Dinosaurs nesting on tidal flats

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Abstract

The discovery of a dinosaur clutch in Upper Cretaceous tidal flat deposits in the Tremp Formation in Biscarri, Isona (Lleida, Spain), provides insight into the mode of life of some dinosaurs. The clutch occurs in grey marls that were originally waterlogged, muddy sediments. The seven elongated eggs are parallel to each other and with their long axis almost vertically oriented. A mound-nest is inferred in contrast to hole-nest described so far for similar eggshell types. The permeability of the eggshell (oospecies Meiolithus straungeri) was highly enhanced by a braided network of multicanaliculate pore canals. The nest style, eggshell porosity and tidal flat environment point to a periaquatic habitat for these dinosaurs. A modern analogue occurs in crocodiles, alligators and some birds living in periaquatic environments, with nesting on plant-mounds above waterlogged soils or even on floating vegetation.

Keywords: dinosaurs; eggs; nests; Spanish Pyrenees; tidal flats; Upper Cretaceous

1. Introduction

Evidence for the dinosaur mode of life is rarely found in the fossil record, and thus habitat hypoth- eses have been supported by indirect arguments. Sauropods were classically regarded as aquatic dinosaurs, having been considered too heavy for walking on land. They were depicted submerged in shallow water with snorkel-like nares (Hay, 1908). Such a ‘hippo-like’ mode of life was widely accepted also for hadrosaurs until the early 1970s. During the ‘Dinosaur Renaissance’ of the 1970s, some authors rejected the aquatic lifestyle hypothe- sis, based on morphofunctional analyses that sug- gested terrestrial habits (Bakker, 1971; Coombs, 1974). This view is now largely accepted by most dinosaur palaeontologists.

Most of the fossil evidence for dinosaur nesting habits is found in deposits from terrestrial habitats (flood-plains, palaeosols, abandoned channel-fills deposits; Erben et al., 1979; Iatzoura et al., 1991; Cousin et al., 1994). However, because the amni- otic egg is adapted for terrestrial conditions, most aquatic amniote vertebrates nest on dry land; thus, the terrestrial settings for dinosaur nests are not necessarily evidence for a terrestrial habitat for dinosaurs.

There are in fact some indications for a periaqu- atic habitat for some dinosaurs. Dinosaur eggs are frequently found in shallow marine and estua- rine deposits (Weishampel and Horner, 1990; Schwimmer, 1997 and references cited therein). They are usually interpreted as being transported from terrestrial habitats, but it may also indicate
a periaquatic habitat. The findings of articulate parts, half-parts and also near-complete dinosaur skeletons preserved in marine shelves, scavenged by sharks and even encrusted with oysters are rather frequent (Horner, 1979; Fiorello, 1990; Schwimmer, 1997; Canudo et al., 1998); this would be exceptional for fully terrestrial organisms. Moreover, some sedimentary environments in which sauropod bones and tracks are found connect this group to aquatic environments (Dodson, 1990; Lockley et al., 1994). There are also some cases of dinosaur clutches in marginal aquatic palaeoenvironments, such as lake shores (Sahni et al., 1994) and beach deposits (Sanz et al., 1995). These discoveries suggest an aquatic mode of life for these dinosaurs, but do not prove it. Some terrestrial species, such as modern terrestrial emus (Dromaeus novaehollandiae), may nest occasionally on lake shores when the substrate is dry (De Molina, 1987).

The dating of the La Posa Grey Marls relies on biostratigraphic data. The Biscarri section is situated between two marine levels containing corals and the rudist Hippuritella castroi (Liebau, 1971). These eggs belong to the osospecies Megalooolithus siruguei, currently found in grey marls from aquatic palaeoenvironments. This discovery is the first autochthonous dinosaur clutch recovered from a water-saturated environment. Nest settings in coastal waterlogged soils are not found in terrestrial species, and thus it constitutes palaeobiological evidence indicating a periaquatic habitat for these dinosaurs.

2. Geological setting

The Biscarri clutch was preserved in its original position at the top of a 80 m thick succession of grey marls, shales, lignites and limestones from the lower Member of the Tremp Formation named La Posa Grey Marls (Isona, Lleida, Fig. 1; Liebau, 1973). Some lignite beds have been exploited by mining works in this section, a few metres below the interval represented in the Fig. 1. The continental red beds of the Tremp Formation in the Isona area reach up to 900 m in thickness, of which La Posa Grey Marls made up the lower 120 m. These deposits formed part of the Late Cretaceous regressive cycle filling the South Pyrenean foredeep basin.

The La Posa Grey Marls are underlain and interbedded with marine sandstones from the Aren Formation exposed along the periclinal closure of the Isona Anticline. Near Isona, the Aren Formation consists of medium to coarse grained tabular sandstones with sharp contacts, and containing large-scale sigmoidal cross-bedding indicative of tidal influence (Nagtegaal et al., 1983; Lópezmartín and Ardevol, 1999). The contact with the overlying La Posa Grey Marls is marked by a rudist bed mainly composed of Radiolitella pulchellus. The two related formations have been interpreted as a lagoon-barrier island depositional system dissected by tidal inlets, receiving significant freshwater inputs from the continent (Fig. 2) (Liebau, 1971, 1973; Nagtegaal et al., 1983; Díaz-Molina, 1987).

The dating of the La Posa Grey Marls relies in biostratigraphic data. The Biscarri section is situated between two marine levels containing corals and the rudist Hippuritella castroi (Liebau, 1971). These rudist levels have been correlated with the lower Gansserina gansseri zone indicating a Late Campanian age (Vicens et al., 1999; Ardevol et al., 2000). The upper rudist-bearing level, near the top of the Biscarri section (Fig. 1) has also yielded planktonic foraminifera (Globotricana havanaensis, Globotruncanana arca and Rosita fornicata) and benthic foraminifera (five species of Bolivinoides, such as B. peterssoni-laevigatus and B. draco miliaris) (Liebau, 1973). Between these two clear marine intercalations, the section contains transitional and freshwater deposits with oyster beds, coal seams and peritidal limestones bearing ostracods and charophyte remains. The presence of the charophytes Amblypychora c. macrornota and Peckychora serrulata in the grey marls of this section and the absence of Septorrela indicate a Late Campanian age (Feist and Colombo, 1983; Riveline et al., 1996) after recalibration of the Campanian-Maastrichtian boundary (Obradovich, 1993; Gradstein et al., 1995). Palynomorphs are abundant along the whole section, with ferns and palms dominant (De Porta et al., 1985). The sediment containing the dinosaur clutch is
Fig. 1. Map and log of the Biscarri section in La Posa Grey Marls, lower Member of the Tremp Formation, along the periclinal contour of the Isona anticline. The Aren Formation is situated about 50 m below. The position of the upper Hippuritella castroi level is indicated on the map. The situation of the clutch at the top of a grey marl bed is shown, perforated by traces of roots probably from aquatic plants. Map modified from Liebau (1973).

a homogeneous grey-coloured marl with iron-encrusted root casts (rhizoliths). The marls around the eggs contained plant debris, crustaceans (decapods, euryhaline ostracods), molluscs, fish bones, ray teeth, an ankylosaur tooth and diverse types of eggshell fragments. The eggs are covered by the same types of grey marl that underlie them, and contain framoidal pyrite encrusting the interior side of the eggshells. The marls bear iron-reduced colours and locally contain bioturbation structures (burrows) preserved as siderite concretions that are diagenetic products of the first-formed hydroxides (Ai-Agha et al., 1995). Leaching features, nodulization or motting (marmorization) are absent, indicating that the profile was within the phreatic nape, below the vadose zone. These features indicate water-saturated sediments in a reducing coastal environment, from peritidal swamp (lignites and limestones) to tidal flats (grey marls), comparable to the modern coastal series of mud flats and salt marsh (Verger, 1968; Pye et al., 1990; Allen and Pye, 1992; As-Agha et al., 1995).

A hydromorphic soil developed after the nest was laid, since the eggs were perforated by the root traces passing through them. The rusty-coloured rhizoliths, reaching about 1 m deep and 1–2 cm in diameter, are evenly spaced and gently curved but do not show frequent branching. They are coated with hydrated iron (limonite) and gypsum crystals, which occasionally also encrust lignites and mollusc shells. The iron-coated root traces, and the absence of nodulization, leaching horizon, shrinkage and brecciation, indicate waterlogged gley pedogenetic processes (Retallack, 1998, 1990). The gypsum is early diagenetic, linked to local pyrite hydration in a gley-type hydromor-
Fig. 2. Palaeogeographic reconstruction and cross-section during the late Campanian in the Southern Pyrenees. La Posa Grey Marls correspond to the deposits of lagoon and peritidal swamps. The Aren Formation corresponds to deposits of mesotidal barrier islands system and ebb-tidal delta [according to Nagtegaal et al. (1983) and Díaz-Molina (1987)]. The position of the Biscarri dinosaur clutch corresponds to tidal mud-flat deposits. After flooding and burial of the nest, plants from the supratidal salt marsh prograded over the area to produce the root traces.

phic soil (hydrated pyrite when combined with calcite produces limonite and calcium sulphate; Nicolas, 1949; Ai-Agha et al., 1995).

From these data it can be inferred that the Biscarri dinosaur clutch was deposited in water-logged tidal flat sediments from a perialgalonal environment. After the eggs were laid in the tidal flat, the nest was flooded and covered with mud and subsequently perforated by the growth of plant roots associated to the prograding salt-marsh (Duchaufour, 1960; Freytet and Plaziat, 1982; Rahmani, 1998). Scattered eggshells from hatched eggs of the same egg-type as those of the Biscarri clutch have been reported in sediments a few metres above the clutch (Moratalla, 1993). A concentration of dinosaur eggshells has been reported 5 km north of Biscarri, in a similar sedimentary facies (Rosell, 1967). Eggshells were also found in the neighbouring Barranco de la Posa lignites (Erben et al., 1979). Moreover, bones of sauropod dinosaurs have been found at the neighbouring Torra mine. In addition, probable dinosaur footprints have been reported in the area (Díaz-Molina, 1987; unpublished findings of Eduard Dominguez, Isona). These discoveries document the existence
of stable dinosaur populations living in La Posa lagoon during the Late Campanian.

3. Description of the clutch

The clutch contains seven incomplete dinosaur eggs. Recent erosion has removed the top of the eggs and we do not know if more eggs were originally present (Fig. 3). The preserved clutch is about 90 cm long and about 40 cm wide. Five out of the seven eggs lie in a single plane, and the other two eggs are in a slightly higher plane (Fig. 3). The eggs are arranged in a longitudinal pattern, with five eggs aligned to each other (numbers 1, 2, 3, 5 and 7 in Fig. 3) and two other eggs lying parallel. This evidence is consistent with either both oviducts laying the eggs or with a single oviduct laying the eggs in two phases. The first hypothesis agrees better with the preserved state of the clutch and with other clutches observed in the area (Sanz et al., 1995).

A sediment of similar texture and composition surrounds and fills the eggs and clutch (Fig. 1). Lack of clear margins around the clutch, and the vertical disposition of the eggs (Fig. 3) suggest that the nesting structure was not a buried hole. Moreover, the sediment indicates deposition in water-saturated conditions, and wet mud is an unsuitable substratum for nest burial because it impedes air conductance. Therefore, this nest setting is not compatible with the burial-nesting pattern attributed to sauropods (Erben et al., 1979; Kerourio, 1981; Williams et al., 1984; Cousin et al., 1994; Sanz et al., 1995). The waterlogged conditions of the ground and the arrangement of the clutch, since the hatchling would have disturb the ordered, vertical, closely packed eggs. Dinosaur clutches similar to that described here have not been reported before. Nests with seven or eight eggs from Southern France (Kerourio, 1981; Cousin et al., 1994; Garcia, 1998) have been attributed to a hole nesting with the eggs arranged at different levels. Other clutches show a circular pattern (Zhao, 1979; Horner, 1987; Sahni et al., 1994).

The eggs are elongated in shape, with an estimated maximum length of about 25 cm and 18 cm width. An exact reconstruction of the egg shape, either ellipsoid or ovoid, is precluded because of the preservation state of the eggs. However, the bimodal distribution pattern of the eggshell thickness (Fig. 4) is more consistent with a symmetrical ellipsoid shape. The thickest eggshell fragments correspond to both poles and the thinnest ones to the equatorial area. Ovoid eggs would present a more continuous distribution of eggshell thickness. An ellipsoid-shaped egg is also congruent with a simultaneous production of eggs. An ovoid, asymmetrical shape would indicate a sequential production of eggs, such as in the case of the avian reproductive apparatus (Varricchio et al., 1997).

The eggshell is well preserved, with no signs of erosion, internal surfaces slightly encrusted with pyrite frambooids, and empty pore channels. Eggshell thickness, external ornamentation and shape of the shell crystalline units (Fig. 5) identify the eggshells as *Megaloolithus siruguei* (Vianey-Liaud et al., 1994). This oospecies name includes both spherical and elongated eggs (Williams et al., 1984; Vianey-Liaud et al., 1994; Garcia, 1998). The surface is nodular, with relatively rounded or polygonal nodes (compacituberculate ornamentation). The pore openings are subrounded in shape, with an estimated pore density of about 120–150/cm². No reabsorption craters have been found in the mammillary cores at the base of the shell units along the inner surface, indicating that these eggs were not hatched. This interpretation is also supported by the preserved structure of the clutch, since the hatching would have disturb the ordered, vertical, closely packed eggs.

The crystal units are spherulitic, high, narrow, well individualized and not fused (dinosauroid-spherulitic basic type, and discretetuberculiferous morphology). They show convex growth lines (Fig. 5b) and a sweeping extinction pattern under the polarizing microscope. The eggshell thickness varies between 2.3 and 3.2 mm. The pore canals are relatively straight, with a relatively constant diameter along their length. A braided network of transverse canals connects the primary canals...
Fig. 3. Apical and lateral views of the preserved lower part of the Biscarri clutch (BIS111, photographs by Uly Martin). The original block embedded in a polyurethane matrix was disturbed by falling down during transport. Restoration was performed by reversing it upside down. Below, an interpretative scheme of the original arrangement of the eggs in lateral view. Five of the seven eggs are in one plane, with the exception of numbers 3 and 6. The eggs were laid almost vertically, with the main axes directed towards the ground. The erosion surface is indicated, shaded areas represent lost material above it. The clutch is housed at the Museo Municipal of Isona i Conca Dellà (Lleida, Spain).
embryos from Patagonia with eggshells attributed to Megaloolithidae (Chiappe et al., 1998), although they are not identical to the Old World Megaloolithus eggshells.

4. Palaeobiological interpretation

The Biscarri egg clutch preserved in tidal flat marls is the first autochthonous discovery that documents an unexpected nesting pattern for dinosaurs on the surface of wetlands. This document indicates a case of using waterlogged soil for dinosaur reproduction. No other cases of dinosaur eggs from reducing environments have yet been reported. All other cases of dinosaur eggs recorded from periaquatic deposits come from oxidized, subaereal sediments and are interpreted as nests on dry land after retreat of the shore (Sahni et al., 1994; Sanz et al., 1995). *Megaloolithus siruguei* eggshell fragments have been recovered from grey marls attributed to palustrine–lacustrine palaeoenvironments (Williams et al., 1984; Moratailla, 1993; Vianey-Liaud et al., 1994; Garcia, 1998), but these findings could be allochthonous fossils and do not allow one to infer the original nest habitat.

A nest setting analogous to the Biscarri dinosaur clutch is found in recent Amniota. Mound-nesting of the eggshell greatly (Williams et al., 1984). This feature has not been described in other dinosaur eggshell types.

Associated embryos or hatchlings have not been found yet for this oospecies. Wesnampel et al. (1991) and Grigorescu (1993) attributed a similar egg type from Rumania to hadrosaurs, although the only well-documented case of hadrosaur eggs until now corresponds to a different eggshell type (prolatospherulitic: Hirsch and Quinn, 1990). Bones from both titanosaur sauropods and hadrosaurs have been documented in the Tremp basin (Lapparent and Aguirre, 1957; Brinkmann, 1984; Casanovas et al., 1987; Mastiera and Ullastre, 1988). The tubospherulitic eggshell type was traditionally attributed to sauropods without direct evidence (Matheron, 1869; Dughii and Sirugue, 1957; Sahni et al., 1994). This hypothesis has been recently supported by findings of titanosaur embryos from Patagonia.
The unusual pore canal system of the Biscari eggshells gives additional support to the hypothesis of a riparian habitat for these dinosaurs. The complex network of pore canals found in *Megaloolithus siruguei* (Fig. 5) enhanced the eggshell permeability greatly, which was already extremely high in tubospherulitic eggs (more than eight times that of avian eggshell: Seymour, 1980;
Williams et al., 1984), and high enough to cause the dehydration of the embryo in terrestrial conditions, even in humid climate. The conductance value for *M. siruguei* has been estimated to be 12–17 times the calculated value for avian eggshells of the same size (Williams et al., 1984). A procedure to avoid the fatal loss of water was the burial of the eggs, documented for other tubopsherulitic oospecies (Williams et al., 1984; Erben et al., 1979; Grigorescu et al., 1994; Sanz et al., 1995; García, 1998). In the case of the Biscarri eggs, a buried hole-nest is excluded, because damp mud is an unsuitable cover for the eggs. The water-saturated sediments would smother the embryos because they could no breathe. This clutch was of necessity lying above the sediment, probably surrounded by vegetation, as suggested by the preserved structure (see above).

However, *Megaloolithidae* eggs with highly porous eggshells are unsuitable for nesting on the surface, because they would be dehydrated unless they lie in a water-saturated atmosphere. Furthermore, *Megaloolithus siruguei* eggshells have an enhanced permeability. Dehydration in surface nests is usually avoided by covering the eggs with a mound of plants or sand, which serves also as a temperature control (Coombs, 1989). Whether or not the Biscarri clutch was covered by a plant mound, the sophisticated pore canal network of its eggshells indicates that suffocation was the main danger for the embryos, not dehydration.

The Biscarri dinosaur clutch thus shows two unusually combined palaeobiological features: evidence of surface nesting and an enhanced permeability of the eggshells. The anastomosing pore canal network in *Megaloolithus siruguei* eggshells could be adaptive in the case that some external pores were filled by sediment: the transverse canals would then serve as auxiliary connections between the main vertical canals, allowing gas exchange in the whole egg, thus improving the probability of embryo survival. The extremely high conductance of this egg type would function best in a habitat with a wet mud soil and water-saturated atmosphere. An extremely permeable egg enhancing the ventilation of the embryo would be adaptive for nesting in sultry, air-stagnant swamp environments.

5. Conclusions

The Biscarri clutch documents that some dinosaurs with *Megaloolithus siruguei* eggshell had paludal nesting habits and inhabited tidal flat environments. The clutch was found in coastal plain sediments interpreted as perilagoonal facies of tidal mud-flats, at the transition with a salt-marsh. The clutch with at least seven eggs was not buried, as is usual in other dinosaur clutches, but laid above the surface of a humid mud substrate, probably with plants supporting the vertical closely packed eggs. *M. siruguei* bears an elaborate, anastomosing pore system that made the eggshell extremely permeable. The high internal connectiv-ity of the pore canal network could be adapted to prevent the accidental closure of pores by mud.

Such a combination of surface nesting and enhanced water conductance of eggshells suggests a water-saturated, air-stagnant atmosphere, where the embryos do not have the risk of dying by dehydration but rather by suffocation. Different localities have yielded fragments of this eggshell-type from paludal sediments, but until now evidence of autochthonous fossils have been lacking. The Biscarri clutch is the first documentation of a dinosaur nesting in waterlogged habitats. Comparison with modern Amniota nesting in such habitats suggests a mode of life linked to water for this type of dinosaur.

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