Biomacromolecules in recent phosphate-shelled brachiopods: identification and characterization of chitin matrix

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ABSTRACT
Phosphate-shelled brachiopods differ in filter-feeding lifestyle, with Lingula anatina an active infaunal burrower, and Discinisca tenuis a shallow marine epibenthic animal. The shells of these animals are built of organophosphatic constituents, the organic fibres/sheets reinforced with calcium phosphate to provide a sophisticated ultrastructural robustness. This investigation examined the nature of the organic fibres in order to improve understanding of how living organisms produce hierarchically structured biomaterials. Unlike powdered samples commonly used in previous studies, organic fibres were isolated for the first time and the shell fractions were purified, in order to study the content and nature of the biopolymer fibres. Biochemical methods including Calcofluor staining revealed a chitin matrix. Ultrastructural analysis, thermal gravimetric analysis, and spectroscopic analyses show that the core polysaccharide framework is composed of layers of β-chitin sheets and/or fibrils that are coated with a fibrous organic matrix. There is more chitin matrix in the L. anatina shells (26.6 wt.%) compared to the D. tenuis shells (12.9 wt.%). Taken together, the data show that the chitin matrix contributes to increased skeletal strength, making L. anatina highly adapted for life as an active burrower. In comparison, D. tenuis contains less chitin and lives as attached epibenthos in a shallow marine environment.
GRAPHICAL ABSTRACT

First spectroscopic evidence of β-chitin sheets in recent organophosphatic brachiopods

Introduction

In nature, living organisms are highly organized and are capable of producing three-dimensional hierarchically structured materials such as teeth, bones, and seashells. These biomaterials are inorganic–organic nanocomposites with exceptional structures and functions. Inorganic components comprise biominerals, that is, calcium carbonates and phosphates or silica [1–3], whereas organic macromolecules constitute a combination of soluble- and insoluble-organic components, primarily peptide-, protein- and polysaccharide-based layers which provide a scaffold for the inorganic phase [4–6]. Water insoluble macromolecules are most frequently built from monomers-based either on sugars or peptides. Sugar monomers and biomolecules based on peptides are present throughout the animal kingdom [5, 7, 8]. The former are chemically modified, most commonly by amination, and are referred to as chitin, whereas the product of subsequent deacetylation of chitin is known as chitosan [8, 9]. As for the biomolecules based on peptides, these can either be functional or structural proteins comprised of a variety of fibrous, tubular, and triple helices in the case of collagen [5, 10].

Chitin is semicrystalline in nature and has three polymorphic forms: α-, β-, and γ-chitin [11–14]. Polysaccharide α-chitin is the most common, in which the chains of α-chitin are arranged in an anti-parallel manner with strong intermolecular hydrogen bonding [12]. β-Chitin chains are parallel to each other, with relatively weak intermolecular hydrogen bonding [11]. Chains of γ-chitin are a mixture of α- and β-chitin, with two parallel chains in one direction and a third one in the opposite direction [13, 14]. Typically, chitin is the structural polymer to be found in three-dimensional architectures of fungi, yeasts, diatoms, sponges, corals, annelids, and mostly in arthropods, including insects, arachnids, and crustaceans [10, 13, 15, 16]. In the layered cuticle of arthropod exoskeletons, chitin is cross-linked with substantial amounts of proteins to form variably oriented yet organized fibres [17, 18]. In the matrix sheets of calcareous bivalve shells, the core polysaccharide framework is composed of interlamellar structure of...
β-chitin, and is frequently reinforced and/or composited with extracellular matrix [4, 19, 20], other biomolecules include glycosaminoglycans [21–23], and/or inorganic components [2, 4]. These composite materials are high-strength yet lightweight, and provide functional flexibility, support ultrastructures, mechanical stability and compressive strength [2, 24, 25]. Living organisms use a combination of organic matrices and inorganic constituents, in various ratios, to produce extraordinary arrays of diverse shell morphologies which eventually facilitate adaptation to a diverse suite of habitats [26, 27].

Brachiopods are a diverse group of marine bivalved, sessile, filter-feeding lophophorate animals currently classified into three subphyla: Linguliformea, Craniiformea and Rhynchonelliformea [28]. Craniiformea and Rhynchonelliformea utilize calcium carbonate and ~2% of an organic matrix to hierarchically assemble ultrastructures of the calcitic shells [29]. In contrast, linguliform brachiopods such as L. anatina and D. tenuis use organic biopolymers reinforced with calcium phosphate (Ca-phosphate) to produce the shells [1, 30]. Linguliform brachiopods are an important ancestral animal group, with an unexcelled fossil record that originated some 530 million years ago in the early Cambrian as part of the emergence of biomineralization across disparate animal clades [28, 31]. Linguliform brachiopods exhibit distinctive features including a ciliated lophophore, chitinous chaetae, as well as a fleshy pedicle for anchoring to (or within) the substrate [28, 32]. Typical D. tenuis uses a pedicle for attachment to hard substrates, so as to survive wave action in the steeper slopes of the shallow seafloor [26]. In contrast, L. anatina has an elongate pedicle to prop itself up with the shell against the sediment. The downward U-shaped burrowing is accomplished by cyclical complex motions of the valves against the sediment, with the pedicle trailing behind the shell [26, 27, 32–34]. Definitive evidence for an active U-shaped burrowing lingulid lifestyle is unknown from the Palaeozoic, where the obolids burrowed pedicle first [32–34]. Lingulides have occurred in a vertical burrowed life position since the early Palaeozoic while the first appearance of U-shaped burrows constructed by lingulids likely occurred during the Mesozoic [35, 36]. Although burrowing has been connected with naturally occurring
demineralization [37] and maceration [35], the survival of Lingulidae over time has been attributed to the depth of burrows [26, 32–34]. For instance, the shell of L. anatina is more flexible and has remained well adapted to deep burrows as a refuge against bioerosion of the substrate and predation since the early Palaeozoic [26, 33–35].

Lingulid shells are used as paleoenvironmental proxies due to their complete fossil record and their distribution in modern oceans [28, 38]. Despite their palaeontological importance, the growth mechanism of brachiopod shells including linguliforms, is not clearly understood, and regarding the organic biopolymers, detailed structural and molecular analyses are topical in order to understand the mechanisms and formations of lingulid shells. Although biochemical studies and spectroscopic analyses on powdered brachiopod samples have identified proteins associated with chitin from linguliform shells [39–41], the composite fibres have not been characterized, so little is known about the nature of the organic sheets. Chemical fixation and scanning electron microscopy (SEM) were used to propose that the organic fibrils in linguliform shells are composed of chitin, a dense mat of fibrils which are probably collagenous, and membranous laminae within the shell layers [21, 42]. However, these previous studies were based on a more or less complete mixture of organic biopolymers and Ca-phosphate mineral. It is necessary to establish an easy and efficient method to reveal the composition of the organic sheet layers in brachiopod shells. The approach of this study is to use field emission gun-SEM (FEG-SEM), light and fluorescence microscopy, and comparison of the imaged region with natural composite biominerals analysed by microRaman spectroscopy and Fourier transform infrared (FTIR) spectroscopy. The goal of this study is to identify and characterize the major organic frameworks in lingulid brachiopod shells. This investigation facilitates a clearer understanding of morphological structures of fossilized, recent phosphate-shelled brachiopods and the biosynthesis of organic–inorganic composites, and the design of new biomaterials.
Materials and methods

Materials

Three phosphatic brachiopod shells of Lingula anatina Lamarck, 1801 and Discinisca tenuis Sowerby, 1847 (SI Fig. 1) were used for this study, including both recently alive and fresh specimens (live). Recently alive shells of D. tenuis and L. anatina were collected from Walvis Bay, Namibia, and from Moreton Bay, Queensland, Australia, respectively. The fresh (live) specimens of L. anatina used in this study were collected in the Bay of Guangxi, China, and were preserved in 10% formalin. While long exposure of biological samples to formalin does not alter the organic composition, it does diminish the fracture toughness due to etching/removal of calcium phosphate. Shells of brachiopods are laminated and are covered with a non-mineralized organic layer, the periostracum, which ranges from 1 to 30 μm thick [43, 44].

Sample preparation and thermal gravimetric analysis

The three shell samples were mechanically cleaned with a scalpel, and then washed with Milli-Q water to remove organic and inorganic contaminants. Samples were soaked in hydrogen peroxide (35%; Chem-Supply, UN2014) for about 2 h to remove surface organic detritus. Varied chemical procedures, extraction temperatures, and different times have been described to achieve optimal demineralization and deproteinization of chitin matrix [45, 46]. For this study, each shell sample was demineralized with 5% acetic acid at ambient temperature for about 48 h. Subsequently, shell samples were washed several times with Milli-Q water, followed by deproteinization and/or depigmentation with 5% sodium hydroxide at ambient temperature for ~48 h. During these procedures the samples were monitored intermittently using Fourier transform infrared (FTIR) spectroscopy. After 24 h in sodium hydroxide, several layers were peeled off the shells using tweezers: these are termed the organic fibres. The remaining shell samples and the organic fibres were collected from the solution, washed in Milli-Q water until a pH of about 6.8 was reached, and then rinsed in cold acetone twice. The shell samples and organic fibres were then dried at room temperature.

Shells were broken into a few mm-sized pieces and heated using a thermal gravimetric analyzer (TGA) model 2050 (TA Instruments, USA) equipped with a differential thermal gravimetric (DTG) analyzer. About 10 mg of sample was measured at a heating rate of 10 °C/min over a temperature interval from 25 to 900 °C. The moisture content, organic biopolymer content, and residual ash content were determined from mass losses. The analyses were recorded twice for each sample.

α-Chitin extracted from shrimps (Sigma-Aldrich) and squid pen β-chitin were used as standards for this study. A squid pen was prepared by degrading any possible inorganic matrix with 5% hydrochloric acid at room temperature for 24 h. After washing with Milli-Q water, the squid pen sample was treated with 5% sodium hydroxide at 65 °C for 6.5 h, and was washed with Milli-Q water until a pH of about 6.8 was reached.

Calcofluor white staining and chitinase digestion test

For biochemical staining, isolated organic fibres from the brachiopod shells were rinsed with Milli-Q water, and then 25 μL of a solution of 10 g sodium hydroxide in 90 mL Milli-Q water and 10 mL glycerine were added to the samples. After 40 s, a 0.1% Calcofluor White M2R (Sigma-Aldrich; Fluorescent Brightener F-3543) solution was added and the samples were kept in the dark for ~30 min. Samples were rinsed twice with Milli-Q water and dried at room temperature. Images of the stained samples were captured using 40 x objective on a Zeiss fluorescence microscope system (Zeiss Axioskop 40 FL system) equipped with the Axiocam 506 mono high-resolution camera and Zeiss ZEN2 lite software. The images were acquired below the saturation level of the camera, and all images shown are representative of results.

For the chitinase digestion process, sodium hydroxide-etched samples and organic fibres were rinsed with phosphate buffer (pH 6.0), then treated with chitinase (EC 3.2.1.14, No. C-6137, Sigma-Aldrich) from Streptomyces griseus in a phosphate buffer (pH 6.0; 1 mg/mL). Control samples were treated with phosphate buffer without chitinase. The
treated and control samples were incubated at 25 °C for ~ 20 h and later rinsed with Milli-Q water. Samples were allowed to dry at room temperature, before imaging without prior coating using a Phenol XL Desktop back-scattered detector scanning electron microscope (SEM).

MicroRaman and Fourier transform infrared spectroscopy

Raman spectra were recorded at room temperature using a Horiba Jobin Yvon LabRAM HR Evolution spectrometer equipped with an Olympus BX41 microscope and an automated x–y stage. A red wavelength helium–neon laser (633 nm) was used and scattered light was dispersed by a grating with 600 grooves/mm (750 nm spacing). Rayleigh radiation was blocked using an edge filter. Spectra of the organic fibres were recorded in the 400–1800 cm\(^{-1}\) range with an acquisition time of 60 s, 10 accumulations, and a delay time of 3 s. Twenty spectra were measured for all samples. The Raman spectrometer was calibrated before and after measurement using the 520.46–520.60 cm\(^{-1}\) peak of a silicon wafer. Data acquisition and spectra treatments were carried out with the commercially available program LabSpec v6 (HORIBA Jobin Yvon GmbH).

FTIR measurements were conducted as described previously [1], with 64 accumulations and a resolution of 2 cm\(^{-1}\). Background spectra were measured at the start of each analysis. Spectra treatments were performed using the software Origin LabPro 2017. A linear baseline was subtracted for the spectra and for the amide I region, over the range of 1580–1720 cm\(^{-1}\). Peak fitting of the spectra was performed to reveal the different components based on Gaussian functions.

Scanning electron microscope imaging

Both organic fibres and shell samples were mounted on aluminium SEM sample holders and gold coated for imaging, prior to analysis with a JEOL JSM-7100F field emission gun SEM at an electron energy of 10 kV and a 10 mm walking distance. For the chitinase digestion test, samples were evaluated without coating with a Phenol XL Desktop back-scattered detector SEM at an electron energy of 15 kV and ~ 5 mm walking distance.

Results and discussion

Results

Vibrational spectroscopy—microRaman and FTIR spectroscopy

Raman spectra of organic fibres from the shell samples (Fig. 1) show absorption bands that are typical of chitin (N-acetyl-D-glucosamine). The band assignments are listed in Table 1, where they are compared with the spectra of standard α-chitin and squid pen β-chitin, and literature data [1, 20, 47–52]. The spectra of the organic fibres reveal the presence of C=O (carbonyl) functional groups based on the amide I stretching vibration at ~ 1654 cm\(^{-1}\), and correspond primarily to the β-form of chitin. The deconvolutional fit of the amide I band in the 1580–1720 cm\(^{-1}\) range...
Table 1 Raman and FTIR band assignment for α-chitin, β-chitin, and chitin (organic) fibre in the 4000–400 cm\(^{-1}\) range

|   | Raman | FTIR |   | Raman | FTIR |   | Raman | FTIR |   | Raman | FTIR |
|---|-------|------|---|-------|------|---|-------|------|---|-------|------|
| α-chitin |       |      | β-chitin |       |      | L. anatine (FL/RL) |       |      | D. tenius |       |      |
| Raman | FTIR | Raman | FTIR | Raman | FTIR | Raman | FTIR | Raman | FTIR |
| 3476 | 3434 | 3260 | 3104 | 2959 | 2935 | 2879 |       |      |      |       |
| 3532* | 3536* | 3424/18 | 3106/1 | 2958/61 | 2932/1 | 2877 |       |      |      | 2853* |
| 3418 | 3282 | Amide A, N–H ass |      | 2952 | CH\(_3\) ass |      | 2851 | CH\(_2\) ss |      |      |
| 3282 | 3095 | Amide B, N–H ss |      | 2928 | CH\(_2\) ass |      |      |      |      |
| 1654 | 1656 | 1656/5 | 1654/5 | 1656 | 1652 | Amide I, C=O s |      |      |      |      |
| 1617 | 1621 | 1620/1 | 1620/5 | 1627* | 1634* | Amide I, C=O s |      |      |      |      |
| 1553 | 1552 |       |      | 1550 |      |      |      |      |      |      |
| 1447 | 1455 |       |      | 1451 |      |      |      |      |      |      |
| 1428 | 1427 |       |      | 1430 |      |      |      |      |      |      |
| 1413 | 1415 |       |      | 1414 |      |      |      |      |      |      |
| 1376 | 1376 |       |      | 1374 |      |      |      |      |      |      |
| 1327 | 1326 |       |      | 1326 |      |      |      |      |      |      |
| 1308 | 1309 |       |      | 1309/5 |      |      |      |      |      |      |
| 1265 | 1266 |       |      | 1265| 1264 | Amide III, C-H s, N–H ib, C=O ib |      |      |      |      |
| 1224 | 1236 |       |      | 1235/9 |      |      |      |      |      |      |
| 1202 | 1204 |       |      | 1204| 1203 | Amide III, C=O-H d |      |      |      |      |
| 1147 | 1154 |       |      | 1150 |      |      |      |      |      |      |
| 1111/04 | 1113 | 1110 | 1108 | 1110 | 1110 | C–O–C s |      |      |      |      |
| 1005 |       |      |      |      |      |      |      |      |      |      |
| 1010 | 1011 |       |      | 1010 |      |      |      |      |      |      |
| 971* | 974* |       |      | 974/3 |      |      |      |      |      |      |
| 953 | 952 | 953 | 949 | 952/1 | 950/49 | 953 | 950 | PO\(_4^{3–}\) s |      |      |
| 915 | 919 | 925 |       |      |      |      |      |      |      |      |
| 895 | 895 | 902 | 902 | 902/3 | 899/7 | 897 | 898 | CH\(_4\) d, C–O–C glycosidic bond |      |      |
| 746 | 760* |       |      | 761/4 |      |      |      |      |      |      |
| 710 | 701 | 709 | 708 | 708/2 | 708 | 702* | C–O op, C–H op, N–H d |      |      |
| 692 | 687 | 692* | 689/91 | 690* | 692 | C–O op, C–H op, N–H d |      |      |
| 648 | 634 | 648 | 633/16 | 642/4 | 635 | 635 | C–C t, C–O b, C–H b |      |      |
| 598 | 598* |       |      | 597/601 | 592 | 581 | C–C t, C–O b, C–H b, PO\(_4^{3–}\) s |      |      |
| 563 | 568 | 568/9 |       |      |      |      |      |      |      |      |
| 530 | 537 | 528/6 |       |      |      |      |      |      |      |      |
| 496 | 504 | 505 |       |      |      |      |      |      |      |      |
| 450/5 | 455 | 458/5 |       |      |      |      |      |      |      |      |

*Shoulder, s = stretching, ass = asymmetrical stretching, ss = symmetrical stretching, b = bending, ip = in-plane-bending, ab = anti-symmetric bending, w = wagging, sb = symmetric bending, d = deformation, t = twisting, r = rocking, op = out-of-plane bending. Band assignments were carried out using data from the literature [1, 20, 47–52].
(SI Fig. 2) has some ubiquitous features with that of standard \(\alpha\)-chitin, but the vibrational bands have different area percentages (SI Table 1). The rocking CH\(_2\) vibrations at 1376 cm\(^{-1}\) and the amide III bands in the 1202–1327 cm\(^{-1}\) range are predominantly assigned to organic biopolymers (Fig. 1). The CH deformation of the \(\beta\)-glycosidic bond band at 897 cm\(^{-1}\) for \(D.\) tenuis and at \(\sim\) 902 cm\(^{-1}\) for \(L.\) anatina is present at 895 cm\(^{-1}\) in standard \(\alpha\)-chitin and at 902 cm\(^{-1}\) in \(\beta\)-chitin. Bands at 1003 cm\(^{-1}\) in \(L.\) anatina and 1005 cm\(^{-1}\) in \(D.\) tenuis are not typically present in standard \(\alpha\)-chitin and \(\beta\)-chitin, and are attributed to the aromatic C–C and/or C–O stretch of fibrous insoluble protein [1, 53]. A similar \(\beta\)-chitin peak was reported from diatom species \(Thalassiosira\) rotula [47] and \(Riftia\) pachyptila tubeworm [52]. The vibrational bands at \(\sim\) 856 cm\(^{-1}\) and \(\sim\) 820 cm\(^{-1}\) are assigned to a combination of C–C and C–O stretching, and appear in \(L.\) anatina only. Ca-phosphate vibrational bands at 964 cm\(^{-1}\), 608 cm\(^{-1}\) and 581 cm\(^{-1}\) are only present in the \(D.\) tenuis spectrum (Fig. 1; SI Fig. 3). Identification of Ca-phosphate components was most likely due to the fact that both Ca-phosphate and the framework of organic biopolymers form complexes to produce shell ultrastructure.

FTIR spectroscopy was used to probe and/or identify the molecular and functional properties of the organic fibres from the shell samples, and for comparison with the \(\alpha\)-chitin standard and squid pen \(\beta\)-chitin (Fig. 2). The amide I of the insoluble matrix of \(L.\) anatina shows a broad band at 1630–1635 cm\(^{-1}\), similar to \(\beta\)-chitin. However, the spectrum of \(D.\) tenuis organic matrix reveals a broad band with a partial splitting pattern in two bands at 1652 cm\(^{-1}\) and 1634 cm\(^{-1}\), while standard \(\alpha\)-chitin has two bands at 1656 cm\(^{-1}\) and 1621 cm\(^{-1}\). The band at 1550 cm\(^{-1}\) is assigned to amide II, while bands in the 1203–1308 cm\(^{-1}\) range are attributed to amide III. The stretching C-O bands in the 1153–950 cm\(^{-1}\) range (Fig. 2) are indicative of chitin. The fingerprint region of the organic fibres of the phosphate-shelled brachiopods are doublet bands at 702–708 cm\(^{-1}\) and 691–692 cm\(^{-1}\), suggesting crystalline chitin. The stretching band of the OH group for shell samples is centred at \(\sim\) 3424 cm\(^{-1}\), with a shoulder near 3532 cm\(^{-1}\). Other spectra bands for amide A (3282–3272 cm\(^{-1}\)), amide B (3106–3095 cm\(^{-1}\)), and the C-H stretching regions in the 2952–2851 cm\(^{-1}\) range are all characteristics of crystalline chitin and/or a protein motif. The bands and functional groups are assigned in detail in Table 1.

![Figure 2](image-url)  
**Figure 2** FTIR spectra of chitin fibre from brachiopod shells: Recent *Lingula anatina* (RL), fresh *Lingula anatina* (FL), recent *Discinisca tenuis* (DT), and pure \(\beta\)-chitin (BC) and \(\alpha\)-chitin (\(\alpha\)C) as a reference. a Shows a larger wavenumber range (4000–600 cm\(^{-1}\)) than the expanded range (1800–600 cm\(^{-1}\)) in (b).

**Biochemical analyses of chitin matrix**

Organic fibres from the shell samples were stained with the blue fluorescent stain Calcofluor White, also known as Fluorescent Brightener M2R (Fig. 3; SI Fig. 4). Calcofluor White forms hydrogen bonds with \(\beta\)-(1–3)- and \(\beta\)-(1–4)-linked polysaccharides [20],
enabling identification and visualization of the chitin matrix [15].

When the shell samples were treated with *Streptomyces griseus* chitinase, no clear evidence of chitinase activity was apparent under a light microscope. However, SEM imaging showed that the chitin-bearing fibres of the lingulid shells were damaged and disorganized (Fig. 4). Unlike the control samples (SI Fig. 5), chitinase targeted the chitin in the shells, leading to degradation and perforation. There was no distinctive difference between the isolated organic fibres and the NaOH-treated samples (not shown).

**Morphology of the organic fibre extracts**

Surface imaging of the organic fibres shows ultrastructural features of biopolymers, although the fibre morphology varies from shell to shell (Fig. 5). Organic fibres in *L. anatina* contain pores between the fibres (Fig. 5a), and closer imaging of the radiating units of the ridges reveals organic granules (Fig. 5a'). The organic fibre extract from *D. tenuis* shows lamellar units (Fig. 5b) that are comparable with the rheomorphic folding discussed by previous authors [21]. This suggests that the polymeric structure of the polysaccharide is preserved after acid (demineralization) and sodium hydroxide (deproteination) treatments. A polymeric structure as rheomorphic...
folding is observed in D. tenuis (Fig. 5b’), unlike the fibrous structure found in the organic fibre extract of L. anatina.

Structural imaging of brachiopod shells after demineralization and deproteination

In the shells of L. anatina, there are apatite-rich lamellae and organophosphatic-rich lamellae (Fig. 6a, b). The thickness of the apatite-rich layers is up to 840 nm, whereas the thickness of the organophosphatic-rich lamellae is up to 1.5 μm (Fig. 6a). In the organophosphatic lamellae, the organic sheet laminations are parallel to the Ca-phosphate sub-lamellae (Fig. 6b), but do not penetrate into the apatite-rich lamellae which are approximately perpendicular (Fig. 6b). Within the apatite-rich laminae of L. anatina, there are nanoscale fibrils that are mineralized with Ca-phosphate nanoparticles (~ 50–70 nm) to form a fibrous biocomposite (Fig. 6c). FEG-SEM imaging of L. anatina reveals composite organic sheets with Ca-phosphate granules (SI Fig. 6a). The organic sheets or lamellae coated with nanoparticle granules (in SI Fig. 6a) represent several layers of organic sheets that were peeled off the shells for the characterization of the chitin matrix (see experimental). Some biomineralized nanofibrils are arranged approximately perpendicular to the micro-laminations (SI Fig. 6a).

After demineralization and deproteination, the shell of D. tenuis reveals only one major and continuous compact layer that is penetrated by canals with diameters of ~ 90–120 nm (Fig. 6d). This conspicuous sets of canals consists of various aggregates of organic-coated granules in the 50–100 nm range (Fig. 6e; SI Fig. 6b). As shown in Fig. 6f, the shell of D. tenuis is composed of ~ 50–60 nm spherical Ca-phosphate nanoparticles, with the aggregates of organic-coated granules naturally compacted to form mosaics (see Fig. 6f), similar to a previous study[21]. Rod-shaped crystallites (Fig. 6f) may perhaps be self-organized chitin fibrils, which are coated with proteins before mineralization. The apatitic components entwine with the organic fibrils (Fig. 6e, white arrows), resulting in a stratiform succession of microscale laminae [42, 44].
Thermal gravimetric and differential thermal gravimetric analyses

TGA and DTG analyses of brachiopod shell samples, standard α-chitin, and standard β-chitin were carried out to detect degradation features in the organic matrix. TGA weight loss and multistage decomposition steps occur for the brachiopod shells, α-chitin, and β-chitin (Fig. 7). The first weight loss of 5.1 wt% to 5.6 wt% for both L. anatina and D. tenuis occurs over the 30–200 °C range, and is attributed to removal of water. The second weight loss of 26.6 wt% for L. anatina and 12.9 wt% for D. tenuis (Fig. 7a) is recorded in the 200–650 °C range and is interpreted to be due to the combustion of a complex mixture of shell macromolecules, including chitin fibre and a protein moiety [54, 55], that is occluded in the apatitic shells. The maximum DTG value for the commercially obtained α-chitin is recorded at 357 °C, whereas the peak shift occurs at 335 °C for L. anatina and 347 °C for D. tenuis, comparable with the squid pen β-chitin standard at 341 °C (Fig. 7b). These maximum DTG values for the shells are in the 250–350 °C range for thermal stability of β-chitin [54, 55]. Lower temperature peaks are considered to be a measure of lower thermal stability [54]. However, the DTG thermogram of L. anatina is similar to α-chitin, and shows a broad peak at around 600 °C (Fig. 7b), which is attributed to the thermal degradation of a pyranose ring and the decomposition of residual carbon [54]. DTG thermograms of the shells reveal apatitic peaks at 840 °C for L. anatina and 740 °C for D. tenuis. The residual mass after heating to 900 °C is 65.9 wt% for L. anatina and 79.6 wt% for D. tenuis. This suggests the presence of inorganic constituents that were not extracted during the acid-treatment stage.

Discussion

The presented data provide a much better characterization of organic fibres in two organophosphatic brachiopod shells that have well established lineages in the fossil record, compared to previous studies [41]. After purification, the thermal degradation results confirm the presence of a chitin matrix in the shells of two representative taxa belonging to separate superfamilies of lingulid organophosphatic brachiopods, L. anatina (Linguloidea) and D. tenuis (Discinoidea). The L. anatina shells are composed of twofold more abundant chitin matrix compared to the D. tenuis shells. Chitin is rarely found in a pure state in nature, and is most commonly linked covalently with proteins [13, 56]. Pure chitin can be difficult to isolate, for example in mollusc shells [4, 19] and brachiopod shells [41]. Nevertheless, modern chitin staining techniques, such as the use of the fluorochrome Calcofluor White which binds to D-glucopyranose polysaccharides enable identification of chitin [15, 22], see Fig. 3. Calcofluor White is widely used for the rapid identification of chitin in chitin-containing organisms, i.e. fungi [15, 57], chytrid parasites on phytoplanktonic eukaryotes and prokaryotes [58], and in the shells of molluscs [20, 22, 59].
In this study, chitinase perforated and/or degraded the studied samples, thus confirming the presence of a chitin matrix in the shells of brachiopods. In the shell of *L. anatina*, the chitin layer was not readily degraded by chitinase [39]. Chitin matrix is covalently linked with other biomolecules, or may be masked by other organic matrices such as glycosaminoglycans and extracellular matrix [1, 39, 42]. Also, when chitin forms complexes with other biomolecules such as melanin, bromotyrosine, and/or inorganic components such as calcium carbonate, chitinase has little or no effect on such chitin-containing biomaterials using light and fluorescent microscope [15, 60]. Still, after chitinase activity, the SEM images (Fig. 4) are comparable to the SEM images of fossilized brachiopod shell structures from the lower Cambrian [61]. The basic effect of chitinase in this study extends our understanding concerning the evolution of biomineral formation in phosphatic small shelly fossils [61, 62].

Chitin can be readily distinguished using vibrational spectroscopy, such as by Raman and FTIR spectroscopy, due to its distinctive bands [1, 47–51]. The analyses presented here using vibrational spectroscopy reveal the biochemical composition of brachiopod shells, with chitin fibres entwined with minor shell proteins, supporting previous work [41]. The spectral bands of the organic fibres compare closely with the chitinous system containing aminopolysaccharide chitin (Figs. 1 and 2). The FTIR and Raman spectra bands associated with the amide I vibrational modes reveal broad and complex bands that are centred at high frequencies and are attributed to β-chitin. The Raman band is centred at ~1655 cm⁻¹ for brachiopod shells, and is commonly assigned to stretching of the C=O group hydrogen bonded to the N–H group of the neighbouring intra-sheet chain [49]. In the FTIR spectra, the amide I position is at 1652 cm⁻¹ for *D. tenuis*. The band shifts by ~19 ± 3 cm⁻¹ towards a lower frequency for *L. anatina* (Table 1). The FTIR spectra 1634 cm⁻¹ band for *D. tenuis* and a 1646 cm⁻¹ band for *L. anatina* are present as a shoulder, and are likely due to defects in the crystalline structure of the chitin fibre, or possibly a shell-associated protein. The deconvolutional fit of the amide I band for the FTIR spectra in the 1580–1720 cm⁻¹ range also reveals a band at ~1622 cm⁻¹, compared with the α-chitin band at 1620 cm⁻¹ (SI Fig. 7). The occurrence of this band may show a specific hydrogen bond of carbonyl (C=O) with the hydroxymethyl group of the next chitin residue of the same chain, or may be due to the presence of the enol form of the amide moiety [49, 63, 64]. A ~1003 cm⁻¹ vibrational ring peak is visible in the Raman spectra of the brachiopod shells, and is commonly observed in the published spectra of chitin [47, 52] and collagen [53, 65]. Recent analyses provide comprehensive data on *D. tenuis* and *L. anatina* shells, with identification of a collagen matrix as the primary protein motif within the inorganic matrix of phosphate-shelled brachiopods [1].

Total organic macromolecules in the shells of *D. tenuis* is 24.6 wt.% and for *L. anatina* is 40.6 wt.% [1]. After demineralization and deproteination, the amount of organic macromolecules reduced to 12.9 wt.% for *D. tenuis* and 26.6 wt.% for *L. anatina*. The extraction methods and calculation from TGA reveal the presence of phosphatic soluble organic macromolecules in *L. anatina* (14.0 wt.%) and *D. tenuis* (11.7 wt.%). Based on the previous work of Lévéque et al. [66], there may be soluble organic matrices in Ca-phosphate shells that greatly contribute and/or influence the nucleation and growth of brachiopod shells. The inorganic matrix provides the structural strength, stiffness and rigidity for many invertebrate shells, including organophosphatic brachiopod shells [67]. However, polysaccharide chitin offers increased strength and flexibility [67, 68].

*L. anatina* is clearly laminated with an alternating chitinous system and biomineralized chitin-phosphate layers. In contrast, lamination of the mineralization framework is less pronounced in the sessile epibenthic *D. tenuis* [67, 68]. Biomineralized laminae provide strength to the shells of *L. anatina*, and allow bending flexibility [68], a factor that may be important in the complex shell-first burrowing of living lingulids [27, 32–34]. However, flexibility is less important for the shallow marine epibenthic phosphate-shelled brachiopods [67]. Further, *L. anatina* shells contain an organic primary layer that protects the secondary layer from incipient cracks that can occur during the movement of the animal through the sediment [42, 43]. The flexible secondary layer of *L. anatina* consists of stacks of alternating mineralized and non-mineralized β-chitin sheets [42], see Fig. 6. Chitin chains lie parallel with the c-axes of the apatite components, and chitin fibrils are usually coated or covered with spherical and/or elongated shapes of Ca-phosphate particles [1, 39, 42]. The organized structure of the *L. anatina* shell helps divert fractures,
and thus provides high flexibility while the animal burrows itself into and through the sediment [67, 68].

The analyses reported here reveal the architectural structures and polysaccharide matrix of two important phosphate-shelled brachiopods and demonstrate further a practical bioanalytical approach for analysing lophotrochozoan shells.

Conclusions

1. Based on FEG-SEM, vibrational spectroscopic analyses, and biochemical studies, phosphate-shelled brachiopods (*Lingula anatina* and *Discinisca tenuis*) contain chitin fibre composites with a fibrous protein concentration, which create suitable microenvironments for inorganic nanoparticle aggregation.

2. The polysaccharide-apatitic matrix composition of the shells of *L. anatina* is distinctly different from that of *D. tenuis*. For *L. anatina*, the polysaccharide fibre component is internally reinforced by a nanoscale Ca-phosphate component and is distinctively laminated. *D. tenuis* used a different shell architecture, because it contains a continuous compact layer with canals and a stratiform succession of microscale laminae.

3. This study highlights the differences in the shell ultrastructure of *L. anatina* and *D. tenuis*. *L. anatina* has a distinctive two-layer structure with apatite-rich lamellae and organophosphatic-rich lamellae. *D. tenuis* has a major and continuous compact layer, with organic fibrils entwined or intercalated with a Ca-phosphate mineral. However, based on the extraction methods and calculation from TGA, the amount of soluble shell macromolecules is similar, 14.0 wt.% for *L. anatina* and 11.7 wt.% for *D. tenuis*.

4. It is proposed that the high polysaccharide chitin matrix content of *L. anatina* may contribute to its complex burrowing mechanism, in which the valves function as flexible digging tools, combined with the injection of water in a shell-first burrowing action that produces U-shaped burrows. The shell of *L. anatina* is composed of chitin sheets or lamellae, forming a laminar shell design that produces a flexible shell. Taken together, a flexible shell gives an advantage for burrowing and prompt movements in the burrow [26, 27, 32–35].

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Authors’ contributions

OBAA designed the study, carried out the analyses, interpreted the data, and prepared the original manuscript. YL and ZZ provided fresh (live) specimens of *Lingula anatina*; LEH provided recently alive *Discinisca tenuis* shells, and GAB provided recently alive *Lingula anatina* shells. KCD carried out the fluorescence imaging. LEH, GAB and SCG coordinated the study. All authors reviewed the manuscript and gave final approval for publication.

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Declarations

Conflict of interest The authors declare no competing interests.

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