Chemical identification of microfossils from the 1.88-Ga Gunflint chert: Towards empirical biosignatures using laser ablation ionization mass spectrometer

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Abstract
In this contribution, we investigated the chemical composition of Precambrian microfossils from the Gunflint chert (1.88 Ga) using a miniature laser ablation ionization mass spectrometer (LIMS) developed for in situ space applications. Spatially resolved mass spectrometric imaging (MSI) and depth profiling resulted in the acquisition of 68,500 mass spectra. Using single mass unit spectral decomposition and multivariate data analysis techniques, we identified the location of aggregations of microfossils and surrounding inorganic host mineral. Our results show that microfossils have unique chemical compositions that can be distinguished from the inorganic chert with high fidelity. Chemical depth profiling results also show that with LIMS microprobe data, it is possible to identify chemical differences between individual microfossils, thereby providing new insights about nature of early life. Analysis of LIMS spectra acquired from the individual microfossils reveals complex mineralization, which can reflect the metabolic diversity of the Gunflint microbiome. An intensity-based machine learning model trained on LIMS Gunflint data might be applied for the future investigations of putative microfossils from silicified matrices, where morphological integrity of investigated structures is lost, and potentially in the investigation of rocks acquired from the Martian surface.

KEYWORDS
Gunflint, Mars, mass spectrometry, microfossils, space instrumentation

1 | INTRODUCTION

In situ research and remote sensing have provided multiple lines of evidence that clement conditions were present on the surface of early Mars.1–3 Moreover, recent radar studies reveal evidence of subglacial liquid water on Mars,4 which supports the hypothesis that microbial life forms (extinct or extant) may be preserved within the Martian subsurface.4,5

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All these observations provide a strong rationale for the search of biosignatures on the Red Planet. The current state of space exploration provides foundation for new measurement techniques and novel analytical approaches to identify and characterize minerals and potential signatures of life, if any, on Mars. However, in situ investigation of rocks on Mars faces multiple technological difficulties, ranging from constraints on instrumentation robustness, size, and power consumption to the quality of the acquired data. Some authors proposed an implementation of multicriteria approach to confirm or reject a biogenic origin of the given sample. Multicriteria approach, thus, requires several instruments onboard of the rover providing a multiplex analysis of the same sample and identification of morphological, molecular, elemental, and isotopic signatures of life. However, traditional methods used in space research, for example, bulk analysis and remote sensing instrumentation, might not be sufficiently sensitive to detect faint features from micrometer-sized (and below) organic material or microbial remains and, in some cases, can alter the chemistry of the sample. There is a growing demand for sensitive, in situ instruments with high spatial resolution and minimal sample processing, providing elemental and organic composition detection, which will enhance the scientific return from the missions to Mars and icy moons of Jupiter.

In addition to the development of analytical methods, development of chemometric tools has also proven to be a field of high importance to the current and future space exploration programs. For example, multivariate curve resolution alternating least squares have been shown to successfully identify various minerals and compounds from Raman hyperspectral images, overcoming the spectral overlap issues. The linear mixture model (LMM) was successfully used to quantify the abundance of major elements using the laser-induced breakdown spectroscopy (LIBS) spectra. Furthermore, a data fusion approach was reported for complementary analytical techniques (Raman and LIBS) that improved the classification limits of investigated binary compounds.

Laser-based mass spectrometry is an emerging and sensitive technique that has shown to be capable of measuring extremely low concentrations (fmol) of amino acids in desorption mode, elemental detection of single microbes in Martian mudstone analog material, and provides chemical (element, isotope) analysis in ablation mode of any solid material. The latter can be conducted with a depth resolution on the scale of tens of nm and with high element detection sensitivity down to the ppb level. This makes laser-based mass spectrometry an attractive method in the field of in situ chemical analysis on planetary surfaces. The upcoming ExoMars mission/Rosalind Franklin Rover contains a Laser Desorption/Ionization (LDI)-Quadrupole Mass Spectrometer in its instrument suite, stimulating further development of LDI instruments. However, laser ablation ionization mass spectrometer (LIMS) capability to detect and identify billion-year-old microfossils has not been shown so far.

Since the discovery of Precambrian microbial communities in the 1.88-Ga Gunflint Formation (Ontario, Canada) in the early 1960s, many more examples of Precambrian life have been found, but the Gunflint Formation retains its place as a premier Precambrian Fossil-Lagerstätten, demonstrating that Paleoproterozoic life was widespread, already complex, and diversified. The Gunflint chert sample in this study has been used as a Martian analog, reflecting the iron-rich nature of the Martian sediments, as well as taking into account that siliceous sediments have high preservation capacity and are of interest to upcoming astrobiological missions. However, despite being among the best example of Precambrian life, phylogenetic affinities and metabolic speciation of the Gunflint microfossils remain largely unknown. Similarly, microfossils of unknown affinities dominate the majority of the Precambrian record. Traditionally, the classification of types of microfossils has relied on morphological features and later advanced to include isotope fractionation and multielement nano-characterization of individual microfossils. However, within the space instrumentation domain, detection of individual microfossils remains a technological challenge.

In this contribution, we present results on mass spectrometric imaging (MSI) and chemical depth profiling from the Gunflint chert using a miniature time-of-flight reflectron-type mass spectrometer developed for in situ space applications, equipped with a femtosecond (fs) UV-258-nm laser ablation ion source. Mass spectrometric studies conducted on microfossils allowed their identification within the inorganic host material. Utilizing the depth profiling approach, we removed the contaminated layer present on the rock surface and probed the original chemical composition of the microfossils. The network-based approach used for the interpretation of hundreds of recorded mass spectra revealed a new topological dimension, where separate mineralogical inclusions present within the same analytical spot (inclusions smaller than the size of the ablation spot) can be readily separated.

The analysis of the mass spectra from the depth profiles revealed the presence of major biorelevant elements (CHNOPS), microscopic inclusions of Cu, Cr sulfides, rare earth element (REE) minerals in addition to the majority of Fe-dominated mineralization associated with the microfossils. These observations can indicate the presence of sulfur-processing and iron-processing species but also can indicate the presence of intracellular bio-mineralization machinery...
(passive mineralization) within the Gunflint microbiome to withstand possible Cr and Cu toxicity within an already highly ferruginous environment. H/C, O/C, Si/C ratios, and principal component analysis (PCA) scores calculated from the depth profiling dataset show ratios and intensity regions in which microfossils can be identified. Large-scale mass spectra sampling allowed the construction of binary classification machine learning (ML) models, which can be used for the identification of microfossils from other Precambrian cherts and other rocks (once calibrated), where morphological integrity of the putative microfossils is lost, thus providing a way to assess biogenicity, by comparing spectra from other cherts to our truly biogenic model data.

Overall, the LIMS microprobe shows an ability to identify micro- and nano-mineralization associated with the microfossils. LIMS imaging combined with accurate depth profiling has the potential to reveal new insights into the distribution, preservation, and elemental speciation of microfossils from Precambrian cherts and a potential to deliver insights into the chemical composition of samples acquired from the Martian surface.

2 | MATERIALS AND METHODS

A standard double-polished thin section of the Gunflint chert (collected from the Schreiber beach locality, Ontario, Canada) with a thickness of ~30 μm was used in the current study. The sample was mounted on a metal sample holder with vacuum-compatible copper tapes to fix the sample on the surface of the steel holder. No additional treatment of the sample was performed. A miniature time-of-flight mass spectrometer (TOF-MS) developed at the University of Bern has been used to study the Gunflint sample. The mass analyzer has small dimensions: Ø 60 mm × 160 mm, which makes it suitable for space exploration programs as part of the lander or rover. The time-of-flight mass analyzer works in the positive ion detection mode and provides a single unit mass resolution and a ppm-level sensitivity. For the detailed characterization of the instrument and method in general, we refer the interested reader to the reviews and technical reports describing figures of merit. In the current laboratory environment, a mass spectrometer is accompanied by an integrated microscopy system and a fs UV-258-nm ionization source. The instrument is designed to have spatial molecular, elemental, and isotopic mapping capabilities of solid samples. The microscopy system utilized in the current laboratory environment is not space qualified; however, a separate space prototype was developed in our group that combines the microscopy system, mass analyzer, and an ion source. High precision XYZ translation stage is used for the accurate sample positioning under the instrument: 1-μm positioning accuracy is typically achieved between the internal microscope and laser focal point positions. Laser power output stability, beam profile, and crater shapes are checked prior to the measurement campaign.

The Gunflint sample studied in this work could be approximated as a dielectric glass with inclusions of dark absorptive features (microfossils). To enhance the ionization efficiency of the quartz mineral, which constitutes most of the sample, and shorten the gap between the ionization efficiency of microfossils and host mineral, we implemented a frequency tripling system (STORC) into the beam delivery line, which allowed us to reach a stable fs UV-258-nm laser radiation. Photon energy, at this point, reached 4.8 eV, which is well suited for the ablation and ionization of any solid materials. Ti:sapphire fs Laser from Clark Inc. generates infrared (IR) 775-nm fundamental wavelength (180 fs), which is guided into the frequency doubling and tripling system. β-BBO crystals are used in the doubling and tripling parts of the STORC system to achieve UV light generation.

In this contribution, we performed two separate data acquisition campaigns: (1) the MSI of the 1.5 × 2-mm area with 10-μm gaps between ablation craters and (2) depth profiling campaign within specified areas—microfossil-rich zone and a host mineral (quartz) area. The MSI campaign yielded 30,000 (150 × 200 pixels) mass spectra, where every pixel represents a histogram of 200 single laser shot spectra. The output mass spectra were processed using a single mass unit spectral decomposition (extraction of consecutive mass peak areas, following the footsteps of Meyer et al., 2017). Overall, 182 single mass unit intensities, retrieved from every pixel, were used to form the isotope intensity maps. Further, the Gaussian process (Kriging) interpolation was used to increase the resolution of the output maps by a factor of 2. In total, $^{28}$Si, $^{16}$O, $^{12}$C, and $^1$H maps were calculated to visualize the distribution of microfossils and filling quartz mineral on the surface of the sample. The second data acquisition campaign was performed to remove the organic contamination layer present on the surface and probe the original chemical composition of the microfossils and subsequently build a binary classification model. The depth profiling campaign resulted in the acquisition of 38,500 spectra from 15 depth profiles acquired from the microfossils-rich zone and a host mineral (see the supporting information for detailed information about data processing and filtering). The output spectral intensity profiles from both locations were log-transformed and z-score standardized.
The PCA of the depth profiling dataset was conducted using the correlation matrix computed on centered data. The first three principal components were extracted from the depth profiling dataset. The weighted correlation networks of inclusions present in the depth profiles were calculated using the direct Pearson pairwise correlation scores of 182 single unit mass intensities. The correlation scores retrieval resulted in the acquisition of 16,380 Pearson correlation pairs for given inclusion. Further, extracted correlation pairs were used as weights defined on pairwise edges in the construction of the undirected network. The force-directed layout (ForceAtlas2) provided with Gephi\textsuperscript{46,49} was used to visualize the network structure. The edge weight threshold was implemented to remove the insignificant correlation values from the network (see further down in the text). The pairwise kernel density estimates of biorelevant ratios and element intensities were used to visualize the density distributions of investigated locations (silicified host mineral and microfossils). Furthermore, 24 binary classification ML models (including classification trees, support vector machines, and ensemble models) were scored using the Matlab ML presets (see supporting information for more details). The isotope ratios and synthetic metrics (e.g., geometric mean values of light masses) were added to the ML dataset, making 196 variables in total. The fivefold cross-validation was used to avoid overfitting of the dataset. Within best-performing models, an additional 30-step Bayesian optimization (search through different learning hyperparameters) procedure was implemented to test for potential improvements in the output performance. For a more detailed description of ML models, information on PCA, and weighted correlation networks, we refer to the supporting information.

3 | RESULTS AND DISCUSSION

3.1 | LIMS imaging and depth profiling

All experimental measurement procedures—surface imaging and depth profiling—performed on the sample with a miniature LIMS system are schematically illustrated in Figure 1. The drawing is out of scale and intended to give a better understanding of the subject of this study. The distribution of microfossils embedded in a quartz matrix is shown with gray lenticular structures. Most of the preserved species studied by nano-microscopy are hollow and represent partially collapsed cell walls. An approximate estimate of the thickness of the cell walls mentioned in the figure is 500 nm; however, actual thicknesses are varying. For detailed morphological studies of these microfossils, we refer to literature\textsuperscript{35,39} The small layer on the surface represents surface contamination with recent organic material. A focused UV-258-nm fs laser beam shown in Figure 1 was used to ablate and ionize material from the Gunflint chert. The produced ions were transmitted into a miniature TOF-MS (LIMS) developed for the operation on planetary surfaces.

To acquire information on the chemical composition of the stromatolitic layering from the Gunflint chert sample, we performed MSI of an area of a thin section containing two morphological features: (1) dense population of microfossils and (2) clean host mineral area. We identified a rectangular area (1.5 mm × 2 mm) (Figure 2A,B) where these features were present. To accurately sample the area under investigation, a 10-μm gap between the ablation craters was chosen for the imaging, resulting in a grid with 30,000 ablation spots (with a single mass spectrum corresponding to each spot—see Figures S5 and S3–S5). To avoid material displacement and crater-to-crater cross-contamination from the ablation processes, a pulse train of 200 laser pulses was applied to each location, yielding a single mass spectrum. Because imaging implies probing material from different parts of the sample with different light absorption properties, suitable pulse energies were determined on preliminary craters from various locations prior to the imaging. Laser pulses with an energy of 0.36 [μJ] per pulse (measured at the sample surface) were found to be appropriate for both the dense microfossil assemblage and the clean host area. The diameter of the analytical spot was determined to be around 4–5 μm for the imaging campaign within dominantly quartz locations and 7–8 μm within microfossils.

Figure 2D,E shows the distribution of \(^{1}\text{H}\) and \(^{12}\text{C}\) signal intensities on the surface of the Gunflint sample, extracted from the mass spectra using Simpson integration (details about integration procedure can be found in literature\textsuperscript{50}) and defined as an output current registered from the detector per unit of time (log\(_{10}\) electrons * ns\(^{-1}\)). As is clear from Figure 2D,E, hydrogen and carbon are spatially correlated with the location of the microfossils identified by the optical microscopy (see Figure 2B). Resolution of the imaging reached a single cell level (see Figures S4 and S5) and could be improved by a factor of 2 in future campaigns without any analytical interference. Figure 2F,G displays the distribution of \(^{28}\text{Si}\) and \(^{16}\text{O}\) intensities recorded on the surface of the sample. Both isotopes show a relatively homogeneous distribution, with a good correlation to each other. The host material in which the microfossils are embedded is diagenetic quartz with varying crystal sizes. It is also possible to observe an enhanced signal of \(^{28}\text{Si}\) and \(^{16}\text{O}\) within the microfossil’s lamination area due to the enhanced absorption of light by microfossils. Imaging of minor and trace elements was not
possible due to isobaric contamination by hydrocarbon clusters in the mass spectra (see Figure S12). The separate depth profiling campaign was performed to remove the contaminated layer on the surface and probe the original chemical composition of the microfossils and host area.

3.2 Depth profiling

Figure 3 shows the locations at which depth profiles were measured within the lamination area containing the population of microfossils and the host quartz area. Depth profiling analysis was conducted on nine spots containing microfossils and six spots of clean host quartz. Spots studied by depth profiling are independent of the grid that was used for MSI. For seven spots from the microfossils and six spots from the host area, 2500 spectra were measured simultaneously on two acquisition cards and summed together (forming a 5000-spectra dataset for a single depth profile before summation), where each spectrum consists of 200 single laser pulse spectra and corresponds to the single ablation layer. Two additional spots on a microfossil-rich area were measured with reduced histogramming down to 64, and 32 single laser shot spectra, in an attempt to obtain an even finer sampling of the microfossils. In total, 3000 spectra were collected from each of these spots (6000 spectra before preprocessing). A data extraction procedure is performed, retrieving intensities of the single mass units, utilizing direct Simpson integration of the time-of-flight windows determined for each mass. In total, 182 single unit masses from each spectrum have been retrieved, including a background signal (noise measurements), which were determined in the time-of-flight window, free of any ion signal.

Figure 3A,B displays location and morphology of exemplary craters acquired during the depth profiling campaign. The dark patches in Figure 3B represent an aggregation of microfossils, where an arrow indicates location of the single lenticular microfossil. Figure 3C shows variation of the $^{12}$C mass peak intensities measured at the selected location. As is clear from Figure 3C, the $^{12}$C mass peak intensity measured within 32,500 spectra reveals clear intensity separation.
boundary between two locations (note that intensities are presented in the log scale). A limited amount of $^{12}\text{C}$ signal, registered from the host region, may originate from the surface of the sample due to the widening of the crater, while ablation is progressing. Figure 3D shows the single depth profile from the microfossil-rich location and reveals increased $^{12}\text{C}$ intensities within specific depth regions, indicating that the measured carbon originates from the inclusions (Figure 3D, bars with location X and Y). In contrast, the depth profiles registered from the host region (Figure 3C) show the presence of a significantly reduced amount of carbon, in comparison with the depth profile acquired from the microfossil-rich lamination area. Localized aggregation of intense peaks of carbon within the bulk of the quartz matrix
is interpreted to be individual bodies of microfossils (see the sketch—Figure 1). Moreover, carbon-enriched inclusions are associated with other biorelevant elements: CHNOPS (see further in the text and Figure 4B,C), which indicates that these inclusions are indeed individual microfossils, located in the distinct depth regions. However, to prove that these localized spectra are acquired from a single source (microfossil), we calculated the correlation networks, which will be presented later in the text. Here, we need to mention that LIMS, being a destructive method, provides sensitive and spatially resolved measurements that are hard to achieve using bulk characterization methods. For example, in the low-biomass simulation of Martian sediments,10,11 the results have shown that LIMS can identify spatially constrained biosignatures in Mars analog environments.

Figure 4 shows spectra measured from different mineralogical inclusions present within the Gunflint subsurface. The spectrum measured at the host area (Figure 4A) reveals the chemical composition corresponding to the diagenetic quartz. Intense peaks of Si and O are readily recognized. Additionally, peaks of H, C, Na, K, and chain of SiO clusters can be identified in the spectrum (Figure 4A). The mass spectrum in Figure 4B was measured within location X (Figure 3D) at depth position 250–500. The spectrum reveals the complex chemical composition of the microfossil body intermixed with the host chemistry. The chemical composition measured within this spot represents a mixture of the
quartz mineral, kerogen from microfossil cell walls, and a polymetallic inclusion associated with the microfossil. Intense mass peaks of transition metals, Ti, Mn, Fe, and Cu, can be identified in the spectrum with a major contribution from Cr and additional minor contribution from La and Ce monoxides. The presence of kerogen is identified from the detection of C, H, N, O, P, and S. Additionally, multiple low-intensity CxHy compounds can be identified in the spectrum.

Figure 4C shows the mass spectrum measured at depth locations 2350–2500 (location Y; see Figure 3D) and reveals a different composition compared with the inclusion described above. In addition to the mass peaks registered from quartz and CHNOPS, there are also peaks of Mg, Al, K, Ca, Cr, Mn, La, and Ce, and mass peaks of Cu and Ag could be noted. The latter are rather unexpected to be found within microfossils because they are known to be elements with high cytotoxicity (i.e., they are toxic to cells).

Although LIMS can yield sensitive measurements of elements and isotopes, it can be challenging to determine mineralogical composition of multiple microinclusions (smaller than the size of the probing laser spot). The pairwise correlation factors between single mass unit intensities are calculated in an attempt to identify mineralogical composition of investigated inclusions. Figure 5 shows two networks calculated from the individual microfossils (Figure 3D) and
visualized using an open-source graph drawing platform Gephi. The mass correlation network shown on the left side was calculated from location X, which is located at the depth region: 250–500. Correlation factors ($\rho$) larger than 0.4 were used to visualize only strongly correlated masses (see full correlation matrices in the supporting information). Node colors are chosen according to the modularity rank, calculated from the network topology (modularity $Q$ is a parameter that measures the density of links in the graph inside communities as compared with links between communities). Blue nodes correspond to the elements registered from the quartz mineral, whereas red nodes represent elements and isotopes registered from the microfossil body. Clear separation of the inclusion from the host chemistry could be observed within this network—host mineral (quartz) colored with blue nodes and a microfossil (red nodes).

The mass correlation network shown on the right side of Figure 5 was calculated from the depth region 2350–2500, at location Y (see Figure 3D), and represents the chemical composition of another microfossil. Due to the lower number of spectra registered from this inclusion (size of the inclusion was considerably smaller than the inclusion described above), the $\rho$ cutoff value was set to 0.2. However, even with low correlation factors, it is possible to obtain the appropriate modularity ranks and to increase the interpretability of the data. Within the analyzed microscopic inclusion, S is interconnected with Cu, implying the presence of covellite within the body of the microfossil (see Figure 5, location Y, top red nodes), or closely attached to it. P measured in the microfossil is more interconnected with the host (quartz) elements (left blue node in network Y). This observation can be attributed to the “shouldering” effect of the intense $^{30}$Si peak, which affects the integration window of P, thus modifying its $\rho$ value and positioning in the graph. One notable feature of this correlation network is that by identifying the least and most interconnected nodes, it is possible to find predictive masses that are unique for each of the given mineralogical classes. The topology of the correlation networks also reveals a centrality measure, which indicates the importance of the element in a network. We can see that in Figure 5 left, there are some elements that are present both within the quartz and a microfossil—H, Na, K, O—these nodes could be characterized with high betweenness centrality. $^{56}$Fe is also among the central elements in a network, due to the presence of the isobaric contribution of Si$_2$. Analyzing the microfossil-related network separately (only red
nodes), one can see that $^{12}$C, $^1$H, $^{41}$K, $^{54}$Fe are among the central isotopes, which also reflects the importance of this isotope in the chemical composition of the microfossil bodies. Any organic matter preserved within the bulk of the host mineral requires the existence of such networks with measurable centrality of C and H and a separate modularity rank. Depending on the chemical integrity (state of decay) of the inclusions, better preserved microfossils also have better connectivity and higher covariance, whereas finely dispersed carbon incorporated into the body of the matrix will not have such metrics and likely to be located within the outer nodes of the network.

The co-occurrence of P, Ce, and La (Figure 4B,C) within studied locations is indicative of the presence of the monazite microscopic inclusion. There are two possible interpretations for the presence of REEs in association with the microfossils: first is the intracellular incorporation and passive mineralization of La and Ce by living organisms and, second, postmortem mineralization (secondary incorporation of these elements). However, because La and Ce have been measured within our dataset only in association with microfossils, intracellular incorporation seems more plausible. The connection of Fe with S (Figure 5 left) indicates the presence of pyrite, which was described as a byproduct of the metabolic activity of sulfate-reducing microbes. However, localization of metallic nodes in the lower part of the network indicated presence of the third inclusion, which likely represents Cr-rich nodule with impurities of Fe, Mn, Cu, and Ti (see bottom red nodes in Figure 5, location X). The complex chemical composition associated with some of the microfossils might also be indicative of the development of tolerance to the polymetallic toxicity. For example, Cu, registered at location Y, is an essential trace element for aerobic organisms; however, Cu might be lethal to microbes if homeostasis is not maintained. Cr has also been reported as a highly toxic and mutagenic element for bacterial colonies. The C and H bearing microfossils reveal close association with Na, K, Mg, Ca also with Fe and S. Overall, analysis of the inclusions from the depth profiles reveals a complex chemical composition indicating presence of chemically distinct microfossils with identifiable hydrothermal mineralization patterns (presence of typical hydrothermal elements like Cu, Fe, and Ag).

### 3.3 Identification of empirical biosignatures

As it was shown in Figures 2, 4, and 5, carbon and hydrogen peak intensities are correlated with the location of the microfossils, and within our sample, carbon might be used as a tracer of the microfossils. The full depth profiling dataset sampled from the microfossil-rich locations was divided into host and microfossil data by thresholding the carbon signal. Depth profiles from the microfossil locations were sorted using a threshold of 5.8 log$_{10}$ el/ns (higher than the noise level) to create a subset of data that represent only microfossil-related spectra, assuming that the C signal originates from the microfossils. The depth profiling dataset was additionally filtered to the depth region 500–2500 within both locations (host mineral area or aggregation of microfossils), to avoid contribution from the surface data. In total, after filtering, we formed a dataset with 12,000 spectra from the host location and 1454 spectra from the microfossil-rich location. Figure 6 presents kernel density estimates (two-dimensional density maps) calculated for two specific regions and represents a probability distribution function of element intensities for two groups: quartz and microfossils (i.e., inorganic or bioorganic intensity regions). Figure 6A–D represents the variation of the signal from the microfossils plotted against the same mass intensities measured from the host region—red kernels are calculated from the microfossil-rich location, and blue kernels from the host area (see Figure 3A,B). Multiple non-overlapping intensity regions associated with the microfossils could be observed. These intensity regions can serve as predictive borderlines for the identification of the organic remnants from other Gunflint-like cherts. In contrast, Figure 6E–G represents the variation of the elements associated with the inorganic host, shows mostly overlapping intensities, and indicates that most of the spectra from the microfossils have a significant contribution from the quartz mineral. Figure 6F shows that the Fe signal registered from the microfossils interferes with the Si$_2$ molecule, and significant parts of it protrude into the higher Fe content area, indicating increased Fe content within the microfossil bodies. Figure 6H displays a perfectly overlapping variation of the Gaussian background signal derived from the two locations.

Figure 7A shows partially overlapping clusters of $^{1}$H/$^{12}$C and $^{16}$O/$^{12}$C ratios measured from the host (blue kernels) and the microfossils (red and orange kernels). Because most of the microfossils are hollow (see Figure 1) and smaller than the LIMS analytical spot size, they will be sampled with the encapsulating host mineral. In addition to the chemical composition of the microfossils, compositional details of the host mineral are likely to be registered. Hence, intensity values of $^{1}$H and $^{16}$O, formerly occurring within microfossils, are interfering with the same isotopes from the quartz mineral; hence, they can be subtracted. The results of these corrections are shown in Figure 7A,B with orange kernels. As can be seen from this figure, the locations of the kernels from the microfossils coincide with empirically determined
regions of organic compounds (lipids, peptides, sugars, and condensed hydrocarbons [kerogen]). The identification of ratio boundaries for different organic compounds was demonstrated in the literature using ultra-high-resolution mass spectrometry. Black boxes shown in Figure 7A schematically represent the location of those boundaries. Empirically determined shapes of the complex organic compounds are derived at overlapping areas and typically have more complex shapes to those presented on the plot.

Data collected from the densely populated microfossil area (Figure 7A red and orange kernels) hint at the presence of lipid and peptide signatures, which aligns with the previously reported identification of amides from the Gunflint microfossils and exceptional preservation capacity of cherts. However, they likely represent a mixing ratio between the original kerogen and quartz ratios. A fraction of the data also intrudes into the area of kerogens, which can be classified as a Kerogen Type I (Algal). However, there is a part of the red kernels that overlap with the host mineral data as well. Data collected from the host area contain mostly quartz mineral, and most of the measured ratios represent signal-to-noise ratios of $^{16}\text{O}$ and $^1\text{H}$, due to the low concentration of $^{12}\text{C}$ within analyzed depth profiles from the host locations. The results presented in Figure 7A indicate that organic hydrocarbons measured at the Gunflint sample are significantly reduced in $^{16}\text{O}$ and $^1\text{H}$, due to the low concentration of $^{12}\text{C}$ within analyzed depth profiles from the host locations. The same correction procedure as in Figure 7A has been applied for $^{16}\text{O}/^{12}\text{C}$ distribution, which is shown with orange kernels.

Figure 7C,D shows the distribution of PCA values obtained from the two distinct groups—measurements from the host area in contrast to the measurements from the microfossil-rich zone (12,863 spectra in total). Spectra from the microfossil location could be separated from the spectra collected within the host area, except for a small number,
which interferes with the host measurements. To increase the separability of the dataset, we further thresholded C intensities to $6 \log_{10} \text{el/ns}$ (see supporting information), which reduced the microfossils dataset from 1454 spectra to 863 spectra. We schematically identified the transition boundary with a light blue transparent line, which shows the location of the estimated transition boundary between the inorganic host (quartz) spectra and spectra from the microfossils. The transition of one class into another could be explained by the ablation of the small portions of the microfossils (nm thick cell walls in the bulk of the host mineral). The thickness of the rims of the collapsed cell walls reported from the Gunflint microfossils varies from the tens of nm$^{39}$ to first micrometers$^{35}$, which might explain statistically...
more prevalent low-intensity regions in KDE plots (Figures 6 and 7). Figure 7C,D indicates that intermediate levels between classes are sampled when spectra from one class gradually turn into the spectra from the other class. It is worth noting that even with only the first two principal components, a clear separation between two main classes can be seen. Additionally, as was demonstrated before (Figures 4B,C and 5), and noting the dispersion of the PCA loadings, it is possible to identify that there are potentially more than one class of microfossils within the Gunflint dataset. A significant part of the dataset is clustered within the dark red areas, representing the majority of simple kerogen containing microfossils. However, we could see from Figure 7C that kernels protrude from the hydrocarbon saturated area towards areas with notable metallic content, pointing towards the presence of uptake of Fe, Mn, and Cr. This observation agrees with the results reported previously on the diversity of the microbiome within Gunflint waters. Cyanobacteria were proposed to be a dominant part of the Gunflint stromatolites31,40,58; however, other interpretations are possible. Presence of Mg in the spectra (Figures 4B,C and 5) can indicate the presence of degradation products of chlorophyll, because all chlorophyll molecules share chlorin magnesium ligand in their structure, supporting the photosynthetic hypothesis. A community of saprophytic heterotrophs was proposed as part of the microbiome,35 which are assumed to have a different set of metallic catalysts and enzymes, hence, identifiable chemical fingerprints. Nevertheless, clear separation of the chemically distinct subclasses of microfossils requires even higher statistics and linkage to the morphological features.

Overall, the broad set of mass spectrometric characteristics, measured from the Gunflint microfossils, can be identified using the full feature space, with the ML classification models (see Table S1). By applying an ensemble classification algorithm (adaptive boosting), we achieved a 99.7% separation rate between classes: inorganic host (quartz) or organic inclusions (microfossils). A small subset of misclassifications is attributed to the transition boundary line, where spectra are protruding from the host region towards microfossils. By extracting the 196 features from the single mass spectra, we created 19,110 unique sets of mass and ratio pairs, in which two classes might be separated (i.e., C vs. H, and C vs. O). It is possible to achieve a full separation score of 100% by further limiting the C intensities; however, this will affect the quality of the learned borderlines between classes. Such models, containing the empirical biosignatures from the known terrestrial samples with a proven biogenic origin, might be used as a deployable solution onboard of the Martian rovers, providing an additional line of evidence towards establishing the biogenicity of a given putative sample by assessing the proximity of the data to the spectra acquired from the Gunflint and/or other model samples. However, future work will be required to identify the capacity of LIMS system for distinguishing between true biological organic material (life) as opposed to organic material that was produced non-biologically, for example, via Fischer-Tropsch-Type (FTT) synthesis.

4 | CONCLUSION

In summary, the chemical composition of Precambrian microfossils from the 1.88-Ga Gunflint Formation was investigated using a laser-based miniature TOF-MS. Locations of microfossils were identified on the surface of the sample utilizing MSI. The composition of individual microfossils embedded within the chert was identified using depth profiling and single mass unit spectral decomposition. Utilizing MSI, weighted mass correlation networks, isotopic ratios, and projection of intensities into the low dimension using PCA, it was demonstrated that the microfossils, and associated with them mineralization, have a unique chemical composition that could be identified from the LIMS microprobe data. High-throughput LIMS imaging combined with depth profiling has been shown to be capable of yielding new insights into the distribution, preservation, and elemental speciation of the microfossils in Precambrian cherts.

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CONFLICT OF INTEREST

No competing interests exist.

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The data that support the findings of this study are available from the corresponding author upon reasonable request.

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