The Volyn biota (Ukraine) – indications for 1.5 Ga old eukaryotes in 3D-preservation, a spotlight on the 'boring billion'

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- 20 Abstract

21 The Volyn biota, fossilized organisms with a minimum age of 1.5 Ga, were found in cavities in 22 granitic pegmatites from the Korosten pluton, NW Ukrainian shield. Fossilization was due to 23 influx of hydrothermal fluorine-rich waters, which silicified the outermost part of the 24 organisms, thus preserving the 3D morphology. Details of the morphology (investigated by 25 scanning electron microscopy) show that the majority of the specimens is filamentous, of a 26 large variety with diameters ranging from $\sim 10 \ \mu m$ to $\sim 200 \ \mu m$, thin filaments with typical 27 branching, thick filaments with ball-shaped outgrowths and dented surface. Filaments can be 28 straight or conical, curvilinear or strongly curved, up to mm in length, some with a central 29 channel. Some filaments show indications for segmentation, are grown as sessile organisms 30 onto substrate; others show both intact ends, indicating a non-sessile, free-living lifestyle. 31 Objects with flaky morphology and agglutinating filaments are interpreted as fossil biofilms. 32 Other objects are hollow and show a large variety of forms; spherical objects are scarce. Infrared 33 spectroscopy indicates the presence of chitosan in one filament type, electron microprobe 34 analysis of nm-sized inclusions in filaments identified the presence of Bi(Te,S) minerals, and

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both observations are compatible with the interpretation as fungi-like organisms. Stable C- and N-isotope data of bulk samples are in the range of -31 to -47 $\& \delta^{13}$ C, and of +3 to +10 $\& \delta^{15}$ N, indicating possible methanogens as part of the subsurface micro-ecosystem. The Volyn biota indicate that at 1.5 Ga complex forms of life existed in the continental deep biosphere, well above the microscopic level, including fungi-like organisms resembling eukaryotes.

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42 **1 Introduction**

43 Precambrian fossils are generally not well preserved because of the absence of skeletal parts. 44 In addition, most Precambrian fossil record is from sedimentary rocks with strong diagenetic 45 or even metamorphic overprint, which destroyed much of the original morphology and in 46 extreme cases of very old organisms left only an isotopic signature (e.g. Alleon et al., 2018; 47 Berbee et al., 2020). Therefore, their biogenicity is often disputed especially when the organic 48 matter (OM) is completely replaced, often by silica or pyrite. A preservation of 3D-morphology 49 is very rare and requires special fossilization conditions, which include first prevention of rapid 50 decay of the OM and then preservation of the space around the fossil in order to preserve its 51 original morphology. These conditions were fulfilled in pegmatites of the Volyn pegmatite 52 field, Ukraine, associated with the Korosten Pluton. These so-called 'chamber pegmatites' 53 contain large miarolitic cavities in which OM named (oxy)-kerite was found and in previous 54 investigations interpreted as an example of a-biogenic formation (Ginzburg et al., 1987; 55 Luk'yanova et al., 1992), later re-interpreted as fossil cyanobacteria (Gorlenko et al., 2000; 56 Zhmur, 2003) from a geyser type deposit. Ginzburg et al. (1987) give a composition of 60-76 wt% C, 5-7 wt% H, 9-23 wt% O, 8-9 wt% N, and 2-3 wt% S and an empirical formula of 57 58 C₄₉₁H₃₈₆O₈₇(S)N. Gorlenko et al. (2000) and Zhmur (2003) mention masses of up to 3 kg of 59 kerite in one of the cavities with an irregular distribution within the pegmatite.

60 The organisms lived in these cavities and provide an example of the Precambrian deep 61 biosphere. Their fossilization conditions included sudden influx of hot hydrothermal waters in 62 the geyser system, where magmatic fluids rich in SiF₄ mixed with meteoric waters (Franz et al., 2022a), infiltration of Si-Al into the outermost layer of the fossils, and formation of dominantly 63 64 clay mineral encrustations. The 1.76 Ga intrusion age of the pegmatites (U-Pb zircon; 65 Shumlyanksyy et al., 2021) provides a maximum age of the fossils; the minimum age of 1.5 Ga is provided by the age of formation of a breccia, which contains degraded OM, brown opal with 66 67 OM, buddingtonite which NH₄-content was provided by the degraded OM, and muscovite (⁴⁰Ar-³⁹Ar laser ablation data; Franz et al., 2022b). An additional argument for this age comes 68

from the brown opal, which cements the breccia and contains organic matter (Franz et al., 2017, and references therein). Therefore, we assume that breccia formation must have occurred when organic matter was already present in the 'chambers'. Although some of the miarolitic chambers collapsed, producing the muscovite-opal-bearing breccia, other chambers are still intact and were mined since the 1930ies for piezo quartz and until now for pegmatite minerals such as beryl and topaz (Ivanovich and Alekseevich, 2007; Lyckberg et al., 2009, 2019).

We report here details about the morphology and the internal structure of the fossils, investigated by scanning electron microscopy (SEM) and electron microprobe analysis (EMPA), and provide stable C-N isotope and infrared spectroscopy (FTIR) data, which allow speculating about the types of organisms. An important point is that these 'micro'-fossils in many cases reach a size well above the microscopic level, with filaments of several mm in length. The age of the fossils of 1.5 Ga in the middle of the 'boring billion' and gives insight into the organisms of the deep continental biosphere.

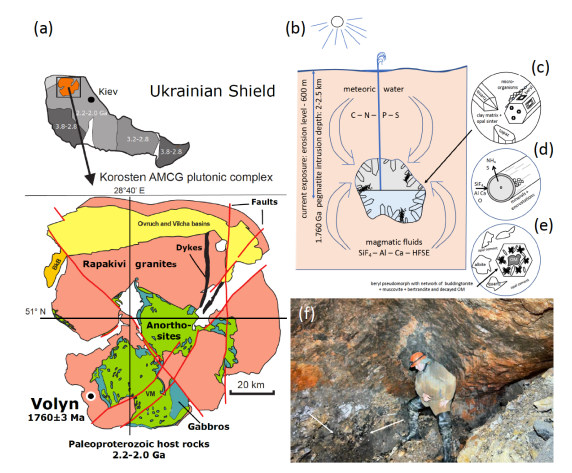


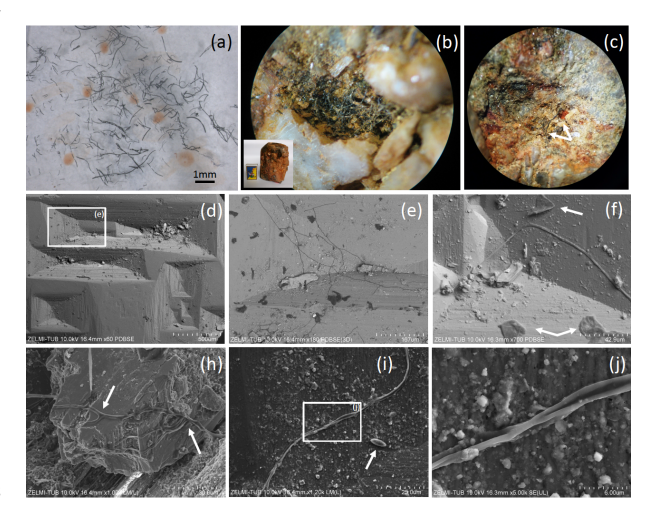
Fig. 1 (a) Location of the Volyn pegmatite field in the Ukrainian shield, which hosts the Volyn biota. (b) Conceptual model for the fossilization in the miarolitic cavities ('chambers') in the pegmatite. Current exposure is from the erosion level to a depth of 600 m. Kerite is attached to the walls, often to feldspar, but also (c) on topaz and beryl crystals. (d) Fossilization produces a thin rim of Si-Al enrichment and encrustations of clay minerals. (e) In collapsed chambers, a breccia formed with pseudomorph of

buddingtonite+muscovite+opal after beryl. Decaying kerite produced NH4 for formation
of buddingtonite. The Ar-Ar-LA age of muscovite is considered as minimum age for the
fossils. (f) View into a chamber, arrows point to black kerite; (all images reproduced from
Franz et al., 2022a).

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94 **2** Geological framework and sample material

95 The locality in the Ukrainian Precambrian shield is associated with the Korosten anorthosite-96 mangerite-charnockite-granite plutonic complex (Shumlyanksyy et al., 2012) (Fig. 1). The 97 samples were recovered from underground in shaft 3 of the mine from a depth of approximately 98 100 m, one sample was obtained from the mineralogical museum of the Academy of Sciences, 99 Kiev, and one beryl sample with kerite on beryl was collected from the mine tailings (Table 1). 100 Two additional samples of topaz from the museum in Kiev with kerite (Fig. 2) were not 101 investigated in detail. The samples from underground could be simply picked up with no need 102 for separation from rock matrix and were stored in plastic sample bags. Kerite has not been 103 found outside the cavities, which are in the core of the pegmatite. It exists as fine fiber masses 104 between fragments of the wall of the cavities and as larger masses hanging freely on the walls, 105 attached to feldspar and often around topaz (Fig. 2b, c) and to beryl (Fig. 2d-j). The bottom of 106 the cavities is covered with soft clay.



109 Fig. 2 (a) Photograph of sample #0, illustrating the pieces of broken, solitary kerite 110 filaments of mm-length. (b) Photograph of kerite filaments on topaz (white crystals with 111 Fe-oxide-hydroxide staining; inset shows topaz crystal with 5-cm large matchbox for 112 scale, image diameter approximately 2 mm). (c) Filaments of different diameter on topaz 113 (arrows; image diameter approximately 3 mm). (d) SEM image (with combined back 114 scattered mode) of beryl prism surface with characteristic etch pits. Rectangle indicates 115 position of (e), which shows filamentous kerite together with kerite in irregular shape (dark contrast indicates organic matter). (f) SEM image, arrows point to kerite with 116 117 irregular shape. (h) Kerite filaments with branching (arrows) in dissolution feature of bervl. (i) Kerite filament and spherical kerite (arrow) in an etch pit of beryl; rectangle 118 119 indicates position of (j), illustrating the irregular diameter of the filament.

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The sample #0 consists of broken filaments of several mm length (Fig. 2a) and it is likely that the original length was much larger on the cm scale. It was also found grown onto a topaz crystal (Fig. 2b, c). On beryl it was found attached to dissolution features on the surface of the crystals, but not only in the common filamentous form, but also in irregular shape (Fig. 2d-j) and rarely in spherical shape (Fig. 2i). Although the previous reports mention mostly filaments

- 126 with smooth surface, our new observations revealed a large variety of different types of
- 127 filaments, described below.

128 **3 Methods**

The samples were investigated by SEM and EMPA. SEM images were obtained with a Hitachi SU8030 instrument, equipped with an EDAX EDS system with a 30 mm² silicon drift detector (SDD) fitted with a silicon nitride window. Samples were coated with an approximately 5 nm thick Ir layer allowing for high-resolution imaging of the filaments' surfaces without the structure of commonly applied Au coating. The kerite samples without further cleaning or preparation were mounted on Al stubs stickered with conductive carbon tabs. The beryl crystals with kerite filaments were dust-cleaned with compressed air and coated with C.

- 136
- 137 Table 1: List of samples

No./GFZ no.	Year of sampling	Material	Location		
0/Museum Ac. Sci. Kyiv	unknown	kerite	unknown		
1/G017809	2018	kerite	shaft 3		
2/G017810	2018	kerite	shaft 3		
3/G017811	2018	kerite	shaft 3		
4/G017812	2018	kerite	shaft 3		
5/G017813	2013	kerite	shaft 3		
6/G017814	2013	kerite	shaft 3		
7/G017815	2013	kerite	shaft 3		

2008-V-10	2008	beryl crystal	mine tailings
		with etch pits	pegmatite #2

139 The JEOL JXA-8530F field emission electron microprobe at TU Berlin was used to investigate 140 mounts embedded in epoxy, but with C-coating, for quantitative results and less absorbance 141 (compared to Ir). EPMA data for element distribution maps of cross sections or of parts of the 142 rim of the filaments and flaky kerite were acquired in the wave-length dispersive mode using 143 an 8 kV, 20 nA beam with a probe diameter of 64 nm. Back-scattered electron images (BSE) 144 were taken to select appropriate sites. Mappings were done in stage scan-modus with pixel 145 resolution between 277 and 360 x 180 and 265, with a pixel size of mostly 80 nm, and a dwell 146 time per pixel of 200 ms. Total scan areas varied between 70 x 36 µm to 33.2 x 31.8 µm.

147 Stable isotope analysis and concentration measurements of nitrogen and carbon were performed 148 simultaneously with a THERMO/Finnigan MAT V isotope ratio mass spectrometer, coupled to 149 a THERMO Flash EA 1112 elemental analyzer via a THERMO/Finnigan Conflo IV- interface 150 in the stable isotope laboratory of the Museum für Naturkunde, Berlin. Isotope ratios are expressed in the conventional delta notation ($\delta^{13}C / \delta^{15}N$) relative to atmospheric N (Mariotti, 151 152 1983) and VPDB (Vienna PeeDee Belemnite standard). Standard deviation for repeated 153 measurements of lab standard material (peptone) is generally better than 0.15 per mill (‰) for 154 both N and C. Standard deviations of concentration measurements of replicates of our lab 155 standard are <3% of the concentration analyzed.

156 FTIR absorption spectra of several small, 40-60 µm wide, translucent dark-brown fragments of 157 kerite (sample #0, which showed the least mineralization crust) were measured in the spectral range 7000 – 700 cm⁻¹ at room temperature using a Bruker IFS 66 spectrometer equipped with 158 159 an IR-microscope. The kerite fragments were selected under a binocular microscope and placed 160 on an IR-transparent KBr plate. Spectra were taken in the transmittance mode at a spectral resolution of 4 cm⁻¹ with a measuring spot diameter of 40 µm. The reference spectra were 161 162 measured through the same KBr plate. The time-averaged signal was collected over 200 scans 163 in both reference and sample spectra. For comparison, absorption spectra of chitin (poly-(1,4)-164 β -N-acetyl-D-glucosamine) and >75% deacetylated chitin, or chitosan (2-amino-2-deoxy-165 (1,4)-β-D-glucopyranan, both produced by Sigma-Aldrich Chemie GmbH (C7170-100G, 166 C3646-10G) from shrimp shells, were measured in several single flattened, 30-50 microns thick 167 transparent flakes of these materials at the same conditions. Band assignments are based on 168 literature comparison (Table 1 Supplement).

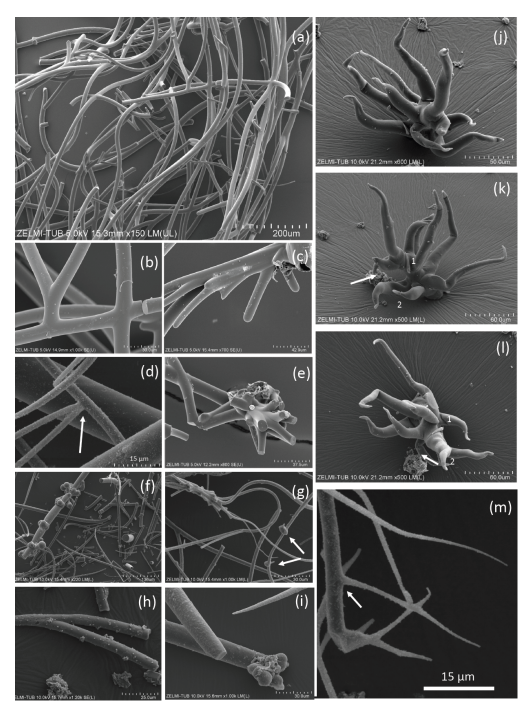
169 **4 Results**

170 **4.1 Morphology**

171 **4.1.1 Filaments**

172 Filaments are curvilinear with smooth surfaces and circular cross section (Fig. 3) with different 173 types of ends (Fig. 4). Other types have a structured surface, some are conical, others strongly 174 curved (Figs. 5, 6). Branching is typical for filaments with smooth surface, and was observed 175 as Y-, T-, and double-T-branching (Fig. 3b, h), as multiple branching (Fig. 3c), and combined 176 Y-T-branching (Fig. 3d). Clear indications for anastomosing filaments were not found. Multiple 177 branching represents the beginning of growth of filaments (Fig. 3e). In others, globular 178 outgrowths possibly mark the beginning of new branches (Fig. 3g). Whereas the diameter of 179 the individual filaments can be homogeneous between approximately 10 µm and 20 µm (sample 180 #0), others (e.g. sample #3; Fig. 3f) show different diameters, between a few µm and several 181 tens of µm. Ball-shaped outgrowths at the end of a filament occur together with a conical thinning-out filament (sample #1; Fig. 3i). Conical, thinning out filaments originate in Y-182 183 branching from a thicker filament with constant diameter (Fig. 3m). One object was identified 184 with multiple conical filaments, with claw-like curved ends (sample #6; Fig. 3j, k, l). The 185 bottom part can be interpreted as beginning of growth of the filaments on a substrate, i.e. the 186 clay mineral assemblage in the miarolitic cavities.

Most filaments are broken pieces of larger filaments, and preserved length is in the order of mm, and it can be assumed that the original length was up to cm. Complete filaments were observed, with one end ball-shaped, the other end thinning out (Fig. 6i, o). Whereas beginning of a filament is rarely observed, ends are frequently preserved (Fig. 4) and can be either simply round (Fig. 4a), ball-shaped (Fig. 4b-f), rarely with oval shape (Fig. 4e), or conical-thinning out (Fig. 4g, 1, m).



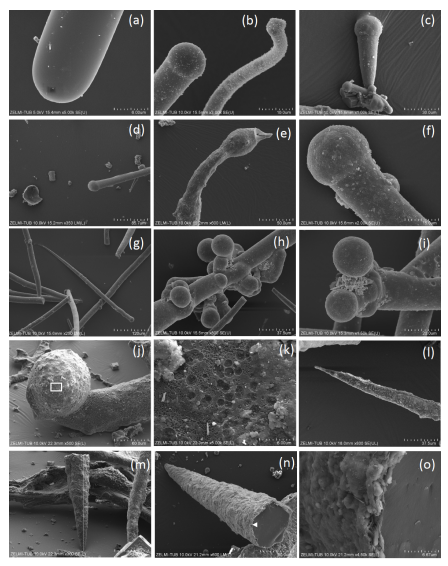
194 Fig. 3 SEM images of curvilinear filaments with smooth surfaces and circular cross 195 section. (a) Overview of sample #0, illustrating the amount of material with homogeneous 196 diameter of approximately 10 µm, length of more than 1 cm, round ends. (b) Branching 197 with Y-, T- and double-T-junctions. (c) Multiple branching and (d) combined Y- and T-198 branching. (e) Possible multiple branching representing the beginning of the filaments. (f) Overview (sample #3) with filaments of variable diameter and (g) multiple branching 199 200 (upper left) and small outgrowths (arrows). (h) Sample #4 with Y-branching. (i) Sample 201 #1 showing 3 filaments, one thinning out (upper left), one with constant diameter with ball-shaped outgrowths on end (below), and a slightly conical one (above). (j, k, l) Image 202 203 of multiple, conical filaments with claw-like ends, growing from a common center; view 204 of the same object (sample #6) in different perspectives. In (k) and (l) numbers 1 and 2 205 identify the same beginning and end of a filament; arrows point to a fluorite crystal. (m) 206 Y-branching of a thinning-out filament (arrow) starting from a filament with constant

thickness. The star-like shape in the center is not branching, it shows different filaments in different heights.

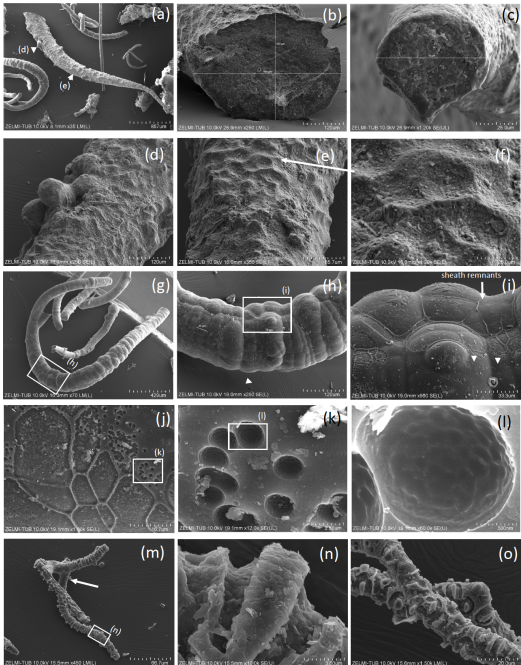
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Ball-shaped outgrowths (Fig. 4h) and multiple ball-shaped ends (Fig. 4i) possibly mark the
beginning of new branches, and balls can be situated asymmetrically at the end of a filament
(Fig. 4j). The structured surface of this ball-shaped end is caused by the fossilization process,
as indicated by the round pores in the surface, together with mineral incrustations (Fig. 4k).
This is also seen on the surface of a 300 µm long conical filament fragment (Fig. 4m, n), which
has a µm-wide rim of mineral incrustations with a homogeneous interior part (Fig. 4o).

- 216 The structured surface is only partly a result of the fossilization process. Figure 5a-f shows a 217 filament with approximately 4 mm preserved length and oval cross section (120x80 µm thick 218 on one end), which has a dented surface and bulbous outgrowths (Fig. 4d). Another example of 219 a strongly curved filament (Fig. 4g-l) with bulbous surface, several mm in length and near to 220 200 µm diameter shows irregular segmentation in distances between 35 µm and 70 µm. On the 221 surface of the filament, relicts of a sheath are visible, partly the sheath is intact. The transition 222 between the intact sheath and the remnants exhibits a polygonal structure and circular 1-2 µm 223 wide holes, probably caused by decay/fossilization. Segmentation is also seen in a branched 224 filament with approximately 3-5 µm wide ridges (Fig. 4m, n, o). This filament has a mineralized 225 outer part of clay minerals with irregular ridges; however, where branching starts, the surface 226 is intact. We interpret these irregular ridges as irregular segmentation of the filament, 227 accentuated and emphasized by fossilization.
- 228 Some samples have joint occurrence of filaments with smooth, slightly, and strongly bulbous 229 surfaces (Fig. 6a, b), and joint occurrence of straight, slightly, and strongly curved filaments 230 with irregular segmentation (Fig. 6c, d). The strongly bulbous filaments are transitional to 231 outgrowths (Fig. 6d). Segmentation is indicated (Fig. 6e) and the surface can be strongly 232 sculptured. The filaments have variable diameters from 75 µm (Fig. 6e) to approximately 250 233 μ m (Fig. 6d, f). Some thin filaments show clear indication for segmentation (Fig. 6g, h). The 234 strongly sculptured surface consists of small ball-shaped outgrowths. Joint occurrence of 235 filaments with strongly sculptured surface and smooth surface and with slight striation 236 perpendicular to filament length, and filaments with strong sculptured surface (Fig. 6k, l, m, n), 237 indicates that these are probably different types of organisms, not different stages of 238 fossilization.

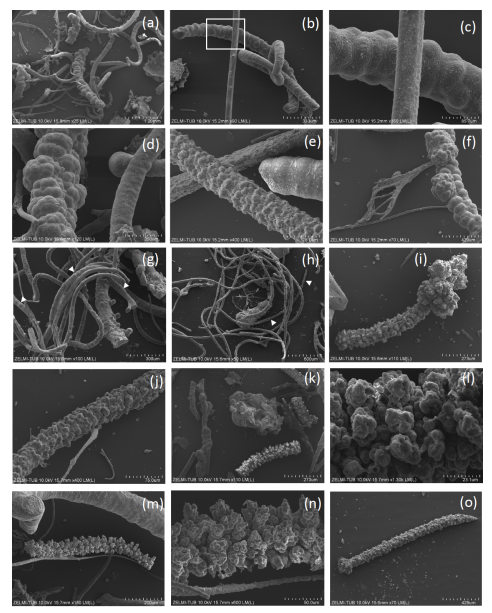


240 Fig. 4 SEM images of ends of filaments with smooth surface. (a) Simple round end (sample 241 #0). (b) Ball-shaped end of straight and curved filament (sample #3). (c) Ball-shaped end 242 of conical filament (sample #1). (d) Ball-shaped end of straight filament (sample #5). (e) 243 Oval-shaped outgrowths near end of filament (sample #7). (f) Ball-shaped end (sample 244 #1). (g) Complete filament with one end thinning out, one with a round end (sample #1). 245 (h) Ball-shaped outgrowths and ends (sample #3). (i) Double ball at end of filament 246 (sample #1) (j) Ball-shaped end; rectangle indicates position of (k), surface of the ball with 247 mineral incrustations and porosity, interpreted as result of decay/fossilization (sample 248 #6). (1) Thinning-out of a filament (sample #5). (m, n) Cone-shaped filament in different 249 perspective, approximately 300 µm preserved length (sample #6); white rectangle 250 indicates position of (o) detail of the 1-2 µm wide rim with mineral incrustations.



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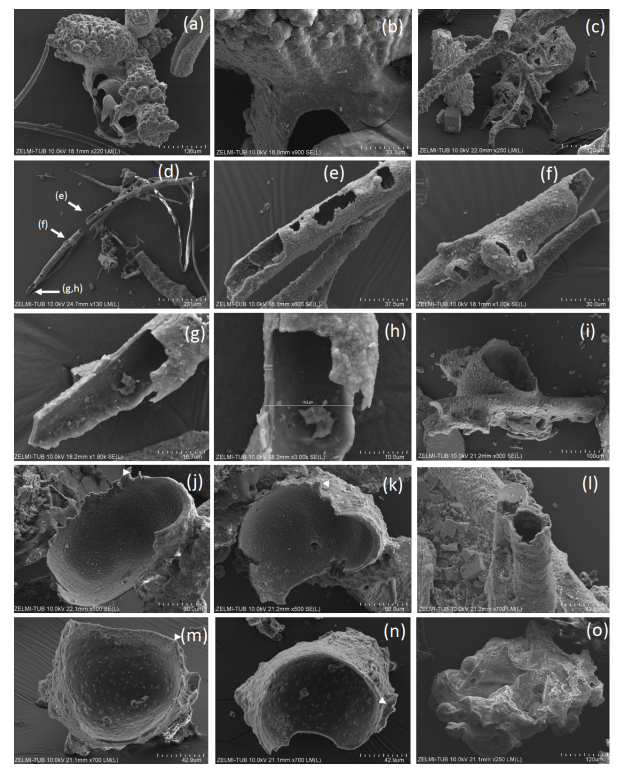
Fig. 5 Filaments with structured, rough surface. (a) Conical filament of approximately 4 254 mm preserved length, upper oval diameter (b) 440 µm x 320 µm, (c) thin end 70 µm 255 (sample #5); triangles point to details shown in (d), bulbous outgrowths, and (e, f) dented 256 surface. (g) Strongly curved filament with bulbous surface, several mm in length and near 257 to 200 µm diameter (sample #5). Rectangle shows position of (h), bulbous surface with 258 irregular segmentation in distances between 35 µm and 70 µm; rectangle indicates 259 position of (i), white triangle to position of (j). (i) In the upper part of the filament, relicts 260 of a sheath are visible (single arrow), in the lower part the sheath is intact (triangles point 261 to the contact). (j) The transition between the intact sheath and the remnants in the lower 262 part of the filament exhibits a polygonal structure and (k, l) circular 1-2 µm wide holes, 263 probably caused by decay/fossilization. (m) Branched filament with approximately 3-5 264 µm wide ridges (sample #2). Note intact surface where branching starts (arrow). (n) Detail 265 of central part of (m). Platy objects are clay minerals. (o) Similar feature of filament 266 surface (sample #4) with irregular ridges, indicating irregular segmentation. 267



269 Fig. 6 SEM images of filaments with structured, rough surface 2. (a) Overview illustrating 270 joint occurrence of smooth, slightly, and strongly bulbous surfaces (sample #5). (b) Joint 271 occurrence of straight, slightly and strongly curved filaments; rectangle indicates detail 272 in (c) with irregular segmentation of the slightly curved filament. The straight filament 273 also shows a slight structure on the surface (lower right). (d) Joint occurrence of slightly 274 bulbous (right) and strongly bulbous filaments, transitional to outgrowths. (e) Filament 275 with indication for segmentation (right) and filament with strongly sculptured surface; 276 note small diameter (75 µm) compared to the large filament in (d). (f) Thick filament with 277 bulbous outgrowths, next to thin agglutinated filaments. (g, h) Thin filaments with 278 indication for segmentation (white triangles). (i) Complete filament of approximately 1 279 mm length with strongly sculptured surface and outgrowths. (j) Part of a filament with 280 strongly sculptured surface. (k) Joint occurrence of filaments with strongly sculptured 281 surface and smooth surface, together with and irregularly shaped object (center). (1) 282 Detail of strongly sculptured surface, which consists of small ball-shaped outgrowths. 283 Note fluorite crystal in upper right, below label (m), which shows joint occurrence of thick 284 filament (top) with slight striation perpendicular to filament length, and filament with 285 strong sculptured surface, detail shown in (n). (o) Almost 2 mm long complete filament, 286 one thin end, one with outgrowths.

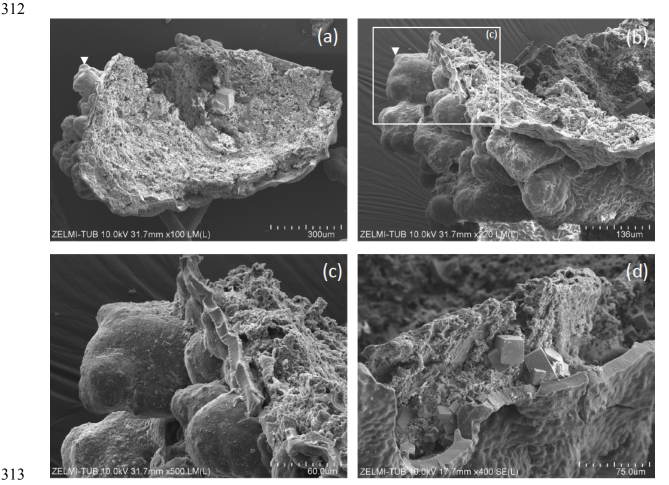
288 4.1.2 Hollow objects

289 Some objects appear hollow (Fig. 7); one object (Fig. 7a, b) has a hollow lower part transitional 290 into a more solid upper, strongly bulbous part. The hollow rather irregular objects (Fig. 7c) 291 occur together with filaments. Filaments can be also hollow (Fig. 7d-h) and the thickness of the 292 outer rim is approximately 2 µm (Fig. 7h). This is the width of the fossilized outer part of 293 filaments, which we documented in the previous study (Franz et al., 2022a) and therefore we 294 interpret the hollow objects as organisms in which the interior part was completely decayed 295 during and after the fossilization process. Some of the hollow objects are bowl-shaped (Fig. 7i-296 n). One such object (Fig. 8) is >1 mm large and from the view in different perspectives is can 297 be seen that it is grown onto mineral substrate; next to the clay minerals fluorite is a 298 characteristic mineral and indicates a high fluorine activity in the fossilizing fluid (Franz et al., 299 2022a). The base of mineral substrate is followed by an approximately 10 µm thick solid rim 300 with bulbous outgrowths.



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302 Fig. 7 SEM images of hollow objects. (a) Irregular-bulbous base of a strongly sculptured 303 object, with (b) detail of the transition (center in (a); sample #5). (c) Irregular hollow 304 object below filaments (sample #6). (d) Hollow filament, approximately 1 mm preserved 305 length; position of enlarged parts in (e-h) is indicated (sample #5). The mineralized rim is 306 1-2 µm wide, diameter near 20 µm. (f) Bulbous outgrowths are also hollow. (i) Filament 307 with an attached hollow form, similar to outgrowths, but much larger (sample #6). (j, k) Same object as in (i), enlarged in two different perspectives; white triangle indicates 308 309 identical point. (1) Hollow filament next to a filament with a central channel (sample #6). 310 (m, n) Isolated hollow bowl-shaped object in two different perspectives; white triangle 311 indicates identical point (sample #6). (o) Irregular object, partly hollow (sample #6).

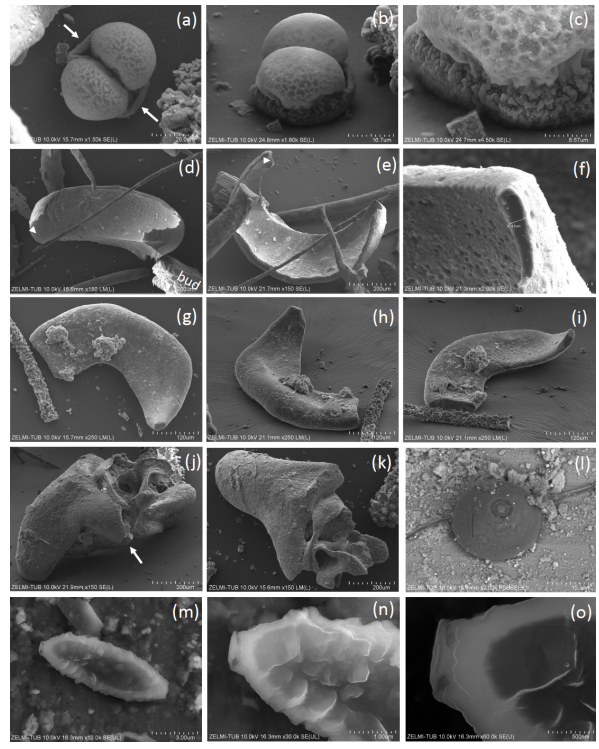


314 Fig. 8 SEM images of >1 mm large bowl-shaped object (sample #5) (a) seen from below, grown onto mineral substrate; euhedral crystal is fluorite, white triangle indicates 315 316 position of (b), enlarged part of the rim. Rectangle indicates position of (c) illustrating the 317 base of mineral substrate (right) followed by an approximately 10 µm thick solid rim with bulbous outgrowths. (d) Detail of the solid rim with several fluorite crystals. 318

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320 4.1.3 Spherical objects

- 321 Most spherical objects (Fig. 9) appear as rather complete, with only some parts broken off. One 322 object with a double-ball shape (Fig. 9a,b) is clearly grown onto the substrate (Fig. 9c). The 323 double-ball with remnants of a sheath points to cell separation. Note the different size of the 324 objects from $< 10 \,\mu\text{m}$ (Fig. 9m) to $> 1 \,\text{mm}$ (Fig. 9g). Two small objects identified on the etched
- 325 beryl surface appear like seeds or spores (Fig. 9l, m).
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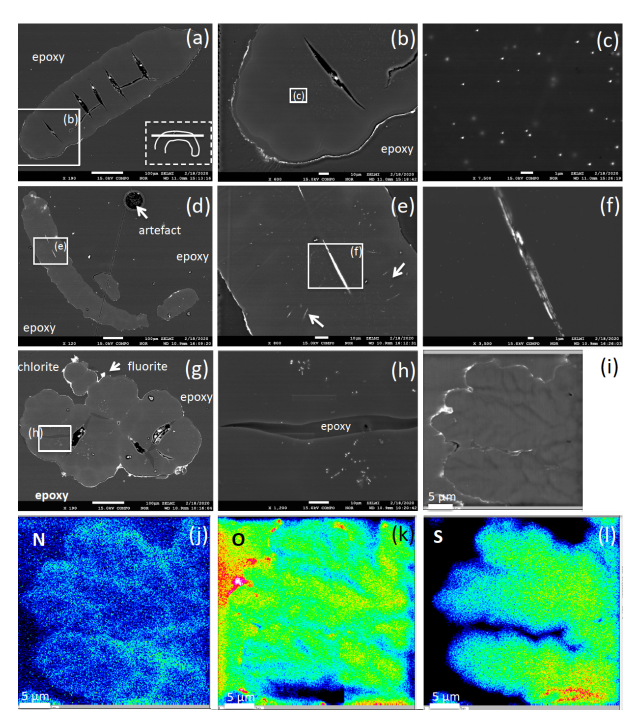
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328 Fig. 9 SEM images of spherical objects. (a, b, c) Same object in different perspective and 329 magnification; arrows in (a) point to a sheath; the euhedral crystal in (c) is fluorite. The 330 object growth from a flat mineral surface into a double-ball with dented surface. (d, e) 331 Same object in different orientation; white triangle indicates identical position; bud = 332 buddingtonite. (c) The thickness measured at one point is approximately 6 µm. (g, h, i) 333 Approximately 0.5 mm large object in different perspective with mineral incrustations. (j, k) Irregular, partly hollow object in different perspective. (l) Perfectly round object, 334 335 sitting on a filament, on etched surface of beryl (compare Fig. 2d); the circular round 336 structure on its top is beam damage. (m, n, o) Oval object on etched surface on beryl 337 (compare Fig. 2i). The lower contrast (dark) in the central part indicates less dense (partly 338 hollow) material.

339 4.1.4 Irregular objects

- 340 Irregular, flaky objects are abundant, especially on the surface of the beryl crystal (Fig. 2e, f),
- 341 but also in many samples (e.g. Fig. 6k, 7a, c, o, 8, 9j, k). They show the same fossilization
- 342 features as the filaments with a thin rim enriched in Si, Al, Ca, and P, loss of N, and oxygenation
- 343 (Franz et al., 2022a). In some samples (Fig. 6f) filaments appear agglutinated by OM and we
- 344 interpret these as well as the irregular objects on the beryl crystals as fossilized biofilm.
- 345 4.2 Internal structure
- 346 For investigation of the internal structure we used SEM images of broken filaments and other
- 347 objects, as well as polished sections embedded in epoxy, investigated by BSE images including
- 348 mapping of element distribution. Data of open-pyrolysis and TEM data (Franz et al., 2022a)
- had shown that the OM is highly mature, amorphous oxy-kerite. Indications for an outer cell
- 350 wall are absent, because the outer rim of the fossils is silicified, partly with formation of mineral
- 351 incrustations.
- 352 Segmentation of filaments, which might be a characteristic phenomenon for certain organisms 353 and is observed in the filaments' morphology (Figs. 5g, h, 6b, c, e, h) is not obvious in cross
- 354 section, but one section shows internal cracks, separating the filament in ~50 µm to 100 µm
- 355 wide segments (Fig. 10a, b). A section of a bulbous fossil shows cracks, which separate the
- 356 individual bulbs from each other (Fig. 10g, h).
- 357 The outer rim of the filament shows the typical enrichment of Si and Al (Fig. 10b), and the
- 358 inner, homogenous and not silicified part shows abundant, nm-sized mineral inclusions (Fig.
- 359 10c). They are located in the central part and thus not related to the fossilization process,
- 360 irregularly distributed or in linear array of several crystals (Fig. 10e, h). The minerals were
- analyzed with the EDS-system and due to their small size in the order of a few nanometers,





364 Fig. 10: BSE images of filamentous (a-f) and bulbous fossils (g, h, i), embedded in epoxy, polished thin section and element distribution (j, k, l). (a) Part of curved filament; 365 366 orientation of section is shown in rectangle (dashed lines), position of enlargement (b) in 367 rectangle (solid lines). Open cracks (black contrast, with impurities from polishing 368 material) indicate approximately 50 µm to 100 µm wide segments. (b) Silicified outer rim 369 (white contrast, irregular) and a narrow, up to 10 µm wide inner rim, are interpreted as 370 effect of fossilization. The homogeneous appearing central part shows in the enlarged 371 image (c) irregularly distributed inclusions, tens of nm in size, of Bi-S-Te minerals. (d) 372 Filament with two, central oriented Bi-S-Te mineral inclusions, approximately 50 µm in 373 length and 1-2 µm wide, enlarged shown in (e) and (f). Arrows in (e) point to straight 374 aligned inclusions, and (f) shows irregular contrast, possibly caused by heterogeneous 375 distribution of Fe and Cu in the Bi-S-Te minerals. (g) Bulbous fossil, with silicified rim and encrustations of chlorite and fluorite. Cracks, partly filled with epoxy, separate individual bulbs from each other. (h) Enlarged part showing irregularly distributed and aligned nm-sized Bi-S-Te mineral inclusions, and epoxy-filled crack. (i) Bulbous fossil with element distribution of N (j), O (k), and S (l), indicating an interior structure with possible former cell walls. The color code goes from cold to warm, blue = low concentration, red = high concentration.

382 much smaller than the excitation volume of the electron beam, only mixed analyses with the 383 organic material could be obtained (Table 2). Recalculation of the analyses without the organic 384 compounds C, O, and N yielded an atomic ratio of Bi:(S,Te) near 1:1, indicating minerals such 385 as ingodite Bi(S,Te) or joseite Bi₄(S,Te)₃. The example of the bulbous filament (Fig. 10g) with 386 inclusions also shows a Bi(S,Te) mineral, located in the central part. The heterogenous BSE 387 contrast is caused by different trace compounds of Fe and Cu. Element distribution of N and O 388 (Fig., 10j, k) in a bulbous fossil, indicated by different BSE contrast (Fig. 10i), show an internal 389 structure, possibly indicating a primary separation into different cells, whereas S (Fig. 10l) 390 shows a systematic decrease towards the rims of the object, as a result of decay and/or

391 fossilization.

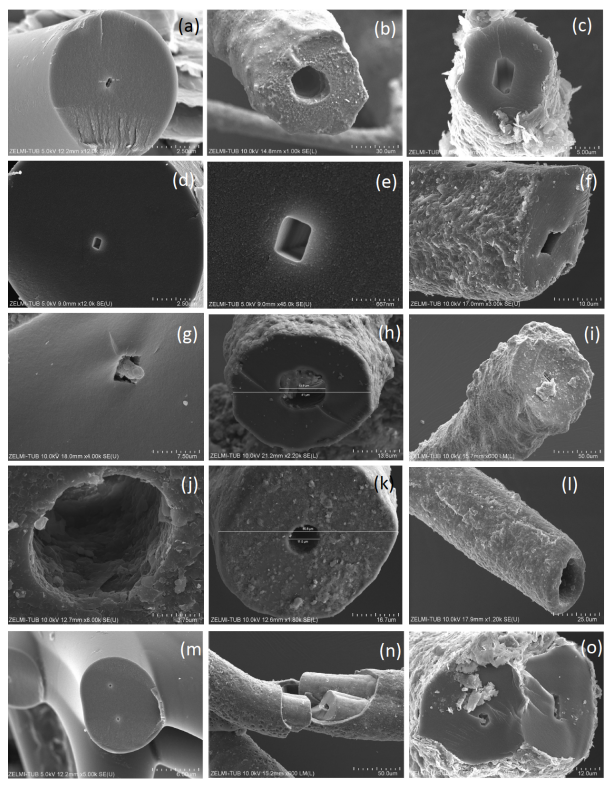
392

Analysis#	15 06 ¹	13 03 ²	13 ³ n=18	Min-max
S atom%	0.27	2.59	0.20	n.d 0.52
Те	0.13	0.06	0.12	n.d. – 0.51
Bi	0.29	2.05	0.24	0.01-0.68
Pb	0.03	n.d.	n.d.	
Fe	n.d.	0.19	n.d.	
Cu	n.d.	0.22	n.d.	
С	86.24	84.86	83.38	80.19-96.15
Ν	5.91	4.89	3.16	n.d7.18
0	7.13	5.14	10.12	2.74-15.78
Sum ⁴	100	100	100	
recalculated	15 06	13 03	13 n=19	Min-max
S atom%	38	51	37	3-55
Те	18	1	25	1-90
Bi	40	40	46	7-68
Pb	4	0		
Fe		4		
Cu		4		
Sum	100	100	100	

393 Table 2 EDS analyses of Bi-sulfide-telluride inclusions

¹ Fig. 10h; ² Fig. 10f inclusion in channel; ³ average of 18 analyses, inclusions in matrix, Fig.

395 10b,c; ⁴ normalized; n.d. = not detected 396



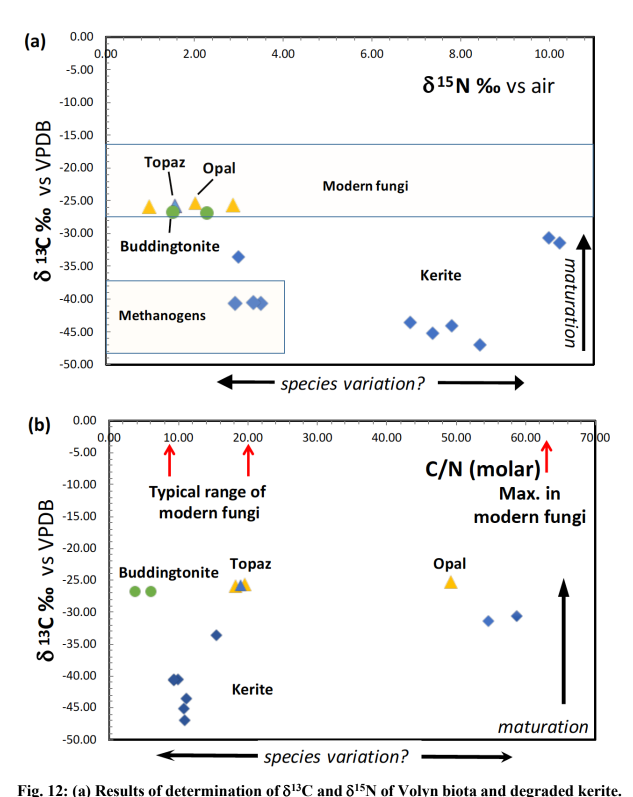


400 Fig. 11: SEM images of broken filamentous fossils, illustrating the central channel. (a,b,c) 401 Six-sided channel in filament with (a) smooth outer surface, (b) dented surface, and (c) 402 strongly mineralized surface. (d, e, f, g) Rectangular channel; (e) is enlarged part of (f). 403 (h) Round, slightly irregular channel. (i) 4 μ m x 6 μ m wide channel on filament with 404 dented surface. (j) Round channel, enlarged from (k), approximately 12 μ m wide in a 405 filament of nearly 70 μ m diameter. (l) Slightly conical end of a filament with large, round 406 channel. (m) Two filaments one with a small μ m-wide channel attached to a hollow

filament. (n) Channel in a filament with sheath-like structure. (o) Two filaments with six sided channels.

- 409 A very characteristic feature of the filaments is a central channel (Fig. 11), observed in many
- 410 but not all of the filaments. The cross section of the channel can be six-sided (Fig. 11a-c,m),
- 411 rectangular (Fig. 11d-f), or round (Fig. 11h-l). The channel diameter is variable and ranges from
- 412 approximately 0.5 μ m to 25 μ m in filaments with an outer diameter between approximately 5
- 413 μ m and 100 μ m; examples in Fig. 11 show 5 μ m with a channel of 260 nm x 550 nm (a), 50
- 414 μ m with a channel of approximately 20 μ m (b), 10 μ m with a channel of 2.5 μ m x 4 μ m (c),
- 415 100 μ m with a channel of 400 nm x 560 nm (d,e), 41 μ m with a channel of 14 μ m (i).
- 416 4.3 Stable isotopes and C/N variation
- 417 Stable isotopes of C and N were obtained from all bulk samples (Table 1); it was not possible
- 418 to determine individual fossilized objects. In addition, we determined OM in black opal and
- 419 OM adherent to topaz (see sample list in Franz et al., 2022a).
- 420 Results of δ^{13} C and δ^{15} N-determination and the molar C/N show a large variation (Fig. 12). All δ^{13} C values are negative, and for kerite fossils vary between -47 (sample 2) and -31 ‰ (sample 421 1); δ^{15} N values vary between ~3 to 4 ‰ (samples kerite 0, 4) and ~10 ‰ (samples 1, 3). OM 422 associated with opal and topaz (considered as 'secondary') and buddingtonite, which obtained 423 424 its N from decayed OM, is less negative and homogeneous in δ^{13} C with values between -25 and -27 ‰. The C-values should be considered as maximum values, since alteration either by 425 deep-seated CO₂ from the mafic magmas or from meteoric waters would have increased δ^{13} C. 426 The close group of δ^{13} C and δ^{15} N values for secondary OM indicates that during maturation 427 428 and decay they all have reached a similar value. The variation of the N-isotopes is not correlated 429 with the C-isotopes, and there is also no correlation with C/N.
- 430
- 431 Table 3 Results of $\delta^{15}N$, $\delta^{13}C$, and molar C/N of bulk kerite samples

Sample#	weight mg	δ ¹⁵ N ‰	mg N/sample	% N	$\delta^{13}C$ ‰	mg C/sample	% C	molar C/N
1	2.76	9.99	0.038	1.37	-30.66	1.91	69.07	58.74
2	2.37	8.44	0.067	2.85	-46.99	0.63	26.52	10.87
3	2.21	10.23	0.027	1.20	-31.38	1.24	56.10	54.58
4	2.52	2.98	0.033	1.31	-33.61	0.44	17.34	15.48
5	4.01	7.37	0.096	2.38	-45.19	0.88	21.98	10.78
6	3.14	7.79	0.037	1.19	-44.06	0.27	8.55	8.39
7	4.29	6.87	0.074	1.73	-43.58	0.71	16.54	11.17
Opal 8	50.15	2.02	0.013	0.03	-25.32	0.55	1.09	49.23
Topaz 9	54.46	1.56	0.023	0.04	-25.73	0.38	0.69	18.89





436 Fig. 12: (a) Results of determination of δ^{13} C and δ^{15} N of Volyn biota and degraded kerite. 437 Symbols: Blue diamonds – dominantly filamentous kerite, with small amounts of flaky

438 and spherical OM; yellow triangle - black opal with OM; blue triangle - OM adherent to

439 topaz; green dots - buddingtonite from breccia (from Franz et al., 2017). Fields of modern

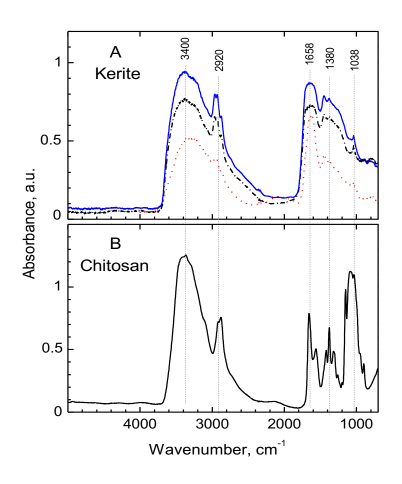
440 fungi from Mayor et al. (2009) and methanogens are summarized in Struck (2012). (b)

441 Molar C/N ratio of kerite fossils and degraded OM. Range of C/N of modern fungi from

442 Mayor et al. (2009).

443 **4.4 FTIR investigation**

444 All measured FTIR spectra of morphologically different kerite fragments in the sample #0 are 445 very similar (Fig 13a) and resemble closely the chitosan spectrum (Fig 13b); both spectra are dominated by two main groups of absorption bands located in the regions of 3500-2500 446 cm⁻¹ and 1800-900 cm⁻¹. The first group consist of overlapping broad bands due to O-H and N-447 H stretching vibrations, with a group of characteristic narrow peaks of C-H stretching vibrations 448 on their long-wavelength wing in the region of 2960-2870 cm⁻¹ (Fig. 13; for detailed band 449 450 assignments and for spectra of chitin see Table 1 Supplement). The peak in vicinity of 1650 451 cm⁻¹ is diagnostic of C=O group (Wanjun et al., 2005; Coates, 2011; Loron et al., 2019), the band at 1560 cm⁻¹ (broad shoulder near 1570 cm⁻¹ in kerite spectra) was assigned to N-H 452





454 Fig. 13: FTIR spectra of filamentous fossil compared to standard materials chitin and 455 chitosan. (a) Complete spectra of three pieces of sample kerite #0, the sample with less mineralization, showing two main regions of absorption: 3500 cm⁻¹ to 2800 cm⁻¹ and 1850 456 cm⁻¹ to 900 cm⁻¹; (b) Standard material chitosan. Compared to chitosan the major 457 absorption bands in kerite spectra are broader, the weak shoulder near 3100 cm⁻¹ in 458 459 chitosan spectrum is not present in kerite. The narrow triplet near 2950 cm⁻¹ is observed as doublet in chitosan, shifted to lower wavenumbers. In the part from 1800 cm⁻¹ to 700 460 461 cm⁻¹, kerite shows only broad absorption, shifted towards higher wavenumbers compared

to chitosan, with three superimposed distinct weak peaks at 1450 cm⁻¹, 1380 and 1038 cm⁻¹
the first is not present in chitosan, which has a number of distinct peaks in this region.

- bending vibrations in amide group. The relatively weak band near 1420 cm⁻¹ (1450 cm⁻¹ in kerite) was attributed to C-H bend (Loron et al., 2019), and the sharp peak at 1380 cm⁻¹, which was reported in cellulose, chitosan, and chitin spectra, was assigned to superposition of O-H bend (pyranose ring; Li et al., 2009) and symmetrical bend of CH₃ group. A band centered near 1315 cm⁻¹ in chitin and chitosan spectra due to C-N stretching vibrations in amide group (Vasilev et al., 2019; Wanjun et al., 2005) is not observed in kerite.
- A broad, weak band at around 2100 cm⁻¹ is present in spectra of kerite and chitosan (Fig. 13), and the same type of weak bands are shown in published chitosan spectra (see Table 1 Supplement), but not mentioned and assigned. It can probably be attributed to overtone or combination bands of pyranose ring vibrations. At lower wavenumbers, in all measured spectra there is a series of strong (1150, 1180, 1030 cm⁻¹) and several weak bands caused by different types of C-O vibrations in polysaccharides (Nakamoto, 1997; Wanjun et al., 2005; Li et al., 2009; Coates, 2011; Loron et al., 2019; Vasilev et al., 2019).
- A general observation is that in kerite spectra, compared to chitosan, all characteristic absorption bands of the amide group and the pyranose ring become broader and weaker, in agreement with earlier studies of spectroscopic changes during chitin/chitosan degradation (Wanjun et al., 2005; Zawadzki and Kaczmarek; 2010; Vasilev et al., 2019). Nevertheless, the main absorption features caused by amide group, diagnostic of chitosan, are still present in kerite spectra.

484 **5 Discussion**

485 **5.1 Interpretation of morphological and internal characteristics**

The Volyn biota show an astonishingly large variation of different types of filaments and other forms, pointing to the interpretation that different organisms were involved. We have already interpreted the flaky objects of OM on the surface of beryl crystals (Fig. 2e,f) as biofilms (Franz et al., 2022a). Agglutinated filaments (Fig. 6f) and the hollow object agglutinated to a filament (Fig. 7i) can similarly be interpreted as fossilized biofilms. The sheath-structure (obvious e.g. in Fig. 5i, j) is also an indication for the presence of a biofilm or extracellular polymeric substances (EPS). 493 Some objects have a base onto which they grew (Figs. 3j-l, 8, 9a-c) and one object shows a 494 hollow lower part, from which bulbous outgrowths originate (Fig. 7a, b), pointing to sessile 495 organisms. Filaments are generally fragmented, but a few filaments have been found with two 496 intact ends (Figs. 4c, g, 6i, o), and we interpret this as non-sessile, free-living organisms.

Thickness of the filaments varies from $\leq 10 \ \mu m$ to $\geq 200 \ \mu m$. In filaments with diameter up to approximately 30 μm , branching with thinning out of the branch clearly show that these are within-species variations (irregular diameters of filaments, Fig. 2i, j, are interpreted as collapse structures during fossilization). However, very thick filaments with diameters in the range of $\geq 200 \ \mu m$ with a structured, bulbous surface (e.g. Fig. 6), or conical objects (Fig. 4m) are interpreted as different species. The length of both types of filaments reaches the mm-range, and since they are fragments possibly up to cm-length.

504 Branching as indication for growth of the organisms is typical in the thin filaments, with Y-, 505 T-, double-T-, and multiple branching (Fig. 3), but anastomosing was not observed. In thick 506 filaments with diameter near 200 µm branching was not found. The ends of filaments also hint 507 to the type of growth. Simple round ends are rare, more typical are ball-shaped ends (Fig. 4). 508 Ball-shaped outgrowths along filaments are interpreted as beginning of a branching (Fig. 4h). 509 In the complete filaments (Fig. 4c, g) with one end thinning out, one with a ball-shaped end, 510 the thinning-out end is possibly the origin, the ball-shaped protrusions the growing end, because 511 ball-shaped ends are rather continuous in shape, from a small protrusion (Fig. 4b) to a more 512 complete ball (Fig. 4f, i). Similar protrusions were found at the end of recent, large bacterial 513 filaments (Volland et al., 2022). However, branched, thinning-out ends of the filaments (Fig. 514 3j-l, m) indicate ends similar to Spitzenkörper, what in modern fungi is described as a 515 continuous and indefinite process of cell extension (Fischer et al., 2008).

Segmentation in thin filaments (Figs. 5m, 6g, h) with distances of a few µm up to tens of µm is 516 517 accentuated by mineralization (Fig. 5n), with irregular ridges caused by mineralization. Thick 518 filaments do not show a clear segmentation; the morphology is more irregular and shows 519 rounded, polygonal structures on the surface with dimensions of approximately 20-30 µm 520 (parallel to filament axis) x 35-70 µm (perpendicular to filament axis) (Figs. 5g, h, i, 6b, c). 521 Between the polygonal structures on the surface, remnants of a sheath are visible. In cross 522 section (Fig. 10) segmentation is clearly visible by cracks with a distance of approximately 50-523 100 µm.

524 Bulbous forms (Figs. 7a, b, 8) mark the beginning of growth of some objects, and bulbous 525 outgrowths are very typical for thick filaments (Fig. 6, d, f), which extend into approximately 526 20 μm large objects, which consist of smaller bulbs (Fig. 6l, n). In thin filaments with typical 527 branching, the outgrowths are rare and more regularly ball-shaped (Figs. 3f, g, 4h), indicating 528 one species with prominent growth by branching of thin filaments, and another species with 529 growth by outgrowths along thick filaments.

Among the spherical objects, only the small ones with a size of a few μ m (Fig. 91-o) resemble spores or other types of seeds/fruit bodies. The irregular, large objects several hundred μ m in size (Fig. 9d-k) do not fit into any scheme of known organisms. Similarly, there is no obvious interpretation for the large bowl-shaped and irregular hollow objects (Fig. 8). The small doubleobject with a partly preserved sheath (Fig. 9a-c) grown on a substrate has some similarities with cell division.

536 The function of the conspicuous central channel (Fig. 11) in many, but not all filaments with 537 different shape in cross section is speculative, likely providing pathways for transport of 538 components for cell extension along the filament axis. In one example we observed a type of 539 filling in the channel (Fig. 11g), so in the original organisms it might have been filled with an 540 easily degradable substance. It is not clear if a hollow form (Fig. 7e, 1) is a different phenomenon 541 or due to special preservation conditions. The width of the preserved rim is in the same order 542 of magnitude as the silicified rim $(1-2 \mu m)$ and therefore it might just be a remnant of a filament, 543 in which the central part was completely degraded.

544 Another special feature of the internal structure are the nanometer-sized mineral inclusions of 545 Bi-S-Te minerals (Fig. 7). The organisms were able to concentrate these elements, either 546 irregularly distributed (Fig. 7c) or rod-like aligned (in a bulbous object; Fig. 7h) or within the 547 channel (Fig. 7e). It is unclear if the relatively large Bi-S mineral with some Cu and Fe contents 548 in the center of a thick filament in the central channel is the original position of the Bi-S 549 concentration or an effect of fossilization. Modern fungi are able to concentrate Te (and Se) as 550 nm-sized crystals (Liang et al., 2020) and could be used in technology for soil mycoremediation 551 (Liang et al., 2019). In black shales, the organophilic element Bi might behave similar as Se 552 (Budyak and Brukhanova, 2012). Biogeochemistry of Te is probably analogous to Se (Missen 553 et al., 2020), but little is known about the link of Bi to S and Te in OM (such as in coal, e.g. 554 Finkelman et al., 2019). The concentration of Bi-S-Te in the organisms of the Volyn biota is 555 another indication for fungi-like organisms, although other organisms such as bacteria are also 556 able to concentrate Te (Missen et al., 2020).

557 Remnants of cell membranes, separating individual cells, could not be identified, and to answer 558 the question if some of the organisms were multicellular is speculative. However, the large size

559 of many objects of the Volyn biota already indicates that possibly they were not single-celled 560 but multicellular, notwithstanding that single-cell bacteria (Thiomargarita magnifica; Volland 561 et al., 2022) can reach the size of cm. These macroscopic single-cell bacteria show a very simple 562 straight filament, whereas the large objects from the Volyn biota show a much more 563 complicated form; the surface of large filaments shows a bulbous structure with sizes in the 564 order of tens of µm (Figs. 5g-i, 6c, f, 9a, b), well visible with a polygonal network (Fig. 5j). In 565 the internal structure we also see phenomena that could be explained as separate cells, such as 566 the gaps in a filament (Fig. 10a) or in a bulbous object (Fig. 10g). The interior structure visible 567 in the element distribution of N (Fig. 7j) might indicate the original distribution in former 568 interior cell walls, in which chitin-like substance was concentrated. Finally, the small spherical 569 object shown in Fig. 9a, b might be taken as two cells, with an envelope of a sheath.

570

571 **5.2 Stable isotopes**

Modern fungi show a very wide variation of δ^{15} N from -5 % to +25 %, with the main cluster 572 573 between - 5 ‰ and +12 ‰, and δ^{13} C is restricted to -19 ‰ to -29 ‰ δ^{13} C, with the main cluster 574 at -22 ‰ to - 28 ‰ δ^{13} C (Mayor et al., 2009; Fig. 12a). Whereas the N-isotopic signature of 575 kerite is consistent with the interpretation as fossil fungi, the C-isotopic signature is much lower 576 than that of modern fungi. However, fungi live from consumption of organic matter, and the C-577 isotopic signature is then transferred to the fungi without strong isotopic effect (Peterson and 578 Fry, 1987). I. e. during incorporation of carbon from modern plants to fungi, the δ^{13} C-signature of -27 ‰ to -30 ‰ in plants changes to - 25 ‰ to -27.5 ‰ δ^{13} C in fungi (e.g. Högberg et al., 579 580 1999). Assuming that the isotope fractionation in the Volyn biota was similar, the consumed 581 organism had a C-isotopic signature of c. -35 % to -50 % δ^{13} C. These very low values are 582 consistent with the interpretation that the primary organisms were methanogens. Another factor, 583 which must be considered, is intracellular heterogeneity as observed in bacteria (Lepot et al., 2013). The membrane (lipids) can have a signature of 10 $\% \delta^{13}$ C lower than the bulk cell, and 584 585 degradation during fossilization of the proteins and polysaccharides can lower the now 586 determined C-signature. It is also possible that the fungi consumed biofilm. Fossil biofilms of the 2.75 Ga Hardey Formation (Australia), probably coexisting with methanogens, 587 588 methanotrophs, and sulfur-metabolizing bacteria have δ^{13} C of -55 ‰ to -43 ‰ (Rasmussen et 589 al., 2009), well in the range of δ^{13} C-values observed here. The biofilms, described by 590 Rasmussen et al. (2009), lived in synsedimentary cavities similar to stromatolites, pointing to

the importance of cavities for the preservation of organic matter, similarly as the biofilms atVolyn in the deep biosphere.

593 Maturation clearly affects the C- and N-isotope ratios, which we see in degraded OM preserved 594 in black opal, in OM adherent to topaz, and buddingtonite which obtained its NH₄ from OM. These samples have much more positive δ^{13} C values around -26 ‰ and more homogeneous 595 δ^{15} N values near +1.5 to +3 ‰ (Fig. 12a). In contrast, the large variation of δ^{15} N between 3 ‰ 596 597 and 10 ‰ in the kerite samples (Fig. 12a) and C/N between 10 and >50 (Fig. 12b) possibly 598 indicates a variation of the species. These values were less influenced by maturation, as there 599 is no correlation between δ^{13} C and C/N in all samples (fossils and degraded OM). Alleon et al. 600 (2018) in their description of the 3.4 Ga old Strelley Pool microfossils (Western Australia) 601 argued that though the fossils experienced heating up to 300 °C, the C/N did not change 602 significantly. Also, for anthracite coal it has been shown that the original C/N did not vary with 603 coalification (Anwita et al., 2020).

604 Loron et al. (2019) reported fossil fungi from the 1 Ga Grassy Bay Fm Canada, and provided 605 proof via chitin remnants (FTIR) and showing the characteristic bilayered fungal cell walls 606 (TEM data). However, the few SEM images for the Grassy Bay biota do not allow a comparison 607 with the Volyn biota. Following their discussion, the FTIR investigation of the filamentous 608 Volyn sample shows good indications for preserved chitosan as part of the OM. Degradation 609 studies of chitosan (Wanjun et al., 2005; Zawadzki and Kaczmarek; 2010; Vasilev et al., 2019) 610 showed that the spectra of kerite has the same characteristic bands as chitosan at approximately 611 250 °C; at lower as well as at higher temperatures these bands disappear. Completely 612 independent temperature estimates for the fossilization based on phase equilibria of Be minerals 613 yielded the same temperature range (Franz et al., 2017).

614 **5.3 Taxonomy and comparison with Precambrian biota**

Film-like microfossils were described from the 3.4 Ga old Strelley Pool (Western Australia; Alleon et al., 2018), the 3.3-3.5 Ga old Onverwacht Group (Australia; Westall et al., 2001), from the 2.75 Ga old Hardey Formation (Australia; Rasmussen et al., 2009) and there is little doubt that biofilms existed for a long time in the Earth's history and are an integral component of the ancient life cyle (Hall-Stoodley et al., 2004). It seems safe to assume that the irregular (Fig. 2f, and images in Franz et al., 2022a) and sheath-like structures (Figs. 5i,j, 6f, 9a) of the Volyn biota were biofilms.

- We have already pointed out that some of the organisms show analogies to fungi. Based on the
- 623 molecular clock technique, Wang et al. (1999) estimated the divergence between the three-way

624 split of the animal-plant-fungi kingdoms at 1.58±9 Ma, much earlier than the 'Precambrian 625 explosion'. This age is in the same range as the minimum age of the Volyn biota. Other 626 molecular clock estimates indicate that the first zygomycetous fungi occurred on Earth during 627 the Precambrian, approximately 1.2–1.4 Ga ago (review in Krings et al., 2013). Diversification 628 of fungi and transition to land was dated at ca. 720 Ma (Lutzoni et al., 2018) and they estimate 629 the origin of fungi at ca. 1240 Ma, similarly as Berbee et al. (2020), who placed the origin of 630 fungi at ca. 1300 Ma. If indeed the Volyn biota contain fungi-like organisms, their origin as 631 well as colonization of land occurred earlier than ca. 1500 Ma.

632 Bengtson et al. (2017) reported fungus-like organisms in the 2.4 Ga Ongeluk Formation (South 633 Africa) from the deep biosphere, which are however not terrestrial but marine. The important 634 fact is that these fossils were found also in open cavities, though of a completely different size, 635 mm-amygdales in low-grade metamorphic basalt, in contrast to the huge cavities of tens of 636 meter size in the pegmatites from Volyn. The filaments from the Ongeluk biota with a diameter 637 of ca. 2 µm to 12 µm are generally thinner than the Volyn biota and show anastomosis, but also 638 Y- and T-branching, and sometimes bulbous protrusions, 5-10 µm in diameter. A special feature 639 is what Bengtson et al. (2017) call 'broom structure', diverging filaments growing from a 640 substrate of clay minerals (chlorite), and the filaments consist also of the same type of chlorite. 641 These structures (shown in 2D in thin sections) could be similar as the object from the Volyn 642 biota (Fig. 3j, k, l), and what we called 'multiple branching' (Fig. 3c, e, g). A significant 643 difference between the two biota is the fossilization process, which resulted in the Ongeluk 644 biota in complete replacement of the filaments by clay minerals, whereas at Volyn fossilization 645 is restricted to the outermost rim and most of the C is preserved (Franz et al., 2022a).

646 Good evidence for fungi-like organisms were reported from the early Ediacaran Doushantuo 647 biota, at approximately 635 Ma (Gan et al., 2021). These fossils are pyritized, but with remnants of organic matter, and consist of branching filaments (Y-, T-branching, but also with A- and H-648 649 type and anastomosis) and associated hollow spheres. Compared to the Volyn biota, the 650 filaments are thinner (two types, one with average 6.8 µm, one with average 2.7 µm), whereas 651 the observable length in thin section with hundreds of µm is possibly in the same range as in 652 the Volyn biota. The spheres of the Doushantuo biota are hollow and coaxially aligned, but also 653 similar to what we described as ball-shaped outgrowths; their size varies from average 16 µm 654 to 20 µm in small ones and large spheres with 36 µm to 102 µm, similarly to the Volyn biota 655 (Fig. 4h, i for the small spheres, Fig. 4j for large spheres). The fact that the spheres of the 656 Doushantuo Formation are hollow is possibly due to the fact that they are mostly pyritized, i.e. 657 most of the organic matter was decomposed. The small spheres were interpreted (Gan et al.,

658 2021) as possible spores, the larger ones were possibly symbiontic organisms living together659 with the fungi.

660 Myxomycetes (slime molds) are other possible eukaryots, which might have existed in the 661 Proterozoic, although Stephenson et al. (2008) considered 50 My as the oldest fossil record. 662 Their diverse morphology during the different stages of their life cycle including amoeboid 663 forms leaves much room for speculation. Filamentous, mm-long sporocaps, such as shown in 664 Fig. 3a in Rikkinen et al. (2019) are similar to what we see in Fig. 4b. The structured surfaces 665 shown in Fig. 6 are somehow similar to what Dagamac et al. (2017) showed in their Figs. 7-9 666 from recent Arcyria complex, though on the µm-scale, whereas those from the Volyn biota are 667 much larger. The image of multiple, conical filaments with claw-like ends, growing from a 668 common center (Fig. 3j, k, l) is similar to *Copromyxa protea* shown by Schnittler et al. (2012) 669 in their Fig. 4-2. Hollow objects (Fig. 7, i-k, m, n) resemble open sporocaps of liceaceae 670 (Schnittler et al., 2012, in their Fig. 5-12). Finally, large objects such as the open, bowl-shaped 671 one with bulbous outgrowths (Fig. 8) could be interpreted as plasmodium of a myxomycete 672 with beginning development of fruiting bodies (e. g. Fig. 2, life cycle of myxomycetes, 673 transition from stage H1 to A; Stephenson and Schnittler, 2016).

Other possible organisms described from the Precambrian are all very different from the Volyn biota and are excluded as possible analogues e.g. palynomorphs, which are among the earliest clear records of terrestrial life (Wellman and Strother, 2015); the 1.67 Ga eukaryotic Changcheng biota, (Miao et al., 2019), or vase-shaped metazoan microfossils, considered as the oldest evidence for heterotrophic protists, e.g. Urucum Formation, Brazil (Morais et al., 2017).

679 Most of the Precambrian biota listed in the literature are considered as photosynthetic 680 organisms, probably not a likely analog for the Volyn biota. E. g. the 770 Ma (Cryogenian) 681 Chichkan Fm. in Maly Karataou, Kazakhstan (Sergeev and Schopf, 2010) contains biota in 682 fine-grained black chert, which were deposited in a mid-shelf and a near-shore environment 683 with stromatolites. Most of the biota listed by Sergeev and Schopf (2010) are cyanobacteria, 684 rather small mostly up to the 10 µm range and thus do not serve as analogues for the Volyn 685 biota. They also list a number of larger protista (incertae sedis) in the 100 µm-range, however 686 with little morphological similarity to the Volyn biota. No similarity was found to eukaryotes 687 (acryitarchs) from 1.1 Ga old Taoudeni basin, Mauretania (Beghin et al., 2017). Red algae 688 (rhodophytae) from the 1.05 Ga Hunting Fm, considered as among the oldest eukaryotes 689 (Butterfield, 2000; Gibson et al., 2018) are photosynthetic organisms and can also be excluded.

690 **5.4 Model for a Precambrian deep biosphere ecosystem**

691 The Volyn occurrence is a well-preserved example of a fossil ecosystem of the deep continental 692 biosphere. We exclude an a-biotic origin as previously postulated (Ginzburg et al., 1987; 693 Lu'kyanova et al., 1992) because of the extremely low δ^{13} C values and the large variation in 694 morphology. A-biotic pseudofossils have been produced experimentally, e.g. by Nims et al. 695 (2021) and references therein, when sulfide is oxidized in the presence of organics. These 696 'organic biomorphs' show a large variety of morphologies, mostly filamentous, but also 697 globular. In a siliceous environment (for many cases chert) such organic biomorphs can be 698 replaced by silica, and their morphology can be well preserved. However, for the Volyn biota 699 such a sulfide rich environment did not exist. Additionally, we take the presence of chitosan as 700 another indication for a true fossil. McMahon (2019) provided another example of pseudo-701 fossils, which is however restricted to an iron-rich environment; these pseudofossils consist of 702 hematite or Fe-oxides/hydroxides, conditions not realized in the highly differentiated 703 pegmatites, which are very poor in Fe. Rouillard et al. (2018) produced another type of 704 pseudofossils with an amazing large variety of morphologies, which might occur in 705 hydrothermal, silica-rich rocks, but requires a high activity of Ba, for which there is no 706 indication in the Volyn pegmatites.

In combination with textural arguments, the age determination of muscovite, formed in
pseudomorphs after beryl, points to a minimum age of 1.5 Ga (Franz et al., 2022b); the
maximum age is restricted by the intrusion of the igneous rocks at 1.760 Ga (Shumlyanksyy et
al., 2021).

711 The geological context argues for a continental, terrestrial environment, because the KPC 712 intruded into continental crust most likely in a within-plate tectonic setting (Shumlyanskyy et 713 al., 2012, 2017). After intrusion uplift to the erosion level occurred, documented by an 714 unconformity, and sedimentation started with sandstones and shales at approximately 1.4 Ga 715 (Zbranki Formation; Gorokov et al., 1981), later than or coeval with the pseudomorph 716 formation and the minimum age of the microfossils. The depth, where the organisms lived, is 717 an open question, but the occurrence in the underground mines indicate a depth of up to at least 718 150 m. The age of 1.5 Ga is much later than the Great Oxidation Event of the Earth's 719 atmosphere, allowing for the evolution of complex species and ecosystems on the land 720 (sub)surface. The supply of organic matter to the underground for the production of the high 721 amounts of kerite is speculative. In a geyser system, which we invoke for the whole geological 722 situation, intense growth of organisms at the surface is a common observation. In such systems 723 continuous exchange between surface and depth is evident. This also excludes very deep (more 724 than several hundreds of meters) biosphere. The biota was more likely located near to the

surface. Unfortunately, no information is available right now, which of the many pegmatites
from the Volyn pegmatite field contains kerite and which – in what depth – are devoid of kerite.
This remains to be investigated in the future.

Drake et al. (2017) reported partly mineralized fungi from the deep continental (granitic) biosphere (up to 740 m). The fossilization process also included maturation of the OM and final mineralization by clay minerals. The source of carbohydrates was living or dead bacterial biofilms, similar to what we speculate about the Volyn biota.

- The large size of the filaments up to cm in length is a-typical for bacteria and archaea. Although Volland et al. (2022) described recent cm-long bacteria, these are still the exception, and it is more likely that some of the Volyn biota were multi-cellular eukaryotes. Their suggested age of 1.5 Ga is the age range given for the first appearance of eukaryotes (see review in Butterfield, 2015). Putative cm-sized Precambrian fossils (different from the Volyn biota) were reported from the 2.1 Ga old Francevillian biota (El Albani et al., 2014); however, they are completely pyritized and occur in diagenetically overprinted black shales, which makes the interpretation
- 739 difficult.

740 The Volyn biota must have been highly radiation resistant, because a U-Th-K-rich granitic-741 pegmatitic system has a high radiation level. There are a number of different organisms, such 742 as bacteria (e.g. Deinococcus radiodurans), archaea (Thermococcus gammatolerans) or 743 microscopic fungi (e.g. Cladosporium sphaerospermum), which fulfill this requirement; see 744 review in Matusiak (2019). During the mining operations in Soviet times, a high Rn content 745 was measured inside cavities, when they were broken into. The general radiation levels, 3000 746 times higher than the allowed limit at that time, were even higher 1.5 billion years ago. Deeply 747 black-colored quartz crystals in the pegmatites are of the 'morion' type and also indicate high 748 radiation. Recent observations at the Tschernobyl power plant have led to the speculation about 749 radiotrophic fungi (e.g. Matusiak, 2019; Prothmann and Zauner, 2014), which produce melanin 750 as a protection against radiation and enhancement of fungal growth via capture of ionizing 751 radiation for energy conversion (Dadachova et al., 2007; Tugay et al., 2017). Mycoremediation 752 is at least a well-documented mechanism for a very effective method of radio nuclides pollutant 753 removal considering the versatility of fungi in terms of their ecology, nutritional modes, 754 adaptability, morphology, physiology, and metabolism (Shourie and Vijayalakshmi, 2022). 755 Fungi are known as extremophylic organisms (e.g. Blachowicz et al., 2019) and we can expect 756 that in the Proterozoic or possibly already earlier in Earth history similar organisms were active

and resistant to a high radiation level, in an epoch when the ozone layer was not yet fullydeveloped.

759

760 **6 Summary and conclusions**

761 The exceptional 3D preservation of the 1.5 Ga Volyn biota is due to the fossilization conditions 762 in open cavities, with SiF₄-rich fluids as the driving agent. There are a number of indications 763 that fungi-like organisms were likely an important part of the microecosystem – hyphen with 764 branching (though not anastomosing), growth in thinning-out ends, and also in bulbous 765 extrusion, both at the end of filaments and along the filaments. Sheath-like structures are clearly 766 visible, and there are good indications for a former biofilm and extracellular proteinic 767 substance. The large size and internal structure of the organisms and the segmentation visible 768 on thick filaments points to multicellular organisms, and the nano-sized inclusions of Bi(S,Te) 769 crystals have an astonishingly good analog in recent fungi. Other organisms, which might have 770 been present in this subsurface micro-ecosystem are myxomycetes or myxomycete-like. The 771 stable N- and C-isotopic signature is in accordance with such an interpretation.

The fungi-like organisms possibly lived from lithotrophic methanogens; alternatively or additionally bacteria such as cyanobacteria were transported from the surface downwards into the cavities. The geyser system of the Korosten Pluton provided an ideal framework for growth of bacterial or algal organisms at the surface. In the deep biosphere, attached as well as freeliving forms of organisms are observed.

The Volyn biota show that fungi-like organisms developed before 1 Ga (Loron et al.; 2019), and support the speculation that the fossils from the 2.4 Ga Ongeluk Formation were fungi-like organisms (Bengtson et al., 2017). Molecular clock data, especially the three-way split of the kingdoms animals-plants-fungi at 1.58±9 Ma (Wang et al., 1999) are still uncertain, but our data indicate that it must have occurred early in the Proterozoic.

The Volyn biota also prove that a deep continental biosphere was already present in the Early Mesoproterozoic/Late Paleoproterozoic. It is known that in the subseafloor environment microbial life existed in the Archean (Cavalazzi et al., 2021), as described from the 3.4 Ga old Onverwacht Group of the Barberton greenstone belt, but from the continental environment this has not yet been reported.

787

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- 801

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