa. 1) is a right patella with minimal erosion along the proximal, distal and lateral margins and the distal part of the medial facet (Figure 1a). Based on comparative size and morphology, Pa. 1 could belong to the same individual represented by the specimen 216.1-Pa. 5 (Trinkaus, 1975, 2000). Krapina 215.3-Pa. 3 (Pa. 3) is also a right patella with minimal damages in the distal margin of its posterior aspect (Figure 1b). Originally suggested to represent the same individual of 215.2-Pa. 2 (Trinkaus, 1975),
not included in this study, Pa. 3 has been later tentatively associated to the specimen 216.9-Pa. 14 (see above; Trinkaus, 2000). The fossil 216.1-Pa. 5 (Pa. 5), which could represent the same individual of Pa. 1 (Trinkaus, 1975, 2000) and which has been previously reported for its general inner conformation (Cazenave et al., 2019), is a left patella with some erosion along the proximal, lateral and medial articular margins and the posterior aspect of the apex (Figure 1c). Krapina 215.5-Pa. 6 (Pa. 6) is a right patella bearing some damages on the proximal and distal portions of the medial articular surface (Figure 1d). The specimen 216.3-Pa. 8 (Pa. 8) is a left patella displaying minor abrasion on the proximal, medial and distal edges (Figure 1e). Finally, the left patella 216.9-Pa. 14 (Pa. 14), perhaps associated to Pa. 3, is the least preserved specimen within the fossil assemblage considered here (Figure 1f). Its mediolateral and anteroposterior diameters are preserved, but it lacks the proximal third of the articular surface and the distal extremity of the apex.
It also shows erosion and damages on the distal third of the posterior aspect.

Descriptive and quantitative information on this fossil assemblage, including height, width, thickness, and metrics of the medial articular and lateral facets, can be found in Gorjanović-Kramberger (1906), Trinka (1975, 2000), Radovčič et al. (1988), Rosas et al. (2020) (for Pa. 5, see also Cazenave et al., 2019). Ten of the 16 patellae forming the entire Krapina assemblage, including five of six among those considered in this study, have been previously imaged in the radiographic atlas of the Krapina’s human remains (Kricun et al., 1999; figure 123). All fossils are housed at the Croatian Natural History Museum, Zagreb.

The EH comparative material consists of 22 finely preserved adult patellae from individuals of both sexes, all from the right side and lacking macroscopic evidence of alteration or pathological changes. The assemblage represents eight individuals of African ancestry (four males and four females aged 22–32 years) and seven of European ancestry (three males and four females aged 21–51 years) selected from the Pretoria Bone Collection stored at the Department of Anatomy of the University of Pretoria, South Africa (L’Abbé et al., 2005), and seven patellae (from two likely male and five likely female individuals whose estimated age at death ranges between c. 30 and c. 50 years) from the Imperial Roman graveyard of Velia, Italy, stored at the “Luigi Pigorini” National Museum of Prehistory and Ethnography (rev. in Beauchesne & Agarwal, 2017).

To test the possible influence of laterality, we also used an additional set of left patellae (representing three males and five females of both African and European ancestry) belonging to eight among the 15 individuals forming the comparative sample from the Pretoria Bone Collection. This additional set was uniquely used for testing laterality. Details on the composition of the EH sample are provided as Table S1.

### 2.2 Methods

The patellae from Krapina were imaged by X-ray microtomography (µXCT) at the Multidisciplinary Laboratory of the International Centre for Theoretical Physics (ICTP) of Trieste, Italy (Tuniz et al., 2013), at an isotropic voxel dimensions of 33 μm. The archeological specimens from Velia were also imaged at the ICTP at an isotropic voxel size ranging from 25 to 29 μm. The specimens from the Pretoria Bone Collection were scanned at the South African Nuclear Energy Corporation (Necsa), Pelindaba, with an isotropic voxel size ranging from 24 to 59 μm (Table S1). Following acquisitions, a virtual transformation of each dataset has been carried out to coherently orient all specimens by using Avizo v.8.0.0. (Visualization Sciences Group Inc.). More specifically, each image stack has been manually rotated, until the most lateral and medial surface junctions between the apex and articular surface in posterior view are aligned with the x-axis, the most lateral and medial points in proximal view are aligned with the y-axis, and the articular surface in medial view is aligned with the z axis. The new image stack orientation has then be saved in Avizo by resampling the data.

In each specimen, we firstly delimited the cortico-trabecular complex (CTC), that is, the component which includes the cortical shell (lamina) and the intimately related adjoining portions of the supporting denser trabecular network (Cazenave et al., 2019, 2020; Mazurier, 2006; Mazurier et al., 2010). By using the routine MPSAKv2.9 (in Dean & Wood, 2003), we measured the CTC mean thickness (CTT) across the anterior (aCTT) and posterior (pCTT) surfaces by using the sagittal and the transversal slices respectively extracted at the maximum anteroposterior thickness and the mediolateral breadth (Figure 2a). Because of damages on their posterior aspects, it was not possible to measure the pCTT across both sagittal and transversal virtual sections in Pa. 6 (Figure 1d), while in Pa. 14 the pCTT was not measured across the sagittal slice (Figure 1f). To conduct size-independent

![Figure 2](https://example.com)
intra- and inter-taxic comparisons, in each specimen the absolute CTT values (in mm) were standardized with respect to the maximum breadth and provided as percent (%) values in each specimen.

Besides the CTC assessment, in all specimens we also virtually extracted four homologous cubic volumes of interest (VOIs) whose individual edge length systematically equals 10% of the mediolateral maximum patellar breadth. The VOIs systematically sample the proximal (pVOI), distal (dVOI), medial (mVOI), and lateral (lVOI) regions (Figure 2b). The geometric centers of the pVOI and dVOI were placed at the ½ of the mediolateral breadth and of the anteroposterior thickness, and at ½ of the proximodistal articular height from the proximal and distal margins, respectively. The mVOI and lVOI were positioned at the ½ of the proximodistal articular height and of the anteroposterior thickness, and at ¼ of the mediolateral breadth from the medial and lateral margins, respectively.

To measure the site-specific structural properties of the trabecular network, the cubic VOIs were binarized into bone and air using the “half maximum height” (HMH) quantitative iterative thresholding method (Spoor et al., 1993) and the region of interest protocol (ROI-Tb; Fajardo et al., 2002). Owing to the biasing effects of a non-spherical VOI by using the mean intercept length (MIL) algorithm in Quant3D (Ryan & Ketcham, 2002), on the largest centered sphere fitting completely within each extracted VOI we measured: (i) the trabecular bone volume fraction (BV/TV, in %); (ii) the trabecular thickness (Tb.Th., in mm); and (iii) the degree of anisotropy (DA).

For all variables, a number of intra- and inter-observer tests for accuracy run by independent observers on both the fossil assemblage and on selected specimens from the comparative sample provided differences less or near 4%.

To assess the possible influence of size, in both fossil and comparative extant samples the Pearson’s correlation coefficient was calculated between each endostructural variable and the mediolateral breadth of the patella, a dimension found to scale with body size in hominoids (Jungers, 1990; Pina et al., 2014).

Statistical analyses were performed in R v3.4.4 (R Core Team, 2018) by RStudio v1.2.5033, while plots were generated using ggplot2 (Wickham, 2009). Due to small sample sizes, only non-parametric statistical tests were performed. The significance of the differences between the taxa for anterior CTT of the sagittal and transversal sections, BV/TV, Tb.Th. and DA of each VOI was tested by the two-sample t test via 

t

By the quantile method (Ryan & Ketcham, 2002) but statistical tests show no significant differences for the anterior CTT of both sagittal and transversal sections. The most marked average differences concern the posterior thickness measured across the sagittal section (7.9% vs. 7.1%) and the anterior thickness across the transversal section (7.3% vs. 6.7%). However, given that the comparative sample shows a wider range of variation for both parameters (4.0%–12.8% and 4.5%–13.1%, respectively, vs. 6.7%–10.6% and 5.9%–9.7% in Krapina), such differences can be considered as negligible. Indeed, even if the Neanderthal patellae show the average thickest cortices (Table S2), for the sagittal and transversal anterior CTT and the transversal posterior CTT the absolute highest values have been measured in humans (Table S2). In terms of cortical thickness topographic distribution, it does not matter the orientation plan, in both Krapina and the EH sample, the CTC is on average, thicker anteriorly (aCTT > pCTT; Table 1), but intra-individual deviation from this pattern has been found in both assemblages (one case in Krapina and 23% and 14% of cases in the comparative sample for the sagittal and the transversal plane, respectively). In all cases and in both samples, CTT variation is size-independent (Table S3).

3.1 | CTT distribution

In both sagittal and transversal planes, the Neanderthal patellae from Krapina show, on average, a slightly thicker CTT compared to the EH sample used in this study (Table 1, Figure S1) but statistical tests show no significant differences for the anterior CTT of both sagittal and transversal sections. The most marked average differences concern the posterior thickness measured across the sagittal section (7.9% vs. 7.1%) and the anterior thickness across the transversal section (7.3% vs. 6.7%). However, given that the comparative sample shows a wider range of variation for both parameters (4.0%–12.8% and 4.5%–13.1%, respectively, vs. 6.7%–10.6% and 5.9%–9.7% in Krapina), such differences can be considered as negligible. Indeed, even if the Neanderthal patellae show the average thickest cortices (Table S2), for the sagittal and transversal anterior CTT and the transversal posterior CTT the absolute highest values have been measured in humans (Table S2). In terms of cortical thickness topographic distribution, it does not matter the orientation plan, in both Krapina and the EH sample, the CTC is on average, thicker anteriorly (aCTT > pCTT; Table 1), but intra-individual deviation from this pattern has been found in both assemblages (one case in Krapina and 23% and 14% of cases in the comparative sample for the sagittal and the transversal plane, respectively). In all cases and in both samples, CTT variation is size-independent (Table S3).

The specimens Pa. 1 and Pa. 5, suggested to represent the same individual (Trinkaus, 1975, 2000), are close for the sagittal anterior and transversal posterior average CTTs, but distinctly distinguishable for the transversal aCTT (7.4% and 5.9%, respectively), the left patella Pa. 5 systematically displaying a slightly thinner shell (Table 1, Figure S1a,c, Table S2). Slightly higher differences in CTC arrangement are recorded between Pa. 3 and Pa. 14, the second possible individual pair of patellae within the Krapina’s assemblage included in this study (Trinkaus, 2000). In this case, the greatest difference concerns the transversal pCTT (5.6% and 3.1%, respectively; Table 1, Figure S1bf, Table S2). However, compared with the homologous fossils from Krapina is suitable for the purposes of a subtle quantitative assessment of their conformation (Figure 3, Supporting Information S1 and Figure S1). As noted in previous studies on the outer morphology of this assemblage (Gorjanović-Kramberger, 1906; Radovči, 1988; Trinkaus, 2000), the anterior aspect of Pa. 3 shows an enthesophyse formation likely resulting from the ossification of fibers of the quadriceps femoris tendon, and the articular surface of Pa. 8 bears minor alterations typical of degenerative changes. However, as previously observed in the case of the Magdalenian patella Chancelade 1 also displaying similar outer degenerative alterations (Cazenave et al., 2019), the μXCT record of both Pa. 3 and Pa. 8 shows that the corresponding underlying cortical and trabecular tissues lack any evident structural change with respect to the surrounding areas, with no evidence of local cortical thinning or thickening and/or trabecular rarefaction or increased density (Figures 3b,e and S1b,e).
values of CTC cross-sectional differences measured for the eight pairs of patellae representing as many individuals from the Pretoria Bone Collection (Table S1), where fluctuating (nondirectional) asymmetry ranges between 0.1% and 3.2%, with no pattern of laterality (Table S4), the differences recorded within the possible pairs Pa. 1-Pa. 5 and Pa. 3-Pa. 14 (range < 2.5%) are fully compatible with intra-individual normal variation. We, therefore, cannot reject the hypothesis that Pa. 1-Pa. 5 are from the same individual as well as Pa. 3-Pa. 14 (Trinkaus, 1975, 2000).

3.2 | Trabecular bone structural organization

The individual and average values of the structural properties of the trabecular network assessed in the patellae from Krapina and the average values measured in the comparative EH sample for the proximal (s), distal (i), medial (m), and lateral (l) VOIs are shown in Table 2 and rendered in Figure 4.

The Neanderthal values do not significantly differ from the EH estimates but for the degree of anisotropy (DA) of the dVOI and for

![Figure 3](https://onlinelibrary.wiley.com/doi/image/10.1002/ajpa.24709)

**Figure 3** Coronal (upper row) and sagittal (lower) virtual sections extracted across the center of the right patella 215.1-Pa. 1 (a), the right patella 215.3-Pa. 3 (b), the left patella 216.1-Pa. 5 (c), the right patella 215.5-Pa. 6 (d), the left patella 216.3-Pa. 8 (e), the left patella 216.9-Pa. 14 (f) from Krapina, and of an extant human (EH) representative (g; a 22 years old female). Scale bar: 1 cm.
TABLE 1 Individual and mean values of the mean cortico-trabecular thickness (C TT, standardized % values) measured for the anterior (aCTT) and posterior (pCTT) aspects across the sagittal and transversal virtual slices, respectively (Figure 2a), in the Neanderthal patellae from Krapina and in the extant human (EH) reference sample used for comparisons (n = 22).

<table>
<thead>
<tr>
<th>Specimen/sample</th>
<th>aCTT sagittal</th>
<th>pCTT sagittal</th>
<th>aCTT transversal</th>
<th>pCTT transversal</th>
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</thead>
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<tr>
<td>Krapina</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>215.1-Pa. 1</td>
<td>8.0</td>
<td>6.7</td>
<td>7.4</td>
<td>5.8</td>
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<tr>
<td>215.3-Pa. 3</td>
<td>7.9</td>
<td>7.2</td>
<td>7.4</td>
<td>5.6</td>
</tr>
<tr>
<td>216.1-Pa. 5</td>
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<td>7.0</td>
<td>5.9</td>
<td>4.5</td>
</tr>
<tr>
<td>215.5-Pa. 6</td>
<td>8.4</td>
<td>-</td>
<td>6.8</td>
<td>-</td>
</tr>
<tr>
<td>216.3-Pa. 8</td>
<td>8.8</td>
<td>10.6</td>
<td>9.7</td>
<td>6.4</td>
</tr>
<tr>
<td>216.9-Pa. 14</td>
<td>9.2</td>
<td>-</td>
<td>6.6</td>
<td>3.1</td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>8.4 (0.5)</td>
<td>7.9 (1.9)</td>
<td>7.3 (1.3)</td>
<td>5.1 (1.3)</td>
</tr>
<tr>
<td>EH (SD)</td>
<td>8.3 (2.3)</td>
<td>7.1 (2.4)</td>
<td>6.7 (2.1)</td>
<td>4.8 (1.7)</td>
</tr>
</tbody>
</table>

the trabecular bone volume fraction (BV/TV) of the mVOI (Figure 4, Table S5; p values of 0.02 and 0.005 for the DA of the dVOI and BV/TV of the mVOI, respectively), the latter reflecting a higher average bulk density in the medial region characterizing the Neanderthal assemblage. Besides the dVOI DA, the differences between Krapina and the comparative sample measured for the other parameters of the medial and lateral VOIs slightly exceed those expressed between the proximal and distal VOIs. However, as already noted for the thickness of the CTC (Table 1), topographic variation of the trabecular network structural properties shown by the comparative human sample is higher than expressed by the Krapina assemblage by itself (Figure 4).

In all Neanderthal specimens, a structural reinforcement of the trabecular network indicated by higher BV/TV and Tb.Th. values is found in the lateral VOI (Table 2), whose values statistically differ from those measured in the other three VOIs (Table S6; p value of 0.03). A relative lateral reinforcement is also variably developed in the EH patellae (including comparing to humans from Cazenave et al., 2019, 2020; Tables 2 and S6), where it is associated with a proximal reinforcement of the network, this latter feature systematically lacking in Krapina (see also Cazenave et al., 2019, 2020; Toumi et al., 2006; Tables 2 and S6). Conversely, the Krapina assemblage and the EH networks share a relatively denser trabecular area approximately positioned at the proximalateral margin; a higher number of radially oriented trabeculae around the medial peripheral area; a vertically oriented anterior bundle; and two obliquely oriented proximal and distal bundle-like structures running from the anterior to the posterior endosteal surfaces (see also Cazenave et al., 2019, 2020).

Among all 12 variables examined for each sample (three structural properties by four VOIs), a significant correlation with the patellar mediolateral breadth has been recorded only for the medial DA in the fossil sample and the proximal BV/TV in the comparative human sample (Table S3).

The possible Krapina’s pair Pa. 1-Pa. 5 shows some closer affinities in trabecular organization compared to the signal from other specimens forming the Neanderthal assemblage, but the two patellae differ for the BV/TV of the distal VOI (22.5% vs. 30.1%), the Tb.Th. of the lateral VOI (0.36 vs. 0.28 mm), and the DA of the distal VOIs (0.50 vs. 0.34). Conversely, for the mVOI DA (0.57 and 0.67, respectively) they show the highest values of the entire fossil assemblage (average of the remaining specimens = 0.44). Structural affinities also exist between the specimens forming the other possible pair, Pa. 3 and Pa. 14, but in this case they differ for the BV/TV (25.8% vs. 38.0%) and the DA (0.24 vs. 0.50) of the dVOI (Table 2). Indeed, while Pa. 3 displays at all sites BV/TV values distal to the average of the whole fossil assemblage, Pa. 14 is among the most endostructurally robust specimens. As recorded for the thickness values of the CTC, even greater differences than those revealed by the comparative analysis of the two possible Neanderthal pairs Pa. 1-Pa. 5 and Pa. 3-Pa. 14 have been measured in our ad hoc sample of eight pairs of EH patellae (Table S7). As recorded for the CTC arrangement, no pattern of laterality between the paired patellae has been found in the EH sample for the structural properties of the trabecular network (Table S7), that can be associated to fluctuating asymmetry (i.e., small, random deviations away from perfect bilateral symmetry).

4 | DISCUSSION

The quantitative endostructural analysis of six patellae from the Neanderthal site of Krapina, Croatia, has revealed a modest degree of variation in cortical bone topographic distribution and site-specific arrangement of the trabecular network, associated with a lower degree of intra-individual organizational heterogeneity than commonly observed in EHs (Cazenave et al., 2019, 2020; Hocchel et al., 2015; Toumi et al., 2006, 2012; this study). This, despite a certain degree of dimensional and morphological variation displayed by the Krapina fossils in a number of outer features (e.g., expression of the vastus lateralis notch, concavity of the lateral articular facet and of the proximodistal profile of the articular crest; Radovčič et al., 1988; Rosas et al., 2020; Trinkaus, 2000; see Table S8). However, given the extremely limited size of the investigated Krapina sample, such observation needs confirmation.

In the Krapina assemblage, for most measures of cortico-trabecular thickness distribution and structural properties of the trabecular bone, a comparable structural signature is expressed by the two pairs of specimens Pa. 1 and Pa. 5, on the one side, and, to a lesser extent, Pa. 3 and Pa. 14, on the other side, originally suggested to likely represent two individuals because of their outer features and proportions (Trinkaus, 1975, 2000). Even though bony endostructure
TABLE 2  Individual and mean values of the bone volume fraction (BV/TV, in %), the trabecular thickness (Tb.Th., in mm) and the degree of anisotropy (DA) of the volumes of interest (VOIs) sampling the proximal (pVOI), distal (dVOI), medial (mVOI), and lateral (lVOI) regions (Figure 2b) measured in the Neanderthal patellae from Krapina and in the extant human (EH) reference sample used for comparisons (n = 22).

<table>
<thead>
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<th>Specimen/sample</th>
<th>pVOI</th>
<th>dVOI</th>
<th>mVOI</th>
<th>lVOI</th>
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</thead>
<tbody>
<tr>
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<td>Tb.Th.</td>
<td>DA</td>
<td>BV/TV</td>
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<td>0.55</td>
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<td>0.59</td>
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</tr>
<tr>
<td>Krapina 216.3-Pa. 8</td>
<td>36.3</td>
<td>0.28</td>
<td>0.66</td>
<td>39.0</td>
</tr>
<tr>
<td>Krapina 216.9-Pa. 14</td>
<td>34.9</td>
<td>0.26</td>
<td>0.52</td>
<td>38.0</td>
</tr>
<tr>
<td>Krapina 217.9-Pa. 16</td>
<td>34.8</td>
<td>0.28</td>
<td>0.57</td>
<td>31.6</td>
</tr>
<tr>
<td>Mean</td>
<td>34.8</td>
<td>0.28</td>
<td>0.57</td>
<td>31.6</td>
</tr>
<tr>
<td>(SD)</td>
<td>(2.5)</td>
<td>(0.01)</td>
<td>(0.05)</td>
<td>(6.6)</td>
</tr>
<tr>
<td>EH (SD)</td>
<td>40.1 (9.9)</td>
<td>0.31 (0.13)</td>
<td>0.52 (0.19)</td>
<td>30.0 (12.4)</td>
</tr>
</tbody>
</table>

Note: Significant differences (p value < 0.05) are shown in bold (see Table S5 for statistical results).
pattern. Differently from the first one, this second prediction is at least partially supported. Indeed, for their structural properties, the Krapina’s patellae tend to differ from the recent human ones by showing a higher medial density (higher BV/TV and Tb.Th.) and by the lack of a proximal cancellous reinforcement, which is instead variably developed in the extant specimens. As a whole, this results into two generically distinct patterns for both the trabecular bone volume fraction and the trabecular thickness (but not for the degree of anisotropy): lateral \(>\) proximal \(\geq\) distal \(\leq\) medial, in Krapina, versus proximal \(\geq\) lateral \(>\) distal \(\geq\) medial, more typical in EHs.

Such reinforcements are interpreted as related to the mechanical stresses acting at the quadriceps tendon, but also to reflect a functional similarity in the action exerted by the central (vastus intermedius) and lateral (vastus lateralis) components of the quadriceps muscles (Toumi et al., 2012). Differently, the medial side of the patella, whose cancellous network is absolutely and relatively denser in the Krapina fossils, undergoes the loads of the medial component of the quadriceps (vastus medialis), which inserts obliquely on the patella to actively resist its lateral pull and contributes the medial stability of the knee-joint (Balcarek et al., 2014; Rajput et al., 2017; Toumi et al., 2012). Accordingly, the endostructural configuration of the Krapina’s patellae suggests a mechanical environment of the Neanderthal knee joint comparable to the EH condition for the loads exerted by the vastus lateralis, but not for those of the vastus medialis, whose functional action in the stabilization of the articulation might have been absolutely and relatively stronger than commonly occurs in EHs.

Besides some differences between Neanderthals and EHs in the relative development, positioning and function of the vastus muscles (e.g., Belcastro et al., 2006; Belcastro & Mariotti, 2017; Rosas et al., 2020), the influence on the Neanderthal knee joint of the greater robusticity of the distal femoral epiphysis remains also to be clarified. With this regard, a comparative study combining linear measurements and a 3D geometric morphometric analysis of the outer patella in a whole sample of 27 late Middle-Late Pleistocene Neanderthals (including four specimens from Krapina) has shown that Neanderthals had a more oblique attachment of the complete quadriceps muscular package expressed by a large size of the medial articular facet and a relative displacements of the medial and lateral masses.
of the patellar body than expressed in EHs. This configuration, compatible with the differences in the endostructural conformation of the trabecular network recorded in our study, has been related to a distinct rotation of the tibia with respect to the femur uniquely characterizing the Neanderthal knee joint, but interpreted as more likely representing a consequence of body form evolution in *Homo* rather than a specific functional adaptation of the knee (Rosas et al., 2020).

It is noteworthy that inter-individual variation in patellar endostructural arrangement expressed by our comparative extant sample is wide and a Krapina-like trabecular conformation pattern is also found in a couple of extant specimens used in this study. On the other hand, a previous analysis of the Regourdou 1’s patella (Madelaine et al., 2008; Trinkaus, 2000; Vandermeersch, 1981), hardly distinguishable for its outer proportions and morphology from the average condition expressed by the Krapina assemblage (Table S8) and, more widely, by the Neanderthal condition (Rosas et al., 2020), has revealed a rather modern human-like cancellous pattern, with a lateral and proximal bone reinforcement (Cazenave et al., 2019, 2020). This suggests a potentially wide range of inter-population variation among Neanderthals. However, in the absence of comparative information from additional assemblages, the extent of such variation and any possible evidence for time- and geographic-related trends remain unknown.

Embedded within tendons and ligaments and acting “to protect the knee joint, to lengthen the lever arm of the quadriceps femoris, and to increase the area of contact between the patellar ligament and the femur” (White et al., 2012, p. 252), the patella adapts its outer and inner structure in response to the mechanical environment and the evolutionary changes in force transmission at the knee joint (Cazenave et al., 2019, 2020; Rosas et al., 2020; Toumi et al., 2006, 2012). In this view, the assessment of the evolutionary patterns of its endostructural conformation should provide subtle information of relevant biomechanical significance. However, the number of hominin fossil specimens examined so far is less than negligible and quantitative information on EH variation is also limited. Accordingly, future anthropological research should pay greater attention to comparatively detail the inner structural evolutionary anatomy of this bone in the hominin fossil record and to develop analyses on representative EH samples from various chrono-geographic contexts aimed at elucidating the influence of variables such as age, sex, body size and shape, physical activity, among others, on the variation patterns of its cortical bone topography and trabecular arrangement.

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**CONFLICT OF INTEREST STATEMENT**

The authors declare no conflict of interest.

**DATA AVAILABILITY STATEMENT**

The data that support the findings of this study are available from the corresponding author upon reasonable request and upon permission of the Croatian Natural History Museum for the scans of the Neanderthal sample from Krapina.

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**REFERENCES**


**SUPPORTING INFORMATION**

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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