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1 **Els Casots (Subirats, Catalonia), a key site for the Miocene vertebrate**  
2 **record of Southwestern Europe**

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87 **ABSTRACT**

88 Els Casots is one of the richest fossil vertebrate sites of the Vallès-Penedès Basin (Catalonia,  
89 Spain). It was discovered in 1989 and excavated briefly during the 1990s, resulting in the  
90 recovery of thousands of remains and the erection of several new mammal species.  
91 Excavations resumed in 2018 and continue to date. Here we provide updated results regarding  
92 the age, stratigraphy, biota and palaeoenvironment of the site. The age of the site is well  
93 constrained to ~15.9 Ma thanks to recent bio- and magnetostratigraphic data, thus coinciding  
94 with the onset of the Miocene Climatic Optimum (MCO). The stratigraphic succession at the  
95 site area indicates lacustrine to palustrine environments with cyclically oscillating water level.  
96 There are several fossiliferous layers that have yielded a vertebrate fauna comprising up to 74  
97 different vertebrate species including amphibians, reptiles, birds and mostly mammals. The  
98 finding of several articulated partial skeletons indicate that the site records an autochthonous  
99 to parautochthonous assemblage. The abundance and completeness of the vertebrate remains  
100 together with a well-constrained age and detailed stratigraphic and palaeoenvironmental data,  
101 make els Casots a key site for understanding wetland ecosystems in southern Europe during  
102 the MCO.

103

104 **KEYWORDS:** Miocene Climatic Optimum, fossil vertebrates, taphonomy,  
105 palaeoenvironment, Vallès-Penedès Basin, Spain

106

## 107 **Introduction and historical overview**

108 The Early to Middle Miocene transition was a crucial period marked by important climatic  
109 and faunal changes. The former notably comprise the Miocene Climatic Optimum (MCO), the  
110 warmest interval since the late Eocene, which peaked between ~17–15 Ma (Zachos et al.  
111 2001). This interrupted the long-term Cenozoic cooling trend and Antarctic ice-sheet build up  
112 (Shevenell et al. 2008; Foster et al. 2012), resulting in estimated mid-latitude atmospheric  
113 temperatures 5–7° C warmer than today (Steinthorsdottir et al. 2021). The MCO coincided  
114 with a series of faunal dispersals, such as those of modern cricetid rodents or bovids which  
115 —contrary to earlier Miocene dispersals (e.g., those between Africa and Eurasia)— were not  
116 facilitated by new continental connections. During the Miocene these newcomers would  
117 replace older groups of Palaeogene origin and ‘modernise’ the mammal faunas. This time  
118 interval is well represented in several Spanish basins, such as the Calatayud-Montalbán Basin  
119 (Aragon; Van der Meulen et al. 2012; García-Paredes et al. 2016), the Magro Basin  
120 (Valencia; Ruiz Sánchez et al. 2003), the Ribesalbes-Alcora Basin (Valencia, Crespo et al.  
121 2019) or the Bardenas Reales (Navarra) in the Ebro Basin (Suárez-Hernando 2017). However,  
122 their record mostly consists of microvertebrates, larger mammals being generally scarce. By  
123 contrast, the Vallès-Penedès Basin (Catalonia) presents a rich record for the Early/Middle  
124 Miocene transition that includes several sites that have yielded both large and small  
125 vertebrates. Amongst these, els Casots stands out as one of the richest sites of the whole  
126 basin, including thousands of specimens, often exceptionally complete.

127 Els Casots (note that the correct toponym according to the Cartographical and  
128 Geological Institute of Catalonia is ‘els Casots’, with ‘els’ not capitalized) pertains to the  
129 municipality of Subirats (l’Alt Penedès), less than 40 km away from Barcelona. During the  
130 late 19<sup>th</sup> century, several short-lived lignite mines were active near els Casots and eventually  
131 the workers found a few fossil bones and teeth that were handled to the priest and geologist

132 Jaume Almera, from the Geological Museum of the Barcelona Seminary. These stand out  
133 amongst the earliest finds of fossil vertebrates in Catalonia. Almera entrusted their  
134 identification to well-known vertebrate palaeontologists of his time, such as Charles Depéret  
135 and Albert Gaudry, who ascribed them to rhinoceroses, ruminants and rodents (Almera 1898).  
136 When the mines closed no further findings were reported and eventually those given to  
137 Almera were lost or destroyed in 1936, at the beginning of the Spanish Civil War. Later, the  
138 renowned palaeontologist Miquel Crusafont, founder of the Institut de Paleontologia de  
139 Sabadell (IPS, predecessor of the Institut Català de Paleontologia Miquel Crusafont, ICP), and  
140 his collaborators surveyed the mine surroundings and only found some fossil plants but not  
141 vertebrates (Crusafont et al. 1955). Finally, in 1989 Toni Adell, an amateur fossil hunter  
142 found several bones on the surface of a vineyard close to the ancient mines (Figure 1a) and  
143 these were reported to S.M.S from the IPS (Moyà-Solà and Rius Font 1993). Systematic  
144 excavations followed between 1989 and 1993, resulting in the recovery of more than 3,000  
145 macrovertebrate remains from different fossiliferous levels (Moyà-Solà and Rius Font 1993).  
146 In addition, samples from some levels were screen-washed and yielded hundreds of  
147 identifiable micromammal teeth. All the recovered fossils were deposited at the ICP and most  
148 of them have been prepared. In 1995, the site was acquired by the municipality of Subirats  
149 and protected as cultural heritage by the Catalan government. However, excavations were  
150 interrupted in 1994, mostly because the discovery or re-excavation of major sites in the same  
151 area (e.g., Can Llobateres) required the limited funding and personnel available.

152           Nevertheless, the study of the fauna continued during the 1990s, mostly focusing on  
153 artiodactyls. New species such as the listriodontine suid *Eurolistriodon adelli* (Pickford and  
154 Moyà-Solà, 1995) or the palaeomerycid *Ampelomeryx ginsburgi* Duranthon et al., 1995 were  
155 erected based on material from els Casots. In the 21<sup>st</sup> century taxonomic studies expanded to  
156 consider crocodylians, rodents and small carnivorans, amongst others (see below). Yet, there

157 are relatively few works devoted to the biota from els Casots and none summarising the  
158 characteristics and significance of the site. In 2018, after more than 20 years of inactivity, the  
159 ICP resumed fieldwork at els Casots. Yearly campaigns are currently conducted, and these  
160 include not only systematic excavation, but also geological survey and sampling to provide  
161 further insights on the age and palaeoenvironment of the site. Furthermore, several actions,  
162 such as building permanent structures to shelter main excavation areas (Figure 1b), have been  
163 undertaken by the Subirats municipality and the site is open for touristic visits upon request.  
164 Here we present an overview of the els Casots site, its geology, age, biota and  
165 palaeoenvironment based on our latest data.

166

## 167 **Geological setting**

168 Els Casots is located at the southern margin of the Vallès-Penedès Basin (VPB), an elongated  
169 half-graben sub-parallel to the Catalan coastline and bounded by the Catalan Coastal Ranges  
170 (Figure 2a), that was formed during the opening of the Western Mediterranean by the late  
171 Oligocene (Roca et al. 1999; Cabrera et al. 2004). The VPB sedimentary record covers most  
172 of the Miocene (from about 20 to 7 Ma) and is mainly continental except for relatively brief  
173 marine episodes that occurred during the late Burdigalian (Early Miocene), Langhian and  
174 early Serravallian (Middle Miocene), and Tortonian (Late Miocene; Casanovas-Vilar et al.  
175 2016). Early to early Middle Miocene continental alluvial deposits interdigitate and are finally  
176 overlain by shallow to deep marine (bay, shelf) and transitional deposits (fan deltas and minor  
177 deltas, shoreline systems and saline lagoons) ranging in age from late Burdigalian to  
178 Langhian. The continental, transitional and marine facies are stacked in transgressive–  
179 regressive sequences that attained their maximum spreading and depth in the basin during the  
180 Langhian sea-level high stand.



181 In general, the Early to early Middle Miocene (Ramblian to early Aragonian mammal  
182 ages) terrestrial vertebrate record of the VPB is comparatively poorer than later intervals (late  
183 Aragonian and Vallesian mammal ages; Casanovas-Vilar et al. 2016). This interval is  
184 recorded by the sequences of the Lower Continental Units, a lithostratigraphical unit that  
185 crops out near the southern basin margins and at the Llobregat and Anoia river valleys. This  
186 assemblage includes intensely red alluvial fan facies sourced from the surrounding reliefs and  
187 minor cyclically arranged alluvial-shallow lacustrine units of various ages (Cabrera Pérez  
188 1981a; Cabrera et al. 1991; de Gibert and Casanovas-Vilar 2011; Casanovas-Vilar et al. 2011,  
189 2016). The younger alluvial-lacustrine assemblage (Subirats alluvial-lacustrine unit; Agustí  
190 and Cabrera 1980; Cabrera Pérez 1981a, 1981b) includes the site of els Casots as well as  
191 other rich fossil sites such as les Cases de la Valenciana (Crusafont et al. 1955; Jovells-Vaqué  
192 et al. 2018), Can Julià and Can Martí Vell (Crusafont et al. 1955; Agustí and Cabrera 1980;  
193 Agustí 1983). This unit unconformably overlays directly the Mesozoic basement (Cabrera  
194 Pérez 1979, 1981a, 1981b) and extends a few kilometres along the present southern basin  
195 margin between the municipalities of Gelida and Subirats (Figure 2). Els Casots is located in a  
196 continental to transitional sequence developed in an extensional faulted zone that affected  
197 both the Mesozoic basement and the Miocene cover.

198

## 199 **Structure, stratigraphy and age of the succession**

### 200 *Structure and arrangement of the basement*

201 The pre-Miocene basement at the area of els Casots consists of Mesozoic, mainly Early  
202 Cretaceous, marine carbonates (Figure 2b). To determine the basement arrangement,  
203 geophysical prospection using ERT (Electrical Resistivity Tomography) methods was  
204 conducted at the site area to infer the thickness of Miocene sediments and preliminarily study

205 the basement morphology. ERT data were collected along two intersecting sections (Figure  
206 2b), and they encompassed three ERT profiles. Geoelectrical results are presented along two  
207 sections built from the 2D models obtained from the ERT profiles (Figure 3). Geoelectrical  
208 section 1 (Figure 3a) shows a preferential stratified behaviour with an alternate of conductive  
209 and resistive layers gently dipping to the centre of the section. It also presents three segments  
210 (A, B, and C) where this stratification is not complete. A similar behaviour is recognized in  
211 section 2 (Figure 3b), with a horizontal stratification and an abrupt change in segment D.

212 The interpretation of section 1 is based on the information provided by two core  
213 samples (CS-A and CS-B, see following section and Figure 4) taken next to the section (see  
214 Figure 2b). The Cretaceous basement is identified as the bottom resistive layer (Figure 3, in  
215 blue), while the different conductive and resistive layers in the Miocene cover can be  
216 associated with the clayey and carbonate sequences described in the cores. Regions A, B, and  
217 C (Figure 3a) may be interpreted as three fractured zones affected by minor normal faults  
218 dipping to the N. In Section 2 (Figure 3b), basement identification is not straightforward  
219 because this section runs along the fractured zone B. However, this section does not present  
220 E-W changes in resistivity except in region D, which coincides with the location of the  
221 inferred NE-SW fault (Figure 2b).

222 Attending to the geological and geophysical data, the pre-Miocene basement at els  
223 Casots area consists of Mesozoic (mainly Early Cretaceous) marine carbonates, which make  
224 up the present surrounding reliefs. During the Early to Middle Miocene these rocks  
225 constituted a weathered, eroded and karstified palaeorelief later affected by NE-SW and  
226 NNE-SSW faults related to extensional processes in the area.

227

228 ***Miocene stratigraphy***

229 The overall stratigraphy of the Subirats alluvial-lacustrine unit at els Casots has been  
230 previously described (Cabrera Pérez 1979, 1981a, 1981b). The lithological succession in the  
231 area nearby the fossil site was further studied by means of geological core sampling next to  
232 the excavation area (core CS-A; Figure 2b) and at the point of maximum thickness of  
233 Miocene sediments as inferred from ERT results (core CS-B; Figure 2b).

234 Core CS-B (Figure 4) reached the Mesozoic basement at 36 m of depth. A 4 m-thick  
235 deposit of clast- to matrix-supported scree breccias overlay the karstified Cretaceous  
236 basement. The often large (up to several cm in diameter) clasts are made of Cretaceous  
237 carbonates and show little rounding or are clearly subangular to angular, thus indicating that  
238 they derived from in situ substrate fragmentation. The matrix consists of mudstone that grades  
239 upwards from ochre/reddish to greenish grey due to reductive conditions associated to the  
240 lacustrine flooding. Grey mudstone including outsized limestone granules and clasts overlay  
241 the clast dominated lowermost deposits. Similar deposits overlay the basement in the  
242 surrounding outcrops (Figure 2b) and include plant, gastropod and vertebrate fossils, badly  
243 broken macrovertebrate remains and abundant micromammals (Cal Sutxet site; Jovells-Vaqué  
244 and Casanovas-Vilar 2021). The lowermost breccia unit is of variable thickness because it  
245 overlaid the pre-existing palaeorelief. This breccia unit is overlain by a 25 m-thick, cyclically-  
246 arranged mudstone/limestone succession that at the bottom includes thin (mm to dm thick)  
247 sub-bituminous coal deposits. Cycles often include a lower dark, organic matter rich term that  
248 grades upwards to grey-bluish massive mudstones with minor, scattered millimetric siderite-  
249 like ferruginous nodules. The upper part of the cycles is a carbonate term that includes a  
250 variety of bioclastic- to micrite-dominated limestones (grainstones, wackestones and  
251 mudstones) that range from a few centimetres to a maximum of a couple of meters thick  
252 (Figure 4). The carbonate facies include bioclastic laminae made up of accumulations of  
253 ostracods, charophytes and micrite-coated algal filaments. Very thin tufa-like carbonates also

254 occur (Cabrera Pérez 1979). Thin ferruginous laminae are associated to some of the bioclastic  
255 facies. Both carbonates and siliciclastic mudstones may include abundant plant fragments and  
256 freshwater gastropod shells or casts. This lacustrine/marshy lacustrine succession has yielded  
257 abundant vertebrate fossils in the excavated area (Figure 5).

258         The lacustrine succession ends rather abruptly and in core CS-B grades along the  
259 uppermost 5 m into intensely reddish and mottled silts and sandstones topped by carbonate-  
260 rich pedogenic to palustrine facies (Figure 4). These palustrine facies crop out W of Cal  
261 Sutxet farmhouse (Figure 2b) and are rich in internal moulds of terrestrial gastropods but are  
262 devoid of vertebrate fossils. They are overlain by red mudstones and green/grey sandstones  
263 that record subaerial distal channelized alluvial fan facies. The continental deposits are in turn  
264 overlain by an oyster coquina and a bioclastic quartzarenite, which marks the early Langhian  
265 transgressive surface in this sector. The Langhian marine transgression and high-stand marine  
266 facies in the area record successive episodes of deposition of bioturbated sandstones overlain  
267 by grey marine bay-shelf mudstones and marls and some episodes of minor reef construction  
268 (Bessedik and Cabrera 1985; Agustí et al. 1990; Domènech et al. 2011; Casanovas-Vilar et al.  
269 2016).

270

## 271 ***Age***

272 Els Casots has been generally correlated to European Neogene Mammal (MN) unit MN4,  
273 which corresponds to the early Aragonian, latest Early Miocene (e.g., Agustí et al. 1990;  
274 Agustí and Llenas 1993; Duranthon et al. 1995; Pickford and Moyà-Solà 1995; Casanovas-  
275 Vilar et al. 2011a, 2011b, 2016; Jovells-Vaqué et al. 2017a). Such correlation was mostly  
276 based on the rodent fauna, which includes the early modern cricetids *Megacricetodon*  
277 *primitivus* and *Democricetodon hispanicus* together with the eomyid *Ligerimys ellipticus*.

278 However, recent studies indicate that els Casots should indeed be correlated to MN5 (Jovells-  
279 Vaqué and Casanovas-Vilar 2021). The MN4/MN5 boundary was previously defined by the  
280 extinction of the eomyid genus *Ligerimys* (e.g., Mein 1999; Kälin and Kempf 2009), which  
281 turned out to be a diachronic event in Western Europe, with the last Iberian species, *L.*  
282 *ellipticus*, persisting for about 0.5 million years after the extinction of the last central  
283 European species, *Ligerimys florancei* (Van der Meulen et al. 2012). However, the extinction  
284 of *L. florancei* is synchronous in both areas (~16.4 Ma), so this has been proposed as the  
285 diagnostic criterion for the definition of the MN4/MN5 boundary (Hilgen et al. 2012; Van der  
286 Meulen et al. 2012). At els Casots, only *L. ellipticus* is present (and in low numbers),  
287 suggesting a correlation to early Aragonian subzone Cb (16.15–15.94 Ma; see Van der  
288 Meulen et al. 2012), corresponding to the earliest MN5, the beginning of the Middle Miocene.

289 Cores CS-A and CS-B were sampled for palaeomagnetism at an average interval of  
290 0.35 cm, collecting 57 samples from core CS-A and 78 samples from core CS-B, and  
291 additional oriented samples were taken from the fossil site on surface and the overlying, non-  
292 lacustrine sediments, up to the Langhian oyster coquina (Figure 2). Bedding dip, when  
293 observable in the cores, was taken as a reference for the orientation of palaeomagnetic  
294 samples. Both stepwise alternating field (AF) and thermal (TH) demagnetization was applied  
295 to all samples, providing comparable results. A viscous component was demagnetized at  
296 temperatures below 200° C and peak AF of 10 mT. Above this range, a stable characteristic  
297 remanent magnetization (ChRM) could be determined up to temperatures of 450° C and peak  
298 AF of 60 mT. The ChRM was ranked as Quality 1 when a linear decay was observed until  
299 complete demagnetization. As the samples from cores lacked azimuthal orientation, the  
300 polarity determination relied only on the inclination of the ChRM, positive (downwards)  
301 inclination representing normal polarity and vice versa. For caution, ChRM components  
302 yielding low inclinations (reference palaeomagnetic inclination at the site is 60°) were not

303 considered as reliable and were ranked as Quality 2. A complementary quality check applied  
304 was the coherency of the angular relationship between the viscous and the ChRM  
305 components. Since the viscous component mostly conform to the north-directed present-day  
306 field, a reversed polarity ChRM should be roughly antiparallel to the viscous component,  
307 while a normal polarity ChRM should tend to align with the viscous component. ChRM  
308 directions meeting this condition were included in Quality 1.

309 Best quality results were obtained from core CS-B, with 66 samples (88%) ranked as  
310 Quality 1, compared to 25 (44%) samples of Quality 1 from CS-A. The whole series shows  
311 reverse polarity except for a normal interval determined at a depth between 12–15 m in CS-A,  
312 but not confirmed in CS-B, which yielded a palaeomagnetic signal of higher quality.  
313 Therefore, we consider this interval of normal magnetization from CS-A possibly resulted  
314 from a secondary magnetization process, unrelated with the time of deposition. Two  
315 additional normal magnetization intervals in CS-B are very thin, marked only by one sample,  
316 and were not considered relevant for correlation purposes. As already exposed, the rodent  
317 assemblage indicates a correlation to early Aragonian subzone Cb, which in the Calatayud-  
318 Montalbán Basin spans the whole normal polarity geomagnetic chron C5Cn.1n and the  
319 earliest part of the inverse polarity chron C5Br (Van der Meulen et al. 2012). Considering  
320 these data, the dominantly reverse polarity interval recorded in the cores is best correlated to  
321 chron C5Br (15.974–15.160 Ma; boundaries after Ogg 2020). Coupled with the  
322 aforementioned correlation to Aragonian subzone Cb (16.15 – 15.94 Ma; Van der Meulen et  
323 al. 2012), this results in an estimated age of ~15.9 Ma for the site of els Casots.

324

325 **Excavated area and taphonomic remarks**

326 The fossiliferous area of els Casots is huge, comprising all the fields around the farmhouses of  
327 Cal Solà and Cal Sutxet, but only a small area of about 6,200 m<sup>2</sup> is protected (Figures 1, 2b).  
328 For the moment being, a small surface of just 400 m<sup>2</sup> has been excavated, most of it during  
329 the 1990s field campaigns. Detailed stratigraphic information from the different excavated  
330 sectors (or ‘spots’) was recorded during the 1990s field campaigns and up to 13 stratigraphic  
331 levels (labelled A to M) were distinguished from surface to bottom (Figure 5). These  
332 represent a ~2 m thick succession consisting of mostly greyish to ochre mudstones, with  
333 occasional thin black mudstone layers. A thin layer of reddish carbonates (L) crops out at the  
334 base of the excavated sections. Mudstones are often laterally discontinuous and of variable  
335 thickness within the site area. They are rich in mollusc shell fragments and may include plant  
336 remains (mostly wood). Carbonate nodules are abundant in some levels.

337         Although all levels have yielded macrovertebrate fossils, level E turned out to be the  
338 most productive in the 1990s campaigns. Field campaigns since 2018 have focused on the  
339 excavation of level J, a bone bed that has yielded partially articulated specimens (Figures 6b–  
340 d, 7). Furthermore, well-preserved carnivoran coprolites and numerous wood fragments have  
341 been found at level J. Taphonomic information was recorded during both recent and older  
342 field campaigns. Macrovertebrate remains occur in high concentrations in all the excavated  
343 area laying subhorizontally, parallel to stratigraphy, and do not show signs of transport.  
344 Therefore, although detailed taphonomic analyses have yet to be conducted, we interpret that  
345 the biota from els Casots represents an autochthonous to parautochthonous assemblage.  
346 Lithostatic pressure has resulted in extensive crushing and deformation of many remains. For  
347 example, the holotype cranium of the suid *Eurolistriodon adelli* (IPS9096; Pickford and  
348 Moyà-Solà 1995: figs. 1–2) is so flattened dorsoventrally that it measures less than 5 cm in  
349 thickness. Almost all the fossils recovered during the 1990s have been prepared for study, but

350 only a few of the most relevant specimens (such as articulated skeletons and crania) found in  
351 the new excavations are prepared.

352         Microvertebrates were recovered during the 1990s field campaigns after screen-  
353 washing sediment samples from three different levels (CS-72, CS-73 and CS-74, certainly  
354 equivalent to excavation level D; Figure 5) and yielded hundreds of identifiable dental  
355 remains, mostly belonging to rodents (Agustí and Llenas 1993). During the new field  
356 campaigns large amounts of sediment have been screen-washed from layers K, J and M  
357 resulting in the recovery of thousands of microvertebrate remains, which include a diverse  
358 palaeoherpetological assemblage, some bird remains and numerous rodent, lagomorph and  
359 insectivoran cheek teeth. Even if only a small part of this collection has been prepared for  
360 study and is considered here, as shown in the following sections, the amount of new  
361 information is so relevant that els Casots can be already considered a key locality for the  
362 Miocene vertebrate record of Southwestern Europe.

363

## 364 **Biota and palaeoenvironment**

### 365 *Flora and invertebrates*

366 Crusafont et al. (1955) reported a single palm leaf found near the abandoned coal mines next to  
367 els Casots. Poorly preserved leaf fragments of helophytic plants, i.e., reeds and sedges (e.g.,  
368 *Typha*) and wood fragments have been recovered during the recent excavations. Sediment  
369 samples next to the excavation area have been screen-washed and yielded gyrogonites which  
370 are currently under study. The charophyte assemblage is dominated by *Chara* cf. *vulgaris* and  
371 *Chara molassica* var. *notata*. Extant *C. vulgaris* inhabits various kinds of freshwater  
372 environments, ranging from small eutrophic temporary ponds to mesotrophic lakes and rivers.  
373 Core samples from the site area (see above) were also analysed in search for palynomorphs, but



374 without success. Invertebrates are represented at the site by fragments of ostracod carapaces,  
375 mollusc shell fragments (such as freshwater gastropods) and, in the non-lacustrine upper layers,  
376 by moulds of terrestrial gastropods. Rare benthic foraminifera (i.e., *Ammonia* sp.) also occur in  
377 the assemblage and were probably transported from nearby marine settings.

378

### 379 *Non-mammalian vertebrates*

380 Highly fragmentary fish dental remains have been recovered by means of screen-washing but  
381 have neither been prepared nor studied. The herpetofauna is especially diverse, including up to  
382 24 different species (Table 1), and the available collection has increased significantly after the  
383 latest field campaigns. Amphibians include anurans and urodeles, the latter represented by just  
384 a single humerus. Anurans are more abundant, but most of the recovered remains do not allow  
385 for a confident taxonomic attribution. Based on ilial morphology, at least three different genera  
386 are present: the discoglossine alytid *Latonia* (already known from the roughly coeval site of  
387 Sant Mamet and other younger sites in the VPB; Villa et al. 2019a), the bufonid *Bufotes* (gr.  
388 *viridis*), and the ranid *Pelophylax*.

389 Chelonian fossils include scarce remains of the geoemydid *Ptychogaster* as well as a  
390 few disarticulated shell plates and some phalanges attributed to the medium-sized tortoise  
391 *Testudo*. The lack of diagnostic shell parts precludes an assignment to the subgenus rank for  
392 both taxa (Luján et al. 2014a, 2016). The most abundant and complete chelonian material,  
393 comprising both shell and postcranial remains, belongs to giant tortoises of the genus  
394 *Titanochelon*. The epiplastron resembles that of *Titanochelon richardi*, common in Middle to  
395 Late Miocene (MN6–MN10) sites of the VPB (Luján et al. 2014b). Squamates are by far the  
396 most diverse group in the palaeoherpetological assemblage. Gekkotans are represented by very  
397 few bones, while lacertids, whose identification is often difficult if not impossible based on

398 isolated and fragmentary bones, are represented by at least a non-amblyodont form together  
399 with a slightly-amblyodont one (cf. *Amblyolacerta* sp.). A scincoid is also present, based on the  
400 tooth morphology of few fragmentary dentigerous elements. Several remains of non-*Anguis*  
401 anguines were recovered, especially osteoderms and vertebrae. A single fragment of an  
402 indeterminate dentigerous bone with an *Ophisaurus*-like dentition as well as at least one trunk  
403 vertebra morphologically similar to those of *Pseudopus* has been recovered. A few vertebrae  
404 and isolated teeth possibly belong to varanid lizards. Two small amphisbaenian premaxillae can  
405 be referred to a member of the western clade of *Blanus* owing to the flat morphology of their  
406 anterior outline in lateral view (Bolet et al., 2014; Villa et al. 2019b). Snakes, in turn, are mainly  
407 identified based on vertebral morphology. The most abundant taxon is a pythonid, but an  
408 indeterminate scolecophidian, an erycine boid, at least one “colubrine” (sensu Szyndlar 1991),  
409 and possibly two viperids (a large and a smaller one) are present.

410 Small-sized crocodylians are extremely abundant at the site, being represented by  
411 countless isolated plates, osteoderms and teeth, as well as by complete skulls. During the 2021  
412 field campaign a complete articulated skeleton was unearthed and is currently under study.  
413 Cranial remains recovered during the 1990s field campaigns were described in detail by Díaz  
414 Aráez et al. (2017) and assigned to the diplocynodontid *Diplocynodon ratelii*. This represents  
415 the youngest record of the genus in the Iberian Peninsula.

416 Avian remains are rare, and most can be attributed to a medium-sized diurnal raptor  
417 showing some morphological similarities with the extant *Circaetus*. There is also evidence of  
418 a larger eagle (cf. *Aquila* sp.), an unknown nocturnal raptor (Strigidae indet.) of the size of an  
419 eagle owl, and the well-known peafowl *Miophasianus altus* and present in other sites of the  
420 VPB (Sánchez-Marco 2021).

421

422 *Small mammals*

423 Small mammals are incredibly abundant in all levels and are represented by isolated cheek  
424 teeth as well as mandibular and cranial material (Jovells-Vaqué et al. 2017b). However, only  
425 some rodent groups have been studied in detail. Els Casots records one of the latest  
426 occurrences in the Iberian Peninsula (see Furió et al. 2012) of the small marsupial  
427 *Amphiperatherium frequens*, represented by a few isolated teeth. Chiropterans are similarly  
428 rare, just a couple of molars of a small-sized taxon have been found to date. Most of the  
429 recovered eulipotyphlan remains belong to the small-sized erinaceid *Galerix symeonidisi*.  
430 Other insectivorans include the heterosoricid *Heterosorex neumayrianus*, represented by just  
431 one fissident upper incisor, a few teeth belonging to the tiny soricid *Paenelimnoecus* cf.  
432 *truyolsi*, and two unicuspid with rather strong cingula that may belong to the dimylid  
433 *Plesiodymylus*. The insectivoran assemblage is somewhat poorer than that of older sites  
434 (MN3–MN4) of the same basin, which usually record up to six different species (Van den  
435 Hoek Ostende et al. 2020). Lagomorphs are represented by the ochotonid *Lagopsis*, common  
436 in Early and Middle Miocene sites of the VPB (López Martínez 1989).

437         Rodents display a remarkable diversity of ground squirrels (Sciuridae) including both  
438 xerins (*Atlantoxerus idubedensis*, *Heteroxerus rubricati*) and marmotins (*Csakvaromys*  
439 *besana*), which were described by Aldana Carrasco (1991, 1992). Cricetids and glirids stand  
440 out as the most abundant rodents. The former, which were thoroughly described by Jovells-  
441 Vaqué et al. (2017a) and Jovells-Vaqué and Casanovas-Vilar (2021), include *Megacricetodon*  
442 *primitivus* and three *Democricetodon* species: *De. hispanicus*, which is the most abundant;  
443 *De. gracilis*, represented by a few small specimens; and a larger indeterminate species  
444 recorded by a single lower first molar. Interestingly, *De. hispanicus* persists for longer in the  
445 VPB than in the Calatayud-Montalbán Basin (the type area of the Aragonian; east-central  
446 Spain), where it is last recorded at 16.63 Ma (Van der Meulen et al. 2003, 2012).

447 *Democricetodon gracilis*, on the other hand, indicates faunal affinities with Central Europe,  
448 whereas it is absent in inner Iberian basins (Jovells-Vaqué and Casanovas-Vilar 2021). Glirids  
449 have only been preliminarily studied (Agustí and Llenas 1993), but up to nine different  
450 species have been identified. Many of these are characterised by their simplified dental  
451 pattern –such as *Peridyromys murinus*, *Pseudodryomys ibericus* and the three species of  
452 *Simplomys* (*S. simplicidens*, *S. robustus*, *S. julii*)– which is generally thought to reflect a  
453 preference for open and arid environments (de Bruijn and Van der Meulen 1982; Van der  
454 Meulen and Daams 1992; Van Dam 2006). *Simplomys simplicidens* and *De. hispanicus* stand  
455 out as the most abundant rodents. The remaining glirids comprise presumably arboreal  
456 species with more complex dental patterns (De Bruijn and Van der Meulen 1982; Van der  
457 Meulen and Daams 1992; Van Dam 2006), namely *Bransatoglis* sp., *Glirudinus modestus*,  
458 *Microdyromys monspeliensis* and *Muscardinus* sp., which are generally scarce. Finally,  
459 eomyids are only represented by less than ten teeth (over more than 700 identifiable rodent  
460 teeth) of *Ligerimys ellipticus*. This endemic Iberian species represents the last survivor of the  
461 genus *Ligerimys*, which disappeared from central Spain at 15.93 Ma (Van der Meulen et al.  
462 2012).

### 463 ***Large mammals***

464 Carnivorans are diverse and comprise multiple families of caniforms and feliforms. Amongst  
465 the former, amphicyonids include two undescribed species of different size, the larger one  
466 being represented by an almost complete skeleton (Figure 7a). Hemycyonids are represented  
467 by *Hemicyon stehlini*. Mustelids include the gulonin *Iberictis buloti*, the oldest member of the  
468 wolverine lineage, which is abundantly represented by dental and cranial material (Figure 6a;  
469 Valenciano et al. 2020). Other mustelids comprise *Paralutra* sp., represented by two  
470 mandibles, which was likely semiaquatic (Ginsburg 1999) as well as two distinct small-sized  
471 marten-like forms (“*Martes*” sp. 1 and 2). Feliforms include herpestids, hyaenids and felids.

472 Herpestids are represented by *Leptoplesictis aurelianesis*, for which only few, but remarkably  
473 complete, craniodental remains have been found at the site. Concerning hyaenids, only a  
474 single mandible tentatively attributed to *Protictitherium* has been recovered. Felids are better  
475 represented by abundant postcranial and dentognathic remains belonging to the lynx-sized  
476 “*Pseudaelurus*” *lorteti*. During the 2021 field campaign, a crushed complete skull was found  
477 and is currently under study.

478 Rhinocerotids stand out as one of the most abundant elements of the fauna, being  
479 recorded by *Dromoceratherium mirallesi*, which was originally described from the  
480 neighboring locality of Can Julià (Gelida, MN4) by Crusafont et al. (1955). This poorly  
481 known rhinoceros is generally included in *Plesiaceratherium* (Yan and Heissig 1986) but  
482 ongoing studies of the type material and additional remains from other VPB sites support the  
483 distinction of *Dromoceratherium* (Sanisidro et al. 2018). Further rhinocerotid remains include  
484 indeterminate juvenile mandibles (Figure 6c), isolated teeth, and postcranial elements pending  
485 detailed study but certainly belonging to another species. In contrast to rhinocerotids, equids  
486 are extremely rare at els Casots, being only represented by lower molar fragment assigned to  
487 *Anchitherium* sp. (Rotgers and Alba 2011).

488 Artiodactyls include the small cainotherid *Cainotherium miocaenicum*, a very  
489 common element in older (MN3–MN4) sites of the same basin (see Crusafont et al. 1955;  
490 Casanovas-Vilar et al. 2016), which is much rarer at els Casots, being represented by cranial  
491 and mandibular fragments. Concerning suoids, els Casots is the type locality of two species: a  
492 small “peccary-like” suoid, included in the family Palaeochoeridae (Van der Made 2020) or  
493 Siderochoeridae (Pickford 2017), and a larger listriodontine suid. The former is best  
494 represented by a crushed skull that is the holotype of *Choeromorus ibericus* (Pickford and  
495 Moyà-Solà 1994; Pickford 2017). Alternatively, Van der Made (2020) synonymized this  
496 species with *Choeromorus primus*, with the material from els Casots representing its last

497 appearance datum. The listriodontine is assigned to *Eurolistriodon adelli*, a species originally  
498 described based on a skull with associated skeleton from els Casots (Pickford and Moyà-Solà  
499 1995) that is alternatively included in *Bunolistriodon* by Van der Made (2020). However,  
500 cladistic analyses indicate that *Eurolistriodon* is the sister taxon of *Listriodon* s.l. (i.e.,  
501 including *Bunolistriodon*; Orliac 2006; Orliac et al. 2010). Although *E. adelli* has been cited  
502 from multiple MN4 and MN5 localities (Van der Made 1997, 2020; Pickford and Morales  
503 2003), in the VPB it is only known from els Casots.

504 Ruminant remains are very abundant, including pecoran and non-pecoran forms.  
505 Among the latter, the tragulid “*Dorcatherium*” *crassum*, which belongs to a stem group  
506 distinct from *Dorcatherium* s.s. (Sánchez et al. 2018), is represented by several dentognathic  
507 fragments (Alba et al. 2014). The pecoran fauna is diverse. Palaeomerycids are abundantly  
508 represented by *Ampelomeryx ginsburgi*, a species erected based on els Casots material  
509 (*Duranthon* et al. 1995). *Ampelomeryx ginsburgi* is abundantly represented by partial  
510 skeletons of various individuals (Figure 6b) including complete crania. It belongs to a  
511 palaeomerycid lineage characterised by possessing flattish non-pneumatised ossicones located  
512 over ‘eyebrow’ supraorbital projections, and flat forked occipital appendages (Sánchez et al.  
513 2015). Lagomerycids are rare faunal elements in the VPB, and only a few specimens  
514 (assigned either to *Lagomeryx rutimeyeri* or *Lagomeryx pumilio*) have been recovered from  
515 Early Miocene sites (Crusafont et al. 1955). The few teeth from els Casots are referred to *L.*  
516 *rutimeyeri* because they are clearly larger than those of *L. pumilio*. Cervids are common and,  
517 although no protoantlers have been recovered, considering some similarities with the Sant  
518 Mamet material (referred by Crusafont et al. 1955 to *Procervulus dichotomus*), we tentatively  
519 ascribe the remains to *P. cf. dichotomus*. Together with palaeomerycids, bovids stand out as  
520 the most abundant ruminants at els Casots, being represented by profuse dental remains and  
521 even a partial skeleton of *Eotragus* (Figure 7b). The horn cores are small, short relative to

522 their diameter, and show no evidence of torsion. All these traits, coupled with the moderately  
523 hypsodont teeth, justify an attribution to *Eotragus noyei*.

524 Proboscideans include both deinotheres and gomphotheres. The former are represented  
525 by a few cheek teeth assigned by Gasamans et al. (2021) to *Prodeinotherium* cf. *bavaricum*, a  
526 species characteristic of the early Middle Miocene (MN5–MN6) of Europe. Gomphothere  
527 remains are more common and include a partial skeleton uncovered during 2019 (Figure 6d)  
528 and additional cranial and dental material from the 1990s that is here assigned to  
529 *Gomphotherium* cf. *subtapiroideum* based on dental morphology.

530

### 531 ***Palaeoenvironment***

532 Els Casots fossil site represents a shallow freshwater lacustrine-palustrine (Figure 8)  
533 environment as evidenced by the presence of charophytes, helophytic plants, ostracods,  
534 freshwater gastropods, fishes and various semiaquatic tetrapods. The latter include a diverse  
535 amphibian fauna, pond turtles (*Ptychogaster*), and abundant remains of small-bodied  
536 crocodylians (*Diplocynodon*). The mammal assemblage also includes the semiaquatic otter  
537 *Paralutra*, the aquophilous tragulid “*Dorcatherium*”, and other taxa likely linked to wetlands.  
538 For example, palaeomerycids, abundantly represented by *Ampelomeryx*, generally are thought  
539 to be associated to boggy forests and would have fed on soft leaves and aquatic vegetation  
540 (Köhler 1993; DeMiguel et al. 2014). Concerning suoids, craniodental morphology suggests  
541 that *Choeromorus* species might have fed on swamp plants too (Pickford 2012). Stratigraphic  
542 data show cyclical fluctuations in the water level, probably related to climatic cycles.

543 The faunal list includes some megathermic taxa, such as crocodylians, giant tortoises,  
544 monitor lizards, pythonid and boid snakes. These indicate warm conditions associated with  
545 the MCO. The small mammal assemblage is overwhelmingly dominated by the cricetids

546 *Democricetodon* and *Megacricetodon* and the terrestrial glirid *Simplomys*. Considering that  
547 the site represents a certainly humid environment, the absence of beavers (castorids) is quite  
548 unexpected. The diversity of arboreal small mammals is surprisingly low, with only four  
549 dormice (glirid) species and no tree or flying squirrels. Insectivorous small mammals are also  
550 rare and only include four insectivorans and the marsupial *Amphiperatherium*. The number of  
551 arboreal and insectivorous small mammal species has been found to be positively correlated  
552 to precipitation and environmental humidity (Van Dam, 2006), so this may indicate that at els  
553 Casots more open and arid woodland environments occurred not far from the wetlands. The  
554 larger-bodied herbivores include only browsing species, many of them of small size (e.g.,  
555 *Choeromorus*, *Lagomeryx*, *Procervulus*), which likely inhabited the more forested  
556 environments (Fortelius et al. 1996) surrounding water masses. The different signal provided  
557 by small and larger mammals may result from taphonomic bias. Whereas larger mammals  
558 represent a (para-)autochthonous assemblage, birds of prey are commonly the main  
559 accumulation agents of small mammal remains, which would then provide a more regional  
560 signal. This palaeoenvironmental reconstruction is congruent with macrofloral (Sanz de Siria  
561 Catalán 1993, 2001) and palynologic (Bessedik and Cabrera 1985; Jiménez-Moreno and Suc  
562 2007) data from nearby Langhian deposits from the Penedès area, which indicate tropical to  
563 subtropical environments with seasonal rainfall. Scattered forests rich in tropical elements  
564 developed at low altitudes near humid areas, whereas far away from the wetlands more open  
565 woodlands with herbs (Poaceae, Asteraceae) and leguminous trees (e.g., *Acacia*) defined the  
566 regional vegetation (Jiménez-Moreno and Suc 2007).

567

## 568 **Conclusions**



569 With up to 74 different vertebrate species represented by thousands of specimens, els Casots  
570 stands out as one of the richest early MN5 sites in Europe. It includes multiple fossiliferous  
571 layers, some of them bone beds, that have yielded remarkably complete remains, including  
572 partial skeletons. Recent field campaigns enable constraining its age combining bio- and  
573 magnetostratigraphic data to ~15.9 Ma (Aragonian subzone Cb, earliest MN5, Middle  
574 Miocene) and have also provided detailed insight on the stratigraphy of the site area and  
575 structure of the pre-Miocene basement. Sedimentologic, palaeobotanic and faunal data  
576 indicate that els Casots corresponds to an ancient freshwater lacustrine or palustrine  
577 environment that developed in a tropical-subtropical climate with rainfall seasonality.  
578 Vertebrate remains represent an autochthonous to parautochthonous assemblage, as evidenced  
579 by the partial articulation of several specimens. Many large mammals appear to have  
580 inhabited the forests surrounding the water masses, while small mammals provide evidence of  
581 drier woodlands further from the wetlands. Finally, it is noteworthy that the site dates to the  
582 onset of the Miocene Climatic Optimum and provides a unique opportunity to study wetland  
583 ecosystems from southern Europe during this exceptionally warm period.

584

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600

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627

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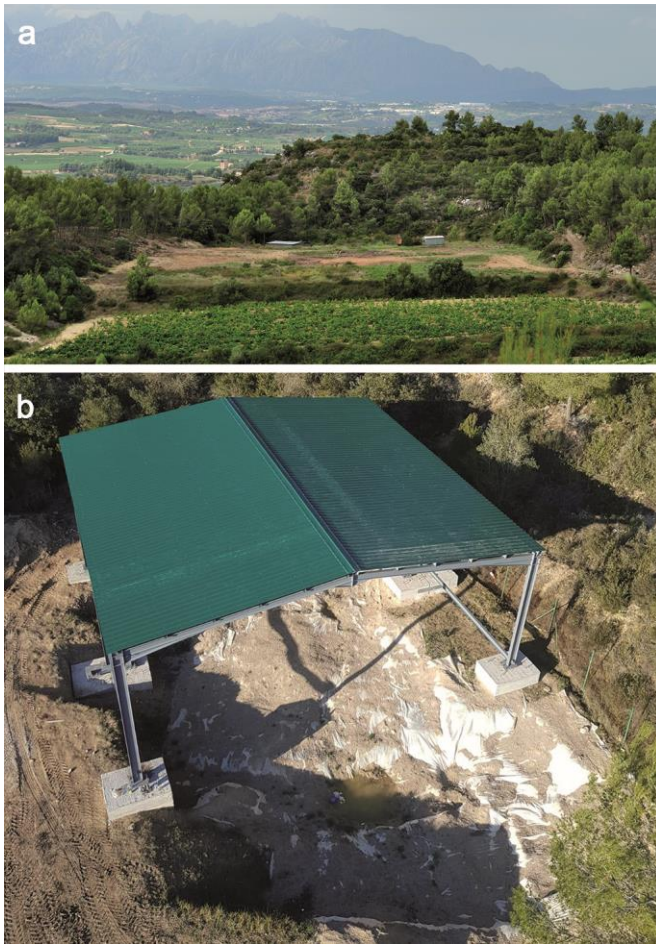
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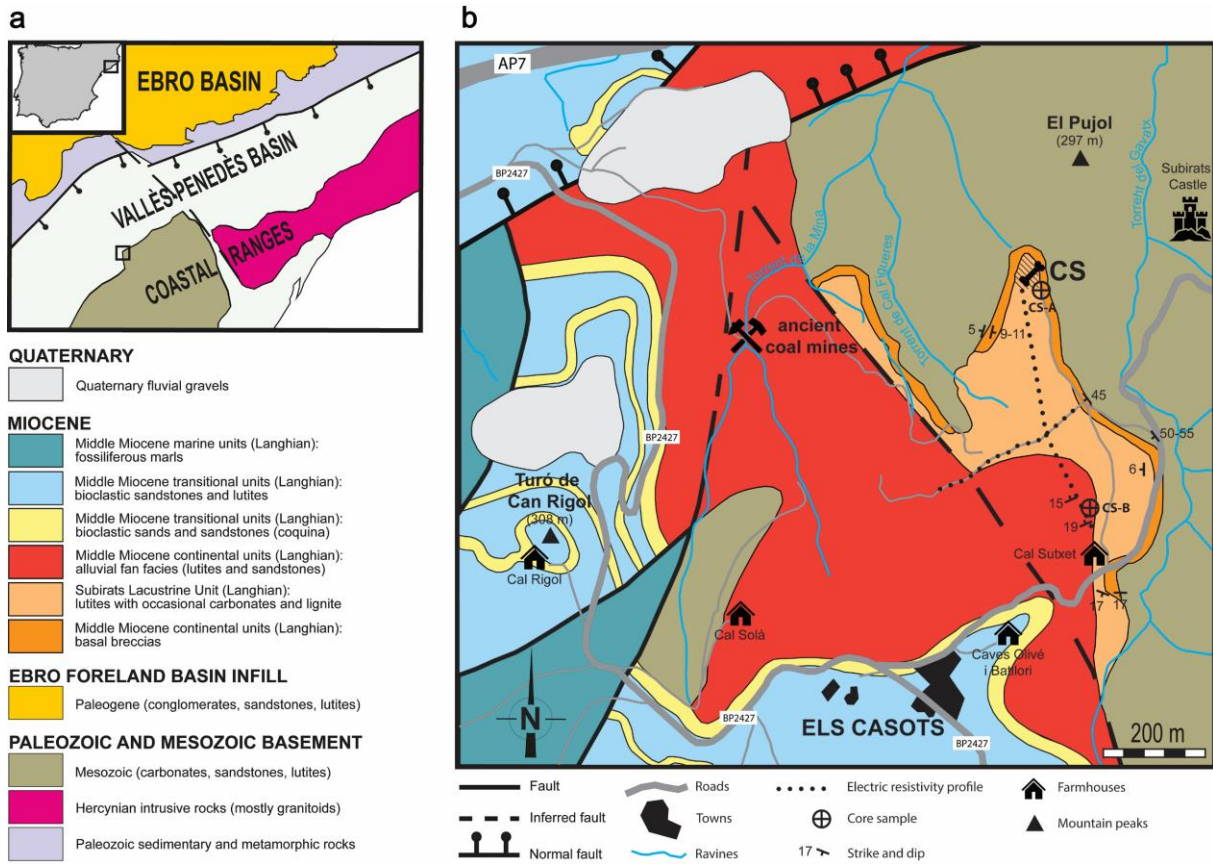
866 **Figures**



867

868 **Figure 1.** a, Panoramic view of the site of els Casots in 2011. Site perimeter is the deforested  
869 field behind the vineyards and corresponds to the hatched area in Figure 2. b, Excavated  
870 surface after the 2019 field campaign. A metal structure has been built over the main  
871 excavation area to protect the site from heavy rains and allow for guided touristic visits.  
872 Photograph courtesy of Ajuntament de Subirats.

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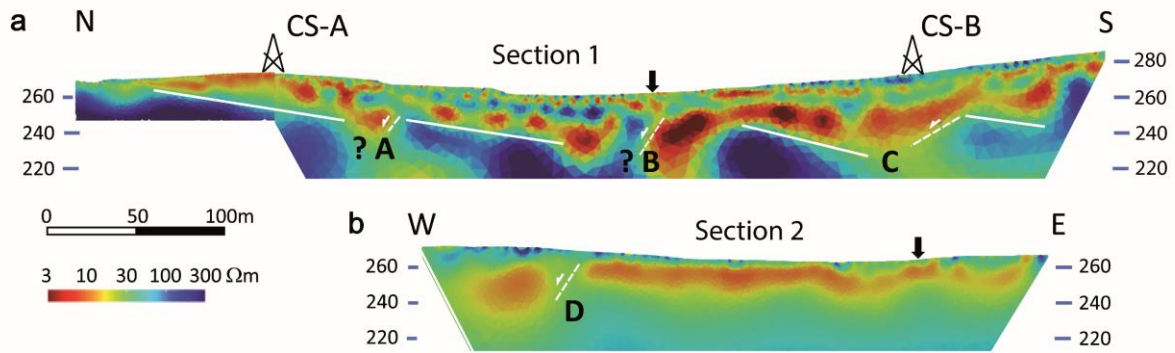


874

875 **Figure 2.** a, Geographical location and general geological context of the Vallès-Penedès  
 876 Basin showing the location of the study area. b, Detailed geological map of the area around  
 877 els Casots, showing the location of the fossil site, the electric resistivity profiles, and core  
 878 samples. Observe the proximity of els Casots to the ancient coal mines as well as its situation  
 879 directly overlying the Mesozoic basement. Also notice the close proximity of the site to the  
 880 first Middle Miocene (Langhian) transitional and marine deposits.

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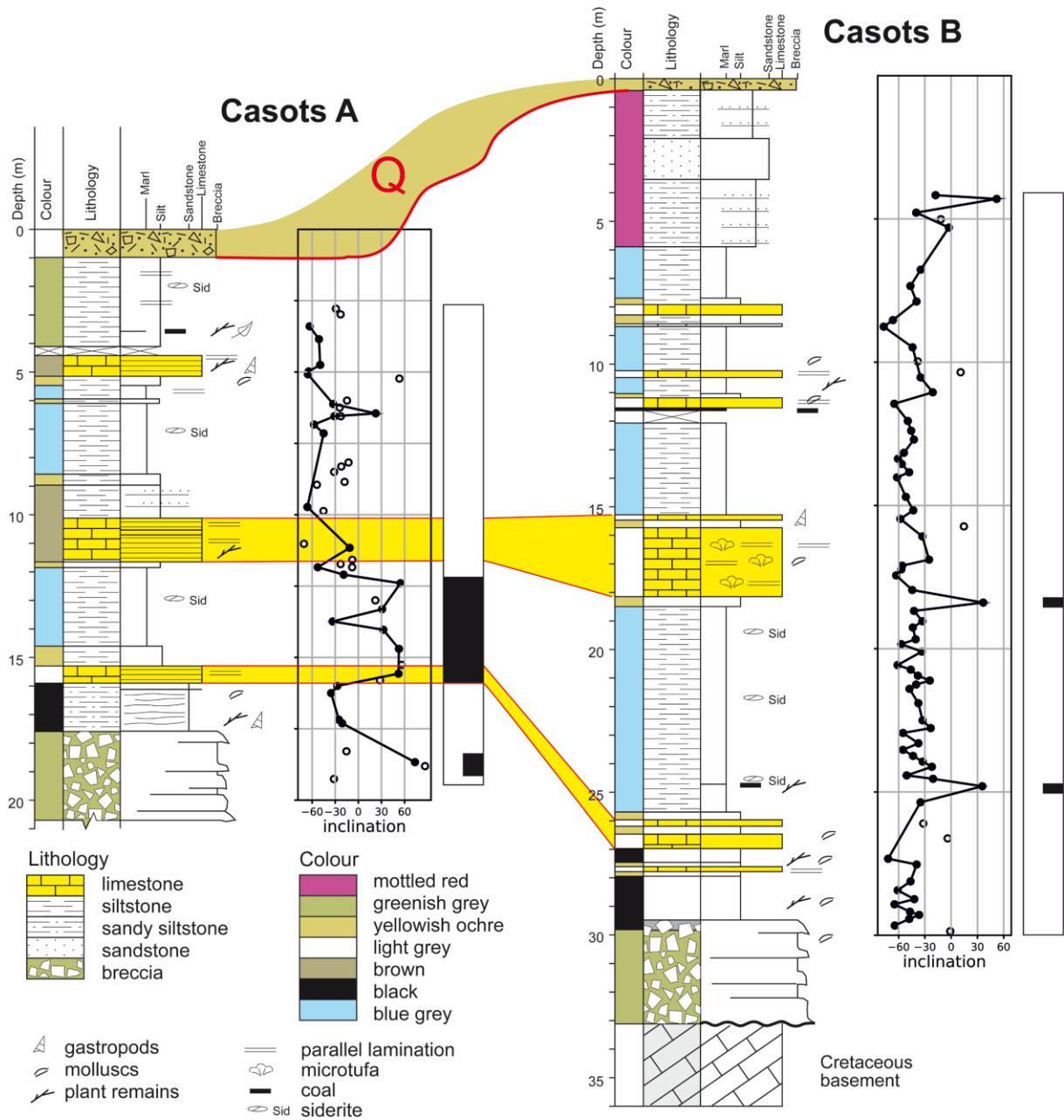




882

883 **Figure 3.** Geoelectric (ERT) sections 1 and 2. a, ERT section 1 is 585 m long with a direction  
 884 S170, overlapping 34 m with two ERT profiles, one of 144 m long with an electrode spacing  
 885 of 2 m, and the other of 475 m and 5 m of electrode spacing. b, ERT section 2 is 355 m with a  
 886 direction N65 corresponding to a unique ERT profile with 5 m electrodes spacing. Data were  
 887 acquired with a Syscal-Pro (Iris-Instruments), and the models were obtained with the ResIPy  
 888 code (Blanchy et al. 2020). Locations of the drilled wells are projected in section 1. White  
 889 lines show the expected contact between Miocene cover and the basement, whereas the black  
 890 arrow indicates the point where both profiles intersect. Segments A–D represent resistivity  
 891 changes interpreted as faulted zones (see text for interpretation). The vertical scale is the  
 892 topographic elevation above sea level in meters. Section location is shown in Figure 2.

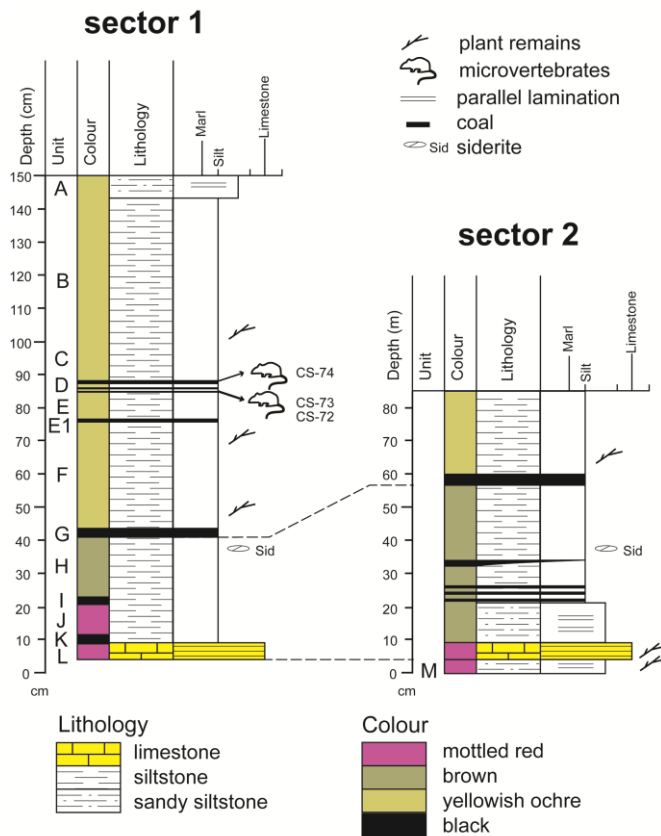
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895 **Figure 4.** Stratigraphic logs of cores Casots A and B (see Figure 2b) and palaeomagnetic  
 896 inclination from studied samples. Positive inclinations represent normal polarity (in black)  
 897 and negative inclinations represent reverse polarity (white). Black circles indicate directions  
 898 of best quality (Quality 1), while white circles indicate directions of lower quality (Quality 2,  
 899 see text for explanation). Note the long reverse polarity interval in CS-B core.

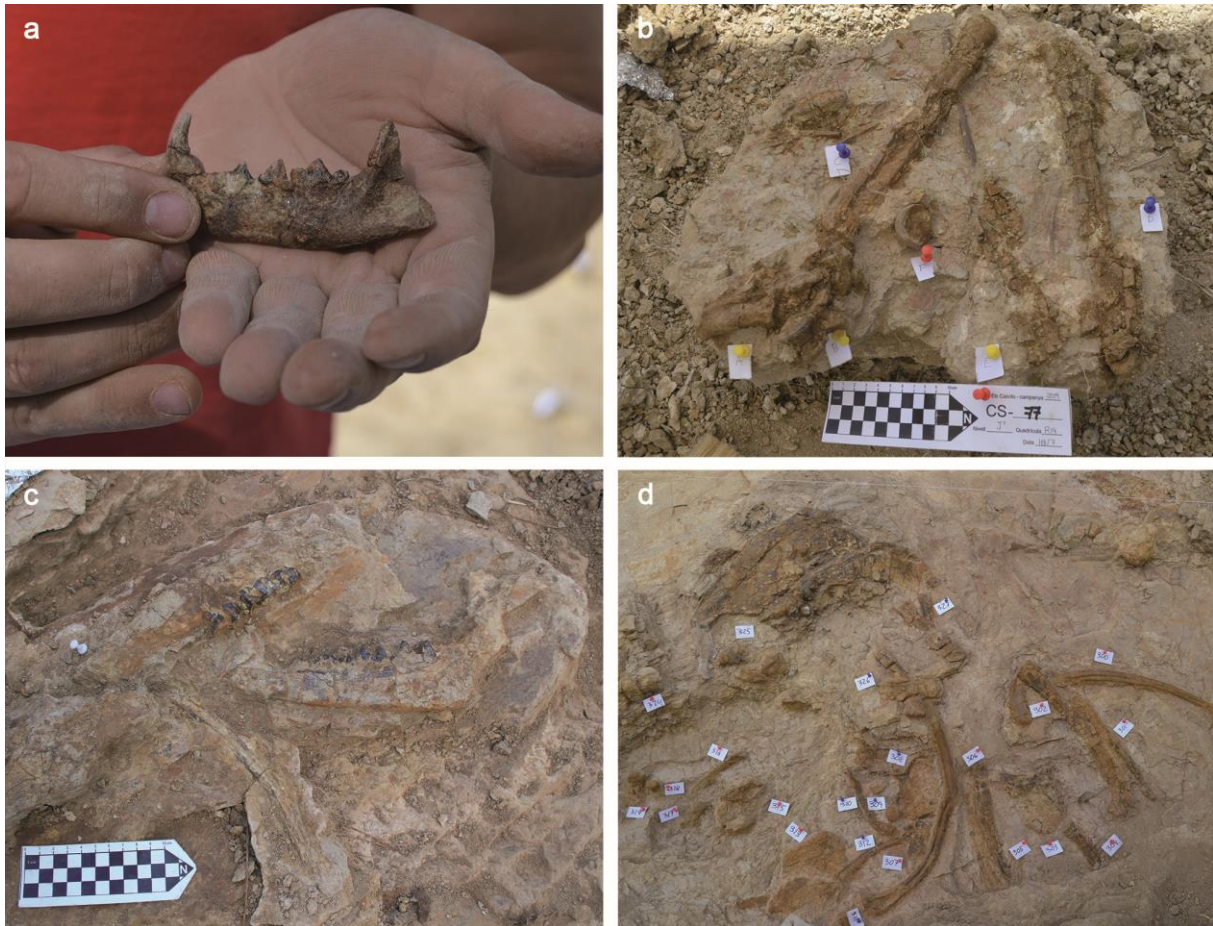
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902 **Figure 5.** Detailed stratigraphic column of the excavation area (sectors 1 and 2). Capital  
 903 letters refer to fossiliferous levels. In this sector, levels A–I were almost entirely excavated  
 904 during the 1990s, so current excavations focus on levels J–M. The levels sampled for  
 905 micromammals during the 1990s (CS-72, CS-73 and CS-74) are also indicated.

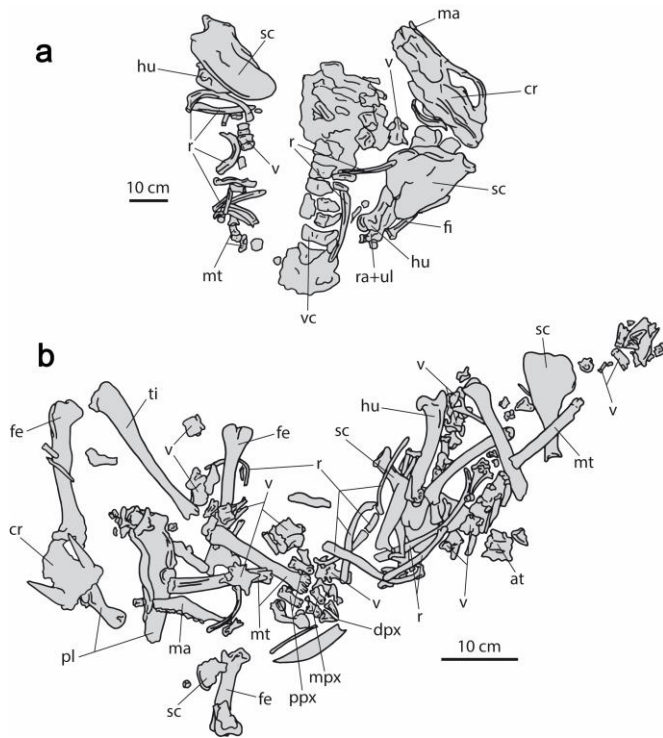
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907

908 **Figure 6.** Some pictures of the recent field campaigns at els Casots. a, Almost complete  
 909 mandible of *Iberictis buloti* (field number CS-150) recovered from level K in 2019. b, Part of  
 910 a disarticulated hindlimb of *Ampelomeryx ginsburgi* (field number CS-77) recovered from  
 911 level J in 2019. From left to right the long bones correspond to a metatarsal, a femur fragment  
 912 and a tibia. c, Two juvenile hemimandibles of a rhinocerotid (field numbers CS-633 and CS-  
 913 634) found in level K in 2021. d, Bone accumulation of different elements (mostly ribs and  
 914 vertebrae) of *Gomphotherium cf. subtapiroideum* found in level K in 2019. A mandibular  
 915 fragment with two molars can be observed in the upper left corner (field number CS-325).  
 916 The whole accumulation of *Gomphotherium* bones covers a surface of 2 m<sup>2</sup>.

917



918

919 **Figure 7.** Excavation plans of two fossil accumulations excavated during the 1990s. a, Partial  
 920 skeleton of *Amphicyonidae* sp. 1 (larger size) extracted during the 1993 campaign (collection  
 921 number IPS11428). Observe the anatomical connection of some elements such as the  
 922 vertebral column (vc) or the cranium and mandible (cr, ma). b, Bone accumulation of two  
 923 different-sized juvenile ruminant individuals found during the 1994 campaign. The smallest,  
 924 and more abundantly represented is *Eotragus noyei*. Although most elements are  
 925 disarticulated, a few are in anatomical connection, such as vertebrae (v) and phalanges (ppx,  
 926 mp, dpx). Collection number for the cranial fragment (cr) of *E. noyei* is IPS11878.  
 927 Orientation and exact position of the bone accumulations within the excavation area was not  
 928 recorded. Anatomical abbreviations: at, atlas; cr, cranium; dpx, distal phalanx; fe, femur; fi,  
 929 fibula; hu, humerus; ma, mandible; mp, middle phalanx; mt, metapodial; pl, pelvis; ppx,  
 930 proximal phalanx; r, rib; ra, radius; sc, scapula; ti, tibia; ul, ulna; v, vertebra; vc, vertebral  
 931 column.

932



933

934 **Figure 8.** Reconstruction of the fauna and palaeoenvironment of els Casots. In the foreground  
935 the felid “*Pseudaelurus*” *lorteti* stares at a large amphycionid that has hunted an individual of  
936 the suid *Eurolestriodon adelli*. Behind them, three individuals of the palaeomerycid  
937 *Ampelomeryx ginsburgi* walk away. To their left and in the background, partly hidden behind  
938 a palm, there is the cervid *Procervulus* cf. *dichotomus*. The crocodylian *Diplocynodon ratelii*  
939 can be seen resting in the lake. In the background, a herd of *Gomphotherium* cf.  
940 *subtapiroideum* approaches the lake shore. Plant remains at els Casots are too fragmentary, so  
941 the reconstructed vegetation is based on that from contemporaneous nearby sites (see Sanz de

942 Síria Catalán 2001). Large plants include the palm *Sabal* and leguminous trees *Acacia* and  
943 *Mimosa*. Bushes in the foreground are *Caesalpinia* and *Cassia*, whereas reeds (*Typha*) grow  
944 abundantly on the lake shore. Palaeoart by Ó. Sanisidro, courtesy of Ajuntament de Subirats.

945

**Table 1.** Updated faunal list of the tetrapod assemblage from els Casots.

| <b>Class</b>   | <b>Order</b>  | <b>Family</b>   | <b>Taxonomic assignment</b>                   |  |                             |
|----------------|---------------|-----------------|---|--|-----------------------------|
| Amphibia       | Anura         | Alytidae        | <i>Latonia</i> sp.                            |  |                             |
|                |               | Bufoinae        | <i>Bufo</i> (group <i>viridis</i> ) sp.       |  |                             |
|                |               | Ranidae         | <i>Pelophylax</i> sp.                         |  |                             |
| Reptilia       | Urodela       |                 | Urodela indet.                                |  |                             |
|                | Testudines    | Testudinidae    | <i>Titanochelon</i> sp.<br><i>Testudo</i> sp. |  |                             |
|                |               | Geoemydidae     | <i>Ptychogaster</i> sp.                       |  |                             |
|                | Squamata      |                 |   | Gekkota indet.   |                             |
|                |               | Lacertidae      |   | cf. <i>Amblyolacerta</i> sp.<br>Lacertidae indet.<br>Scincoidea indet.   |                             |
|                |               | Anguillidae     |   | cf. <i>Ophisaurus</i> sp.<br><i>Pseudopus</i> sp.  |                             |
|                |               | Varanidae       |   | cf. Varanidae indet.   |                             |
|                |               | Blaniidae       |   | <i>Blanus</i> sp.  |                             |
|                |               | Incertae sedis. |   | Scolecophidia indet.   |                             |
|                |               | Boidae          |   | Erycinae indet.  |                             |
|                |               | Pythonidae      |   | <i>Python</i> sp.  |                             |
|                |               | Colubridae      |   | “Colubrinae” indet.  |                             |
|                |               | Viperidae       |   | <i>Vipera</i> sp. (“Oriental vipers”)<br>Viperidae sp. 2 (cf. “European vipers”)   |                             |
|                |               | Aves            | Crocodylia                                    | Diplocynodontidae  | <i>Diplocynodon ratelii</i> |
|                |               |                 | Galliformes                                   | Phasianidae  | <i>Miophasianus altus</i>   |
|                | Falconiformes |                 | Accipitridae                                  | cf. <i>Circaetus</i> sp.<br>cf. <i>Aquila</i> sp.  |                             |
|                | Mammalia      | Strigiformes    | Strigidae                                     | Strigidae indet.   |                             |
| Incertae sedis |               | Herpotheriidae  | <i>Amphiperatherium frequens</i>              |  |                             |
| Chiroptera     |               |                 | Chiroptera indet.                             |  |                             |
| Eulipotyphla   |               | Erinaceidae     |   | <i>Galerix symeonidisi</i>   |                             |
|                |               | Dimylidae       |   | cf. <i>Plesiodimylus</i> sp.   |                             |
|                |               | Heterosoricidae |   | <i>Heterosorex neumayrianus</i>  |                             |
|                |               | Soricidae       |   | <i>Paenelimoecus</i> cf. <i>truyolsi</i>   |                             |
| Lagomorpha     |               | Ochotonidae     |   | <i>Lagopsis</i> sp.  |                             |
| Rodentia       |               | Sciuridae       |   | <i>Atlantoxerus idubedensis</i><br><i>Heteroxerus rubricati</i><br><i>Palaeosciurus</i> sp.<br><i>Csakvaromys besana</i>                           |                             |
|                |               |                 | Gliridae                                      | <i>Bransatoglis</i> sp.<br><i>Glirudinus modestus</i><br><i>Microdyromys monspeliensis</i><br><i>Muscardinus</i> sp.<br><i>Peridyromys murinus</i> |                             |



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|                |                 |  |
|----------------|-----------------|--|
|                |                 | <i>Pseudodryomys ibericus</i>                  |
|                |                 | <i>Simplomys julii</i>                         |
|                |                 | <i>Simplomys robustus</i>                      |
|                |                 | <i>Simplomys simplicidens</i>                  |
|                | Eomyidae        | <i>Ligerimys ellipticus</i>                    |
|                | Cricetidae      | <i>Democricetodon hispanicus</i>               |
|                |                 | <i>Democricetodon gracilis</i>                 |
|                |                 | <i>Democricetodon</i> sp. 3                    |
|                |                 | <i>Megacricetodon primitivus</i>               |
| Carnivora      | Amphicyonidae   | Amphicyonidae sp. 1                            |
|                |                 | Amphicyonidae sp. 2                            |
|                | Hemicyonidae    | <i>Hemicyon stehlini</i>                       |
|                | Mustelidae      | <i>Iberictis buloti</i>                        |
|                |                 | <i>Paralutra</i> sp.                           |
|                |                 | “ <i>Martes</i> ” sp. 1                        |
|                |                 | “ <i>Martes</i> ” sp. 2                        |
|                | Herpestidae     | <i>Leptoplesictis aurelianensis</i>            |
|                | Hyaenidae       | cf. <i>Protictitherium</i> sp.                 |
|                | Felidae         | “ <i>Pseudaelurus</i> ” <i>lorteti</i>         |
| Perissodactyla | Rhinocerotidae  | <i>Dromoceratherium mirallesi</i>              |
|                |                 | Rhinocerotidae sp. 2                           |
|                | Equidae         | <i>Anchitherium</i> sp.                        |
| Artiodactyla   | Cainotheriidae  | <i>Cainotherium miocaenicum</i>                |
|                | Siderochoeridae | <i>Choeromorus ibericus</i>                    |
|                | Suidae          | <i>Eurolistriodon adelli</i>                   |
|                | Tragulidae      | “ <i>Dorcatherium</i> ” <i>crassum</i>         |
|                | Palaeomerycidae | <i>Ampelomeryx ginsburgi</i>                   |
|                | Lagomerycidae   | <i>Lagomeryx rutimeyeri</i>                    |
|                | Cervidae        | <i>Procervulus</i> cf. <i>dichotomus</i>       |
|                | Bovidae         | <i>Eotragus noyei</i>                          |
| Proboscidea    | Deinotheriidae  | <i>Prodeinotherium</i> cf. <i>bavaricum</i>    |
|                | Gomphotheriidae | <i>Gomphotherium</i> cf. <i>subtapiroideum</i> |

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