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Evidence for molecular structural variations in the cytoarchitectures of a Jurassic plant

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ABSTRACT

In this study, we investigate the molecular structural characteristics of organic remains in various cellular organelles from a 180 Ma Jurassic royal fern belonging to the Osmundaceae family of ferns, and compare their carbon isotopic compositions to a now-living species of royal fern (Osmunda regalis). We discovered molecular structural variations indicated by Raman and infrared spectral parameters obtained from various fossilized cellular organelles. The organic remains preserved in the chromosomes and cell nuclei show marked structural heterogeneities compared to the cell walls during different stages of the cell cycle. The fossil and extant ferns have similar δ13C values obtained from bulk samples, supporting evolutionary stasis in this plant lineage and an unchanged metabolic pathway of carbon assimilation since the Jurassic. The organic remains in the cellular organelles of the fossil seem to be less heterogeneous than those in the extant fern, likely due to the preferential preservation of certain cellular compounds during fossilization. Taphonomic processes appear to have diminished the subcellular isotopic heterogeneities. Our research sheds light on the functioning of ancient plant cellular organelles during mitosis, provides insights to the taphonomic processes operating at molecular and isotopic levels, and shows the practicability of in situ techniques in studying the evolution and behaviors of ancient cells.

INTRODUCTION

Exceptional preservation of fossilized cellular organelles is extremely rare but highly significant as these structures provide details of the evolution of life and functioning of ancient cells. The oldest fossilized cells with preserved ultrastructures can be traced to the putative planktonic organisms preserved in the 3.2 Ga Moodies Group, South Africa (Javaux et al., 2010). Complex intracellular organelles such as pyrenoids, nuclei, and lipid vesicles have been reported from Neoproterozoic strata (e.g., Hagadorn et al., 2006). Some of them, however, are still controversial and argued to be encysting protists (Huldtgren et al., 2011) or mineral inclusions such as apatite (Schiffbauer et al., 2012). More-abundant and highly diverse cells containing clear subcellular structures have been found within fossilized Phanerozoic land plants (e.g., Remy et al., 1994). In particular, a petrified Jurassic fern belonging to the family of royal ferns (Osmundaceae) from the Korsaröd locality, southern Sweden, contains exceptionally preserved cytoarchitectures, including cytoplasm, cytosol granules, nuclei, and chromosomes (Bonfleur et al., 2014).

Previous studies on the preservation of fossilized cells largely report morphological observations using imaging techniques such as scanning electron microscopy (e.g., Bonfleur et al., 2014), transmission electron microscopy (e.g., Hagadorn et al., 2006; Javaux et al., 2010), and X-ray transmission microscopy (Huldtgren et al., 2011; Schiffbauer et al., 2012), but fewer studies use a combination of Raman spectroscopy, Fourier-transform infrared spectroscopy (FTIR), and secondary-ion mass spectrometry (SIMS) to investigate the molecular structural and isotopic characteristics of organic remains preserved in fossils (Qu et al., 2018). Here we take the petrified royal fern fossil from the Korsaröd site in southern Sweden as an example, and further expand the investigation of molecular structural and isotopic characteristics of organic remains in various cytoarchitectures. We aim to explore the behaviors of organelles during the cell cycle of ancient fern, and test the use of in situ techniques for investigating the molecular structural and isotopic characteristics of organic remains in various fossilized cytoarchitectures. We further aim to better understand the taphonomic processes and their effects on the preservation of organic carbon, and in particular cellular components in volcanic “Konservat-Lagerstätten” deposits.

METHODS

One fossil specimen (Osmundaceae) was provided by the Swedish Museum of Natural History, Stockholm. The comparative extant fern Osmunda regalis was obtained from the Botanical Garden in Lund, Sweden. Two polished thin sections (~40 μm thick) from the fossil were prepared for petrographic and Raman analyses at the University of Bergen, Norway, and the PARI analytical platform at the Institut de Physique du Globe de Paris. The FTIR analyses were performed on two doubly polished wafers of the fossil at Beamline D7 in the MAX III ring in Lund, Sweden. The same Raman spectroscopic and FTIR analyses were performed on the extant royal fern, but unfortunately the signal bands cannot be distinguished from the high background noise and/or baseline absorption. The SIMS analysis was performed on the gold-coated...
surface of three polished samples (one fossil and two extant samples) at the Swedish Museum of Natural History. The carbon isotopic analysis on two bulk samples (one fossil and one extant) was carried out at the University of Bergen. More details of sampling, geological background, analytical methods, and strategies of data acquisition are presented in the GSA Data Repository1.

RESULTS
The fossil royal fern was entirely permineralized by calcite (Figs. 1A–1C) directly following its burial by volcanic lahar deposits at ca. 180 Ma, capturing cells in various stages of the cell cycle (Figs. 1E–1N). The Raman spectra of its molecular structural order, reflecting the molecular structural order of carbonate material (Qu et al., 2015). Some spectra contain an additional sharp peak at 1086 cm\(^{-1}\) (Fig. 1D), representing the calcite mineral matrix. Raman mapping performed on the fossil fern shows that the relative abundance of calcite (indicated by the intensity of the 1086 cm\(^{-1}\) band [I-Cal]) is complementary to that of organic matter (indicated by the intensity of the 1600 cm\(^{-1}\) band [I-1600]) (Figs. 1E–1N). The I-1350/I-1600 values, indicating molecular structural order of the organic matter, vary markedly and span larger ranges in the nuclei and cytoplasm than in the cell walls (Figs. 1C–1N and 2). Their variations seem to correspond to the morphology of various cytokinetries during different stages of the cell cycle (Figs. 1C–1N).

The FTIR spectra obtained from the fossil have distinct absorbance bands at 2850 cm\(^{-1}\), 2925 cm\(^{-1}\), and 2955 cm\(^{-1}\) (Fig. 3A), corresponding to the stretching vibrations of symmetric CH\(_2\) and asymmetric CH\(_2\) and CH\(_3\), respectively. The intensities of the bands at 2925 cm\(^{-1}\) and 2955 cm\(^{-1}\) show the relative abundance of organic matter in the cytoarchiteutres (I-2925 and I-2955 in Fig. 3). The IR parameter R\(_{2925}\), defined as the intensity ratio of the 2955 cm\(^{-1}\) band versus the 2925 cm\(^{-1}\) band, is widely used to describe the branching index of carbon chains (Igisu et al., 2009; Qu et al., 2017). The R\(_{2925}\) values obtained from the cell walls are higher than those from the cytoplasm (Fig. 2), which is further illustrated by IR maps (Fig. 3B).

Organic matter in the bulk samples of the fossil has a \(\delta^{13}C_{org-bulk} = -27.7\%e\) (Vienna Pee Dee belemnite), and in the extant specimens has a \(\delta^{13}C_{org-bulk} = -29.5\%e\) (Fig. 2). More-pronounced \(\delta^{13}C\) variations (up to 7.0%e) in the extant fern (Fig. 2). More-pronounced \(\delta^{13}C\) variations (up to 7.0%e) in the extant fern occur within different cytokinetries as measured in situ by SIMS (Figs. 2, 4A, and 4B). Both the fossil and the extant fern have lower absolute values of \(\delta^{13}C_{org-SIMS}\) than the respective bulk samples (Fig. 2). The counts of \(^{13}C\text{N}\) ions in the ion images (Fig. 4) provide a semiquantitative representation of the intracellular abundance and spatial distribution of organic matter, which are consistent with the Raman and FTIR maps (1600 in Figs 1E–1N; I-2925 and I-2955 in Fig. 3B).

DISCUSSION
Distinguishing Primary Biosignatures from Secondary Alteration
The nature of sedimentary organics is determined by the molecular structure of their precursor biomass and by subsequent alteration.
(Qu et al., 2015; Vajda et al., 2017, and references therein). The primary components of the plant cell wall are polysaccharides, including cellulose, hemicellulose, pectin, and in some cases lignin (Buchanan et al., 2015). These compounds contain few functional groups of CH₂, as indicated by higher R₃/₂ values (Figs. 2 and 3). The structural order of organic remains derived from polysaccharides in cell walls did not change markedly during the cell cycles as implied by the relatively constant I-1350/I-1600 of cell walls compared to that in the nucleus and cytoplasm (Figs. 1 and 2). In contrast, the cytoplasm and membrane contain abundant CH₂ on the long carbon chains (e.g., lipids and proteins), as indicated by lower R₃/₂ values of the cytoplasm from both the inner and outer cortex compared to their cell walls (Figs. 2 and 3). This observation is consistent with a previous study on Devonian (410 Ma) fossilized plants from Rhynie chert (Scotland) (Qu et al., 2015). The nucleus and cytoplasm contain abundant heteroatoms (e.g., P, N, S, O, H, in the protein and nucleic acid, and other metal elements such as Ca, Mg, K). Moreover, the molecular structures of nucleic acids (DNA and RNA) vary significantly during mitosis (Buchanan et al., 2015). Therefore, the diverse chemical compositions of the carbon precursors could explain the marked molecular structural variations in the organic remains of the fossilized nucleus and cytoplasm, as recorded by larger variations in I-1350/I-1600 relative to that of the cell walls (Figs. 1 and 2).

The carbon isotopic composition of an organism is determined by its assimilated carbon source and the metabolic pathways utilized (Hayes, 2001). The similarity in carbon isotopic values between the fossil (δ¹³Corg-bulk ~ -27.7%o) and extant (~29.5%; Fig. 2) fern implies the same metabolic pathway and coincides with evolutionary stasis as proposed for the fossil Osmundaceae (Bomfluer et al., 2014). The minor difference (1.8%o) in δ¹³Corg-bulk observed between the fossil and extant ferns (Fig. 2) could perhaps reflect slightly ¹³C-enriched Jurassic atmospheric CO₂ globally and/or locally, or different water-use efficiency (Farquhar and Richards, 1984) induced by dissimilar Jurassic atmospheric CO₂ concentration. Alternatively, the higher δ¹³Corg-bulk (by 1.8%o) of the fossil may result from the preferential loss of isotopically light compounds combined with the preservation of ¹³C-enriched organic remains during post-depositional processes (Clayton, 1991), which is consistent with the higher average δ¹³Corg-SIMS values from the fossil fern (Fig. 2).

Concerning the in situ carbon isotope measurements, the low δ¹³Corg-SIMS values from the extant fern could be the result of a matrix effect related to the presence of water. House (2015) showed that hydrated organic materials cause a matrix effect during SIMS analysis, leading to an apparent observed ¹³C depletion. Because we used graphite as a standard (virtually free of water) during the analysis of the hydrous extant fern sample, we likely caused this water-related matrix effect. In the fossil fern, this effect may have been less, because the material is less hydrous than in the extant fern. It is therefore important to note that we do not interpret the absolute δ¹³Corg-SIMS values, but only the internal variations within the extant and fossil ferns, respectively.

The variations of δ¹³Corg-SIMS in different cellular organelles (Fig. 2) possibly reflects diverse organic compounds with heterogeneous isotopic compositions. Indeed, isotopically heterogeneous cellular compounds from modern terrestrial plants with ¹³C variations >10%o have been reported (e.g., Collister et al., 1994). However, terrestrial plant compounds preserved in sediments generally have less carbon isotopic heterogeneity (variations in δ¹³C of a few per mil; e.g., Naraoka and Ishiwatari, 2000). In this study, the narrower ranges of δ¹³Corg-SIMS (Fig. 2) from the fossil fern could imply preferential decomposition and preservation of certain compounds (Igisu et al., 2009) during post-depositional processes, and thus the taphonomic processes seem to have diminished the subcellular isotopic heterogeneities.

**Ultrastuctural Variations in Chromosomes during Mitosis**

Here we characterize specifically the molecular structural variations of organic matter in the nuclei and chromosomes preserved in the cell cycle of the fossil Jurassic royal fern (see more details in Figs. 1E–1N; and in the Data Repository). The nucleus is a dynamic organelle whose morphology and chemical composition change across different stages of the cell cycle. The nuclear envelope and matrix, predominantly consisting of lipids, fibrin, protein, and minor RNA, decompose and reconstruct throughout mitosis (Buchanan et al., 2015). The nucleolus, consisting of abundant RNA and protein, is also highly dynamic and experiences transformation, decomposition, and regeneration in the cell cycle (Buchanan et al., 2015). The chromatin...
and chromosome have a chemical composition of mainly DNA and protein, with minor RNA. During mitosis, their chemical compositions and sequences of nucleotide generally remain unchanged, but molecular structures of the DNA chains experience three-dimensional packaging, folding, and twining with the involvement of various enzymes (Buchanan et al., 2015). This could be a reason for more-pronounced molecular structural variations in the nuclei, as is indicated by large ranges of $\delta^{13}C$ for the extant fern compared to those of the Jurassic fern during the cell cycle and their preservation during post-depositional processes. Our study provides unprecedented insights into the characteristics and behaviors of cellular organelles of a Jurassic fern during the cell cycle and their preservation during post-depositional processes. This study also demonstrates the use of in situ techniques of Raman spectroscopy, micro-FTIR, and SIMS in studying ancient cellular organelles.

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