1 Conserved and derived expression patterns and positive selection on dental genes reveal

2 complex evolutionary context of ever-growing rodent molars

3

4 AUTHORS:

- 5 Zachary T. Calamari^{1,2,3,4,*}, zachary.calamari@baruch.cuny.edu
- 6 Andrew Song^{1,5}, ajs557@cornell.edu
- 7 Emily Cohen^{1,6}, ec4744@nyu.edu
- 8 Muspika Akter¹, muspika.akter@baruchmail.cuny.edu
- 9 Rishi Das Roy⁷, rishi.dasroy@helsinki.fi
- 10 Outi Hallikas⁷, outi.hallikas@helsinki.fi
- 11 Mona M. Christensen⁷, mona.christensen@helsinki.fi
- 12 Pengyang Li^{3,8}, pengyang.li@cshs.org
- 13 Pauline Marangoni^{3,8}, pauline.marangoni@cshs.org
- 14 Jukka Jernvall^{7,9}, jernvall@fastmail.fm
- 15 Ophir D. Klein^{3,8,*} ophir.klein@cshs.org
- 16
- ¹⁷ ¹Baruch College, City University of New York, One Bernard Baruch Way, New York, NY
- 18 10010, USA
- ¹⁹ ²The Graduate Center, City University of New York, 365 Fifth Ave, New York, NY 10016, USA
- 20 ³Program in Craniofacial Biology and Department of Orofacial Sciences, University of
- 21 California, San Francisco, San Francisco, CA 94158, USA
- ⁴Division of Paleontology, American Museum of Natural History, Central Park West at 79th
- 23 Street, New York, NY, 10024, USA

- ⁵Cornell University, 616 Thurston Ave, Ithaca, NY 14853, USA
- ⁶New York University College of Dentistry, 345 E 34th St, New York, NY 10010
- ⁷Institute of Biotechnology, University of Helsinki, FI-00014 Helsinki, Finland
- ⁸Department of Pediatrics, Cedars-Sinai Guerin Children's, 8700 Beverly Blvd., Suite 2416, Los
- Angeles, CA 90048, USA
- ⁹Department of Geosciences and Geography, University of Helsinki, FI-00014 Helsinki, Finland
- 30 *Corresponding authors
- 31

32 ABSTRACT

33 Background: Continuously growing teeth are an important innovation in mammalian evolution, 34 yet genetic regulation of continuous growth by stem cells remains incompletely understood. 35 Dental stem cells are lost at the onset of tooth root formation, but this loss of continuous crown 36 growth is difficult to study in the mouse because regulatory signaling overlaps with signals that 37 pattern tooth size and shape. Within the voles (Cricetidae, Rodentia, Glires), species have 38 evolved both rooted and unrooted molars that have similar size and shape. We assembled a de 39 novo genome of Myodes glareolus, a vole with high-crowned, rooted molars, and performed 40 genomic and transcriptomic analyses in a broad phylogenetic context of Glires (rodents and 41 lagomorphs) to assess differential selection and evolution in tooth forming genes. 42 Results: Our de novo genome recovered 91% of single-copy orthologs for Euarchontoglires and 43 had a total length of 2.44 Gigabases, enabling genomic and transcriptomic analyses. We 44 identified six dental genes undergoing positive selection across Glires and two genes undergoing 45 positive selection in species with unrooted molars, Dspp and Aqp1. Transcriptomics analyses

46	demonstrated conserved patterns of dental gene expression with species-specific variation likely
47	related to developmental timing and morphological differences between mouse and vole molars.
48	Conclusions: Our results support ongoing dental gene evolution in rodents with unrooted molars.
49	We identify candidate genes for further functional analyses, particularly Dspp, which plays an
50	important role in mineralizing tissues. Our expression results support conservation of dental
51	genes between voles and model species like mice, while revealing significant effects of overall
52	tooth morphology on gene expression.
53	
54	3-10 Keywords: Evolution, selection, Glires, molar, root, dental, development, genome, rodent,
55	tooth
56	
57	DECLARATIONS
58	Ethics approval: The University of California, San Francisco (UCSF) Institutional Animal Care
59	and Use Program and the Finnish national animal experimentation board approved protocols for
60	humane euthanasia and collection of tissues for animals used in this study under protocols
61	AN189916 (UCSF) and KEK16-021, KEK19-019, and KEK17-030 (University of Helsinki).
62	
63	Consent for publication: Not applicable.
64	
65	Availability of data and materials: The datasets supporting the conclusions of this article are
66	available in the GenBank repository under [GenBank reference number to be added upon
67	acceptance] and in the article's additional files.
68	

69 Competing interests: The authors declare that they have no competing interests.

71	Funding: This research was supported by National Science Foundation grants CNS-0958379,
72	CNS-0855217, OAC-1126113, and OAC-2215760 through the City University of New York
73	High Performance Computing Center at the College of Staten Island; OAC-1925590 through the
74	MENDEL high performance computing cluster at the American Museum of Natural History;
75	Academy of Finland to JJ; Doctoral Programme in Biomedicine, University of Helsinki to
76	MMC; and National Institutes of Health NIDCR R01-DE027620 and R35-DE026602 to ODK.
77	
78	Authors' contributions: ZTC and ODK designed the study. ZTC and PM performed animal
79	husbandry. ZTC performed and oversaw tissue sampling, sequencing, genome assembly and
80	annotation for Myodes glareolus. ZTC, AS, JR, EC, and MA performed genome computational
81	analyses. PL performed qPCR analyses. OH, MMC, RDR, and JJ designed and implemented
82	RNA sequencing experiments. ZTC wrote and all authors contributed to and approved the
83	manuscript.
84	
85	Acknowledgements: The authors thank A. Joo, N. Ahituv, G. Amato, A. Narechania, S. Singh,
86	A. Scott, and A. Paasch for advice on methods and access to cluster computing resources.
87	
88	BACKGROUND
89	Hypselodonty, or the presence of unrooted and thus ever-growing teeth, has evolved
90	multiple times in mammals. Glires-the clade containing rodents, rabbits, and their relatives-
91	have hypselodont incisors (1), and multiple Glires have also evolved hypselodont molars (Fig.

92 1). At least in rodents, molar hypselodonty evolved considerably later than hypsodont molars, 93 which are high crowned but rooted, which in turn evolved later than hypselodont incisors. In 94 Glires, molars appear to increase in crown height from low-crowned brachydont (low-crowned, 95 rooted), through hypsodonty (high-crowned, rooted), toward hypselodonty (high-crowned, 96 unrooted) (2). Mice (*Mus musculus*), the primary mammalian model species of dental research, 97 have highly derived hypselodont incisors while retaining brachydont molars. Because of this, 98 mice do not provide information about the hypsodont teeth that likely preceded hypselodonty. 99 Mammalian teeth sit in bony sockets, held in place by soft tissue (periodontal ligament) 100 attached to cementum-covered tooth roots (3). Ligamentous tooth attachment may have arisen 101 along with a reduction in the rate of tooth replacements, providing greater flexibility for 102 repositioning the teeth as the dentary grows (4,3). Consequently, the limited replacement of 103 mammalian teeth (two sets of teeth in most mammals and one in Glires) may have spurred the 104 evolution of hypsodont and hypselodont teeth, both with high crowns that compensate for tooth 105 wear from gritty or phytolith-heavy diets (5,6), and resulted in further modification of the 106 anchoring roots. The convergent evolution of unrooted molars in Glires presents an opportunity 107 to identify whether consistent developmental and genomic changes underlie the formation of 108 hypselodont teeth, in turn revealing the mechanisms that must remain unchanged to produce 109 tooth roots. Furthermore, the relatively recent evolution of molar hypselodonty, starting in the 110 Middle Miocene (approximately 16-12 Ma) (2), should provide molecular evidence for the steps 111 required to make a continuously growing organ. 112 Dental development proceeds from the tooth germ, composed of epithelium and

mesenchyme, through phases known as the bud, cap, and bell (7). Multipotent enamel epithelium
differentiates into the cells that form the tooth crown (8–11). As development progresses in

115	rooted teeth, the epithelium at the tooth apex transitions first to a tissue called Hertwig's
116	epithelial root sheath (HERS), and eventually cementum-covered roots (9,10). Studies have
117	identified numerous candidate genes and pathways with various roles during root development,
118	such as $Fgf10$, which decreases in expression at the beginning of root formation (12–18).
119	Although research on mouse molars has identified genetic signals of root formation, a number of
120	the key genes studied have broad developmental roles, such as Wnt family members (14), or
121	overlap considerably with genes also involved in patterning the size and shape of the tooth
122	(17,19–22). This overlap between shape and root expression patterns confounds our ability to
123	identify a clear signal initiating root formation.
124	Evolutionary novelties such as high-crowned hypsodont and hypselodont molars can
125	arise from differences in gene expression and regulation (23–26). Evolutionarily conserved gene
126	expression levels produce conserved phenotypes, and changes in gene regulatory networks have
127	long been linked to morphological evolution (27,28). The order of genes along a chromosome
128	(synteny) can affect gene expression and regulation, as regulatory sequences are often located
129	near their target genes (cis-regulatory elements) (29-31). Genome rearrangements that place
130	genes near new regulatory elements may result in changes of the expression levels and selective
131	environment of those genes; these small-scale rearrangements of genes may be common in
132	mammals (32-34). Genes involved in molar development are not syntenic in the mouse genome
133	nor are genes with organ-specific expression (35), and thus the regulatory effects of co-
134	localization need not apply to all dental genes at once. Changes in genome architecture between
135	Glires species thus may result in different selective and expression environments for dental genes
136	that could result in the evolution of hypselodont molars.

137 To establish a model rodent species with hypsodont molars for close comparison to 138 hypselodont molars, we sequenced and annotated a highly-complete *de novo* genome of *Myodes* 139 glareolus, the bank vole. The bank vole is increasingly used in medical and environmental 140 research, ranging from studying zoonotic diseases (36) to immune responses (37,38), and even 141 assessing environmental remediation efforts through heavy metals that accumulate in vole teeth 142 (39,40), thus our efforts may be of use beyond dental research. The bank vole's hypsodont 143 molars bridge the gap between low-crowned mouse and hypselodont prairie vole (*Microtus* 144 ochrogaster) molars. We performed a suite of genomic and transcriptomic tests of our new bank 145 vole genome in a broad phylogenetic context to test the hypothesis that dental genes are 146 undergoing positive selection and exhibit different expression patterns in species with unrooted, 147 hypselodont molars. We predicted that genes without conserved syntenic relationships in these 148 species would be more likely to have sites under positive selection or significantly different 149 expression. Our analyses revealed positive selection among two dental genes in Glires with 150 unrooted molars compared to those with rooted molars and demonstrated strong conservation of 151 dental gene expression patterns between bank voles and mice, with key differences related to the 152 timing and patterning of tooth morphology.

153

154 RESULTS

155 Orthology and synteny analyses

To identify which sequences in our bank vole (*Myodes glareolus*) genome and annotation had the same evolutionary history as dental genes identified in other Glires and assess genome rearrangements, we performed orthology and synteny analyses in a broad phylogenetic context. OrthoFinder identified 20,547 orthogroups representing 97.9% of the genes across all 24 analyzed genomes (including the human outgroup). Of the orthogroups, 6,158 had all species

161 present. In our *de novo* bank vole genome, there were 27,824 annotated genes, of which 84.2% 162 were assigned to an orthogroup. Bank vole genes were present in 16,250 orthogroups. On 163 average, the genomes included in the OrthoFinder analysis had 19,814 genes, with 98.2% of 164 those assigned to orthogroups. 165 The completeness and large scaffold N50 (4.6 Megabases) of our bank vole assembly 166 supported its inclusion in generating a Glires synteny network. Using the infomap clustering 167 algorithm, we produced 19,694 microsynteny clusters from this overall synteny network. We did 168 not expect dental genes to share the same microsynteny cluster, and instead examined whether 169 each gene was in the same microsynteny cluster in species with rooted or unrooted molars. 170 Among the microsynteny clusters containing dental genes, 28 networks lacked synteny in at least 171 half the species with unrooted molars or did not have a one-to-one relationship with an 172 orthogroup (Fig. 2).

173

174 *Positive selection analysis*

175 We hypothesized that dental genes in species with unrooted molars are undergoing 176 positive selection. Our positive selection analyses in PAML (phylogenetic analysis by maximum 177 likelihood (41)) identified 6 dental gene orthogroups undergoing site-specific positive selection 178 across Glires (Table 1). One of these genes, *Col4a1*, also was largely not syntenic in species with 179 unrooted molars (Fig. 2). Another orthogroup consisted mainly of predicted sequences similar to 180 *Runx3* but only had sequences from four species; both showed site-specific positive selection. 181 We then assessed 24 genes (those with site-specific positive selection or which lacked synteny in 182 at least half of the species with unrooted molars) for site-specific positive selection in species 183 with unrooted molars compared to species with rooted molars (branch-and-site specific positive

184 selection (42)). Two genes, *Dspp* and *Aqp1* were undergoing this branch-and-site specific 185 positive selection. Both genes had a single highly supported site (posterior probability > 0.95) 186 under positive selection in species with unrooted molars based on the Bayes Empirical Bayes 187 method for identifying sites under selection implemented in PAML (43). Dspp also had multiple 188 sites with moderate support (posterior probability > 0.75). The overall selection patterns on each 189 gene differed. Maximum likelihood estimates of selection for *Dspp* showed the percentage of 190 sites under purifying and neutral selection on all branches were nearly equal (47% and 44%, 191 respectively). Percentages of sites under positive selection in the species with unrooted molars 192 (foreground branches) were nearly evenly divided as well, with 5% of sites from branches where 193 the species with rooted molars (background branches) were undergoing purifying selection and 194 4% of sites from branches where the species with rooted molars were under neutral selection. For 195 Aqp1, nearly all sites were under purifying selection on all branches (91%), and few sites were 196 under neutral selection on all branches (7%). Few sites were undergoing positive selection in the 197 foreground branches and their distribution also was unevenly split between sites under purifying 198 and neutral selection on background branches (0.6% and 0.04%, respectively). The complete list 199 of dental genes with hierarchical orthogroups, microsynteny clusters, and positive selection test 200 results are available in Additional file 1.

Because genes under positive selection are often expressed at lower levels than genes under purifying selection (44–47), we also compared expression levels of *Dspp* and *Aqp1* in postnatal first molars (M1) at postnatal days 1, 15, and 21 (P1, P15, and P21) in bank voles (rooted molars) and prairie voles (unrooted molars) using quantitative PCR. Prairie vole molars expressed *Aqp1* at significantly lower levels in all three ages than bank vole molars (Fig. 3). Prairie vole P1 molars expressed significantly lower levels of *Dspp* than bank vole molars; at

P15 and P21, their molars expressed *Dspp* at lower, but not statistically significantly different,
levels than their bank vole equivalent. For both genes, the prairie vole had consistent expression
levels across three biological replicates, while the bank vole had greater variation in expression
levels across replicates.

211

212 Sequence and secondary structure evolution

213 To detect whether substitutions at sites under positive selection influenced protein 214 structure and evolution, we analyzed ancestral states and secondary structure across Glires. We 215 first reconstructed ancestral sequences along the internal nodes of the Glires phylogeny for the 216 genes undergoing branch-and-site specific positive selection to assess potential secondary 217 structural changes in their protein sequences. At the best-supported site in *Dspp* (position 209 in 218 the gapped alignment, Additional file 2), there were three major amino acid changes. The 219 ancestral Glires sequence started with an asparagine (N) in this position. Two of the three species 220 with unrooted molars represented in the *Dspp* dataset had amino acid substitutions at this 221 position, with Oryctolagus cuniculus substituting a leucine (L) and Dipodomys ordii substituting 222 an aspartic acid (D) at this position (Fig. 4A). All muroids (the clade including the voles in 223 family Cricetidae and mice and rats in family Muridae) in our phylogeny substituted histidine 224 (H) for the asparagine at this position. The secondary structure predicted at this position was a 225 coil for most sequences but a helix for the D. ordii sequence (Fig. 5). Appl sequences varied 226 greatly at the position under putative positive selection in species with unrooted molars (position 227 294 in the gapped alignment, Additional file 3). The ancestral state reconstruction showed twelve 228 changes of the amino acid at this position across Glires (Fig. 4B), yet these changes did not

affect the predicted secondary structure of the protein near this residue, which was a coil for allsequences tested. All secondary structure predictions are available in Additional file 4.

231

232 Developmental gene expression

233 We also assessed differential gene expression between mouse and bank vole molars 234 across early development to study the effects of morphology on expression levels of dental 235 genes. Our gene expression analysis focused on keystone dental gene categories, which are based 236 on the effects null mutations of each gene are reported to have during embryonic dental 237 development (48): "shape" genes cause morphological errors, "eruption" genes prevent tooth 238 eruption, "progression" genes stop the developmental sequence, "tissue" genes cause defects in 239 tissues, "developmental process" genes are annotated with the "GO:0032502" gene ontology 240 term, and "dispensable" genes, while dynamically expressed in developing teeth, have no 241 documented effect on phenotype. The group "other" is composed of the remaining protein 242 coding genes (48). Our bank vole genome was like the mouse and rat genomes in terms of the 243 numbers and expression patterns of genes annotated from these keystone categories (Table 2). 244 Ordination of gene expression results from the bank vole and mouse data at embryonic day 13, 245 14, and 16 (E13, E14, E16) (48) by principal components analysis showed a distinct separation 246 between the mouse and bank vole along the first principal component (PC1) of the 500 most 247 variable genes (Fig. 6A). PC1 explained 82.81% of the variance in these genes; there are distinct, 248 species-specific expression patterns in these tissues. Along PC2 (7.47% of variance explained), 249 E13 and E14 samples differ from the E16 samples, although the difference in time points is 250 much greater in bank voles. Ordination of just the keystone dental genes showed clear 251 separations between tissues based on species and age (Fig. 6B). Within this focused set of genes,

252 however, PC1 and PC2 explain less variance (44.8% and 28.84% respectively), and how species 253 and age relate to the PCs is less clear. There are two distinct, parallel trajectories for the mouse 254 and bank vole. Although within each species there is separation by age along PC1 and PC2, 255 mouse E16 and bank vole E13 occupy a similar position along PC1, and mouse E13 and bank 256 vole E16 occupy a similar position along PC2. 257 Examining individual genes underlying the differences between mouse and vole molars, 258 we note several upregulated genes in our vole molars are broadly expressed in developing molars 259 of other vole species (49,50). Relative to the mouse molars, vole molars overexpressed genes 260 related to forming tooth cusps, including Bmp2, Shh, p21 (also known as Cdkn1a), and Msx2, a 261 difference explained by the faster patterning and larger number of cusps in the vole molar 262 compared to the mouse molar (50). Another gene upregulated in the patterning stage vole molar 263 is Fgf10, which is associated with delayed root formation later in vole molar development (9). 264 Nevertheless, developing bank vole molars at E13, E14, and E16 expressed keystone 265 dental genes in overall proportions like those observed at analogous stages of mouse and rat 266 molar development (Fig. 7). Permutation tests within each bank vole sample showed that log 267 counts for the set of genes related to the progression of dental development were significantly 268 higher than those in the tissue, dispensable, developmental process, and "other" categories at E14 269 and E16. The progression gene counts in E13 molars were higher for all of these except the 270 dispensable category. Shape category genes also were significantly higher than "other" category 271 genes in the E14 tissue. Overall, even though we observed conserved expression patterns of 272 dental genes at the system level, individual genes involved in cusp patterning and morphology 273 differed between the mouse and the vole. 274

275 DISCUSSION

276 Our two goals in sequencing the genome of *Myodes glareolus* were to support the 277 development of a comparative system for studying tooth root development and to investigate the 278 evolution of dental genes in Glires, a clade in which ever-growing molars have evolved multiple 279 times (1). Our new *M. glareolus* assembly and annotation captured nearly all of the single-copy 280 orthologs for Euarchontoglires and provided scaffolds with sufficient length for synteny 281 analyses. It was well represented in ortholog groups and microsynteny clusters across Glires. We 282 tested the hypothesis that dental genes are undergoing site-specific positive selection in species 283 with unrooted molars (branch-and-site specific positive selection (42)) and exhibit differential 284 expression patterns. We predicted that lack of conserved syntenic relationships in species with 285 unrooted molars could place dental genes in regulatory and selective environments that promote 286 changes among genes relevant to tooth root formation. Our analyses revealed that most dental 287 genes have conserved syntenic relationships across Glires, yet two dental genes, *Dspp* and *Aqp1*, 288 were undergoing positive selection in species with unrooted molars. We also demonstrated 289 conserved patterns of gene expression among dental keystone genes between bank voles and 290 mice during early embryonic development, and deviations from these conserved patterns likely 291 related to differences in molar morphology between the two species.

We identified 13 genes which were not syntenic in at least half of the species with unrooted molars, and 6 genes undergoing site-specific positive selection across all Glires. Only one gene, *Col4a1*, lacked synteny and had evidence of positive selection. The two genes undergoing positive selection in species with unrooted molars, *Dspp* and *Aqp1*, both maintained their synteny relationships across the Glires studied. Although we predicted loss of synteny for dental genes in Glires with unrooted molars could result in sequence evolution by placing genes

298 in new selective contexts, our analyses did not support a relationship between non-syntenic genes 299 and positive selection. Maximum likelihood estimates of sites under different types of selection 300 for the genes with branch-specific positive selection did reveal different selective pressures on 301 Dspp and Aqp1 overall; Dspp sites on background branches (i.e., branches with species that have 302 rooted molars) were under a mix of purifying and neutral selection, while nearly all Aqp1 303 background branch sites were under purifying selection. These selection regimes suggest there is 304 greater conservation for Aqp1 function across Glires than for Dspp function. Gene duplication 305 can result in functional redundancy and evolution toward a novel function in some genes (51– 306 54), which may explain positive selection in Aqp1, as there are other aquaporin family genes 307 present. *Dspp* has no paralogs, but overlaps functionally with other SIBLING family proteins 308 (e.g., Opn, Dmp1) (55,56).

309 Aqp1 and Dspp play different functional roles during dental development. Under the 310 keystone dental development gene framework, Aqp1 is a "dispensable" gene: developing teeth 311 express it, but tooth phenotypes do not change in its absence. Appl is expressed in endothelia of 312 microvessels in the developing tooth (57,58). Dspp may be particularly relevant for the 313 formation of an unrooted phenotype if its expression domain or function have been modified in 314 species with unrooted molars. Dspp is a "tissue" category keystone dental gene, meaning the 315 main effects of a null mutation occur during the tissue differentiation stage of dental 316 development (48). Null mutations of *Dspp* cause dentin defects in a condition called 317 dentinogenesis imperfecta (59,60); in some patients, teeth form short, brittle roots (60,61). Dspp 318 knockout mice also exhibit the shortened root phenotype, among a variety of other defects in 319 both endochondral and intramembranous bone, due to the disruption of collagen and bone 320 mineralization (62-64).

321 Our ancestral sequence reconstructions and estimated secondary protein structures 322 allowed us to assess whether nonsynonymous substitutions at sites under positive selection 323 resulted in structural differences, thus potentially affecting protein function. Although unrooted 324 molars are a convergent phenotype across Glires, the sites under positive selection did not 325 converge on the same amino acid substitution in species with unrooted molars, and Aqp1 326 appeared particularly labile at this residue. The non-synonymous substitutions at these sites often 327 resulted in changes of properties of the amino acid in the sequence, for example in *Dspp*, polar 328 asparagine was replaced with non-polar leucine in O. cuniculus. Only one of these substitutions 329 changed the predicted secondary structure. Nevertheless, single amino acid substitutions do 330 produce phenotypes for both Dspp (65) and Aqp1 (66), thus we cannot rule out functional 331 changes in these genes in species with unrooted molars.

332 Although the exact relationship between gene expression and sequence divergence 333 remains unclear (67), studies of genome evolution across small numbers of mammal species 334 show correlations between gene sequence divergence and levels of expression (68). In particular, 335 highly-expressed genes are more likely to experience purifying selection (44-47), while lowly-336 expressed genes and tissue-specific genes may experience positive selection (45). The decreased 337 expression of *Dspp* and *Aqp1* in prairie vole M1 compared to that of the bank vole M1 thus 338 supports our finding of positive selection in these genes in species with unrooted molars. If all 339 species with unrooted molars also exhibit decreased expression levels of *Dspp* and *Aqp1*, it could 340 suggest a strong link between lower levels of the genes and the unrooted phenotype.

Without analyses of functional variation caused by positive selection at these coding sites, or spatial sampling to determine where these genes may be expressed during development, we are limited from exploring the specific effects of *Dspp* and *Aqp1* on root formation.

Nevertheless, we found evidence for evolution of these genes in Glires with unrooted molars, and *Dspp* especially has clinical relevance for tooth root formation. Future studies should explore the spatial distribution of *Dspp* expression, which could be relevant to functional changes in Glires with unrooted molars. If *Dspp* is relevant to the lack of root formation in hypselodont Glires incisors, the positive selection identified here may modify its expression domain or its interaction with yet-unidentified root formation co-factors, thus serially reproducing the unrooted incisor phenotype in molars.

351 Our RNA sequencing results supported the bank vole as a suitable system for studying 352 dental development. Although molar morphology differs considerably across mammals, 353 candidate-gene approaches have identified numerous conserved genes involved in tooth 354 development and morphological patterning (69). Studies of single genes or gene families have 355 identified shape-specifying roles common to multiple species (50,70-72), and high-throughput 356 sequencing of mouse and rat molars demonstrate that both species express sets of dental 357 development genes in similar proportions during early stages of tooth development (48). The 358 similarity of our high-throughput RNA sequencing results to the mouse and rat results in 359 previous studies suggest overall expression patterns of keystone dental development genes 360 within each stage may be conserved in Glires. Our principal component analyses and differential 361 expression analyses measuring changes between mouse and bank vole molars, however, showed 362 that several dental genes' expression levels differed significantly by species and age. Previous 363 research has documented organ expression patterns that are conserved across species early in 364 development and diverge over time, with some major organs displaying heterochronic shifts in 365 some species (73). If the major source of variation in keystone dental gene expression patterns 366 between mice and bank vole molars were solely attributable to species, we might expect to see

367 clear separation between the species along the first or second principal component (PC1 or PC2), 368 like that observed in PC1 of the 500 most variable genes (Fig. 6). If molar development follows 369 the diverging expression patterns observed in other organs, we might expect just the earliest age 370 classes to align on one, or multiple, PCs. Instead, we found two trajectories that were nearly 371 parallel across PC1 and PC2 and multiple keystone dental genes that were significantly 372 differentially expressed with respect to species and age. This variation between species is likely 373 driven by the larger number of cusps in the vole molar, and corresponding upregulation of genes 374 regulating cusp formation. The overall acceleration of patterning in vole molars likely explains 375 the significance of the age variable in our expression results, causing a heterochronic shift in the 376 expression patterns.

377 Our analyses were limited by the small number of rodent species with sufficiently 378 annotated genomes to be included in synteny and positive selection analyses. This limitation left 379 us with a small phylogeny for our ancestral state reconstructions, which thus did not encompass 380 the full diversity of Glires tooth roots, and potentially weakened model-based genomic analyses. 381 Although positive selection analyses using the Bayes Empirical Bayes criterion are robust to 382 smaller sample sizes (43), including fossil species in ancestral state reconstructions can change 383 estimations of ancestral characteristics (74). Innovations in paleoproteomics also offer the 384 opportunity to compare fossil species' dental gene sequences directly to living and estimated 385 ancestral sequences (75,76). By incorporating data for extinct Glires in both morphological and 386 molecular analyses, we can further elucidate links between dental gene evolution and unrooted 387 teeth.

388

389 CONCLUSIONS

390 Our genomics and transcriptomics analyses, based on our newly sequenced, high-quality 391 draft bank vole genome assembly and annotation, showed that bank vole early tooth 392 development is comparable to other commonly used rodent models in dental development 393 research. We identified 6 dental gene orthogroups that were undergoing site-specific positive 394 selection across Glires and two genes, *Dspp* and *Aqp1*, that were undergoing site-specific 395 positive selection in Glires with unrooted molars. *Dspp* appears particularly relevant to root 396 formation, as loss-of-function mutations cause a dentin production defect that can result in 397 shortened tooth roots. Future research must explore the functional role that *Dspp* plays in tooth 398 root formation in Glires and other clades. The rodent dentary is an exciting system for 399 understanding tooth development; it provides an easily manipulated set of tissues that can be 400 produced quickly and features a lifelong population of stem cells in the incisor with genomic 401 mechanisms that are potentially replicated across other teeth in species with unrooted molars. 402 Our results identify candidate genes for future analyses, and our draft bank vole genome and 403 annotation improve the utility of this species for comparative dental research that can uncover 404 the genetic mechanisms of tooth root formation.

405

406 METHODS

407 *Tissue collection and sequencing*

To assemble the bank vole genome, we sequenced tissues from a single adult male specimen housed in a colony at the UCSF Mission Center Animal Facility. We euthanized the animal according to UCSF IACUC protocol AN189916 and harvested muscle, kidney, heart, and liver tissue, which were immediately frozen at -80°C. Tissues were sent to a third-party sequencing service, where they were combined and homogenized to achieve appropriate mass

413	for high molecular weight DNA extraction. We targeted 60x coverage with 150 base pair (bp)
414	reads using 10X Chromium linked-read chemistry (77,78) and sequenced on the Illumina
415	platform. We also targeted 10x coverage with Pacific Biosciences SMRT long-read chemistry.
416	For genome annotation and gene expression analyses, we collected seven biological replicates
417	each of first molars at embryonic days 13-16 (E13, E14, E15, E16), second molars at E16, and
418	jaw tissues at E14 under University of Helsinki protocols KEK16-021, KEK19-019, and KEK17-
419	030 and stored them in RNAlater at -80°C for RNA sequencing, following a tissue harvesting
420	protocol established for mice and rats (48). We extracted RNA from these tissues using a
421	guanidium thiocyanate and phenol-chloroform protocol combined with an RNeasy column
422	purification kit (Qiagen) based on the keystone dental gene protocol (48). Single-end 84 bp RNA
423	sequencing was performed using the Illumina NextSeq 500 platform.
424	

425 *Genome assembly and quality control*

426 We first assembled only the 10X Chromium linked reads using the default settings in 427 Supernova 2.1.1. (77,78). We selected the "pseudohaplotype" (pseudohap) output format, which 428 randomly selects between potential alleles when there are two possible contigs assembled for the 429 same region. This option produces two assemblies, each with a single resolved length of the 430 genome sequence (77–79). We used our lower-coverage, long-read data for gap filling and 431 additional scaffolding. First, we estimated the genome's length using the raw sequence data in 432 GenomeScope (80), which predicted a length of 2.6 gigabases. We then performed error 433 correction of the long reads using Canu (81), removing reads shorter than 500 bp and 434 disregarding overlaps between reads of fewer than 350 bp. We kept only those reads with 435 minimum coverage of 3x for scaffolding. Following long read error correction, we used Cobbler

and RAILS (82) with a minimum alignment length of 200 bases to accept matches for gap fillingand scaffolding of both pseudohap assemblies.

- 438 For quality control, we assessed both unscaffolded and long-read scaffolded pseudohap 439 assemblies by standard assembly length statistics with QUAST (83) and presence of single-copy 440 orthologs with BUSCO v3 (84). Both scaffolded assemblies were approximately 2.44 Gigabases 441 long, with an N50 (the length of the shortest scaffold at 50% of the total assembly length) of 4.6 442 Megabases; we refer to them as Pseudohap1+LR and Pseudohap2+LR. The Pseudohap1+LR 443 assembly had 17,528 scaffolds over 1000bp (base pairs) long, and the Pseudohap2+LR assembly 444 had 17,518 scaffolds over 1000bp long (Table 3). BUSCO searched for universal single-copy 445 orthologs shared by Euarchontoglires, recovering 89.4% of these genes in the scaffolded 446 Pseudohap1+LR assembly and 92.8% of the single-copy orthologs in the scaffolded 447 Pseudohap2+LR assembly (Fig. 8). The two assemblies were similar length and contiguity, but 448 because the scaffolded Pseudohap2+LR assembly recovered more single-copy orthologs, we 449 based annotation and downstream analyses on it. 450 451 *Genome annotation* 452 We annotated the genome using three rounds of the MAKER pipeline (85–87). MAKER 453 combines multiple lines of evidence to annotate a genome. For evidence from gene transcripts, 454 we assembled a *de novo* transcriptome assembly based on the single-end RNA sequencing of all 455 molar and jaw tissues using Trinity (88). We also included cDNA sequences from the Mus
- 456 musculus assembly GRCm38. We used SwissProt's curated protein database to identify protein

457 homology in the genome. Two libraries of repeats provided information for repeat masking: the

458 Dfam Rodentia repeat library (89–91) and a custom library specific to the bank vole estimated

459	based on the modified protocol of Campbell et al. (86). The custom library features miniature
460	inverted-repeat transposable elements identified with default settings in MiteFinder (92), long
461	terminal repeat retrotransposons extracted with the GenomeTools LTRharvest and LTRdigest
462	functions (93) based on the eukaryotic genomic tRNA database, and <i>de novo</i> repeats identified
463	with RepeatModeler (94). We combined elements identified by these programs into a single
464	repeat library, then removed any elements that matched to a custom SwissProt curated protein
465	database with known transposons excluded; this custom repeat library is available in Additional
466	file 5. We trained a custom gene prediction model for MAKER as well. The first iteration of the
467	model came from BUSCO's implementation of augustus (95). Between each round of MAKER
468	annotation, we updated the gene prediction model with augustus.
469	MAKER considered only contigs between 10,000-300,000 bp long during annotation.
470	Our second and third iterations of MAKER used the same settings but excluded the
471	"Est2genome" and "protein2genome" functions, as recommended in the MAKER tutorial. We
472	included a SNAP (96) gene prediction model based on the output of the first round of annotation
472 473	
	included a SNAP (96) gene prediction model based on the output of the first round of annotation
473	included a SNAP (96) gene prediction model based on the output of the first round of annotation during the second and third iterations of MAKER annotation. Annotation quality (i.e., agreement
473 474	included a SNAP (96) gene prediction model based on the output of the first round of annotation during the second and third iterations of MAKER annotation. Annotation quality (i.e., agreement between different lines of evidence and the MAKER annotation) was assessed visually in
473 474 475	included a SNAP (96) gene prediction model based on the output of the first round of annotation during the second and third iterations of MAKER annotation. Annotation quality (i.e., agreement between different lines of evidence and the MAKER annotation) was assessed visually in JBrowse after each iteration and using <i>compare_annotations_3.2.pl</i> (97), which calculates the
473 474 475 476	included a SNAP (96) gene prediction model based on the output of the first round of annotation during the second and third iterations of MAKER annotation. Annotation quality (i.e., agreement between different lines of evidence and the MAKER annotation) was assessed visually in JBrowse after each iteration and using <i>compare_annotations_3.2.pl</i> (97), which calculates the number of coding and non-coding sequences in the annotation in addition to basic statistics about
473 474 475 476 477	included a SNAP (96) gene prediction model based on the output of the first round of annotation during the second and third iterations of MAKER annotation. Annotation quality (i.e., agreement between different lines of evidence and the MAKER annotation) was assessed visually in JBrowse after each iteration and using <i>compare_annotations_3.2.pl</i> (97), which calculates the number of coding and non-coding sequences in the annotation in addition to basic statistics about sequence lengths. Our MAKER annotation covered 2.41 Gb of the scaffolded Pseudohap2

the different types of evidence for an annotation, where scores closer to zero represent better-annotated genes (98).

483

484 Orthology and synteny analyses

485 We analyzed orthology and synteny of the bank vole genome to understand gene and 486 genome evolution related to dental development across Glires with rooted and unrooted molars. 487 We obtained genomes from Ensembl for 23 Glires species and one phylogenetic outgoup, Homo 488 sapiens (Table 4). These genomes all had an N50 over 1 Mb, which improves synteny 489 assessment (99). We first analyzed all 24 genomes for groups of orthologous genes (orthogroups) 490 in OrthoFinder (100), providing a tree topology based on the Ensembl reference tree (Fig. 1) to 491 guide orthology detection. Because we would not analyze the human outgroup in downstream 492 analyses, we implemented the OrthoFinder option that splits orthogroups at the root of Glires 493 (hierarchical orthogroups), thus any group of orthologs studied here represents only genes with 494 shared, orthologous evolutionary history within Glires. We selected MAFFT (101) for multiple 495 sequence alignment and fastme (102) for phylogenetic tree searches within OrthoFinder; we 496 retained the gene trees estimated for each orthogroup for downstream analyses. 497 Although dental development genes are spread throughout the genome, we were 498 interested in whether each gene remained in the same local arrangement across species of Glires. 499 We prepared each genome annotation and sequence file for synteny analysis using the 500 reformatting functions of Synima (103) to extract each peptide sequence associated with a gene 501 coding sequence in the Ensembl annotation. Collinear synteny blocks estimated by MCScanX 502 (104) formed the basis for microsynteny network analyses using the SynNet pipeline (105-107).

503 We inferred networks from the top five hits for each gene, requiring any network to have a

504 minimum of 5 collinear genes and no more than 15 genes between a collinear block, settings that 505 perform well for analyzing mammal genomes (107). Using the infomap algorithm, we clustered 506 the synteny blocks into microsynteny networks, from which we extracted network clusters 507 corresponding to the list of keystone dental genes (48). For each dental gene microsynteny 508 network, we assessed whether genes of species with unrooted molars were not syntenic with the 509 other Glires species' sequences.

510

511 *Positive selection analysis*

512 We aligned protein sequences for each dental gene orthogroup with clustal omega (108) 513 using default settings. Based on universal translation tables, we obtained codon-based nucleotide 514 alignments with pal2nal (109), removing sites in which any species had an indel (i.e., ungapped) 515 and formatting the output for analysis in PAML (41). We pruned and unrooted the orthogroup 516 gene trees from OrthoFinder to contain only tips representing the genes in each synteny network 517 or orthogroup under analysis in PAML. We tested whether any of the genes were undergoing 518 positive selection using a likelihood ratio test comparing site-specific models of "nearly neutral" 519 and positive selection. In these models, ω , the ratio of nonsynonymous to synonymous 520 nucleotide substitutions (also known as dN/dS), can vary at each codon site. In the "nearly 521 neutral" model, ω can take values between 0 and 1, while the positive selection model allows 522 sites to assume ω values greater than 1 (43,110). We allowed PAML to estimate κ (the ratio of 523 transitions to transversions) and ω from initial values of 1 and 0.5, respectively, for both tests. 524 Dental genes with significant site-specific positive selection or those for which over half 525 the unrooted species' sequences were not in the same synteny block as sequences for species 526 with rooted molars formed the basis for our second set of positive selection tests using a branch-

527 and-site model of positive selection. This model allows ω to vary not only among codon sites, 528 but also between "foreground" and "background" lineages (43). We marked the species with 529 unrooted molars as foreground lineages, then ran the model twice: once with ω unconstrained to 530 detect sites undergoing positive selection only on foreground branches, and a second time and 531 with ω fixed to 1, or neutral selection. A likelihood ratio test of the two models determined 532 whether the lineage-specific positive selection model was more likely than a neutral model, and 533 Bayes Empirical Bayes analyses (43) produced posterior probabilities to identify sites under 534 positive selection.

535 Genes under positive selection also tend to have lower expression levels (45), thus we 536 compared expression of the genes with branch-and-site specific positive selection between the 537 prairie (unrooted molars) and the bank vole (rooted molars) to provide further support for 538 selective differences. We collected three biological replicates of first molars from both species at 539 three postnatal stages (P1, P15, and P21) and immediately preserved them at -80°C in lysis 540 buffer (Buffer RLT; Qiagen) supplemented with 40 µM dithiothreitol. RNA was extracted from 541 homogenized tissues using a RNeasy column purification kit (Qiagen). We assessed 542 concentration and purity of extracted RNA using a NanoDrop 2000 spectrophotometer 543 (ThermoFisher Scientific). Using 1 µg of RNA, we synthesized cDNA using a high-capacity 544 cDNA reverse transcription kit (ThermoFisher Scientific). We used 1 µL diluted cDNA (1:3 in 545 ddH₂O) and iTaq Universal SYBR Green Supermix (Bio-rad) in the Bio-rad CFX96 real-time 546 PCR detection system for qPCR experiments, producing three technical replicates for each 547 biological replicate. We normalized cycle threshold (CT) values of genes of interest to GAPDH 548 expression levels and calculated relative expression levels as $2^{-\Delta\Delta CT}$. A two-tailed unpaired t-test 549 calculated in Prism 9 measured whether expression of these genes significantly differed between

550 bank voles and prairie voles. The oligonucleotide primers for each species and gene are in

- 551 Additional file 6.
- 552
- 553 Sequence and secondary structure evolution

554 We performed ancestral sequence reconstruction on the codon sequences of the genes 555 that had evidence of branch-and-site specific positive selection to understand how the sequence 556 has changed through time. The gapped clustal omega alignments were the basis for ancestral 557 sequence reconstruction on the Glires species tree (Fig. 1) using pagan2 (111). For each gene, we 558 plotted amino acid substitutions at the site with potential positive selection. Finally, we predicted 559 secondary structures (i.e., helices, beta sheets, and coils) for each unrooted species' protein 560 sequence and the reconstructed ancestral sequence prior to the change at the site under positive 561 selection using the PSIPRED 4.0 protein analysis workbench (112,113). Comparing these 562 predictions across the phylogeny, we assessed how these substitutions at the site under selection 563 may affect the structure of each protein. 564 565 Developmental gene expression

We used performed quality control and filtering of the short reads for the seven replicates of first molar tissues at E13, E14, and E16 using the nf-core/rnaseq v. 3.11.2 workflow (114) for comparability to previous mouse and rat analyses (48). RNAseq reads were evaluated and adapter sequences were filtered using FastQC v. 0.11.9 (115) and Cutadapt v. 3.4 (116), and ribosomal RNA was removed using SortMeRNA v. 4.3.4 (117). We then aligned trimmed sequences to our bank vole annotation using Salmon v. 1.10.1 (118). Counts were then normalized by gene length. We categorized gene count data into functional groups based on their

573	established roles in tooth bud development (48) using the one-to-one orthology list between our
574	bank vole genome and the mouse GRCm39.103 genome annotation generated from our
575	OrthoFinder output. Using the rlog function of DESeq2 (119), we normalized gene counts within
576	each functional group on a log2 scale. A permutation test assessed whether the mean counts of
577	the progression, shape, and double functional groups were significantly different from genes in
578	the tissue, dispensable, and "other" groups (which are potentially relevant later in development)
579	based on 10,000 resampling replicates of the dataset (48).
580	We also assessed differential expression between the bank vole first molar and published
581	mouse M1 data at the same three time points (GEO accession GSE142199 (48)), combining the
582	data based on the one-to-one orthology relationships used in the functional permutation analysis.
583	Using the mouse E13 molar as the reference level, we modeled expression as a response to
584	species (mouse or vole), embryonic day (E13, E14, or E16), and the interaction between species
585	and day. We considered as significant any gene with a log fold change greater than 1, log fold
586	change standard error less than 0.5, and false discovery rate adjusted p value less than 0.05.
587	
588	
589	
590	
591	
592	
593	
594	
595	

596 TABLES

597 Table 1 – Genes undergoing site-specific and branch-and-site-specific positive selection

Gene	Mus transcript	Myodes transcript	Site	Branch-and-site
Aqp1	ENSMUST0000004774	Mglareolus_00011822	Yes	Yes
Col4a1	ENSMUST0000033898	Mglareolus_00032740	Yes	No
Dspp	ENSMUST00000112771	Mglareolus_00014030	Yes	Yes
Fgf20	ENSMUST0000034014	Mglareolus_00013079	Yes	No
Runx3	ENSMUST00000056977	Mglareolus_00033992	Yes	No
similar to <i>Runx3</i>	_	_	Yes	No*

598 Table 1 Legend: *HOG only contained four genes with one unrooted species' sequence, could

- 599 not be tested for branch-and-site specific selection.
- 600

601 Table 2 – P-values of permutation tests between keystone gene categories in bank vole M1

602 at embryonic days 13, 14, and 16

		Tissue	Dispensable	Dev. Process	Other
E13	Progression	0.0310	0.0942	0.0436	0.0402
	Shape	0.6431	0.9041	0.2289	0.0995
	Double	0.1292	0.1521	0.0716	0.0655
E14	Progression	0.0136	0.0383	0.0437	0.0401
	Shape	0.3115	0.4725	0.0922	0.0454
	Double	0.1288	0.0945	0.0709	0.0630
E16	Progression	0.0140	0.0401	0.0303	0.0274
	Shape	0.3770	1	0.1831	0.0662
	Double	0.1343	0.1099	0.0638	0.0596

603 Table 2 Legend: Italicized values are statistically significant (p < 0.05)

604 Table 3 – QUAST assembly statistics for *de novo* bank vole (*Myodes glareolus*) genome

605 assemblies

	Pseudohap1	Pseudohap1+LR	Pseudohap2	Pseudohap2+LR*
Largest contig	27939478	32658832	27937749	32657565
Total length	2434151515	2441426554	2434099357	2441472313
GC (%)	41.88	41.89	41.88	41.89
N50	4187179	4579815	4187179	4558134
N75	1689669	1818134	1687188	1810460
L50	170	153	170	154
L75	388	357	388	358
Ns per 100 kbp	1151.99	1030.75	1151.96	1030.48

606 Table 3 Legend: *assembly used for annotation and downstream analyses in this paper.

607

608 Table 4 – Genomes used in orthology, synteny, and positive selection analyses

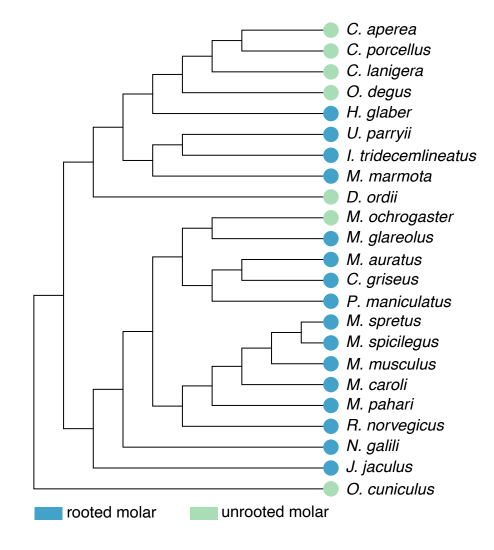
Species	Assembly	Citation
Myodes glareolus	CUNY_Mgla_1.0	This paper
Cavia porcellus*	Cavpor3.0	(120)
Cavia aperea*	CavAp1.0	(121)
Marmota marmota	marMar2.1	(122)
Microtus ochrogaster*	MicOch1.0	(123)
Mus musculus	GRCm39	(124)
Oryctolagus cuniculus*	OryCun2.0	(120)
Dipodomys ordii*	Dord_2.0	(120)
Jaculus jaculus	JacJac1.0	(125)

Rattus norvegicus	Rnor_6.0	(126)
Mus pahari	PAHARI_EIJ_v1.1	(127)
Mus caroli	CAROLI_EIJ_v1.1	(127)
Mus spretus	SPRET_EiJ_v1	(128)
Mus spicilegus	MUSP714	(129)
Cricetulus griseus	CHOK1GS	(130)
Mesocricetus auratus	MesAur1.0	(131)
Peromyscus maniculatus	HU_Pman_2.1	(132)
Nannospalax galili	S.galili_v1.0	(133)
Octodon degus*	OctDeg1.0	(134)
Heterocephalus glaber (F)	HetGla_female_1.0	(135)
Chinchilla lanigera*	ChiLan1.0	(136)
Urocitellus parryi	ASM342692v1	(137)
Ictidomys tridecemlineatus	SpeTri2.0	(138)
Homo sapiens**	GRCh38	(139)

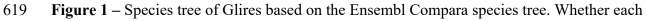
609 Table 4 Legend: *Species with unrooted molars; **Peptide annotation used as outgroup only in

- 610 OrthoFinder analysis.

617 FIGURES



618



620 species has rooted or unrooted molars is indicated at the tip of each branch. Note that unrooted,

- or hypselodont, molars have evolved multiple times across Glires. This topology was the basis
- 622 for our orthology analysis.

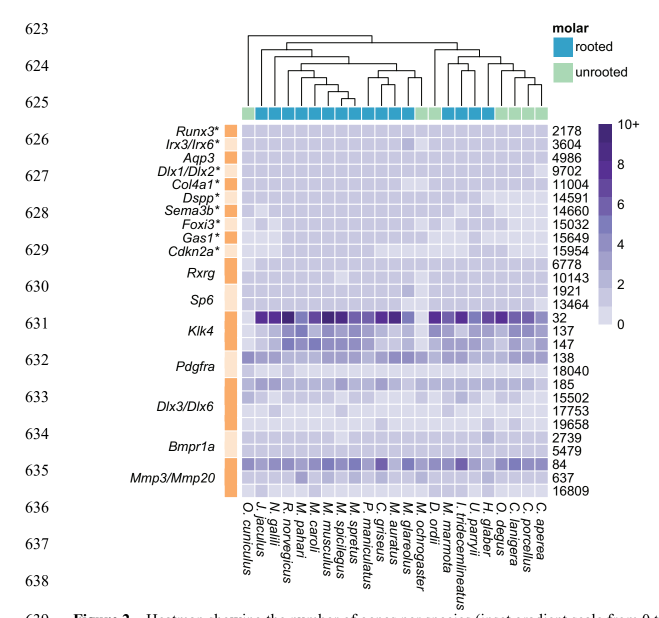
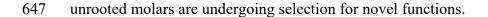


Figure 2 – Heatmap showing the number of genes per species (inset gradient scale from 0 to 10+) in each synteny cluster. The figure shows clusters where species with unrooted molars had no representation or did not have sequences in all microsynteny clusters associated with a single gene, clusters where more than one gene mapped to the same cluster, or a single gene mapped to multiple clusters. Microsynteny cluster number is noted on the right side of the heatmap (one row per cluster), and corresponding genes are noted on the left with alternating bands showing rows to which those genes mapped. * = genes where hierarchical orthogroup did not contain

646 genes for all 23 species. We found little evidence that non-syntenic genes in species with



648

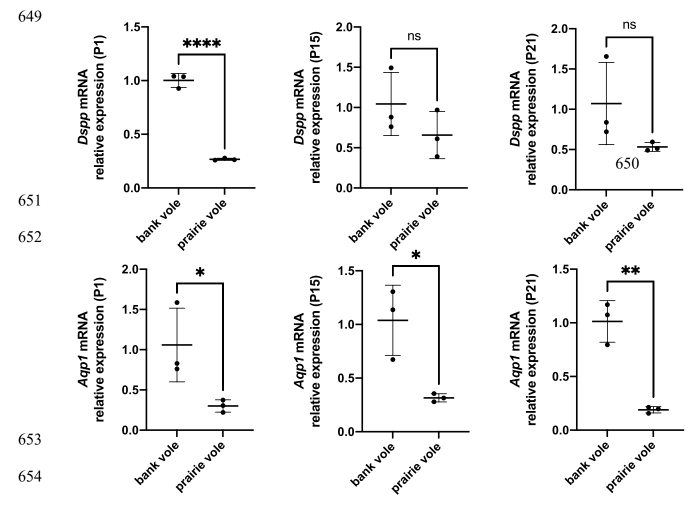
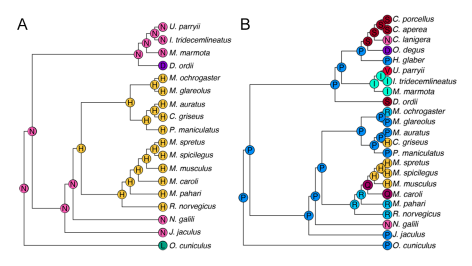


Figure 3 – Quantitative PCR comparisons of *Dspp* and *Aqp1* expression between bank vole and
prairie vole M1 at postnatal days 1, 15, and 21 (P1, P15, P21). Expression levels for both genes
are lower in the prairie vole (unrooted molars), which supports the positive selection detected for
these genes in species with unrooted molars.

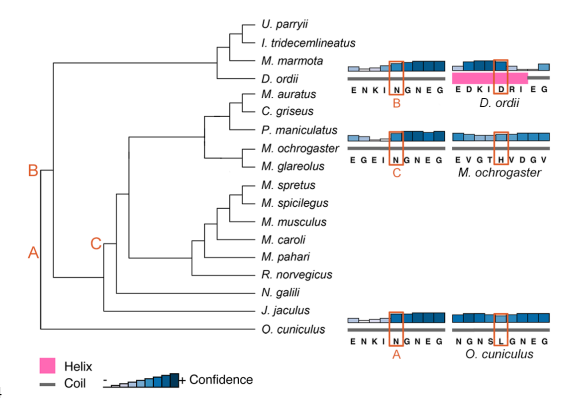
659



668 **Figure 4** – Ancestral state reconstructions of the residue under positive selection in PAML tests.

669 Letters at tips and internal nodes represent IUPAC codes for amino acids. A Dspp; B Aqp1.

- 670
- 671
- 672
- 673





675 **Figure 5** – PSIPRED secondary structure predictions for the three species with unrooted molars

676 represented in the *Dspp* sequences. Letters correspond to the most recent ancestor of each tip

677 species that had a different amino acid at the position under positive selection.

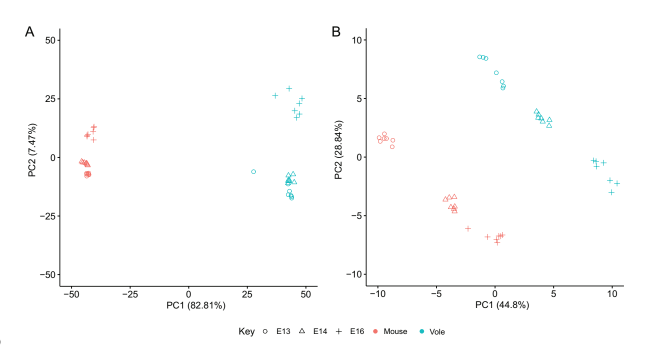




Figure 6 – Principal component (PC) analyses of differentially expressed genes in mouse and
bank vole M1. A PC1 and PC2 of the 500 most variable genes, showing a clear differentiation
between species along PC1 and differentiation between age classes along PC2. B PC1 and PC2
of the keystone dental genes. Both PC1 and PC2 separate age classes within, but not between,
the species, likely due to differences in developmental timing and molar morphology between
mice and voles.



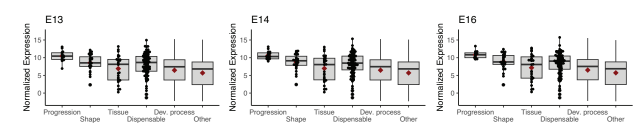
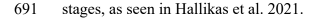
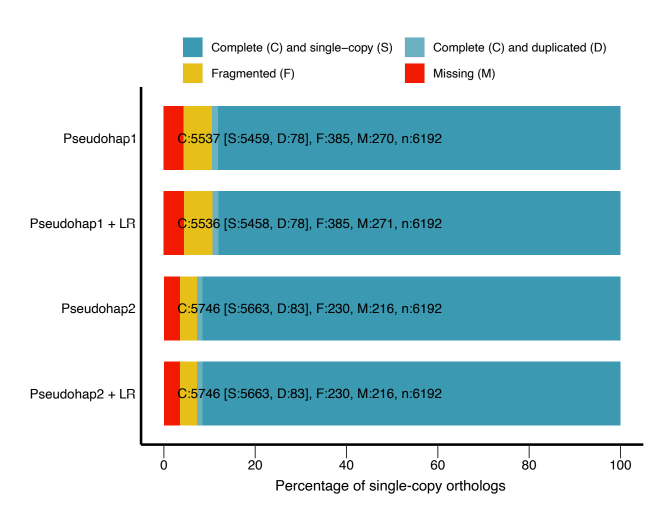


Figure 7 – Box and whisker plots showing normalized log base 2 expression levels for each
keystone gene category in bank vole M1 at embryonic days 13, 14, and 16. Gene expression

690 profiles at these stages are comparable to mouse and rat molars at analogous developmental



692





694 Figure 8 – BUSCO single-copy ortholog recovery for each "pseudohaploid" version of our draft

bank vole genome assembly and these version after long-read scaffolding (denoted by "+ LR").

Each bar represents the cumulative proportion of the 6,192 single-copy orthologs for

697 Euarchontoglires identified by BUSCO represented by complete single-copy, complete-

698 duplicated, fragmented, and missing orthologs. The Pseudohap2 and Pseudohap2 + LR

assemblies had the best single-copy ortholog recovery.

701 ADDITIONAL FILES

- 702 Additional file 1 [.xlsx] Dental gene results Full table of orthology, synteny, and positive
- selection test results for all dental genes assessed.
- 704 Additional file 2 [.txt] Dspp gapped alignment Gapped codon-based alignment for Dspp in
- 705 fasta formatted sequences.
- 706 Additional file 3 [.txt] Aqp1 gapped alignment Gapped codon-based alignment for Aqp1 in
- 707 fasta formatted sequences.
- 708 Additional file 4 [.pdf] Structure predictions PSIPRED Secondary structure predictions for
- each ancestral node and unrooted molar tip species for *Dspp* and *Aqp1*.
- 710 Additional file 5 [.txt] Custom repeat library Custom repeat library of fasta formatted
- 711 sequences used in annotation of the draft Myodes glareolus genome. See Methods for description
- of the process used to generate the library.
- 713 Additional file 6 [.pdf] Oligonucleotide primers List of oligonucleotide primers for *Dspp*,
- 714 *Aqp1*, and *GAPDH* used in bank vole and prairie vole qPCR experiments.
- 715

716 REFERENCES

- Renvoisé E, Michon F. An Evo-Devo perspective on ever-growing teeth in mammals and
 dental stem cell maintenance. Front Physiol. 2014;5 AUG(August):1–12.
- 2. Tapaltsyan V, Eronen JT, Lawing AM, Sharir A, Janis C, Jernvall J, et al. Continuously
- growing rodent molars result from a predictable quantitative evolutionary change over 50
- 721 million years. Cell Rep. 2015;11(5):673–80.

722	3.	LeBlanc ARH, Brink KS, Whitney MR, Abdala F, Reisz RR. Dental ontogeny in extinct
723		synapsids reveals a complex evolutionary history of the mammalian tooth attachment
724		system. Proc R Soc B Biol Sci. 2018 Nov 7;285(1890):20181792.
725	4.	Saffar JL, Lasfargues JJ, Cherruau M. Alveolar bone and the alveolar process: the socket that
726		is never stable. Periodontol 2000. 1997;13(1):76–90.
727	5.	Davit-Béal T, Tucker AS, Sire JY. Loss of teeth and enamel in tetrapods: Fossil record,
728		genetic data and morphological adaptations. J Anat. 2009;214(4):477-501.
729	6.	Damuth J, Janis CM. On the relationship between hypsodonty and feeding ecology in
730		ungulate mammals, and its utility in palaeoecology. Biol Rev. 2011;86(3):733-58.
731	7.	Miletich I, Sharpe PT. Normal and abnormal dental development. Hum Mol Genet. 2003 Apr
732		2;12(suppl_1):R69–73.
733	8.	Harada H, Kettunen P, Jung HS, Mustonen T, Wang YA, Thesleff I. Localization of putative
734		stem cells in dental epithelium and their association with Notch and FGF signaling. J Cell
735		Biol. 1999;147(1):105–20.
736	9.	Tummers M, Thesleff I. Root or crown: A developmental choice orchestrated by the
737		differential regulation of the epithelial stem cell niche in the tooth of two rodent species.
738		Development. 2003;130(6):1049–57.
739	10.	Thesleff I, Tummers M. Tooth organogenesis and regeneration. In: StemBook. Cambridge,
740		MA: Harvard Stem Cell Institute; 2008.

741	11. Krivanek J, Buchtova M, Fried K, Adameyko I. Plasticity of dental cell types in
742	development, regeneration, and evolution. J Dent Res. 2023 Jun 1;102(6):589-98.
743	12. Luan X, Ito Y, Diekwisch TGH. Evolution and development of Hertwig's epithelial root
744	sheath. Dev Dyn. 2006;235(5):1167–80.
745	13. Kumakami-Sakano M, Otsu K, Fujiwara N, Harada H. Regulatory mechanisms of Hertwig's
746	epithelial root sheath formation and anomaly correlated with root length. Exp Cell Res.
747	2014;325(2):78–82.
748	14. Wen Q, Jing J, Han X, Feng J, Yuan Y, Ma Y, et al. <i>Runx2</i> regulates mouse tooth root
749	development via activation of WNT inhibitor NOTUM. J Bone Miner Res.
750	2020;35(11):2252–64.
751	15. Yang S, Choi H, Kim TH, Jeong JK, Liu Y, Harada H, et al. Cell dynamics in Hertwig's
752	epithelial root sheath are regulated by β -catenin activity during tooth root development. J
753	Cell Physiol. 2021;236(7):5387–98.
754	16. Yamashiro T, Tummers M, Thesleff I. Expression of bone morphogenetic proteins and Msx
755	genes during root formation. J Dent Res. 2003;82(3):172-6.
756	17. Yokohama-Tamaki T, Ohshima H, Fujiwara N, Takada Y, Ichimori Y, Wakisaka S, et al.
757	Cessation of Fgf10 signaling, resulting in a defective dental epithelial stem cell
758	compartment, leads to the transition from crown to root formation. Development.
759	2006;133(7):1359–66.

760	18. Ota MS, Vivatbutsin P, Nakahara T, Eto K. Tooth root development and the cell-based
761	regenerative therapy. J Oral Tissue Eng. 2007;4(3):137-42.
762	19. Jernvall J, Thesleff I. Reiterative signaling and patterning during mammalian tooth
763	morphogenesis. Mech Dev. 2000;92:19–29.
764	20. Harada H, Toyono T, Toyoshima K, Yamasaki M, Itoh N, Kato S, et al. FGF10 maintains
765	stem cell compartment in developing mouse incisors. Dev Camb Engl. 2002;129(6):1533-
766	41.
767	21. Tapaltsyan V, Charles C, Hu J, Mindell D, Ahituv N, Wilson GM, et al. Identification of
768	novel <i>Fgf</i> enhancers and their role in dental evolution. Evol Dev. 2016;18(1):31–40.
769	22. Christensen MM, Hallikas O, Das Roy R, Väänänen V, Stenberg OE, Häkkinen TJ, et al.
770	The developmental basis for scaling of mammalian tooth size. Proc Natl Acad Sci. 2023
771	Jun 20;120(25):e2300374120.
772	23. Chen ZJ. Genetic and epigenetic mechanisms for gene expression and phenotypic variation
773	in plant polyploids. Annu Rev Plant Biol. 2007;58(1):377-406.
774	24. Stranger BE, Forrest MS, Dunning M, Ingle CE, Beazley C, Thorne N, et al. Relative impact
775	of nucleotide and copy number variation on gene expression phenotypes. Science. 2007 Feb
776	9;315(5813):848–53.
777	25. Romero IG, Ruvinsky I, Gilad Y. Comparative studies of gene expression and the evolution
778	of gene regulation. Nat Rev Genet. 2012 Jul;13(7):505–16.

779	26. de Montaigu A, Giakountis A, Rubin M, Tóth R, Cremer F, Sokolova V, et al. Natural
780	diversity in daily rhythms of gene expression contributes to phenotypic variation. Proc Natl
781	Acad Sci. 2015 Jan 20;112(3):905–10.
782	27. Erwin DH, Davidson EH. The last common bilaterian ancestor. Development. 2002 Jul
783	1;129(13):3021–32.
784	28. Irie N, Kuratani S. Comparative transcriptome analysis reveals vertebrate phylotypic period
785	during organogenesis. Nat Commun. 2011;2:248.
786	29. Koonin EV. Evolution of genome architecture. Int J Biochem Cell Biol. 2009 Feb
787	1;41(2):298–306.
788	30. Wray GA. The evolutionary significance of cis-regulatory mutations. Nat Rev Genet. 2007
789	Mar;8(3):206–16.
790	31. Acemel RD, Maeso I, Gómez-Skarmeta JL. Topologically associated domains: A successful
791	scaffold for the evolution of gene regulation in animals. WIREs Dev Biol. 2017;6(3):e265.
792	32. Coghlan A, Eichler EE, Oliver SG, Paterson AH, Stein L. Chromosome evolution in
793	eukaryotes: A multi-kingdom perspective. Trends Genet. 2005 Dec 1;21(12):673-82.
794	33. Swenson KM, Blanchette M. Large-scale mammalian genome rearrangements coincide with
795	chromatin interactions. Bioinformatics. 2019 Jul 15;35(14):i117-26.
796	34. Long HS, Greenaway S, Powell G, Mallon AM, Lindgren CM, Simon MM. Making sense of
797	the linear genome, gene function and TADs. Epigenetics Chromatin. 2022 Jan 29;15(1):4.

798	35.	Das I	Rov R	Hallikas O	Christensen	MM.	Renvoisé E.	Jernvall J	Chromosomal
170	$\mathcal{I}\mathcal{I}$		\mathbf{x}	1 Iumab O		TATTAT'		Joinvan J.	Chiomosoniui

- neighbourhoods allow identification of organ specific changes in gene expression. PLOS
 Comput Biol. 2021 Sep 10;17(9):e1008947.
- 801 36. Torelli F, Zander S, Ellerbrok H, Kochs G, Ulrich RG, Klotz C, et al. Recombinant IFN-γ
- 802 from the bank vole *Myodes glareolus*: A novel tool for research on rodent reservoirs of
- 2003 zoonotic pathogens. Sci Rep. 2018;8(1):1–11.
- 804 37. Kloch A, Babik W, Bajer A, Siński E, Radwan J. Effects of an MHC-DRB genotype and
- allele number on the load of gut parasites in the bank vole *Myodes glareolus*. Mol Ecol.

806 2010;19(SUPPL. 1):255–65.

- 38. Migalska M, Sebastian A, Konczal M, Kotlík P, Radwan J. *De novo* transcriptome assembly
 facilitates characterisation of fast-evolving gene families, MHC class I in the bank vole
- 809 (*Myodes glareolus*). Heredity. 2017;118(4):348–57.
- 810 39. Appleton J, Lee KM, Sawicka Kapusta K, Damek M, Cooke M. The heavy metal content of
- 811 the teeth of the bank vole (*Clethrionomys glareolus*) as an exposure marker of

812 environmental pollution in Poland. Environ Pollut. 2000;110:441–9.

- 40. Gdula-Argasińska J, Appleton J, Sawicka-Kapusta K, Spence B. Further investigation of the
 heavy metal content of the teeth of the bank vole as an exposure indicator of environmental
 pollution in Poland. Environ Pollut. 2004;131(1):71–9.
- 41. Yang Z. PAML 4: Phylogenetic Analysis by Maximum Likelihood. Mol Biol Evol. 2007
 Aug 1;24(8):1586–91.

818	42. Zhang J, Nielsen R, Yang Z. Evaluation of an improved branch-site likelihood method for
819	detecting positive selection at the molecular level. Mol Biol Evol. 2005 Dec;22(12):2472–9.
820	43. Yang Z, Wong WSW, Nielsen R. Bayes Empirical Bayes inference of amino acid sites under
821	positive selection. Mol Biol Evol. 2005 Apr 1;22(4):1107–18.
822	44. Drummond DA, Bloom JD, Adami C, Wilke CO, Arnold FH. Why highly expressed proteins
823	evolve slowly. Proc Natl Acad Sci. 2005 Oct 4;102(40):14338-43.
824	45. Kosiol C, Vinař T, Fonseca RR da, Hubisz MJ, Bustamante CD, Nielsen R, et al. Patterns of
825	positive selection in six mammalian genomes. PLOS Genet. 2008 Aug 1;4(8):e1000144.
826	46. Martincorena I, Luscombe NM. Non-random mutation: The evolution of targeted
827	hypermutation and hypomutation. BioEssays. 2013;35(2):123–30.
828	47. Martincorena I, Roshan A, Gerstung M, Ellis P, Van Loo P, McLaren S, et al. High burden
829	and pervasive positive selection of somatic mutations in normal human skin. Science. 2015
830	May 22;348(6237):880–6.
831	48. Hallikas O, Das Roy R, Christensen MM, Renvoisé E, Sulic AM, Jernvall J. System-level
832	analyses of keystone genes required for mammalian tooth development. J Exp Zoolog B
833	Mol Dev Evol. 2021;336(1):7–17.
834	49. Keränen SVE, Åberg T, Kettunen P, Thesleff I, Jernvall J. Association of developmental
835	regulatory genes with the development of different molar tooth shapes in two species of
836	rodents. Dev Genes Evol. 1998;208(9):477-86.

837	50 Jornvall I	Karönan SVF	Theoloff I	Evolutionary	modification	of development in
03/	JU. Jernvan J.	, Keranen SvE.	I nesien I.	Evolutionary	mounication	of development in

838 mammalian teeth: Quantifying gene expression patterns and topography. Proc Natl Acad

839 Sci. 2000;97(26):14444–8.

- 840 51. Hughes AL. The evolution of functionally novel proteins after gene duplication. Proc R Soc
- 841 Lond B Biol Sci. 1997 Jan;256(1346):119–24.
- 842 52. Wagner A. Selection and gene duplication: A view from the genome. Genome Biol. 2002
 843 Apr 15;3(5):reviews1012.1.
- 844 53. David KT, Oaks JR, Halanych KM. Patterns of gene evolution following duplications and

speciations in vertebrates. PeerJ. 2020 Mar 31;8:e8813.

- 54. Copley SD. Evolution of new enzymes by gene duplication and divergence. FEBS J.
 2020;287(7):1262–83.
- 848 55. Fisher LW. DMP1 and DSPP: Evidence for duplication and convergent evolution of two

849 SIBLING proteins. Cells Tissues Organs. 2011 Aug;194(2–4):113–8.

- 850 56. Bouleftour W, Juignet L, Bouet G, Granito RN, Vanden-Bossche A, Laroche N, et al. The
- role of the SIBLING, bone sialoprotein in skeletal biology Contribution of mouse
- experimental genetics. Matrix Biol. 2016 May 1;52–54:60–77.
- 853 57. Felszeghy S, Módis L, Németh P, Nagy G, Zelles T, Agre P, et al. Expression of aquaporin
- 854 isoforms during human and mouse tooth development. Arch Oral Biol. 2004 Apr
- 855 1;49(4):247–57.

856	58. Yoshii T, Harada F, Saito I, Nozawa-Inoue K, Kawano Y, Maeda T. Immunoexpression of
857	aquaporin-1 in the rat periodontal ligament during experimental tooth movement. Biomed
858	Res. 2012;33(4):225–33.
859	59. Zhang X, Zhao J, Li C, Gao S, Qiu C, Liu P, et al. DSPP mutation in dentinogenesis
860	imperfecta Shields type II. Nat Genet. 2001 Feb;27(2):151–2.
861	60. de La Dure-Molla M, Philippe Fournier B, Berdal A. Isolated dentinogenesis imperfecta and
862	dentin dysplasia: Revision of the classification. Eur J Hum Genet. 2015 Apr;23(4):445–51.
863	61. Shields ED, Bixler D, El-Kafrawy AM. A proposed classification for heritable human
864	dentine defects with a description of a new entity. Arch Oral Biol. 1973 Apr 1;18(4):543-
865	IN7.
866	62. Sreenath T, Thyagarajan T, Hall B, Longenecker G, D'Souza R, Hong S, et al. Dentin
867	sialophosphoprotein knockout mouse teeth display widened predentin zone and develop
867 868	sialophosphoprotein knockout mouse teeth display widened predentin zone and develop defective dentin mineralization similar to human dentinogenesis imperfecta type III. J Biol
868	defective dentin mineralization similar to human dentinogenesis imperfecta type III. J Biol
868 869	defective dentin mineralization similar to human dentinogenesis imperfecta type III. J Biol Chem. 2003 Jul 4;278(27):24874–80.
868 869 870	 defective dentin mineralization similar to human dentinogenesis imperfecta type III. J Biol Chem. 2003 Jul 4;278(27):24874–80. 63. Verdelis K, Ling Y, Sreenath T, Haruyama N, MacDougall M, van der Meulen MCH, et al.
868 869 870 871	 defective dentin mineralization similar to human dentinogenesis imperfecta type III. J Biol Chem. 2003 Jul 4;278(27):24874–80. 63. Verdelis K, Ling Y, Sreenath T, Haruyama N, MacDougall M, van der Meulen MCH, et al. DSPP effects on <i>in vivo</i> bone mineralization. Bone. 2008 Dec 1;43(6):983–90.
868 869 870 871 872	 defective dentin mineralization similar to human dentinogenesis imperfecta type III. J Biol Chem. 2003 Jul 4;278(27):24874–80. 63. Verdelis K, Ling Y, Sreenath T, Haruyama N, MacDougall M, van der Meulen MCH, et al. DSPP effects on <i>in vivo</i> bone mineralization. Bone. 2008 Dec 1;43(6):983–90. 64. Chen Y, Zhang Y, Ramachandran A, George A. DSPP is essential for normal development

070		1 .	, ,·	· · · ·	1 .1	1	• •	C (11 /
876	(DSPP)	cause domii	nant negative	e effects 1	n both	dentinoge	nesis ii	mperfecta a	and dentin

- dysplasia by entrapping normal DSPP. J Bone Miner Res. 2012;27(6):1309–21.
- 66. Smith BL, Preston GM, Spring FA, Anstee DJ, Agre P. Human red cell aquaporin CHIP. I.
- 879 Molecular characterization of ABH and Colton blood group antigens. J Clin Invest. 1994
- 880 Sep 1;94(3):1043–9.
- 67. Jordan IK, Mariño-Ramírez L, Koonin EV. Evolutionary significance of gene expression
 divergence. Gene. 2005 Jan 17;345(1):119–26.
- 68. Warnefors M, Kaessmann H. Evolution of the correlation between expression divergence
 and protein divergence in mammals. Genome Biol Evol. 2013;5(7):1324–35.
- 69. Jernvall J, Thesleff I. Tooth shape formation and tooth renewal: Evolving with the same
 signals. Development. 2012;139(19):3487–97.
- 70. Mitsiadis TA. Role of Islet1 in the patterning of murine dentition. Development.
 2003;130(18):4451–60.
- 71. Charles C, Pantalacci S, Peterkova R, Tafforeau P, Laudet V, Viriot L. Effect of *eda* loss of
 function on upper jugal tooth morphology. Anat Rec. 2009;292(2):299–308.
- 891 72. Zurowski C, Jamniczky H, Graf D, Theodor J. Deletion/loss of bone morphogenetic protein
- 892 7 changes tooth morphology and function in *Mus musculus*: Implications for dental
- evolution in mammals. R Soc Open Sci. 2018 Jan 3;5(1):170761.
- 894 73. Cardoso-Moreira M, Halbert J, Valloton D, Velten B, Chen C, Shao Y, et al. Gene
- 895 expression across mammalian organ development. Nature. 2019 Jul;571(7766):505–9.

	896	74. Finarelli JA	, Flynn JJ.	Ancestral	state reconstr	ruction of	f body	size	in the	Caniforn	nia
--	-----	------------------	-------------	-----------	----------------	------------	--------	------	--------	----------	-----

897 (Carnivora, Mammalia): The effects of incorporating data from the fossil record. Syst Biol.
898 2006;55(2):301–13.

899 75. Welker F, Collins MJ, Thomas JA, Wadsley M, Brace S, Cappellini E, et al. Ancient proteins
900 resolve the evolutionary history of Darwin's South American ungulates. Nature. 2015

- 901 Jun;522(7554):81–4.
- 902 76. Warinner C, Korzow Richter K, Collins MJ. Paleoproteomics. Chem Rev. 2022 Aug
 903 24;122(16):13401–46.
- 904 77. Zheng GXY, Lau BT, Schnall-Levin M, Jarosz M, Bell JM, Hindson CM, et al. Haplotyping

905 germline and cancer genomes with high-throughput linked-read sequencing. Nat
906 Biotechnol. 2016 Feb;34:303.

- 907 78. Marks P, Garcia S, Martinez A, Belhocine K. Resolving the full spectrum of human genome
 908 variation using linked-reads. 2017;
- 909 79. Weisenfeld NI, Kumar V, Shah P, Church DM, Jaffe DB. Direct determination of diploid

910 genome sequences. Genome Res. 2017;27(5):757–67.

- 911 80. Vurture GW, Sedlazeck FJ, Nattestad M, Underwood CJ, Fang H, Gurtowski J, et al.
- 912 GenomeScope: Fast reference-free genome profiling from short reads. Bioinformatics. 2017
- 913 Jul 15;33(14):2202–4.

914	81. Koren S, Walenz BP, Berlin K, Miller JR, Bergman NH, Phillippy AM. Canu: Scalable and
915	accurate long-read assembly via adaptive k-mer weighting and repeat separation. Genome
916	Res. 2017 May 1;27(5):722–36.
917	82. Warren RL. RAILS and Cobbler: Scaffolding and automated finishing of draft genomes
918	using long DNA sequences. J Open Source Softw. 2016 Nov 17;1(7):116.
919	83. Gurevich A, Saveliev V, Vyahhi N, Tesler G. QUAST: Quality assessment tool for genome
920	assemblies. Bioinformatics. 2013 Apr 15;29(8):1072-5.
921	84. Simão FA, Waterhouse RM, Ioannidis P, Kriventseva EV, Zdobnov EM. BUSCO: Assessing
922	genome assembly and annotation completeness with single-copy orthologs. Bioinformatics.
923	2015 Oct 1;31(19):3210–2.
924	85. Cantarel BL, Korf I, Robb SMC, Parra G, Ross E, Moore B, et al. MAKER: An easy-to-use
925	annotation pipeline designed for emerging model organism genomes. Genome Res.
926	2008;18:188–96.
927	86. Campbell MS, Law M, Holt C, Stein JC, Moghe GD, Hufnagel DE, et al. MAKER-P: A tool
928	kit for the rapid creation, management, and quality control of plant genome annotations.
929	Plant Physiol. 2014 Feb 1;164(2):513–24.
930	87. Campbell MS, Holt C, Moore B, Yandell M. Genome annotation and curation using

931 MAKER and MAKER-P. Curr Protoc Bioinforma. 2014 Dec 12;48:4.11.1-4.11.39.

022	00 $C = 1.1 \dots M$	C II DI	VN	I	T1	A A A . 1	T
932	88. Grabherr M	О. пааѕ БЈ.	I assour IVI.	Levin JZ.	1 nondson D_A	A. Amit I. et al.	I finity:

- 933 Reconstructing a full-length transcriptome without a genome from RNA-Seq data. Nat
- 934 Biotechnol. 2011 May 15;29(7):644–52.
- 89. Wheeler TJ, Clements J, Eddy SR, Hubley R, Jones TA, Jurka J, et al. Dfam: A database of
- 936 repetitive DNA based on profile hidden Markov models. Nucleic Acids Res. 2013
- 937 Jan;41(Database issue):D70-82.
- 938 90. Caballero J, Smit AFA, Hood L, Glusman G. Realistic artificial DNA sequences as negative
- 939 controls for computational genomics. Nucleic Acids Res. 2014 Jul;42(12):e99.
- 940 91. Hubley R, Finn RD, Clements J, Eddy SR, Jones TA, Bao W, et al. The Dfam database of

941 repetitive DNA families. Nucleic Acids Res. 2016 Jan 4;44(D1):D81–9.

- 942 92. Hu J, Zheng Y, Shang X. MiteFinder: A fast approach to identify miniature inverted-repeat
- 943 transposable elements on a genome-wide scale. In: 2017 IEEE International Conference on
- Bioinformatics and Biomedicine (BIBM). 2017. p. 164–8.
- 945 93. Gremme G, Steinbiss S, Kurtz S. GenomeTools: A comprehensive software library for
- 946 efficient processing of structured genome annotations. IEEE/ACM Trans Comput Biol
- 947 Bioinform. 2013 May 1;10(03):645–56.
- 948 94. Smit A, Hubley R. RepeatModeler Open-1.0. 2008.
- 949 95. Keller O, Kollmar M, Stanke M, Waack S. A novel hybrid gene prediction method
- 950 employing protein multiple sequence alignments. Bioinformatics. 2011 Mar 15;27(6):757–
- 951 63.

- 952 96. Korf I. Gene finding in novel genomes. BMC Bioinformatics. 2004 May 14;5(1):59.
- 953 97. Campbell MS. compare annotations 3.2.pl [Internet]. 2015. Available from:
- 954 https://github.com/mscampbell/Genome_annotation/blob/master/compare_annotations_3.2.
- 955 pl
- 956 98. Eilbeck K, Moore B, Holt C, Yandell M. Quantitative measures for the management and

957 comparison of annotated genomes. BMC Bioinformatics. 2009 Feb 23;10(1):67.

- 958 99. Liu D, Hunt M, Tsai IJ. Inferring synteny between genome assemblies: A systematic
- evaluation. BMC Bioinformatics. 2018 Jan;19(1):26.
- 100. Emms DM, Kelly S. OrthoFinder: Solving fundamental biases in whole genome
 comparisons dramatically improves orthogroup inference accuracy. Genome Biol. 2015
 Aug 6;16(1):157.
- 101. Katoh K, Misawa K, Kuma K ichi, Miyata T. MAFFT: A novel method for rapid multiple
 sequence alignment based on fast Fourier transform. Nucleic Acids Res. 2002
 Jul;30(14):3059–66.
- 102. Lefort V, Desper R, Gascuel O. FastME 2.0: A comprehensive, accurate, and fast distancebased phylogeny inference program. Mol Biol Evol. 2015 Oct 1;32(10):2798–800.
- 968 103. Farrer RA. Synima: A synteny imaging tool for annotated genome assemblies. BMC
- 969 Bioinformatics. 2017 Nov 21;18(1):507.

970	104.	Wang Y, Tang H, DeBarry JD, Tan X, Li J, Wang X, et al. MCScanX: A toolkit for
971		detection and evolutionary analysis of gene synteny and collinearity. Nucleic Acids Res.
972		2012 Apr;40(7):e49.
973	105.	Zhao T, Schranz ME. Network approaches for plant phylogenomic synteny analysis. Curr
974		Opin Plant Biol. 2017 Apr 1;36:129–34.
975	106.	Zhao T, Holmer R, de Bruijn S, Angenent GC, van den Burg HA, Schranz ME.
976		Phylogenomic synteny network analysis of MADS-Box transcription factor genes reveals
977		lineage-specific transpositions, ancient tandem duplications, and deep positional
978		conservation. Plant Cell. 2017 Jun 1;29(6):1278-92.
979	107.	Zhao T, Schranz ME. Network-based microsynteny analysis identifies major differences
980		and genomic outliers in mammalian and angiosperm genomes. Proc Natl Acad Sci. 2019
981		Feb 5;116(6):2165–74.
982	108.	Sievers F, Higgins DG. Clustal Omega. Curr Protoc Bioinforma. 2014;48(1):3.13.1-
983		3.13.16.
984	109.	Suyama M, Torrents D, Bork P. PAL2NAL: Robust conversion of protein sequence
985		alignments into the corresponding codon alignments. Nucleic Acids Res. 2006 Jul
986		1;34(suppl_2):W609–12.
987	110.	Wong WSW, Yang Z, Goldman N, Nielsen R. Accuracy and power of statistical methods
988		for detecting adaptive evolution in protein coding sequences and for identifying positively
989		selected sites. Genetics. 2004 Oct 1;168(2):1041-51.

990	111. Löytynoja A, Vilella AJ, Goldman N. Accurate extension of multiple sequence alignments
991	using a phylogeny-aware graph algorithm. Bioinformatics. 2012 Jul 1;28(13):1684–91.

- 992 112. Jones DT. Protein secondary structure prediction based on position-specific scoring
- 993 matrices. J Mol Biol. 1999 Sep 17;292(2):195–202.

- 994 113. Buchan DWA, Jones DT. The PSIPRED protein analysis workbench: 20 years on. Nucleic 995 Acids Res. 2019 Jul 2;47(W1):W402-7.
- 996 114. Ewels PA, Peltzer A, Fillinger S, Patel H, Alneberg J, Wilm A, et al. The nf-core
- 997 framework for community-curated bioinformatics pipelines. Nat Biotechnol. 2020

998 Mar:38(3):276-8.

999 115. Andrews S. FastQC: A quality control tool for high throughput sequence data. [Internet].

1000 2010. Available from: http://www.bioinformatics.babraham.ac.uk/projects/fastqc

- 1001 116. Martin M. Cutadapt removes adapter sequences from high-throughput sequencing reads.
- 1002 EMBnet.journal. 2011 May 2;17(1):10-2.
- 1003 117. Kopylova E, Noé L, Touzet H. SortMeRNA: Fast and accurate filtering of ribosomal RNAs 1004 in metatranscriptomic data. Bioinformatics. 2012 Dec 1;28(24):3211–7.
- 1005 118. Patro R, Duggal G, Love MI, Irizarry RA, Kingsford C. Salmon provides fast and bias-
- 1006 aware quantification of transcript expression. Nat Methods. 2017 Apr;14(4):417-9.
- 1007 119. Love MI, Huber W, Anders S. Moderated estimation of fold change and dispersion for 1008 RNA-seq data with DESeq2. Genome Biol. 2014 Dec 5;15(12):550.

1009	120. Lindblad-Toh K, Garber M, Zuk O, Lin MF, Parker BJ, Washietl S, et al. A high-resolution
1010	map of human evolutionary constraint using 29 mammals. Nature. 2011
1011	Oct;478(7370):476–82.
1012	121. Weyrich A, Schüllermann T, Heeger F, Jeschek M, Mazzoni CJ, Chen W, et al. Whole
1013	genome sequencing and methylome analysis of the wild guinea pig. BMC Genomics. 2014
1014	Nov 28;15(1):1036.
1015	122. Gossmann TI, Ralser M. Marmota marmota. Trends Genet. 2020 May;36(5):383-4.
1016	123. Di Palma F, Alföldi J, Johnson J, Berlin A, Gnerre S, Jaffe D, et al. The draft genome of
1017	Microtus ochrogaster. Broad Inst [Internet]. 2012; Available from:
1018	https://www.ncbi.nlm.nih.gov/bioproject/72443
1019	124. Mouse Genome Sequencing Consortium, Waterston RH, Lindblad-Toh K, Birney E,
1020	Rogers J, Abril JF, et al. Initial sequencing and comparative analysis of the mouse genome.
1021	Nature. 2002 Dec 5;420(6915):520-62.
1022	125. Di Palma F, Alföldi J, Johnson J, Berlin A, Gnerre S, Jaffe D, et al. The draft genome of
1023	Jaculus jaculus. Broad Inst [Internet]. 2012; Available from:
1024	https://www.ncbi.nlm.nih.gov/bioproject/72445
1025	126. Gibbs RA, Weinstock GM, Metzker ML, Muzny DM, Sodergren EJ, Scherer S, et al.
1026	Genome sequence of the Brown Norway rat yields insights into mammalian evolution.
1027	Nature. 2004 Apr;428(6982):493–521.

1028	127. Kolmogorov M, Armstrong J, Raney BJ, Streeter I, Dunn M, Yang F, et al. Chromosome
1029	assembly of large and complex genomes using multiple references. Genome Res. 2018 Nov
1030	1;28(11):1720–32.

- 1031 128. Lilue J, Doran AG, Fiddes IT, Abrudan M, Armstrong J, Bennett R, et al. Sixteen diverse
- 1032 laboratory mouse reference genomes define strain-specific haplotypes and novel functional
- 1033 loci. Nat Genet. 2018 Nov;50(11):1574–83.
- 1034 129. Couger MB, Arévalo L, Campbell P. A high quality genome for *Mus spicilegus*, a close
- 1035 relative of house mice with unique social and ecological adaptations. G3
- 1036 GenesGenomesGenetics. 2018 May 24;8(7):2145–52.
- 1037 130. Chinese hamster CHOK1GS assembly and gene annotation. Horiz Eagle [Internet]. 2017;
- 1038 Available from: https://www.ensembl.org/Cricetulus_griseus_chok1gshd/Info/Annotation
- 1039 131. Di Palma F, Alföldi J, Johnson J, Berlin A, Gnerre S, Jaffe D, et al. The draft genome of
- 1040 *Mesocricetus auratus*. Broad Inst [Internet]. 2012; Available from:
- 1041 https://www.ncbi.nlm.nih.gov/bioproject/77669
- 1042 132. Lassance JM, Hopi Hoekstra. Improved assembly of the deer mouse *Peromyscus*
- 1043 *maniculatus* genome. Harv Univ Hughes Med Inst [Internet]. 2018; Available from:
- 1044 https://www.ncbi.nlm.nih.gov/bioproject/494228
- 1045 133. Fang X, Nevo E, Han L, Levanon EY, Zhao J, Avivi A, et al. Genome-wide adaptive
- 1046 complexes to underground stresses in blind mole rats *Spalax*. Nat Commun. 2014 Jun
- 1047 3;5(1):3966.

- 1048 134. Di Palma F, Alföldi J, Johnson J, Berlin A, Gnerre S, Jaffe D, et al. The draft genome of
- 1049 *Octodon degu*. Broad Inst [Internet]. 2012; Available from:
- 1050 https://www.ncbi.nlm.nih.gov/bioproject/74595
- 1051 135. Keane M, Craig T, Alföldi J, Berlin AM, Johnson J, Seluanov A, et al. The naked mole rat
- 1052 genome resource: Facilitating analyses of cancer and longevity-related adaptations.
- 1053 Bioinforma Oxf Engl. 2014 Dec 15;30(24):3558–60.
- 1054 136. Di Palma F, Alföldi J, Johnson J, Berlin A, Gnerre S, Jaffe D, et al. The draft genome of
- 1055 *Chinchilla lanigera*. Broad Inst [Internet]. 2012; Available from:
- 1056 https://www.ncbi.nlm.nih.gov/bioproject/68239
- 1057 137. V. Federov, Dalen L, Olsen RA, Goropashnaya AV, Barnes BM. The genome of the Arctic
- 1058 ground squirrel *Urocitellus parryii*. Inst Arct Biol [Internet]. 2018; Available from:
- 1059 https://www.ncbi.nlm.nih.gov/bioproject/477386
- 1060 138. Di Palma F, Alföldi J, Johnson J, Berlin A, Gnerre S, Jaffe D, et al. The draft genome of
- 1061 *Ictidomys tridecemlineatus*. Broad Inst [Internet]. 2012; Available from:
- 1062 https://www.ncbi.nlm.nih.gov/bioproject/61725
- 1063 139. Schneider VA, Graves-Lindsay T, Howe K, Bouk N, Chen HC, Kitts PA, et al. Evaluation
- 1064 of GRCh38 and de novo haploid genome assemblies demonstrates the enduring quality of
- 1065 the reference assembly. Genome Res. 2017 May 1;27(5):849–64.

1066