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New observations on test architecture and construction of *Jullienella foetida* Schlumberger, 1890, the largest shallow-water agglutinated foraminifer in modern oceans

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We present new observations on *Jullienella foetida* Schlumberger, 1890, a giant agglutinated foraminifer with a leaf- or fan-like test reaching a maximum dimension of 14 cm, that is common on some parts of the west African continental shelf. The test wall comprises a smooth, outer veneer of small (<10 µm) mineral grains that overlies the much thicker inner layer, which has a porous structure and is composed of grains measuring several hundreds of microns in size. Micro-CT scans reveal that much of the test interior is filled with cytoplasm, while X-ray micrographs reveal an elaborate system of radiating internal partitions that probably serve to channel cytoplasmic flow and strengthen the test. *Jullienella foetida* resembles some xenophyophores (giant deep-sea foraminifera) in terms of test size and morphology, but lacks their distinctive internal organization; the similarities are therefore considered to be convergent. Based on micro-CT scan data, we calculated an individual cytoplasmic biomass of 3.65 mg wet weight for one specimen. When combined with literature records of seafloor coverage, this yielded an estimate of >7.0 g wet weight m⁻² for the seafloor biomass of *J. foetida* in areas where it is particularly abundant. The relatively restricted distribution of this species off the north-west African coast at depths above 100 m is probably related to the elevated, upwelling-related surface productivity along this margin, which provides enough food to sustain this high biomass. This remarkable species appears to play an important, perhaps keystone, role in benthic ecosystems where it is abundant, providing the only common hard substrate on which sessile organisms can settle.
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Abstract

We present new observations on *Jullienella foetida* Schlumberger, 1890, a giant agglutinated foraminifer with a leaf- or fan-like test reaching a maximum dimension of 14 cm, that is common on some parts of the west African continental shelf. The test wall comprises a smooth, outer veneer of small (<10 µm) mineral grains that overlies the much thicker inner layer, which has a porous structure and is composed of grains measuring several hundreds of microns in size. Micro-CT scans reveal that much of the test interior is filled with cytoplasm, while X-ray micrographs reveal an elaborate system of radiating internal partitions that probably serve to channel cytoplasmic flow and strengthen the test. *Jullienella foetida* resembles some xenophyophores (giant deep-sea foraminifera) in terms of test size and morphology, but lacks their distinctive internal organization; the similarities are therefore considered to be convergent. Based on micro-CT scan data, we calculated an individual cytoplasmic biomass of 3.65 mg wet weight for one specimen. When combined with literature records of seafloor coverage, this yielded an estimate of >7.0 g wet weight m⁻² for the seafloor biomass of *J. foetida* in areas where it is particularly abundant. The relatively restricted distribution of this species off the north-west African coast at depths above 100 m is probably related to the elevated, upwelling-related surface productivity along this margin, which provides enough food to sustain this high biomass. This remarkable species appears to play an important, perhaps keystone, role in benthic ecosystems where it is abundant, providing the only common hard substrate on which sessile organisms can settle.
Introduction

In 1890, Schlumberger described a new and gigantic agglutinated foraminifer from the western coast of Africa (Liberia) and named it *Jullienella foetida* after its collector, the French bryozoan specialist Jules Jullien (*Schlumberger, 1890*). When first discovered, during a French expedition off Liberia in front of “Poor River” at 12.6 meters water depth (Wedabo Beach), Jullien noted that the specimens exuded a particularly "foul-smelling odour", leading Schlumberger to name it *foetida* (from *lat. foetidus* meaning fetid, foetid or malodorous). The species was initially considered to be a bryozoan, but Schlumberger recognized its true character and correctly described it as a single-chambered (monothalamous) agglutinated foraminifer with a large, flat or slightly undulating plate-like test, leaf-like, or fan-like in overall shape and with the chamber interior subdivided by longitudinal partitions (*Schlumberger, 1890*).

*Jullienella foetida* is currently placed within the Schizamminidae, a family established by Nørvang (*1961*) for species of large and somewhat bizarre monothalamous foraminifera that includes the genus *Schizammina* Heron-Allen & Earland, 1929, in addition to *Jullienella* Schlumberger, 1890. This family belongs to the class Monothalamea (‘monothalamids’), a paraphyletic group of single-chambered (monothalamous) foraminifera that encompasses the orders Allogromiida and Astrotrhizida, and includes freshwater as well as marine species (*Pawlowski, Holzmann & Tyszka, 2013*). The monothalamids are subject to ongoing genetically-based revisions and species are currently grouped into a series of clades (*Voltski & Pawlowski, 2015; Gooday et al., 2020*). Unfortunately, there are no genetic data for any species of the Schizamminidae and the relationship of these unusual foraminifera to other monothalamids is therefore unclear.

Since it was first described by Schlumberger (*1890*), *J. foetida* has been widely reported from depths of 14 to 89 m across the West African continental shelf from Western Sahara to Ghana (Fig. 1) (*Longhurst, 1958; Buchanan, 1958; 1960; Nørvang, 1961; Le Calvez, 1963; 1972; Manning & Holthuis, 1981*). It occurs on fine sandy and muddy substrates at densities of up to 200 individuals per m² and covering up to 10% of the sandy seafloor (*Le Loeuff & Intes, 1968; Thiel, 1982; Tendal & Thiel, 2003*). Schlumberger (*1890*) reported that the largest specimens from off Liberia were 6 cm in maximum dimension, and more recent records suggest that it can reach more than twice this size. In situ images of *J. foetida* have shown the thin, plate-like test lying horizontally on the sediment surface with only the lower side partially buried (*Thiel, 1982; Tendal & Thiel, 2003*). These large agglutinated structures often constitute the only available hard substrate on which sessile organisms can settle (*Cook, 1968; 1985*). Its great size, abundance and ecological importance make *J. foetida* an important species in some continental shelf ecosystems of the west African shelf. In this study, we have applied a suite of non-destructive methods, including light microscopy, SEM, X-ray and high-resolution X-ray micro-computed tomography (micro-CT), to reveal new aspects of the internal and external test characteristics of this remarkable foraminifera.
Material and Methods

The new material of *Jullienella foetida* originates from sediment samples collected in 1971 during Meteor Cruise M25 at a water depth of 68 m (sample station # 74/1; Seibold, 1971; 1972). The sample site is located off the coast of Mauritania, north of the capital of Nouakchott at 18°52'N and 16°31'W (Fig. 1). A total of 12 tests was examined (Fig. 2). Images were taken using light microscopy and Scanning Electron Microscopy (SEM, *CamScan MV 2300*, *Vegascan*) and arranged into plates using Adobe Photoshop CS6. X-ray pictures were obtained using a Radifluor 360 generator (Philips Electronics). All specimens analyzed are stored in the micropaleontology collection at the Institute of Geoscience, University of Bonn (LA-2021-Jf-1-14).

To investigate the internal structure further, two individuals of *J. foetida* (Fig. 2A and Fig. 2F) were scanned using the micro-CT scanner *v|tome|xs 240 kV* (GE Sensing & Inspection Technologies GmbH phoenix|x-ray) at the Institute of Geosciences, University of Bonn. During the scans, a total of 1,000 X-ray projections was collected through a 360° rotation of the sample. The specimens were scanned dry at 120 kV and 120 µA (voxel size 0.016 mm). The micro-CT scanner is equipped with a detector panel that produces isotropic voxels (single size image 2024 x 2024 pixels) and a maximum resolution (voxel size) of 1 µm. For all scans, the same shutter speed of 200 ms per capture was used. This generated a stack of grayscale JPEG slice images that were imported into the visualization and analysis program Avizo light 9.2 (ThermoFisher Scientific) for the segmentation of individual architectural elements based on grayscale values (relative X-ray absorption). 3D-reconstructions of the test and volumetric calculations for the test and chamber lumina were then generated, again using Avizo. The raw CT scans and reconstructed 3D models are available for viewing and download on MorphoSource (http://www.morphosource.org/) in Project 000393778.

Micro-CT imaging, which has been used only occasionally with agglutinated foraminifera, was used here to visualize the distribution of the cytoplasm and the test simultaneously. To calculate the area occupied by remnants of dried cytoplasm in vertical and horizontal micro-CT stack images, grey-scale images were analysed using ImageJ software (*Rasband, 1997-2018*). The resulting image analysis provides novel information about the relationship between the cytoplasm and the test.

Results

Overall test morphology

Our specimens of *Jullienella foetida* from Mauritania have large, hard, rigid, leaf-like, fan-like, and plate-like agglutinated tests (Fig. 2), up to ~ 3 cm in size. The surface is wrinkled, often gently undulating and interrupted by more or less distinct arcuate or crescentic ridges (Figs. 2C, G), spaced at intervals of about 1.5 mm. Total test thickness ranges between 800 µm and 1.2 mm (n = 8), with lowest values in smaller individuals and within the slightly depressed areas between
ridges. The wall thickness does not vary significantly during growth stages and remains almost constant throughout the test. In one specimen (Fig. 3A), the plate-like test has overgrown three finger-like projections of what appears to have been an earlier outer margin. Possibly, the growth of this specimen had been interrupted and then redirected as a result of some damage or trauma.

The earliest part of the test appears to be missing in all our specimens of *J. foetida*. Two show a bifurcation in the proximal "stem" (Figs. 2B, E), suggesting that this may be the remains of a link between two lobes of the test (see also Nørvang, 1961), rather than the initial part. Towards the distal end of the test, flattened tubular processes, usually fairly short, extend from the margin. They may branch dichotomously (Figs. 2A, F, I; Fig. 4A, B). The apertures are multiple and consist of numerous rounded, elliptical or slit-like openings at the ends of these tubular extensions of the test periphery.

Test structure

Examination of *Jullienella foetida* tests by SEM revealed an external veneer, comprising small (typically 5-10 µm) angular mineral particles (Fig. 5C, D). The layer is very thin (~20 µm) with a smooth outer surface, but is interrupted by numerous shallow bumps where the much larger underlying grains protrude through it (Fig. 3B). This arrangement, and the transparency of the protruding grains, creates a finely speckled appearance when the wall is viewed at high magnifications under a light microscope. The underlying wall is much thicker (250–350 µm) and composed of subrounded grains, measuring several 100s of microns in size (Figs. 5C, D). It has a very irregular inner surface with deep pits that communicate with open spaces within the wall, creating a porous, labyrinthic structure (Fig. 3D). This is reflected in the micro-CT scans, which show (in reverse view) the interface between the test lumen and inner surface of the wall covered in tiny projections (Fig. 4B, D). To some extent the spaces between the grains are occupied by fine particles similar to those in the outer veneer (Fig. 5E, F). The internal partitions are extensions of the inner layer of the wall and have a similar structure.

Internal structure

The volume of the test and of the internal cavity could be derived from micro-CT data. In two specimens, the agglutinated test alone occupied 67% and 73% (mean 70%) and the test cavity 33% and 27% (mean 30%) of the total volume. Scanning electron, X-ray and micro-CT images show that the test cavity of *Jullienella foetida* is subdivided by a series of discontinuous radiating walls (internal partitions) that have no external expression on the outer surface of the test (Figs. 2, 4, 5). These interior walls are aligned almost in parallel and are spaced at regular distances, subdivide the test lumen into elongated sections. Their radial arrangement reflects the fan-shaped and leaf-like growth form of the large agglutinated test. As the lateral fanning-out of the test increases with growth, new partitions are added (Figs. 2D', J'). Almost all these internal walls are discontinuous, with interruptions often occurring at approximately the same growth stage, allowing efficient protoplasmic communication in both longitudinal and lateral directions. From
an architectural point of view, the longitudinal arrangement of the intermittent partitions may
serve to strengthen the test and prevent the otherwise unsupported “roof” from collapsing, in
addition to channelling cytoplasmic streaming. However, they are not equally developed in all
specimens. In some (Figs. 2D’, I’, J’), they appear consistently strong in the X-ray images, but in
others they are weaker and more intermittent (Figs. 2A’, C’, G’).

Cytoplasm
Along broken edges of the test, remnants of dark brown cytoplasm are present within the lumen
(Fig. 5D) or attached to the inner surface of the wall (Figs. 3E, F). This material contains a
number of diatoms (Fig. 3F). The cytoplasm is also visible in micro-CT scans, where it stands
out as a light-grey, low-density component within the test lumen. In the CT-scan slices shown in
Fig. 6A-B, it is present within the central area as well as the finger-like projections and marginal
openings, but is patchily distributed. Grey-scale image analysis using ImageJ showed that the
dried cytoplasm occupies ~19.2% of the test area in cross section (Fig. 6A), and ~36.4% in
vertical sections (Fig. 6B). However, different horizontal CT-scan slices of the same specimen
(Supplementary Fig. S1) reveal the presence of cytoplasm in other parts of the test lumen. This
suggests that cytoplasm is more widely distributed within the test lumen than is apparent in Fig.
6B, although shrinkage during drying will have created gaps.

Discussion
Comparison with previous observations
The largest specimens of *Jullienella foetida* documented in the literature were observed at ~60 m
depth off the coast of Mauritania (station 192, Meteor expedition 44; Tendal & Thiel, 2003).
Here, they reached a maximum dimension of 14 cm and included a range of morphological
types, including thin, leaf-, fan- and kidney-shaped forms, most of them more or less flat. Other
published illustrations show the leaf-like growth pattern extending from a central juvenile stage
in opposite directions to create a dumbbell-shaped test, or in one direction to form a subcircular
feature (Schlumberger, 1890; Buchanan, 1958; Longhurst, 1958; Nørvang, 1961; Tendal &
Thiel, 2003; see also our Figs. 2A, G). Finger-like tubular processes extending from the test
margin, are characteristic of the species (Schlumberger, 1890; Buchanan, 1960; Nørvang, 1961).
Altenbach et al. (2003) described these features as extending laterally or at an angle of 90° into
the seafloor, occasionally branching at some distance from the main part of the test (Tendal &
Thiel, 2003). Some specimens collected off the coast of Ghana (“Unpublished record” in Fig. 1),
also incorporate tubular processes that project at various angles to the main plane of the test
(Gooday, unpublished observations).

Our specimens of *J. foetida* from off the coast of Mauritania have maximum dimensions of only
~3 cm and are therefore much smaller than many of those illustrated in the literature, including
the specimens of Tendal & Thiel (2003), referred to above, which were also from the
Mauritanian margin. None appears to be intact, and indeed, complete specimens of this species
have rarely been recovered (Buchanan, 1958). They are all plate-like (Fig. 2), and one incorporates a single open space (Fig. 3A), but there is no tendency to form a reticulated structure, as seen in some examples illustrated by Nørvang (1961) and Tendal & Thiel (2003). The marginal processes are also generally less well developed in our material. However, in other respects, our specimens resemble published illustration of *J. foetida* and we have no doubt that they belong to this large and distinctive species.

The discontinuous, fine-grained surface veneer with protruding large grains is reminiscent of the pattern of agglutination seen in *Astrammina rara* Rhumbler, 1931 (Bowser & Bernhard, 1993), although the surface layer in *J. foetida* is more distinct and the protruding grains occupy a smaller area. In common with many other agglutinated foraminiferal species (Heron-Allen, 1915; Lipps, 1973; Armynot du Châtelet, Recourt & Chopin, 2008; Makled & Langer, 2009), *J. foetida* appears able to select particular kinds of grains according to both size and composition. We assume that these are bound together by the kind of organic cement found in monothalamous and some multichambered agglutinated foraminifera (Bender, 1989; 1995; Loeblich & Tappan, 1989; Kaminski, 2004). The typical brownish colour of the test probably reflects the presence of iron chemically bound to an organic cement (Hedley, 1963), and this may be responsible for the strength of the test wall, which is remarkably difficult to break. However, we could not observe any obvious cement in our SEM micrographs (Figs. 5E, F). Berthois and Le Calvez (1966) mention that the test of *J. foetida* is formed from quartz grains bound together by siliceous cement (‘ciment siliceux’) that is secreted by the organism. However, no further details of the cement are given and it is unclear what the authors were referring to. Loeblich and Tappan (1989) conclude that there is no evidence for siliceous cement in any agglutinated foraminifera.

**The possible contribution of *Jullienella foetida* to seafloor biomass**

SEM micrographs of *J. foetida* demonstrate the presence of cytoplasm within at least some of the tests. The dried cytoplasm has a granular appearance and contains scattered diatoms (Fig. 3F). It somewhat resembles the cytoplasm of the large tubular foraminifera *Bathysiphon filiformis* Sars, 1872 from the North Carolina margin (Plate 3, Fig. 2 of Gooday et al., 1992), although in that case biogenic particles were more abundant and diverse. Our micro-CT data provide further information, suggesting that the cytoplasm is distributed throughout the test. These scans allow us to make rough estimates of the individual biomass of the specimen illustrated in Fig. 4A, B. The absolute volumes of the test and lumen were 190.2 mm$^3$ and 71.56 mm$^3$, respectively. If we assume that 50% of the lumen was occupied by cytoplasm (a very conservative estimate given that patches of cytoplasm were present in most parts of the test; see Fig. 6B and Supplementary Fig. S1), and that the density of the cytoplasm is 1.02 g ml$^{-1}$ (= 0.102 mg mm$^{-3}$), then the individual biomass of this specimen would be 3.65 mg wet weight. Gooday et al. (2018) report cytoplasm (‘granellare’) volumes for 4 abyssal Pacific xenophyophore specimens varying from 9.45 mm$^3$ in *Galateammina* sp. to 72.6 mm$^3$ in their specimen 1 of *Psammina* aff. *limbata* Kamenskaya, Gooday & Tendal, 2015. Xenophyophore cytoplasm is packed with barite crystals. If we assume that these occupied 50% of the granellare volume in these xenophyophores, then...
their individual biomass values were between 0.48 to 3.70 mg wet weight, respectively. According to these calculations, our scanned *J. foetida* specimen therefore had a biomass comparable to that of a slightly larger xenophyophore (the maximum dimension of *P. aff. limbata* specimen 1 was ~3.5 cm compared to ~2.2 cm for *J. foetida*), and greater than that of three other xenophyophore specimens.

If similar assumptions are applied to the much larger specimens of *J. foetida* photographed by Tendal & Thiel (2003) at 17° N off Mauritania, then their individual biomass would be much greater than the estimate for our specimen. Thiel (1982) estimated that *Jullienella* covered up to about 10% of the seafloor area. Based on these data, and assuming that the average test thickness is 1 mm, the cytoplasm occupies 50% of the lumen, and again that the wet weight of 1 ml of cytoplasm is 1.02 g (Levin & Gooday, 1992, Gooday et al., 2018), we calculate the maximum possible seafloor biomass of *J. foetida* in Thiel’s (1982) study area to be 15.3 g wet weight m⁻². This estimate would be less if, as seems likely, a proportion of the specimens was dead. For the sake of argument, we will again assume that this proportion is 50%, which would reduce the seafloor biomass to 7.65 g wet weight m⁻².

We emphasise that these estimates are based on very limited data and involve several major assumptions, particularly regarding the extrapolation from our study to that of Thiel (1982). The actual figures should therefore not be taken too seriously. However, they are probably the right order of magnitude and give some indication of the contribution that *J. foetida* could make to seafloor biomass on parts of the NW African shelf (Fig. 1) where it is abundant. The value of 7.65 g wet weight m⁻² is comparable to maximum foraminiferal biomass estimates, in most cases derived from whole assemblages of smaller species, from different settings (Murray, 2006). For shelf seas around Europe and North America it is higher than almost all of those (maximum 2.99 g m⁻²; in one case 16.3 g m⁻²) compiled by Murray & Alve (2000).

Korsun (2002) concludes that at shelf and upper bathyal depths in parts of the Eurasian Arctic, foraminiferal biomass may be dominated by large agglutinated species. Our estimate is higher than that for the St. Anna Trough in the Kara Sea (0.06–1.7 g m⁻²), where biomass in the >500-µm sieve fraction was dominated by *Reophax pilulifer* Brady, 1884 (Korsun et al., 1998). However, an earlier Russian study cited by Korsun et al. (1998) gives values (1 to 10 g m⁻²) that are comparable to ours for large astrorhiziid foraminifera (*Hyperammina subnodosa* Brady, 1884, *Rhabdammina abyssorum* Sars in Carpenter, 1869, *Pelosina variabilis* Brady, 1879) in the Barents Sea. At a 230-m-deep site in the Barents Sea, Kuznetsov (1996) recorded a biomass of 6.2 g m⁻² for *Hormosina globulifera* Brady, 1879 (although his illustration shows a unilocular test resembling *Saccammina sphaerica* Brady, 1871). We therefore believe that our estimates provide plausible maximum values for the seafloor biomass of *J. foetida*.

Comparison with xenophyophores
In terms of test morphology, *Jullienella foetida* resembles some xenophyophores, a group of large monothalamous foraminifera (suborder Xenophyophoroidea) that are common in the deep sea below about 550 m depth (Tendal, 1972; 1996; Gooday et al., 2017). These similarities were first noticed by Goës (1892) who considered that *J. foetida* 'has much in common' with *Neusina agassizi* Goës, 1892, a species he described from the tropical eastern Pacific that is synonymous with the xenophyophore *Stannophyllum zonarium* Haeckel, 1889. He concluded that the two species 'stand much isolated' from other agglutinated foraminifera and 'justly claim to be placed in a family by themselves'. Later, Cushman (1927) established the family Neusinidae to accommodate *Neusina* Goës 1892 and *Botellina* Carpenter, Jeffreys & Thompson, 1870, to which he later added *Jullienella*, and *Schizammina* (Cushman, 1948), apparently unaware of the synonymy between *N. agassizi* and *S. zonarium*. *Jullienella* is in fact quite different from *Stannophyllum*, which has a soft, flexible test ramified by fine proteinaceous fibres (Tendal, 1972; Gooday et al., 2020). As pointed out by Schulze (1907), these fibres ('dünne Chitinfäden') are not present in *Jullienella*.

There is a closer morphological similarity between *J. foetida* and some plate-like species of *Psammina*, such as *P. zonaria* Tendal, 1994, in which a proximal tube widens to become flat and plate-like (Tendal, 1994). The arrangement of cytoplasmic strands in the fan-shaped *P. aff. limbata* (Gooday et al., 2018) is reminiscent of the system of partitions in *J. foetida*, and the corresponding shape of the cell body. *Nazareammina tenera* Gooday, Aranda da Silva & Pawlowski, 2011 is another *Jullienella*-like xenophyophore. Photographs of this abyssal species taken on the surface of a box core resemble in situ images of *J. foetida* (compare Fig. 12A of Gooday, Aranda da Silva & Pawlowski, 2011, with photographs in Tendal & Thiel, 2003). Like some specimens of *J. foetida* illustrated in Plate VIII, Figs. 1,2,5 of Nørvang (1961), *N. tenera* also has a tendency for the plate-like test to break into bar-like elements that may form a reticulated structure. Despite these morphological similarities, all xenophyophores have a distinctive internal organization, comprising light-coloured strands of cytoplasm enclosed within an organic tube ('granellare') and dark accumulations of waste pellets (stercomata), that distinguish them from *J. foetida*. These features are often immediately obvious when a xenophyophore test is broken open (Gooday et al., 2018) but have never been reported in *J. foetida*. The similarities in test morphology between these two taxa are very likely to be convergent, although in the absence of genetic data for *J. foetida*, a phylogenetic relationship between them cannot be entirely ruled out.

We are somewhat less confident about *Jullienella zealandica* Hayward & Gordon, 1984. Some specimens of this species illustrated by Hayward & Gordon (1984) are remarkably similar in their overall external test morphology to *Psammina zonaria*. It is important to note that *J. zealandica* lives at 950 to 1400 m, well within the known depth range of xenophyophores (Tendal, 1972; 1989; 1996) but much deeper than other members of the Schizamminidae. However, the internal test structure is apparently rather different, being subdivided by transverse
partitions in *P. zonarium* but undivided in *J. zealandica*. An examination of the cellular
organization of this species would be helpful in determining whether or not it is a true *Jullienella*
species.

**Distribution**

Since Schlumberger’s original description from off Liberia (*Schlumberger*, 1890), *Jullienella*
*foetida* has been found to be widespread along the western coast of Africa, including between
Western Sahara to Ghana, Mauritania, Senegal, Gambia, French Guinea, Sierra Leone, Liberia,
Ghana, and Côte d’Ivoire (Fig. 1). Our material adds an additional record from off Mauritania.
The species occurs across this range in fairly shallow waters on sandy sediment at depths
between 12 and 89 m and within a temperature range from 16 to 25°C. Abundance is maximal
(up to 200 individuals per m$^2$) at 19°C, a temperature that also corresponds to the occurrence of
the largest specimens (*Tendal & Thiel, 2003*). Although the bathymetric distribution of this and
other schizamminid species may be influenced by sediment grain size (*Buchanan, 1960*), the
need for a high food supply seems to the main factor controlling its overall range. *Tendal &
Thiel (2003)* hypothesized that *J. foetida* is restricted to regions where seasonal upwelling
occurs, which would be consistent with its large test size and likely high individual biomass.
Other large agglutinated foraminifera are reported to occur in areas of organic matter flux to the
seafloor (e.g. *Gooday et al., 1992*). The overall distribution coincides well with large parts of the
Canary Current Upwelling System (CCUS), an area that extends from the Iberian Peninsula to
Guinea, and constitutes one of the most productive coastal upwelling systems in the world
(*Demarcq & Somoue, 2015; Kämpf & Chapman, 2016*). In addition, the CCUS area is situated
adjacent to the Sahara Desert and exposed to one of the highest rates of airborne dust, a major
source of nutrients, in particular iron (*Neuer et al., 2004*). Towards the southern part of the range
(Gulf of Guinea), river runoff becomes the main source for the organic matter deposited on the
continental shelf (*Kämpf & Chapman, 2016*). Despite extensive studies on the shallow benthic
foraminiferal assemblages from reefs, shallow coastal habitats, lagoons and mangrove
environments, *J. foetida* has not yet been recorded from Gabon, Sao Tomé, Príncipe or Nigeria
(*Langer, Fajemila & Mannl, 2016; Fajemila & Langer 2016; 2017; Fajemila, Sariaslan &
Langer, 2020*).

**Concluding remarks**

*Jullienella foetida* is probably the largest agglutinated foraminiferal species occurring in
relatively shallow water (<100 m depth). The thin, basically fan-shaped test can reach lengths of
up to ~14 cm, a size only matched among continental-shelf foraminifera by the discoidal
calcareous nummulitiid (*Globothalamea Cycloclypeus carpenteri* Brady, 1881 (*Briguglio et al.,
2016*). Some deep-sea xenophyophores are larger in terms of test size (up to 20 cm or more;
*Tendal, 1972*), but only a small part (a few percent at most) of this volume is occupied by
cytoplasm. In contrast, our new observations of the internal structure of *J. foetida* suggests that
the cell body probably fills much of the test interior, which would mean that this species is
possibly one of the largest of all foraminifera in terms of biomass. X-ray images of the test reveal an elaborate system of radial partitions that subdivides the test interior into channels (also shown in pl. 17, figs. 7,8 of Loeblich & Tappan, 1988). These may serve to direct the flow of the cytoplasm, and perhaps increase its surface to volume ratio, as suggested recently for the much smaller calcareous foraminifera Chilostomella ovoidea Reuss, 1850 (Nomaki et al., 2020).

Jullienella foetida occupies a restricted geographical range around part of the NW African margin at water depths above 100 m (Fig. 1). It is found in eutrophic settings and on sandy, sometimes rippled substrates, suggesting a preference for energetic environments. Like some other large agglutinated foraminifera (Gooday et al., 1992), the cytoplasm contains diatoms, suggesting that it feeds on detritus. It seems likely that this species fulfils an important, perhaps keystone, ecosystem role by providing the only extensive firm substrate on which sessile organisms can settle (Cook 1968; 1985), thereby increasing local biodiversity, as well as by processing organic matter at the base of the benthic food chain. However, much remains to be learnt about the ecology and biology of J. foetida. It will also be important to obtain DNA sequences from fresh material in order to clarify the place of this giant species, and others currently assigned to the Schizamminidae, within the radiation of monothalamous foraminifera.

Acknowledgements

We are grateful to Hjalmar Thiel (Hamburg) and Alexander Altenbach (Munich)† for providing additional information on the life position of Jullienella and to John Murray for the collection of material from off Ghana. We also thank Georg Oleschinski and Martha Berens for assistances with the light microscopy, X-ray and SEM images and Rico Schellhorn for help with ImageJ.

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Figure 1

Distribution of *Jullienella foetida*. 

**Map showing all known localities where *Jullienella foetida* has been recorded.** The unpublished record from off Ghana is based on a sample in the collections of the National Oceanography Centre, Southampton, of uncertain provenance. The label in the bottle reads ‘Plant material. Agazziz Trawl No 3. 2-5-51 (i.e., 2\textsuperscript{nd} of May 1951). Gold Coast. R. Barrindale’.
Figure 2

Jullienella foetida

**Jullienella foetida; light photographs and corresponding X-ray photographs of 10 specimens.** The radiating linear structures in the X-ray images are interpreted as internal partitions. In some cases, these features are strongly developed along their entire length, but in others they resemble dashed lines, with prominent sections separated by gaps where they are weakly developed or absent.
Figure 3

SEM electron micrographs of Jullienella foetida.

**Jullienella foetida; scanning electron micrographs.** (a) Complete specimen. (b) Test surface showing smooth, fine-grained outer layer with larger grains projecting through it from the underlying wall. (c) Detail of area enclosed by rectangle in figure (a) showing area where the wall has been removed to show internal features and remnant of dried cytoplasm (smaller rectangle). (d) Detail of area indicated by the larger rectangle in figure (c) showing inner surface of test wall with complex pattern of pits and upstanding areas. (e) Detail of area indicated by smaller rectangle in figure (c) showing surface of cytoplasmic remnant. (f) Detail of area indicated by rectangle in figure (e) showing surface of cytoplasm with diatoms.
**Figure 4**

*Jullienella foetida*; micro-CT scans.

*Jullienella foetida*; micro-CT scans of the two specimens shown in Fig. 2a, f. (a, c). Test surface; note the protruding grains. (b, d) Test lumen showing the interface between the test wall and the inner cavity. In effect, this is a view of the inner surface of the wall in reverse. The interface is covered with small-scale irregularities reflecting the labyrinthic nature of the test wall. Note that the internal partitions in (b, d), indicated by open spaces, are developed intermittently, particularly in (b).
Figure 5

Scanning electron micrographs of Jullienella foetida.

**Jullienella foetida; scanning electron micrographs.** (a) Complete specimen. (b-d) Progressively closer views of a broken edge with the coarsely agglutinated test wall overlain by a very thin, fine-grained surface veneer; the test lumen is interrupted by internal agglutinated grains that form, either cross-sections of partitions or more isolated columnar structures. (e) Detail of broken test wall showing large agglutinated grains with intervening spaces filled by fine-grained mortar. (f) Detail of fine-grained mortar.
Figure 6

Micro-CT images of *Jullienella* specimen with coloured cytoplasm.

**Greyscale micro-CT images of the Jullienella specimen illustrated in Fig. 2a,a' and 4a,b with two density components.** (a) Cross section. (b) Section in the plane of the test. The agglutinated test wall (=aw) and internal test partitions (=itp) are well defined as dense and bright white in greyscale scan images. The cytoplasm (=cy) occurs as low-density, material (coloured yellow) and is patchily distributed throughout the test. See also Supplementary Fig. S1.
