Marginal marine agglutinated foraminifera: affinities for mineral phases

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ABSTRACT – The major agglutinated constituents in test material of marginal marine foraminifera are identified as α-quartz and clay particles using complementary spectroscopic techniques. Electron dispersive scattering analysis, micro-laser Raman and Fourier transform infrared (FTIR) spectroscopic techniques revealed detail about elemental and mineral polymorph constituents in test walls. Additionally, FTIR identifies the existence of organic cements and lining materials in wall structures. Micro-laser Raman specifically characterized the titanium oxide mineral, anatase, as a distinctive fraction of agglutinate in Ammobaculites balkwilli Haynes. The mineral represents ~10% of the test material, but comprises a minor component of the sediment and is identified in sediments only after heavy mineral separation. The enhanced concentration of anatase in the test of A. balkwilli suggests that there is a preferential selection for anatase. This provides further evidence that certain foraminifera can select grains specifically, which implies that there exists a selective mechanism and interaction between the organic (secreted) phases in the test walls and inorganic (grain surface) materials. J. Micropalaeontol. 18(2): 183–191, December 1999.

INTRODUCTION

Few biochemical and mineral microprobe investigations have been completed on agglutinated foraminifera, although previous microprobe studies have successfully revealed information about the mineral content and the nature of mineral polymorphs present in test walls (Towe, 1977; Lefurgey, 1978; Véneç-Peyré & Jaeschke-Boyer 1978, 1979; Weiner & Erez, 1985; Gooday et al., 1994; Roberts & Murray, 1995). Accounts of test morphotypes and features in agglutinated foraminifera prior to the introduction of electron dispersive scattering (EDS) analysis, micro-laser Raman and Fourier transform infrared (FTIR) microprobes, were unable to provide precise information about the components in wall structures. The development of more sophisticated analytical techniques provides high spatial resolution to ~2 μm, allowing the identification of the precise inorganic polymorphs used and secreted by foraminifera in test construction (Roberts & Murray, 1995). These data are of crucial importance as the microstructural nature of wall structures – for example, the cement morphology – continues to be one of the most important classification characteristics (Bender & Hemleben, 1988; Hohenegger, 1990). The acquisition of more precise geochemical data further enhances the information already available.

A combination of the above techniques was used to analyse the nature and composition of test components of Ammobaculites balkwilli Haynes, Trochammina inflata (Montagu), Jadammina macrescens (Brady) and Miliammina fusca (Brady), agglutinated foraminifera from marginal marine areas near Southampton, UK. The three spectroscopic techniques applied vary in the way that they interact with sample surfaces, enabling the acquisition of a variety of data (Fig. 1). These analytical techniques allowed the identification of the nature of inorganic and organic components in the test walls of foraminifera. Such data may reveal whether the foraminifera are specifically selecting material from their immediate environment. A detailed description of the organic component analysis is given in Allen et al. (in press). This paper characterizes material in the agglutinated walls and discusses evidence for specific particle selection on the basis of mineralogy by foraminifera.

ANALYTICAL PROCEDURES

Surface sediment samples were collected from marginal marine environments (marsh and intertidal mudflats) at Swanwick and Warsash near Southampton, washed on a 63 μm sieve, and dried at 50°C.

For scanning electron microscopy (SEM), the specimens were mounted onto aluminium stubs and coated with carbon before introduction into the vacuum chamber of the Jeol JSM-6400 scanning electron microscope with a Trarcor Series II energy dispersive X-ray analysis system. Photographs were taken so that a point correlation with EDS analysis could be made. Semi-quantitative point analysis was performed on individual grains and the results collected.

Raman spectroscopy was carried out with a Jobin-Yvon S3000 microprobe using an Olympus BH-2 microscope fitted with a Nikon x80 objective. The 514 nm line of an Ar ion laser (coherent Innova Model 70) was used with a source power of 2–4 mW. Selected specimens were placed loose on a glass slide and analysed using the Raman laser microprobe in the 100–1000 cm⁻¹ region of the spectrum. In this study, integration times of 10–300 s with a single accumulation were used with each analysis taking about 6 min to complete. Calibration was achieved through monitoring plasma lines from the laser and through the use of quartz and sulphur standards. The spectra were processed and manipulated with ISA PRISM software. The peak position was determined at maximum intensity and the line-width measured at half-maximum intensity (Roberts et al., 1995).

Fourier transform infrared analysis was performed with a Nicolet 460 Protégé, magna IR technology FTIR microprobe with a Nic-plan IR microscope. The data were acquired and manipulated with Omnic software. The IR bench conditions were as follows throughout the analysis: acquisition time 200 s, band resolution 4 cm⁻¹ and gain 2–4. Background subtraction
Noise levels were monitored throughout the experiments and were found to be very low. Sections of foraminifera test were placed on a salt window for analysis by FTIR.

X-Ray diffraction (Philips) analyses were performed on sediments collected from the same area as the foraminifera. A 1.00 mass of bulk sediment was ground and the resultant powder was scanned from 2 to 70° 2θ at a scanning velocity of 1.2°/min using Ni-filtered CoKα radiation. Peaks were identified using the associated computer software. Additionally, heavy minerals and clays were separated from the sediment fraction and analysed under the same conditions.

RESULTS

Test coloration of selected agglutinated foraminifera was initially examined using a binocular microscope. _Jadammina macrescens_ and _T. inflata_ have tests which are light grey to brown in colour. Younger chambers can often be seen to be lighter in coloration compared with the rest of the test. _Miliammina fusca_ has a white to brown test which appears to be finely agglutinated and _A. balkwilli_ has a test which consists of angular, medium to coarse, white and dark grains.

Scanning electron microscopy

Scanning electron microscopy of the foraminifera indicates that all the specimens selected have detrital grains as the main constituent of the wall structure. A fibrous cement phase, holding the grains together, is observed at high magnifications. _Jadammina macrescens, T. inflata_ and _M. fusca_ have abundant cement interstitial to the detrital grains, whereas _A. balkwilli_ has coarser grains with a lower proportion of cement (Plate 1). _T. inflata_ and _J. macrescens_ have very similar wall morphology, with outer and inner linings surrounding the test wall. Detrital grains are fairly angular, occasionally tabular. The latter are commonly orientated with the long axis parallel to the test linings and often produce grain imprints in the outer lining (Plate 1, figs 1, 2, 5, 6). _M. fusca_ has abundant medium clay/silt-sized grains, with additional epipelic diatom frustules. The test has no clear outer lining, rather a dense layer of organic material coating the grains on the outer surfaces (Plate 1, figs 3, 7). Detrital grains in the test walls of _T. inflata, J. macrescens_ and _M. fusca_ are approximately <1–4 μm in size and are representative of the clay- to silt-sized portion of the sediment, whereas specimens of _A. balkwilli_ have coarser grains in the size range 10–50 μm, which are representative of the silt–sand-sized portion of the sediment (Plate 1, figs 4, 8).
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Explanation of Plate 1.
Scanning electron photo micrographs of the four marginal marine foraminifera investigated by the microprobe techniques. Test morphology and wall sections of *Jadammina macrescens* (figs 1, 5), *Trochammina inflata* (figs 2, 6), *Miliammina fusca* (figs 3, 7) and *Ammobaculites balkwilli* (figs 4, 8).

The elemental compositions of the test walls were acquired through EDS analysis. *T. inflata, J. macrescens* and *M. fusca* show spectra indicating that quartz is the main detrital grain held in the test (Fig. 2A). In the spectral analysis, the additional presence of Al, along with lower proportions of Mg and Na, suggests that feldspars and clays may also be present as the levels of Mg, Na and Al are in excess of that reported from quartz analysis (Deer *et al.*, 1966) (Fig. 2B). *A. balkwilli* shows similar spectra, indicating the presence of quartz with minor aluminosilicates, i.e. clays and feldspar particles. Certain spectra of *A. balkwilli* indicate that Ti oxides, possibly anatase and rutile, are present in the test (Fig. 2C).

**Raman spectroscopy**
Results from the Raman laser microprobe analysis are outlined in Fig. 3. As a result of the presence of a brown layer on the outside of the tests (thought to be an Fe-rich coating), the specimens were initially difficult to analyse due to the high...
levels of fluorescence. Short acquisition times were used (120 s), with the spectra showing intense bands in the lower wavenumber region. *J. macrescens*, *M. fusca* and *A. balkwilli* show spectra confirming α-quartz as the main constituent of their agglutinated material with characteristic bands at 128, 393 and 464 cm⁻¹.

Fig. 2. Spectra obtained from electron dispersive scattering analysis on agglutinated foraminifera shells. Spectrum A is representative of the analysis of quartz grains (SiO₂) from test agglutinate. Spectrum B represents a mixed elemental composition arising from aluminosilicate mineral phases. Dark grains held within the test of *A. balkwilli*, represented by spectrum C, have TiO₂ compositions, indicative of the presence of rutile or anatase.

Fig. 3. Combined Raman spectra (100–1000 cm⁻¹) obtained from analysis of all agglutinated foraminifera shells. The intensity of the 150 cm⁻¹ peak in *J. macrescens*, *M. fusca*, is significant and occurs in spectra obtained from clay minerals due to the vibration of the AlO₆ groups in kaolinite. The *A. balkwilli* spectrum also has a strong absorption at 150 cm⁻¹ arising from the bending of O–Ti–O structures, and others at 197, 400, 515 and 640 cm⁻¹, representative bands from the analysis of anatase (Frost, 1997). Bands at 128, 379 and 464 cm⁻¹ visible in spectra are representative of quartz.

An intense band at ≈150 cm⁻¹ is observed in certain Raman spectra from *J. macrescens* and *M. fusca* which is in addition to the bands consistent with α-quartz. Kaolinite spectra typically show bands at 150, 461, 512 and 637 cm⁻¹ (Johnston et al., 1985). The 150 cm⁻¹ band is characteristic for the symmetrical bend of the AlO₆ group in kaolinite (Frost, 1997), indicating that this clay mineral is a constituent of agglutinated material, correlating with the EDS spectral results with mixed aluminosilicate composition (Frost, 1997; Johnston et al., 1985). Certain *T. inflata* spectra also have medium-intense absorption at ≈150 cm⁻¹, with additional vibrational bands at 175, 219, 289 and 488 cm⁻¹, although other spectra possess absorptions at 92, 456 and 841 cm⁻¹. The spectra are difficult to characterize as the signal appears to be highly mixed, despite a collection spot size average of ≈2 μm. The spectral bands identified in *T. inflata* spectra are not directly correlatable with published pure mineral values, indicating that a mixed suite of minerals may be present in the agglutinate. Certain bands are comparable with those in montmorillonite (450/467, 521, 621, 843; Johnston et al., 1985) and mica (100, 160, 195, 220, 240 cm⁻¹; Loh, 1973), indicating the potential presence of these minerals.

*Ammobaculites balkwilli* spectra are different as an intense absorption at ≈150 cm⁻¹ occurs with others at 128, 201, 397, 464, 518 and 637 cm⁻¹. Bands at 128, 397 and 464 cm⁻¹ suggest that α-quartz and kaolinite are present as detrital material (128 and 464 cm⁻¹), agreeing with EDS analysis determining grains with mixed aluminosilicate composition. Analysis of darker grains with laser Raman acquired spectra with an intense peak at c. 150 cm⁻¹, attributed to the O–Ti–O symmetrical bend (Frost, 1997), with additional low intensity bands at 197, 400,
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Fig. 4. Combined Fourier transform infrared spectra (400–4000 cm⁻¹) obtained from the analysis of all agglutinated foraminifera shells. Significant bands are evident in the regions around 700 cm⁻¹ (Si-O-Si, Si-O-Al and Ti-O molecular vibrations) and 800–1000 cm⁻¹, representative of silicate and aluminosilicate minerals with Si-O and Si-O-Al stretches. Overtone absorptions from quartz (Shimoda & Aikawa 1994) are identified at 1796, 1883 and 1991 cm⁻¹. Absorptions in the regions 1500–1600 and 2800–3000 cm⁻¹ arise from the vibration of organic materials in the test wall. The broad absorption between 3000 and 3800 cm⁻¹ is produced from fundamental OH-stretching vibrations in inorganic and organic materials (Graetsch et al., 1985), and N-H bond stretches from organic materials (Kokot et al., 1994).

515 and 640 cm⁻¹, characteristic of the mineral anatase (Beattie & Gibson, 1968). This correlates with EDS analysis indicating the presence TiO₂ mineral phases.

To summarize, the laser Raman microprobe has confirmed the presence of α-quartz and clays (kaolinite) in the test agglutinate of the studied foraminifera. The laser Raman microprobe additionally confirms anatase as a TiO₂ component recognized by EDS analysis of the test walls of A. balkwilli.

Fourier transform infrared spectroscopy

Results from the FTIR microprobe analysis are outlined in Fig. 4. Absorptions occur in specific regions of the acquired spectra due to the characteristic stretching and bending of particular molecular moieties. Quartz, clay (kaolinite) and anatase were identified as agglutinating materials with EDS and laser Raman microprobe analysis and thus should be detectable with the FTIR microprobe, although the analytical window for FTIR analysis is slightly larger, lowering the spatial resolution of the microprobe technique.

Trochammina inflata, J. macrescens, M. fusca and A. balkwilli show similar FTIR spectra for agglutinated grains. Absorptions identified at 700 and 798 cm⁻¹ correspond with Si-O-Si bond stretching vibrations for crystalline quartz (700, 785 and 798 cm⁻¹) (Farmer, 1974; Frölich, 1989). The strongest absorptions in all the spectra are in the region 995–1140 cm⁻¹, varying slightly in bandwidth and position. In the analysis of quartz, this broad absorption is regarded as the envelope of several poorly resolved bands assignable to Si-O bond stretches (Frölich, 1989). The variation in band frequencies can be influenced by variations of crystal alignment and orientation, and additionally by composition (Farmer, 1974). The 1796, 1883 and 1991 cm⁻¹ absorptions in T. inflata, J. macrescens and M. fusca spectra, and 1796, 1875 and 1989 cm⁻¹ in A. balkwilli spectra are additionally characteristic of overtone vibrational bands intrinsic to the α, β-, and γ- spectra of quartz (Shimoda & Aikawa, 1994).

Table 1. Reported FTIR spectral band absorbances from proteinaceous materials identifying secondary structures and also specific amino acids.

| Secondary structures | Le Gal et al. (1991) | Scórko-Glonek et al. (1995) | Trouillet et al. (1996) |
|----------------------|---------------------|----------------------------|------------------------|
| Turns + bends          | 1694               | 1691                       | 1691                   |
| Beta                  | 1688               | 1682                       | 1682                   |
| Turns + bends          | 1683               | 1673                       | 1670                   |
| Beta sheet parallel    | 1675 ± 4           | 1665                       | 1660                   |
| Turns and bends        | 1670 ± 2           | 1654 ± 3                   | 1653                   |
| Turn, alpha            | 1663 ± 4           | 1637 ± 3                   | 1635                   |
| Alpha helix            | 1645 ± 4           | 1624 ± 4                   | 1628                   |
| Alpha, random coil     | 1643               | 1625                       | 1625                   |
| Beta sheet antiparallel| 1635               | 1612                       | 1612                   |
| Beta sheet             | 1635               | 1605                       | 1605                   |
| Tyrosine               | 1605               | 1545                       | 1532                   |

The influence of Al-O, Si-O-Al and Ti-O bond vibrations arising from aluminosilicate phases (kaolinite) and titanium oxides (anatase) are also represented in the spectra. Absorption
bands between 700 and 800 cm\(^{-1}\) are enhanced by the stretching vibration of Al-O bonds (around 770 and \(\approx 729-694 \text{ cm}^{-1}\)) (Handke & Mozgawa, 1993) and Ti-O bond vibrations (800–650 cm\(^{-1}\)) (Farmer, 1974) causing band frequency variation around those identified for quartz. Both Si-O-Si and Si-O-Al bridges are present in aluminosilicates and the presence of these bonds influences the region 950–1150 cm\(^{-1}\), producing absorption variation (frequency and intensity) in the foraminiferal spectra.

Fourier transform infrared analysis is ideal for the identification of organic materials, revealing additional information about the test composition (Allen et al., in press). FTIR analysis does not induce sample fluorescence, enabling highly organic materials to be analysed. Two broad absorptions are evident at 1545 and 1647 cm\(^{-1}\) in all the spectra. On closer investigation, smaller absorptions at 1617 and 1669 cm\(^{-1}\) can be detected in spectra from A. balkwilli and M. fusca. Proteinaceous substances in the test walls of the agglutinated foraminifera would cause absorptions to occur in these spectral regions. In all spectra, carbonyl (C=O, stretch) and N–H (bending) vibrations are identified at around 1650 cm\(^{-1}\) relating to the amide I unit, the main peptide link, of a protein (Kokot et al., 1994). Within the amide I region, small absorptions can identify features of the secondary structure of the protein i.e. \(\alpha\)-helices, \(\beta\)-sheets and turns. The intensities of such peaks may indicate the relative amounts of each type of secondary structure (Smith et al., 1994). Absorptions at 1617 and 1669 cm\(^{-1}\) may relate to the presence of tyrosine (a common amino acid) and beta turns/bends of the protein chain. The absorption at 1545 cm\(^{-1}\) in all the spectra arises from the C–N stretch and the N–H in-plane bending absorption in the amide 2 unit of the proteinaceous material (Kokot et al., 1994) (Fig. 5, Table 1). Organic aliphatic molecular moieties (CH\(_2\) and CH\(_3\)) produce spectral absorptions at 2936 and 2972 cm\(^{-1}\) visible in all the spectra from J. macrescens, T. infilata and M. fusca, and at 2858, 2875, 2926 and 2966 cm\(^{-1}\) in the spectra from A. balkwilli. The identification of organic molecular entities identifies organic cements and lining material in all the foraminifera. Bands characteristic of organic material also occur in the lower frequency region (800–1300 cm\(^{-1}\)) of the FTIR spectra, but the presence of inorganic material tends to mask these vibrations and hence prohibit further characterization.

All the spectra have a broad absorption from 3200–3700 cm\(^{-1}\), centring at 3400 cm\(^{-1}\), due to the fundamental OH-stretching vibrations of molecular water and SiOH (Graetsch et al., 1985). This region provides additional evidence that both proteins and carbohydrate components exist within foraminiferal lining materials. N–H bond stretches within proteinaceous components (Kokot et al., 1994) and vibrations arising from O–H bond stretches within carbohydrate units held within the structure will additionally form absorptions in this region.

To summarize, FTIR analysis in the investigation of test components confirms the presence of the major mineral polymorphs present in the agglutinate, but compared with the laser Raman microprobe technique, the spatial resolution was reduced due to the increase in the sampling window. FTIR analysis success fully reveals the nature of the organic cementation phase in all the tests, which the other techniques were not able to detect.

**X-Ray diffraction**

The major constituents of the sediment were found by XRD analysis to be predominantly quartz, together with kaolinite and a small amount of mica and calcite. Anatase was not detected in the bulk sediment analysis, but was detected as a minor component in the analysis of the heavy mineral fraction. The percentage of this mineral in bulk sediment analysis is presumably below the limit of detection and hence is less than 10% of the overall composition of the sediment. Clay fraction analysis of the sediment identified the main mineral type as kaolinite with additional mica. Anatase was not detected, suggesting that less than 10% is present in the clay fraction of the sediment.

**DISCUSSION**

Agglutinated foraminifera from marginal marine areas near Southampton agglutinate quartz and aluminosilicates such as kaolinite for the construction of their test walls. The use of anatase grains for agglutination by A. balkwilli is distinctive. The initial observation of dark grains in the A. balkwilli test was followed with EDS analysis identifying high levels of Ti; Raman analysis characterized the actual polymorph of TiO\(_2\) as anatase. None of the other foraminifera analysed select these grains for incorporation into their test, and hence it appears that A. balkwilli selects the anatase grains specifically. The operative forces of selection are poorly understood and, although most agglutinated foraminifera appear not to select, reports of grain selection have been published (Avnimelech, 1952; Hofker, 1953; Sliter, 1968; Salami, 1976; Jorgensen, 1977; Höttinger, 1986; Bender, 1989). Certain agglutinated foraminifera have been documented to preferentially select quartz particles in high carbonate (chalk) environments where the sediment contains only negligible amounts of quartz (Jorgensen, 1977). The use of anatase in A. balkwilli suggests that particle differentiation is taking place.

**Size selectivity**

Size differentiation might be a factor influencing the process of test construction. A. balkwilli utilizes particles of clay to silt size (4–50 \(\mu m\)) for test construction, whereas T. infilata and J. macrescens select grains of the clay-sized fraction (1–4 \(\mu m\)) from the sediment. Anatase was not detected by bulk XRD sedimentary analysis and hence is presumably present in only very small quantities in the sediment (<10%). However, on heavy mineral separation, anatase was detectable, but remained a minor mineral phase of the sediment compared with haematite and magnetite. Grains of anatase and quartz present in the test walls of A. balkwilli occur in the silt–sand, slightly coarser size fraction of the sediment, whereas J. macrescens and T. infilata do not include anatase in their agglutinated material. Anatase may simply not be available in smaller size categories, hence excluding the mineral from their test walls. Agglutinated foraminifera are reported to select agglutinated grains according to chamber size and can vary the grain distributions depending on the wall area (Avnimelech, 1952; Hofker, 1953; Slama, 1954; Hedley, 1964; Salami, 1976; Bowser & Bernhard, 1995).

**Grain density**

Heron-Allen & Earland (1920) noted that certain foraminifera
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select particles with higher specific gravities than sand grains. Experiments were later performed using minerals of varying specific gravity. The results indicated that no preference was made, leading to the conclusion that agglutinated foraminifera did not have the ability to detect variations in the specific gravity of minerals, although the reasoning behind the specific agglutination of dense minerals still remains unanswered (Salami, 1976). The specific gravity of anatase (3.9) is high compared with quartz and kaolinite (2.65 and 2.6, respectively); however, in an aqueous environment, the effect of a small percentage of heavy mineral in addition to other minerals in the test will make no significant difference to the overall specific gravity of the test, so there would be no advantage to infaunal foraminifera to agglutinate heavier particles.

Heron-Allen (1915) speculated that dense minerals would settle in the lower layers in a sediment profile after wave/current agitation. Depending on the movements of the foraminifera in the sediment, this could increase the probability of denser mineral agglutination. Perhaps *A. balkwilli* pervades these layers during movement through the sediment, enabling the agglutination of anatase along with quartz.

**Pseudopodial/secreted phase interaction**

Pseudopodia possess typical cell membranes (with a lipid bilayer, glycoprotein islands, surface glycoproteins and carbohydrate residues) which are the surfaces involved in sediment interactions (Fig. 6). The carbohydrate entities act as feelers to attach, adhere to substrates (Warren, 1987) and also become involved in cell-cell, cell-matrix recognition processes (Alberts *et al*., 1989). Secretory adhesive products that arise from pseudopodia are also suggested to be involved in prey capture, locomotion, and test construction (Stuart, 1866; Schaudinn, 1893; Sandon, 1934; Jepps, 1942; Nyholm, 1956; Buchanan & Hedley, 1960; Christiansen, 1971). The glycoprotein (glycosaminoglycan) secretions comprise the cement and lining materials present in foraminifera walls (Anderson & Bé, 1976; Langer, 1992). Glycoprotein entities will also be capable of interacting with similar surfaces as they possess functional groups with a high surface charge (Bowser *et al*., 1984). Pseudopodia are capable of interacting with external interfaces, enabling the collection of and adhesion to specific materials. Polystyrene microspheres of various sizes, with net positive, negative or neutral surface charges, with varying degrees of hydrophobicity, were used in an experiment to see how the materials interacted with foraminiferal pseudopodia. The spheres were bound to, and moved along, pseudopodial surfaces, whereas bound bacteria were phagocytosed. This suggests that the pseudopodia are not specific in the surface selection of agglutinate; however, the foraminifera were able to differentiate between mineral and bacterial surfaces (Bowser & Bloodgood, 1984; Bowser *et al*., 1984; Bowser & Rieder, 1985, 1986). Experiments highlighting electrostatic interactive mechanisms between bacterial membranes and quartz interfaces have been reported previously (Beveridge & Murray, 1980; Mera & Beveridge, 1993). Surfaces of quartz, aluminosilicates and TiO₂ materials have highly electrostatically charged regions which will attract sites with oppositely charged interfaces, such as organic foraminiferal materials, forming electrostatic and hydrogen bonds.

The results from the investigation indicate that the agglutinated foraminifera take advantage of materials available in the immediate surroundings, but that *A. balkwilli* is agglutinating materials that are not agglutinated by other foraminifera in the area. Additionally, *A. balkwilli* agglutinates silt-sized grains, whereas the other foraminifera agglutinate clay-sized particles, suggesting that a certain amount of selection may be taking place. Further investigations need to be carried out to determine which of the known mineral properties (size or chemotaxic properties) most influence selection.

**CONCLUSIONS**

This study investigated the nature of the tests of characteristic marginal marine agglutinated foraminifera. These forms definitely use particles of α-quartz and kaolinite as major construction materials in their test formation. This is the first clear microprobe identification of the clay mineral type used in the tests of agglutinated foraminifera. The sediment in the surrounding area is composed of similar constituents to those in the test, so it reasonably follows that they include the most frequently available material in test construction. The more surprising feature is the occurrence of the mineral anatase as a constituent of the material agglutinated by *A. balkwilli*. This mineral is not common in the sediment and was below the detection limit of bulk XRD analysis (<10%), but yet is detected as a minor phase on heavy mineral separation. Anatase is not detected in the clay-size fraction, suggesting that it is absent in this fraction or again exists in amounts below the levels.

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Fig. 6. Diagram of the type of interaction occurring between the glycoproteins or carbohydrates in secreted in cell membranes forming electrostatic or hydrogen bonds with the mineral surfaces.

Glycoprotein chain

Hydrogen bonding

Electrostatic attraction

![Diagram of the type of interaction occurring between the glycoproteins or carbohydrates in secreted in cell membranes forming electrostatic or hydrogen bonds with the mineral surfaces.](image-url)
of detection (<10%). A. balkwilli appears to preferentially select silt-sized particles compared with the agglutination of clay-sized particles by the other foraminifera. This suggests that the organisms are able to differentiate between fractions of sediment. Additionally, silicate surfaces, aluminosilicate surfaces and TiO2 surfaces all have very similar properties and will readily react, through electrostatic attraction or hydrogen bonding, with oppositely charged surfaces, such as pseudopodial membranes and glycosaminoglycan secretory products.

Further research involving the investigation of foraminiferal pseudopodial interactions between different sedimentary size categories and various electrostatically charged surfaces should be performed to determine the controlling factors in the process of agglutination.

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