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1. Introduction

Laser induced breakdown spectroscopy (LIBS) is a form of atomic emission spectroscopy growing in popularity for elemental analysis of soil, sediment and related materials.¹ The sample surface is ablated by a focused laser beam, creating a plasma plume, and the elemental composition of the sample is determined from the wavelengths and intensities of the atomic emissions from the plasma.²⁻⁷ Advantages of using LIBS for elemental analysis are that results are available in real time and that measurements can be performed directly on a solid sample. It is also widely reported that LIBS can be performed without sample preparation. Whilst this is true for some types of sample, the analysis of loosely aggregated heterogeneous material such as soil is facilitated by at least some degree of

Effects of sample pretreatment and particle size on the determination of nitrogen in soil by portable LIBS and potential use on robotic-borne remote Martian and agricultural soil analysis systems

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Field determination of nitrogen in soil is of interest for both terrestrial and Martian applications. Improved management of soil nitrogen levels on Earth could benefit global food production, whilst the determination of soil nitrogen on Mars is required to assess the planet's future habitability. In this study, a mobile laser induced breakdown spectroscopy (LIBS) system with a 1064 nm Nd:YAG laser delivering 25 mJ per pulse was used to assess the effects of sample pretreatment on the measurement of nitrogen in soil. Although pelletisation was preferred, simply milling the sample to <100 mm particle size – which may be more feasible on a remote rover-based analytical platform – improved the spectra obtained. Ablation craters formed in targets prepared from different particle size fractions of the same commercially-available topsoil showed a clear trend in morphology, with smaller particles yielding more uniform craters with fewer fractures. The LIBS emission intensity at 746.83 nm followed a similar trend to results obtained for total nitrogen content in the soil particle size fractions by microanalysis (Perkin Elmer CHN Elemental Analyser) and was well-correlated ($R^2 = 0.94$) with soil nitrate determined by ion chromatography (Dionex DX-100). Although correlations were less good when analysing field soil samples collected from central Scotland ($R^2 = 0.82$ for comparison between LIBS and microanalysis) the study nevertheless demonstrates the potential of portable LIBS for measurement of soil nitrogen content.

> sample manipulation before exposure to the laser, and it has been highlighted in the literature⁸ that further research is needed in this area.

> In addition to applications for the determination of potentially toxic elements, LIBS can allow direct observation of key nutrients in soil.⁹ One element of particular interest because of its role as an essential element for plant growth is nitrogen. The ability to determine nitrogen directly, in soil, in the field, is challenging.¹⁰ However, it would be of enormous benefit to agriculture as it would allow more targeted application of fertilizers¹¹⁻¹⁴ and more effective management of crop nitrogen utilisation.15,16

> The in situ measurement of nitrogen in soils is also of interest in space science.¹⁷ It has been suggested that N_2 was abundant in the early Martian atmosphere but was then depleted by hydrodynamic escape. Nitrates that were produced in the early history of Mars by photochemistry may still be present: indeed, it is estimated that the Martian surface could contain soil nitrates at average levels of 0.3% N (w/w). One of the key goals of research being conducted on Mars is to determine whether the planet has the ability to sustain microbial life, with the hope of creating a soil-based sustainable agriculture system for habitability in future missions. Nitrates are a fundamental

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source for nitrogen for terrestrial microorganisms. Therefore, the detection of soil nitrates is important to assess habitability in the Martian environment.¹⁸⁻³⁰

The detection of nitrogen in soil using laboratory-based LIBS instruments has been reported by several authors.^{14,31,32} However, for agricultural and space science applications, it will be necessary to determine – and ideally map – the soil nitrogen distribution *in situ* using a field-portable instrument.

This is challenging because, as mentioned above s the quality of LIBS results obtained is enormously influenced by the chemical and physical properties of the sample, which in turn depend on how the sample was pretreated prior to analysis.³³⁻⁴⁰ Various methods of soil sample preparation have been developed to achieve the homogeneity required for LIBS analysis, including mechanical pressure,⁴¹ grinding and sieving⁴² and a combination of these.^{39,43-45} However, these potentially may be difficult to replicate in a field setting, especially using an autonomous rover analytical platform.

The aim of the current study is to gain a better understanding of how factors such as sample preparation and particle size are likely to affect the quality of nitrogen determination in soil using a prototype portable LIBS system designed for use with the four-wheel mobile AgriRover platform.⁴⁶ Results were compared with those obtained by well-established laboratory techniques: a CHN Elemental Analyser (for total soil N), and ion chromatography and colorimetry (for soil nitrate concentrations).

2. Experimental

2.1 Soil sampling and sample preparation

Five surface soil samples were collected from agricultural land in central Scotland, UK. The field-moist soil was oven-dried at $40\text{ }^\circ\text{C}$ for two days and sieved to a particle size of 1 mm. Test

portions of one of the samples were then subjected to different pretreatment regimes before analysis by LIBS:

- (a) no further treatment.
- (b) milled to \leq 100 µm particle size.
- (c) milled then pelletised.
- (d) milled then frozen.
- (e) milled then pelletised then frozen.

Milling was performed using a planetary ball mill. Pelletisation was carried out using a manual laboratory press: 7 tonnes of pressure was applied to 2 cm^3 soil samples for 30 seconds using a $1/2$ ["] die. No binders were added. To produce frozen material, soil was placed in a freezer at a temperature of approximately -18 °C for two days.

To investigate the effects of soil particle size, a commercially processed topsoil was purchased from B&Q, Glasgow, UK. The soil was dried; ground in the ball mill; then passed through a set of stainless steel sieves with mesh sizes 1000, 500, 250, 125, 90, 63, 53 and 45 μ m. Material in each particle size fraction (<45, 45–53, 53–63, 63–90, 90–125, 125–250, 250–500 and 500–1000 mm) was pelletised using the apparatus and procedure described above.

2.2 LIBS instrumentation and measurement procedure

A mobile LIBS system, from StellarNet Inc, (Fig. 1) was used in a laboratory setting to measure the concentrations of nitrogen in soil samples. The system consisted of a laser, powered and controlled by a laser drive and control system, a soil sample holder, a sample chamber, and a spectrometer connected to a computer system for data collection and processing.

A Nd:YAG laser operating at 1064 nm, with a beam diameter of 3 mm, pulse repetition rate of 1 Hz, nominal pulse width of 4 ns and energy of 25 mJ per pulse was used to sample the targeted soil. The laser pulse was guided and focused onto the soil

Fig. 1 A schematic diagram of the portable LIBS system and its major components; spectrometers covering the spectral range from 400– 800 nm; sample chamber; vertical laser head; laser trigger unit; and computer system.

Fig. 2 Images of the variation in craters produced by laser ablation in (a) dried & sieved soil, (b) milled (to <100 µm particle size) soil, (c) milled and pelletised soil, (d) milled then frozen soil and (e) milled, pelletised and then frozen soil. Data for 20 laser pulses of 25 mJ of laser power.

surface by a focussing lens to form a high-temperature plasma for vaporization and atomization of the soil surface material. Plasma emissions were measured with fibre optic cables mounted at \sim 45 $^{\circ}$ to the soils surface and connected to four spectrometers, spanning the spectral range 400–800 nm. The spectrometers have an optical resolution of 4 cm^{-1} (equivalent to 1 nm with 25 μ m slit). Background spectra were collected before the experiment and subtracted during data analysis.

Results obtained from LIBS can vary markedly across a single sample pellet.⁴⁷ To obtain as representative a result as possible, the spectra obtained from 20 laser shots (five shots at each of four different locations across the target) were averaged. One to two preliminary shots per location were used to remove any dust adhering to the surface.

2.3 LIBS data analysis

The spectrum from the LIBS system contained complex information which described the elemental composition of the soil. A MATLAB script was written to remove the background spectrum and to highlight and identify peaks at wavelengths of interest.

2.4 Investigation of ablation effects

Soil (3 cm^2) and pellets prepared from one of the agricultural soils following the different sample pretreatment procedures described in Section 2.1 were placed in a specimen holder and then positioned under the laser head. The ablation craters formed on samples subjected to different pretreatments were

Fig. 3 LIBS spectra obtained from (a) dried and sieved soil and (b) dried, sieved and pelletised soil.

imaged and any differences noted. Pellets formed from different size fractions of the commercial topsoil were ablated and the ablation craters imaged using scanning electron microscopy (SEM). Samples were mounted using carbon tabs and coated with gold using a Polaron SC500A sputter coater, under argon. The scanning electron microscope used was a Stereoscan 90, from Cambridge instruments.

2.5 Determination of soil nitrogen and nitrate

Total soil nitrogen content was determined using a Perkin Elmer 2400 CHN Elemental Analyser and manufacturer's recommended conditions. Samples (0.01 g) were analysed in duplicate.

Soil nitrate determination was carried out by either ion chromatography (IC) or using a colorimetric method. For IC, 1 g test portions were shaken in 5 mL deionized water for 4 hours, filtered, and the filtrate collected for analysis. Extracts were analysed in triplicate (giving RSD values < 3.2%) using a Dionex DX-100 instrument with an AS22 column and 5.5 M sodium bicarbonate mobile phase. In the colorimetric method, 8 g test portions were stirred in 25 mL deionized water for 10 minutes, filtered, and 5 mL of filtrate reacted with a chromogenic reagent to give a pink colour. The nitrate content was determined at $\lambda_{\text{max}} = 530$ nm using an S.I. Photonics Model 420 spectrophotometer.

3 Results and discussion

3.1 Effect of sample pre-treatment

The ablation craters formed in soil pretreated in different ways were notably different (Fig. 2). The crater formed in the soil that had only been dried and sieved was relatively large and poorly defined compared to that of the other pretreatments. The raw spectrum obtained was extremely noisy and it was observed that, on contact with the laser, substantial amounts of sample material were ejected leading to contamination of the sample

Fig. 4 SEM images of the craters produced by laser ablation of soil pellets with particle size ranges shown. Data for 20 laser pulses of 25 mJ of laser power. Results for the $90-125 \mu m$ size fraction are not available.

chamber. This highlights that simply scooping soil into a specimen holder – whether in the terrestrial or Martian context – is likely to lead to measurements with high analytical uncertainty that are prone to cross-contamination between test portions.

Milling to \leq 100 μ m gave a better-defined crater and reduced spectral noise. When the soil was frozen its texture changed, with particles adhering together more readily, perhaps due to the presence of a small amount of residual moisture. Regardless of whether the soil had previously been frozen, samples that had been pressed into pellets – as expected – gave the smallest and most well-defined ablation craters and the least spectral noise. The incorporation of suitable apparatus to compress a soil sample is technically feasible on an agricultural rover. However, payload weight restrictions may preclude this option for Martian deployment.

As shown in Fig. 3, the relative intensities of the nitrogen spectral lines differed between samples pretreated in different ways. In particular, the 746.83 nm line increased in prominence following pelletisation. This wavelength has been used previously for quantification of nitrogen by laboratory-based LIBS, using internal calibration with spiked soil¹⁴ and sand.³²

This investigation suggests that, although pelletisation would be preferred, even milling a soil sample prior to analysis improves the spectral quality and is therefore to be recommended.

3.2 Effect of soil particle size

When the ablation craters produced in pellets of the commercial topsoil were imaged by SEM, the crater diameter appeared generally to increase with average soil particle size (Fig. 4).

There was also a clear change in crater morphology. Pellets prepared with particles in the ranges 125–250, 250–500 and $500-1000$ µm had a more jagged profile, relative to the more uniform aspect of those prepared with particles in the <45 and 45–53 mm size fractions. Larger particles also led to an increase in the number and size of fractures emanating from the ablated crater.

The general trends observed agree with expectations from literature. In a study of plant materials, de Carvalho et al. found that after 20 consecutive laser pulses there were notable changes in the crater morphology for targets formed from different particle size fractions of the bulk sample. For the largest particle size (mean diameter $121 \mu m$) the ablation crater was non-reproducible and disordered. Smaller particle sizes (mean diameter \leq 32 µm) produced a crater hole without indication of fractures.³⁹ Capitelli et al. found that ablation craters on sand and sludge differed greatly. They too related this to the form of the sample, including its preparation and texture, as well as its thermal properties, all of which may affect the interaction between the laser beam and the sample surface, hence in turn the LIBS performance.⁴⁸

3.3 Comparison between LIBS and elemental analysis for total nitrogen determination

The total nitrogen content was determined in the different particle size fractions of the commercial topsoil using the CHN Analyser. Results varied between 1.02 and 0.32% – similar to for example the range (0.1–1.2%) recently reported for a suite of Canadian agricultural soils 49 – and general decreased with increasing particle size (Fig. 5). The trend in nitrogen concentration correlated well with the total carbon content determined by microanalysis $(R^2 = 0.905)$ and the organic matter content estimated by loss-on-ignition ($R² = 0.904$) suggesting that the majority of nitrogen present was, as

Fig. 5 Comparison between response from LIBS system at 746.83 nm and nitrogen concentration determined by microanalysis for pellets containing soil particles in different size ranges.

expected,⁵⁰ associated with soil organic matter. The LIBS emission intensities at 746.83 nm followed a remarkably similar trend, indicating that LIBS provided a reliable estimate of the relative soil nitrogen content in the pellets and that – although the sample chamber was not evacuated or flushed with inert gas – the presence of atmospheric nitrogen did not give rise to significant interference.

Comparison between total nitrogen content obtained by microanalysis and LIBS emission intensity at 746.83 nm for the five agricultural soil samples is shows in Fig. 6. A reasonably strong correlation ($R^2 = 0.82$) was observed. Previous workers have shown that it is possible to calibrate the LIBS response by spiking sand³² or soil¹⁴ with extraneous nitrogen compounds. Although stand-alone quantification of nitrogen by portable LIBS was beyond the scope of the current study, Fig. 6 indicates that the approach can at least discriminate between intrinsic nitrogen levels in real environmental samples.

Fig. 6 Relationship between LIBS emission intensity at 746.83 nm and nitrogen concentration determined by microanalysis in five agricultural soils.

Fig. 7 Comparison between response from LIBS system at 746.83 nm and nitrate concentration determined by ion chromatography for pellets containing soil particles in different size ranges.

Fig. 8 Comparison between response from LIBS system at 746.83 nm and nitrate concentration determined by colorimetry for four agricultural soils.

3.4 Comparison between LIBS, ion chromatography and colorimetry for the determination of nitrate

Since a key objective of Mars LIBS deployment is the search for soil nitrates, comparisons were also made between nitrogen determination by field portable LIBS and nitrate concentration determination by established laboratory techniques.

Ion chromatography indicated that $58-179$ mg kg⁻¹ of the nitrogen present in the commercial topsoil was in the form of nitrate (<2% of total soil nitrogen content). This is as expected: nitrate typically constitutes a small fraction of total soil nitrogen in terrestrial systems because the species is soluble and readily lost to the hydrosphere through leaching. The chromatography and LIBS results were found to be highly correlated for the different particle size fractions (Fig. 7) demonstrating that portable LIBS has the sensitivity to differentiate soil nitrate concentrations.

Finally, the nitrate levels in four of the agricultural soil samples were determined by colorimetry and results compared

with LIBS. Broadly similar trends were obtained (Fig. 8) but the correlation between the results of the two techniques $(R^2 =$ 0.596) was much poorer than when different size fractions of a single soil were analysed (Fig. 7). Given that the agricultural soils were collected from adjoining fields and likely to be geochemically similar, further work is needed to study the applicability of LIBS for nitrogen determination in soils that differ markedly in matrix composition (mineral and organic matter content).

4 Conclusions

In this study the effect of sample preparation and particle size on the feasibility of using a portable LIBS system to measure the concentrations of nitrogen in field soil samples has been evaluated. Spectral quality was improved when the samples were pelletised, but even milling to reduce particle size and increase sample homogeneity was beneficial. When benchmarked against well-established chemical analysis techniques, it was found that the relative nitrogen concentrations determined by LIBS followed similar trends to quantitative results for total soil N content obtained by microanalysis (using a CHN Elemental Analyser) and for soil nitrate content obtained using ion chromatography and colorimetry.

Further work is required to develop suitable calibration models for stand-alone nitrogen determination, to establish the limit of detection of the method, and to evaluate the stability and repeatability of field measurement with the LIBS instrument mounted on the AgriRover platform. For terrestrial applications it would be useful to investigate soils with different geochemical characteristics, and to assess whether any scope exists for LIBS-based nitrogen speciation analysis (since it is not the total, but the phytoavailable nutrient content that affects crop yield). This can be investigated by spiking samples with different forms of nitrogen e.g. nitrate, ammonium, and organic species, prior to LIBS analysis. It would also be of interest to carry out LIBS on soil that had been treated to more closely simulate samples likely to be encountered in a Martian setting e.g. by removal of organic matter.

Conflicts of interest

There are no conflicts to declare.

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