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Experimental Perspective on Fallback Foods and Dietary Adaptations in Early Hominins

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Short title: Fallback Foods and Hominin Evolution
The robust jaws and large, thick-enamed molars of the Plio-Pleistocene hominins *Australopithecus* and *Paranthropus* have long been interpreted as adaptations for hard-object feeding. Recent studies of dental microwear indicate that only *Paranthropus robustus* regularly ate hard items, suggesting that the dentognathic anatomy of other australopiths reflects rare, seasonal exploitation of hard fallback foods. Here we show that hard-object feeding cannot explain the extreme morphology of *Paranthropus boisei*. Rather, analysis of long-term dietary plasticity in an animal model suggests year-round reliance on tough foods requiring prolonged postcanine processing in *P. boisei*. Increased consumption of such items may have marked the earlier transition from *Ardipithecus* to *Australopithecus*, with routine hard-object feeding in *P. robustus* representing a novel behavior.
1. Introduction

The australopiths (genera *Australopithecus* and *Paranthropus*) represent the earliest well-documented diversification of the hominin lineage and include the ancestor of our own genus, *Homo* (Fig. 1) [1]. Understanding the paleobiological and phylogenetic implications of phenotypic variation in this group is therefore a critical step in the study of human origins [2-4]. Australopiths were characterized by numerous apomorphic craniodental features—including robust jaws and large postcanine teeth with thick enamel caps—that have long been interpreted as adaptations for countering powerful masticatory stresses associated with a diet of mechanically challenging foods, particularly hard objects [5-8]. However, this scenario has been challenged by recent studies of dental microwear, which have failed to detect signs of routine postcanine processing of hard items in most australopiths [9-11].

One hypothesis invoked to explain this discrepancy between anatomy and microwear is that australopith craniodental adaptations reflect hard fallback foods critical to surviving seasonal periods when easier-to-process preferred resources were scarce [9-10]. Accordingly, the dearth of fossil australopith teeth preserving evidence of hard-object consumption simply reflects the rarity of such fallback items [12] in the diets of these hominins. However, the role of dietary seasonality in shaping the masticatory apparatus of primates and other mammals is unclear. Indeed, there is little evidence that the highly robust jaws of the australopiths *Paranthropus robustus* and especially *P. boisei* would be required of an organism that relies only seasonally on mechanically challenging foods [13]. Here we report the results of a long-term diet-manipulation experiment conducted using an animal model that examines adaptive plasticity [14,15] in cranial and mandibular development vis-à-vis temporal variation in food mechanical properties. Our naturalistic, longitudinal data provide a novel perspective on debates over early hominin
paleoecology and have significant implications for understanding phenotypic variation in extant and fossil species that experience resource seasonality.

2. Materials and Methods

White rabbits (*Oryctolagus cuniculus*) exhibit several characteristics that make them excellent models for investigating questions regarding masticatory biomechanics in primates, including a vertically deep face; a temporomandibular joint situated high above the occlusal plane, capable of rotational and translational movements; intracortical bone remodeling; and a similar pattern of covariation among jaw-muscle activity, jaw loading, and dietary properties [16-20]. Our sample contained 30 five-week-old weanling male rabbits divided equally into three dietary cohorts and raised for 48 weeks. Control subjects were fed a diet consisting solely of rabbit pellets. Annual rabbits were given hay in addition to pellets throughout the experiment, starting with two hay cubes daily for the first 18 weeks and then six hay cubes daily for the next six weeks. This 24-week schedule was then repeated. Seasonal rabbits received pellets and three hay cubes daily for the first six weeks and were then switched to an all-pellet diet for the subsequent 18 weeks, mimicking seasonal reliance on fallback foods. This schedule was repeated in the final 24 weeks.

The mechanical properties of these diets fall within the range of values for foods ingested by wild primates [21,22]. Hay and pellets result in similar levels of bone strain along the rabbit mandibular corpus [16]. However, hay is stiffer than pellets [22] and therefore presents a greater masticatory challenge characterized by longer loading durations and greater cyclical loading. Compared to pellets, hay takes rabbits approximately three times as many chewing cycles and, correspondingly, three times longer to process (Ravosa *et al.*, unpublished data). Dynamic
alterations in dietary behaviors related to differences in the properties of experimental foods are posited to induce osteogenic responses and corresponding ontogenetic changes in the proportions of the feeding apparatus [17,22,23].

Each animal was imaged using micro-CT (Bioscan/Mediso X-CT; settings: 70 kVp, 100 μA, with 71-μm reconstructed isometric voxel size) upon arrival and every two weeks thereafter until Week 24, when they became too large to image. Rabbits were imaged a final time following sacrifice at Week 48. This data set allowed us to track skull development longitudinally from weaning to mature adult stages. Using the segmenting tools available in the program PMOD, we quantified bone cross-sectional areas at three mandibular sites (symphysis, condyle, corpus) and one on the cranium (palate) involved in load resistance during chewing. To control for differences in organismal size that may confound our ability to detect a dietary signal among cohorts at a given age, shape ratios were computed by dividing the square root of each subject’s cross-sectional area at a given time point by its cranial length. These ratios were logged for analysis. Statistical comparisons were performed using the bootstrap to generate confidence intervals for differences between groups [24]. A more detailed description of the measurements and procedures is available in the electronic supplementary material.

3. Results

Mean shape ratios for the three dietary cohorts were statistically indistinguishable at the onset of the experiment (Week 0). By Week 6, seasonal and annual rabbits had significantly greater shape ratios at all sites—indicating relatively larger cross-sectional areas—than control rabbits, while being similar to each other (Table 1, Fig. 2). Annual and seasonal rabbits diverged rapidly following the latter’s shift to a less-challenging all-pellet diet after Week 6. By Week 12,
means for the *annual* cohort were significantly larger than those for the *seasonal* and *control* cohorts at three of the four sites. Notably, *seasonal* rabbits differed from *control* animals only in relative palatal cross-sectional area. This general pattern persisted, with slight differences, through the first half of the experiment (Week 24) and characterized the final set of comparisons: at Week 48, *annual* rabbits had significantly larger symphyseal, palatal, and corporal relative cross-sectional areas than *seasonal* and *control* groups; the latter two cohorts differed only at the symphysis (*seasonal* > *control*). Ratios for the *annual* group at this stage were 6–17% and 4–11% larger than the ratios for the *control* and *seasonal* groups, respectively. Such differences fall within the range of variation observed in closely related primate species that differ in diet (electronic supplementary material, table S4), indicating that phenotypic plasticity is likely an important source of interspecific adaptive variation.

### 4. Discussion

Seasonal hay consumption resulted in adult phenotypes clearly distinct from those of animals that ate hay throughout the experiment, but only minimally differentiated from those associated with the less-challenging all-pellet diet. This finding indicates that the relationship between dietary properties and craniomandibular morphology is highly dependent on loading history, specifically the extent to which a structure is exposed to a behavioral stimulus during development. With respect to linking australopith jaw robusticity to seasonal consumption of hard objects, this observation implies that the more extreme australopiths would have relied on such foods for a greater portion of the year than the more generalized species. It is notable, therefore, that *P. boisei*, the apex of the australopith trend toward increased jaw robusticity [1,6], presents molar microwear suggesting that it processed hard foods *less* frequently than the closely
related but less-specialized *P. robustus* [10,25], the only australopith with microwear consistent
with at least seasonal hard-object feeding [9]. Our results therefore suggest that the apomorphically
masticatory apparatus of *P. boisei* cannot be explained by a scenario in which this species fed
mainly on relatively easy-to-process foods throughout the year while relying on hard objects
during fallback episodes that were shorter in duration than was the case for *P. robustus*. Instead,
the remarkable jaws of *P. boisei* probably reflect regular consumption of items that required
intensive postcanine processing, resulting in masticatory stresses that exceeded those
experienced by *P. robustus*.

Considered within the broader context of australopith variation, the link between
morphology and markedly seasonal hard-object feeding in *P. boisei* appears even more tenuous.
In jaw robusticity, species of *Australopithecus* fall between *Paranthropus* and the geologically
older and more plesiomorphic *Ardipithecus ramidus* [1,3,4]. If the morphological differences
between *Ardipithecus* and basal australopiths signal an adaptive shift to seasonal exploitation of
hard objects in the latter, then our experimental evidence suggests that exaggeration of
australopith craniodental features in *Paranthropus* implies increased reliance on such items.
Microwear results for *P. robustus* fit this scenario, whereas those for *P. boisei* contradict it
[10,25]. Indeed, the microwear signatures of *P. boisei*, *Au. anamensis*, and *Au. afarensis* are
striking in their uniform lack of evidence for consumption of very hard or very tough items
[10,11].

The functional significance of this microwear signature remains enigmatic, but it may
result from prolonged milling and grinding, which would have been necessary when
australopiths consumed tough foods due to the fact that their low-cusped molars are not as well-
suited for shearing such items as are the high-cusped molars of extant folivorous primates [11].
Although speculative, this interpretation is supported by our results: because the microwear data reject frequent hard-object feeding in *P. boisei*, this species must have masticated exceptionally tough foods on a regular basis. The properties of foods eaten by species of *Australopithecus* are more difficult to infer, but differences between these hominins and *Ar. ramidus* in jaw robusticity, megadontia, and microwear [3,4], combined with the absence of a hard-object microwear signature [11], suggests tough-object feeding [26], but not to the degree inferred for *P. boisei*. We posit, therefore, that increases in jaw robusticity from *Ardipithecus* to *Australopithecus* to *P. boisei* reflect progressively greater reliance on tough, probably $^{13}$C-enriched [27] foods and concomitantly elevated masticatory stresses due to higher repetitive loading and longer load durations resulting from extended bouts of milling and grinding [11]. Under this scenario, the hard-object feeding evident in the microwear of *P. robustus* represents a novel feeding strategy, perhaps indicating adoption of a broader niche facilitated by a masticatory apparatus initially shaped by a diet of tough foods but nevertheless capable of processing objects with a wide range of mechanical properties.

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Table 1. Comparisons among dietary cohorts. Significant differences in shape ratios among dietary cohorts are indicated by different type styles; entries in the same style are not significantly different. Alpha levels were adjusted for multiple comparisons using the sequential Bonferroni method within each variable-week combination (i.e., each row). $P$-values and raw data are provided in electronic supplementary material (tables S1, S2).

Figure 1. Cladistic relationships of selected australopiths. Only the five best-represented species are shown here for simplicity, along with *Ardipithecus ramidus* and genus *Homo*.

Figure 2. Jaw growth in dietary cohorts. Logged (base $e$) square roots of bone cross-sectional area versus mean cranial length: (a) condyle, (b) palate, (c) corpus, (d) symphysis. Each datum represents a given week. Data are available in the electronic supplementary material (table S3).
### Table 1

<table>
<thead>
<tr>
<th>Time point</th>
<th>Variable</th>
<th>Direction of observed differences</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Smallest</td>
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<tr>
<td>Week 0</td>
<td>Symphysis ratio</td>
<td>Annual</td>
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<td>Palate ratio</td>
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<td>Week 6</td>
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<td>Palate ratio</td>
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<td>Condyle ratio</td>
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<td></td>
<td>Corpus ratio</td>
<td>Control</td>
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<td>Week 12</td>
<td>Symphysis ratio</td>
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<td></td>
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<td>Week 48</td>
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<td></td>
<td>Corpus ratio</td>
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</tbody>
</table>
Figure 1

Indicates increase in jaw robusticity and postcanine megadontia

Indicates decrease in jaw robusticity and postcanine megadontia