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Dylan J. Malenfanta University of Windsor

Alexandra E. Paulick University of Windsor

Steven J. Rehse University of Windsor

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A Simple and Efficient Centrifugation Filtration Method for Bacterial Concentration and Isolation Prior to Testing Liquid Specimens with Laser-Induced Breakdown Spectroscopy

Dylan J. Malenfant^a, Alexandra E. Paulick^a, Steven J. Rehse^{a,*}

^aDepartment of Physics, University of Windsor, 401 Sunset Avenue, Windsor, Ontario, Canada N9B 3P4

*Corresponding author at: Department of Physics, University of Windsor, 401 Sunset Avenue, Windsor, Ontario, Canada N9B 3P4. Tel.: +1 519 253 3000 E-mail address: rehse@uwindsor.ca (S.J. Rehse).

HIGHLIGHTS

- Bacteria are readily trapped on sub-micron pore size filter media
- A prototype device to hold multiple filters was designed and constructed.
- Centrifuging bacterial suspensions using the device easily collects bacterial cells.
- Removal of the filters allows access to the surface for laser spectroscopy.

GRAPHICAL ABSTRACT



ABSTRACT

An inexpensive filtration device was designed and constructed to rapidly concentrate bacteria in a liquid suspension on the surface of a disposable filter medium while at the same time separating the bacterial cells from larger contaminants in the suspension on the basis of their size. The device consists of a two-stage insert that is held rigidly in a standard tube during bacterial suspension centrifugation. The filters can be easily removed from the insert for subsequent testing with laser-induced breakdown spectroscopy in a process that takes only three minutes. Filter media of 0.45 micron pore size was found to capture approximately 90-95% of the cells in suspension. A limit of detection of 90,000 cells per laser shot was calculated by constructing a calibration curve from multiple suspensions of varying concentration. Deposition of the bacterial lawn across the surface of the 9.5 mm diameter filters was found to be uniform to within +/- 20% of the mean as determined by the total measured optical breakdown emission. Use of 5 micron pore size filter media in the first stage of the insert was found to remove close to 100% of a 12 micron grain size tungsten contamination from the suspension while removing 10% of the bacterial cells. This mounting protocol provides a very convenient method for sample preparation that makes use of common techniques, apparatus, and procedures that would be familiar to clinicians or microbiological pathology laboratory personnel.

KEYWORDS Laser-induced breakdown spectroscopy Bacteria Filter

1. Introduction

The last thirty years have seen numerous experiments performed which have attempted to exploit the speed, utility, and flexibility of the elemental analysis technique known as laserinduced breakdown spectroscopy (LIBS) to detect, identify, or elementally characterize bacterial specimens [1,2,3,4,5]. The goal of these experiments has primarily been to obtain a high signalto-noise ratio LIBS spectrum which can be used to rapidly determine the elemental composition of the bacterial cells ablated by the LIBS laser pulse. This spectrum is dominated primarily by elemental emission from trace inorganic metals such as calcium, magnesium, sodium, and phosphorus, as well as other elements such as carbon and potassium [6]. Molecular emissions are also commonly observed in such spectra [7].

It has been previously noted that great care must be taken to determine how best to prepare, deposit, and sample bacterial cells to optimize the bacterial classification or discrimination, and that the choice of substrate upon which the cells are deposited may strongly impact the success the LIBS-based experiment [8]. In the majority of experiments conducted previously, the bacterial cells were prepared or "mounted" on a carefully selected target substrate prior to LIBS ablation to provide a stable and reproducible platform. Unfortunately, in most clinical or other real-world scenarios in which LIBS might be used in such applications, the number of bacteria, or bacterial titer, may be small, and the contributions to the LIBS spectrum from the underlying substrate which supports the deposited bacterial cells may be significantly larger than the bacterial signal, precluding detection or accurate identification. This is particularly problematic with the use of nanosecond laser pulses which do not control the axial ablation as precisely as femtosecond laser pulses [9].

Two exceptions exist which circumvent this spectral contamination problem. The first utilizes freeze-dried or lyophilized bacterial cells that are then compacted into a solid "pellet" which is an approach typically used for the LIBS testing of unknown powders or residues [10,11,12]. These pellets produce high intensity LIBS sparks due to their density and the absence of water

[13,14,15,16] and can even be mixed with a chemically inert binder to enhance the durability of the pellet during laser ablation [17]. Unfortunately, this lyophilization procedure requires a quantity of bacteria that would not be present in a clinical specimen and requires entirely too much time to make it a realistically competitive diagnostic technology.

The other exception to this is the testing of bacterial cells or spores in aerosol form as either a dense cloud [18], sparse cells dispersed in a dilute gas stream [19], or even as single cells levitated individually [20]. The amount of sample preparation effort this entails, the complexity of the apparatus, as well as the additional biohazards introduced also makes this inappropriate for use in a clinical setting.

While many groups have ablated live bacterial cells from culture directly on the surface of a nutrient medium [21,22,23,24,25] this approach is complicated by a lack of control over cell number and contamination of the cells and spectrum with growth medium contributions. In addition, the colony itself, a loose conglomeration of cells, is not a desirable target for laser ablation, lacking mechanical rigidity and good thermal conductivity. Splashing and target disruption are common outcomes which tend to reduce shot-to-shot repeatability.

A superior choice to all of these is the use of inexpensive, disposable filter media. Other groups have made extensive use of such filters, which are fabricated from nitrocellulose paper, nylon, or other material, due to their widespread availability and their familiarity to clinical microbiologists [7,26,27,28,29]. The ability to quickly separate bacterial cells in a liquid suspension from the liquid itself using the micropores of the filter in their intended role is an obvious advantage.

Recently we have utilized such filter media which provide a highly uniform, extremely flat mounting surface upon which bacterial cells can be deposited to demonstrate rapid and efficient bacterial discrimination of liquid aliquots [30]. These experiments were conducted by creating a loose suspension of cells of known concentration which were then vortex-mixed to maximize the distribution of cells within the suspension. A custom-fabricated circular metal jig with three 4.7

mm diameter holes drilled through it was placed on top of the circular filters and 30 microliters of suspension was deposited in each well with a micropipette. After 20 minutes of settling through the filter in a biosafety cabinet, three smooth lawns of bacteria were created in the shapes of the holes into which the liquid was pipetted [31]. While this method of deposition provided a uniform bacterial deposition, it was still relatively slow and limited the absolute volume of liquid sample that could be tested. Thirty microliters tended to fill the hole in the jig with liquid and subsequent depositions formed a layer of bacteria which effectively stopped up the pores of the filter, preventing the excess water from draining through.

To circumvent these problems, a new method was developed which utilized a custom-built centrifuge-tube insert to hold these filter media during centrifugation. The unique aspect of this insert is that the filter could be safely and trivially removed from the insert after centrifugation for testing in our LIBS apparatus. Utilizing this device and method, much larger volumes of bacteria-containing liquid could be processed for quick and easy separation of the bacterial cells from the liquid filtrate in a shorter period of time. In addition, the insert possessed a two-stage filtration assembly which allowed for the efficient separation of the bacterial cells from larger sized contaminations in the filtrate [32]. Thus, utilizing this insert and process has allowed us to achieve both isolation of the cells (from non-desired contaminants) and concentration of the cells (upon the surface of a filter medium) from a dilute suspension in a very rapid, convenient, and easy-to-use protocol that utilizes apparatus and methodologies highly familiar to microbiological or clinical personnel.

2. Material and Methods

2.1 Centrifuge insert

A centrifuge tube insert capable of holding and filtering at least 1.5 mL of liquid though two filters of varying pore sizes was designed using the SolidWorks 2010 CAD software (Dassault Systémes Solidworks Corporation) and two prototypes were generated via 3-D printing. The insert design and assembly is shown both disassembled and assembled in Fig. 1. The insert consisted of either two or three pieces, depending on whether single-stage or two-stage filtration was desired.



Fig. 1. Centrifuge insert design and prototype. (a) The top or "body" of the insert is threaded to accept the bottom or "base" of the insert (b). A circular filter is placed on the base and a seal is produced by the pressure generated by the threads as the base is screwed into the body. Pedestals under the filter prevent it from resting directly on a flat surface, allowing water to freely pass through the filter and through a hole in the base. Only the first filtration stage of the insert is shown. The base shown in (b) is itself threaded to allow a second identical base to screw into it holding a second filter with a different pore size. (c) Photograph of the disassembled centrifuge insert showing a filter resting in place on the base. (d) Photograph of the assembled centrifuge insert) and (e) the insert secured in place by the cap of a Millipore 10 mL capacity centrifuge tube equipped with a hinged plastic cap.

The lightweight composite insert consisted of a cylindrical tube-shaped main body with a 14 mm outer diameter. The top of the main body was slightly wider at 17 mm, allowing it to rest on the lip of a standard centrifuge tube without falling to the bottom (Fig. 1(a) and 1(e).) This centrifuge tube insert was designed to fit inside a standard 10 mL capacity centrifuge tube equipped with a hinged plastic cap (specifically the filtrate collection tube from an Ultrafree-CL centrifugal filter device, Millipore). The bottom of the body was threaded to allow the second piece of the insert (the base shown in Fig. 1(b)) to securely screw into it. The base of the insert served as a platform upon which the filter was placed, and screwing it into the main body created a seal around the

edge of the filter through applied pressure. A photograph of the disassembled insert with a filter placed upon the base is shown in Fig. 1(c).

To ensure the liquid in the suspension could freely flow through the filter, the filter was supported by a solid ring around the outer circumference of the base and several small pedestal columns (visible in Fig. 1(b)) placed in three concentric rings in order to prevent the filter from resting directly on the base. A hole in the middle of the base allowed the liquid to then drain out of the insert and into the centrifuge tube in which the insert was placed.

When the base was screwed into the body, the filter was securely held on the ring of the base, forming a fairly tight seal, forcing the liquid through the filter. Fully assembled (Fig. 1(d)), the insert was 40 mm in height. The bottom of the base was also threaded to allow a second base piece to screw directly into it. This second base could hold a filter with a smaller pore size allowing for two-stage tiered centrifugation. This mode of operation is discussed in section 3.4.

During operation, approximately 1.5 mL of bacteria-containing liquid suspension was placed within the insert which was then placed within the 10 mL capacity Millipore filtrate collection tube and held firmly in place by a hinged cap attached to the centrifuge tube. (Fig. 1(e)). Due to the small pore size of the nitrocellulose filters, no liquid would come through the insert until centrifugation.

2.2 Nitrocellulose filters

The insert was designed to hold modified nitrocellulose filters. Typical experiments were conducted with 13 mm diameter filters (HAWP01300, 0.45 μ m pore size and GSWP01300, 0.22 μ m pore size, Millipore-Sigma). Due to the diameter of the centrifuge tube, the inner diameter of the insert could not be designed to hold these 13 mm diameter filters. To properly fit on the base of the insert, the 13 mm diameter filter papers were cut with a punch and die set, resulting in a 9.5 mm diameter filter. In subsequent two-stage filtration experiments described in section

3.4, a larger pore size filter was placed on the first-stage filtration base (SMWP01300, 5 μ m pore size, Millipore-Sigma) and the 0.45 micron pore size filter was placed on the second stage filtration base.

2.3 Bacterial sample preparation

All the experiments described in this work were conducted with non-pathogenic *Escherichia coli*, a rod-shaped Gram-negative microorganism of length approximately 1-2 μ m. The bacterial specimens were grown on plates containing tryptic soy agar (TSA) nutrient media. Following incubation, a repeatable quantity of bacteria was harvested from the surface of the nutrient plates and suspended in 1.5 mL of deionized water. The bacterial suspensions were stored in a refrigerator until an experiment was ready to be performed. The suspension was then thoroughly vortex-mixed to create as uniform a suspension as possible.

Concentrations of all bacterial suspensions were obtained by removing a representative quantity of suspension and transferring that volume to an optical quality cuvette for optical densitometry (absorbance) concentration measurements. This absorbance measurement was performed for every new concentration and measured the suspended cell concentration in units of colony forming units (CFU)/mL. All samples used for absorbance measurements were discarded to avoid potential contamination.

2.4 Centrifugation

For all experiments, 1.5 mL of *E. coli* suspension of known concentration was vortexed to achieve a nominally uniform suspension and pipetted into the centrifuge insert, which was secured in the 10 mL Nalgene centrifuge tube. This sample was then then centrifuged for 3 minutes at 5,000 rpm (2,500 g's of force), sufficient to pull all the liquid through the filter. The insert was then removed from the centrifuge tube, disassembled, and the filter was removed from the base. The

filter was secured to a 25 mm x 25 mm piece of steel using double-sided sticky tape for LIBS testing.

2.5 LIBS experiments

The details of the apparatus used to obtain LIBS spectra from the highly flat and uniform filters have been previously described elsewhere [30]. 1064 nm laser pulses of approximately 10 ns duration and 8 mJ pulse energy were focused to approximately 75 µm diameter circular spots on the surface of the filter. Ablation was performed in an argon over-pressure environment. A single shot was always sufficient to ablate all bacteria trapped on the filter as well as a layer of underlying filter media. This ablation was verified spectroscopically as subsequent laser pulses yielded spectra that were not distinguishable from blank filter spectra. It was also verified by scanning electron microscopy and high-resolution optical microscopy performed after the ablation experiments.

After every laser shot the filter was translated 0.635 mm to expose a new surface to the laser pulse. Three single-shot spectra were acquired and digitally averaged to minimize shot noise. Three spectra were chosen for averaging, as opposed to a larger number, to maximize the total number of spectra available for later chemometric analysis (both training and testing) which is not described in this work and also to allow a calculation of the standard deviations of subsequent measurements which were used to construct the error bars in the concentration curve presented in section 3.2. The averaging of three locations allowed the acquisition of over 50 total spectra per filter.

LIBS spectra were acquired with an Echelle spectrometer (ESA 3000, LLA instruments, Inc.) equipped with an intensified CCD camera. Spectra were acquired with a delay time of 2 μ s after plasma formation using an ICCD gate window of 20 μ s for maximum signal acquisition. Once the

spectra were acquired, the integrated area-under-the-curve intensities of 19 backgroundsubtracted emission lines from carbon, phosphorus, magnesium, calcium, and sodium were measured. Table 1 provides the identification of these 19 emission lines used for bacterial measurements, the average LIBS emission intensities of those lines when a high concentration of bacteria was ablated upon the filter (in arbitrary units), and the LIBS emission intensities of those lines when only a blank filter (with no bacteria deposited upon it) was ablated.

Table 1

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Identification and center wavelengths of the nineteen LIBS emission lines measured in this study. The emission intensities (in arbitrary units) for the nineteen lines are shown for bacterial spectra and blank filter (no bacterial deposition) spectra. The bacterial values are the averages of 40 bacterial spectra and the blank filter values are the averages of 20 blank filter spectra.

Element	Wavelength (nm)	Average LIBS Emission Intensity (AU) - Bacteria	Average LIBS Emission Intensity (AU) - Blank Filter
ΡI	213.618	1649	83
ΡI	214.914	879	98
CI	247.856	40342	38082
ΡI	253.398	81	33
ΡI	253.560	387	46
ΡI	255.326	146	35
ΡI	255.491	79	37
Mg I	277.983	141	42
Mg II	279.079	722	37
Mg II	279.553	23476	318
Mg II	279.806	1306	34
Mg II	280.271	12769	163
Mg I	285.213	3343	112
Ca II	317.933	3698	61
Ca II	393.366	41316	722
Ca II	396.847	19527	380
Ca I	422.673	4275	237
Na I	588.995	2751	837
Na I	589.593	1829	572

Numerical values of the emission line intensities were obtained from the spectrometer software (ESAWIN, LLA GmbH) which performed a numeric integration of the full-width at half-maximum region of a background-subtracted peak identified by an automated peak-finding routine. Anticipated linecenters in this peak-finding routine were obtained from the NIST spectral database. Spectral background subtraction was performed prior to integration by fitting a linear function to the background region, which was defined to be 30 pixels on either side of the anticipated linecenter pixel. For these stronger peaks the linear function described the local background well and the calculated background intensity was then subtracted from the data pixel by pixel. The integrated ESAWIN software performed all of this autonomously with no user-defined input and returned spectrally-calibrated integrated areas for all nineteen lines, which were used as the emission intensities, and also the FWHM of all the emission peaks found, which were used to check for any evidence of Stark broadening, line blending, or any other unexpected deviation from the detector-limited FWHM.

3. Results and Discussion

3.1 Filter bacterial retention

Limit-of-detection (LOD) calculations required a knowledge of the number of cells ablated per laser shot. This was calculated from the known initial concentration of the bacterial suspension in the insert, the area over which the bacteria were distributed during centrifugation, and the diameter of the ablated spot. To perform this calculation, the fraction of cells caught on the filter during the centrifugation process needed to be known. Leakage through or around the pressure seal was investigated using optical densitometry on the initial suspension and the filtrate collected from the bottom of the centrifuge tube after centrifugation. Ideally all cells in the suspension should have been caught by the filter and none would have been observed in the filtrate. A calibration curve for optical densitometry measurements was constructed from known concentrations to enable these measurements, and a linear fit to these measurements over eight concentrations gave an r^2 value of 0.999.

Using this fit to calculate concentrations before and after centrifugation, it was determined that 11% of the cells in the initial suspension somehow bypassed the filter and were found in the filtrate. An initial suspension concentration of 8×10⁸ CFU/mL resulted in a lawn on the filter that provided 2.5×10⁶ CFU per ablation event. This was approximately the bacterial lawn density observed and used in previously described experiments [30]. This experiment was repeated for titers at one-sixth the dilution and one-ninth the dilution to ensure that the fractional loss measured was not a strong function of initial titer. These experiments yielded a loss of 7.3 and 5.3% respectively. A measurement of the scatter in repeated optical densitometry measurements on identical bacterial specimens convinced us that this scatter was within the uncertainly of those measurements due to repeatability and did not signify a significant dependence on initial concentration. It did place an uncertainty on an absolute knowledge of the number of cells ablated, however. In all cases, a loss of 11% of the cells from the initial tilter was used as a conservative estimate in all concentrations.

3.2 Concentration curve

A concentration curve was constructed by calculating the total observed LIBS intensity for spectra acquired from five concentrations of bacterial suspensions which spanned approximately two decades. Using a suspension with a known initial concentration, serial dilutions were created prior to centrifugation. These dilutions were centrifuged through the insert utilizing 0.22 μ m filter media and 35 spectra (each the average of three locations) were acquired at each concentration. The plot of these data is shown in Fig. 2, which shows the average total measured LIBS intensity as defined in section 2.5 as a function of the calculated actual number of cells ablated per laser pulse, not the suspension concentrations. Error bars represent the one-sigma standard deviation of the 35 spectra acquired at each concentration. Ablated material from the nitrocellulose filter medium always provided a strong carbon background at all concentrations as

can be seen in the last column of Table 1, resulting in a total measured LIBS intensity for the 19 background-subtracted emission lines which never decayed to a zero reading even for a blank filter sample (the absence of any cells).

A linear fit was performed on these data with a calculated r^2 value of 0.975. An LOD was calculated from this plot using the standard $3\sigma/m$ definition. The LOD was determined to be 60,000 ± 5,000 CFU per ablation event, which was no improvement over previously utilized methods [30]. Obtaining an LOD similar to what had been measured previously with the three-hole circular jig was anticipated, as the concentration of bacterial cells on the filter utilized the same physical mechanism. The concentration of the liquid suspension used to generate the centrifuge-tested sample, however, was significantly lower than when the jig was used and the sample preparation time was greatly reduced. When the uncertainties of the data points obtained from the scatter of the 35 measurements was given a direct weighting on the fit, the goodness of fit improved to an r² of 0.993, but the LOD was then calculated as 90,000 ± 9,000 CFU per ablation event.



Fig. 2. Calibration curve for *E.* coli spectra acquired using specimens prepared at five different concentrations then collected on 0.22 micron filter media using the centrifuge insert.

3.3 Deposition uniformity

During centrifugation, the filter and the insert were held at an angle off from the vertical (as is typical for a "fixed angle" centrifuge). To determine if the deposition across the surface of the filter was uniform, single shot LIBS spectra were collected from across an entire filter with sampling locations spaced in a 0.6825 mm (approximately 1/40 of an inch) square grid. This experiment served as a check that the total observed LIBS intensity was uniform spatially over the surface of the filter which would indicate that the deposition was regular. In addition this allowed for a comparison of the standard deviation of those measurements with the standard deviation of measurements acquired using a previously described method [30].

A false-color map was constructed from the measurements across the surface of the filter. Fig. 3 shows the variation in the total measured LIBS intensity for single-shot LIBS spectra acquired across the surface of the nitrocellulose filter after centrifugation of *E. coli* through the centrifuge insert. The color map indicates the percent difference of the total measured LIBS intensity at any one spot from the average of all measurements as a function of position on the nitrocellulose filter. Red-orange data indicate a measured intensity higher than the average. Blue-green data indicate a measured intensity lower than the average.

No strong patterns emerged in the intensities of the LIBS spectra acquired across the filter. Some systematic increase was observed with motion in the positive x-direction which may indicate a preferential deposition on the half of the filter that hangs lower in the centrifuge, but this increase spanned only approximately a -20% to +20% difference from the mean, which is comparable to the uncertainty of any one measurement. When the fractional standard deviation of the measured spectral lines was compared to previous data, it was seen that the signal was similarly reproducible. In previously published data, the deviation of measured LIBS intensities for lines observed in the bacterial spectra was normally measured to be approximately 20% for small lines and 5-10% for the more intense lines. When using the centrifuge filter, these values

were measured to be 35% and 25%, respectively, before any averaging. Data shown in Fig. 3 is single-shot data, while bacterial data were typically acquired by averaging the measured emission from three ablation events as described in section 2.5. Since the standard deviation is inversely proportional to the square root of *N*, where *N* is the number of measurements, this averaging can be expected to reduce these standard deviations to approximately 20% and 15%, or even lower if more spectra are averaged.



Fig. 3. The variation in the total measured LIBS intensity for single-shot LIBS spectra acquired across the surface of a nitrocellulose filter after centrifugation of E. coli through the centrifuge insert. Laser shots were spaced 0.6825 mm apart. The color map indicates the percent difference of the total measured LIBS intensity from the average of all measurements as a function of position on a nitrocellulose filter. Red-orange data indicate a measured intensity higher than the average. Blue-green data indicate a measured intensity lower than the average.

3.4 Two-stage centrifugation

Bacterial cells are small, approximately 1 μ m in size, compared to red blood cells which are one of the smallest human cells and are ~ 6 – 8 μ m, and typical eukaryotic cells which are about 10 – 100 μ m. This difference in size can be used advantageously to isolate bacteria from clinical specimens based on their smaller size using two filters of varying pore size. To accomplish this, the centrifuge tube insert was assembled with the first-stage base holding a 5 μ m filter and the second-stage base holding a 0.45 μ m filter. This is shown in Fig. 4.

To test the efficacy of the insert for the purpose of separating unwanted material from a bacterial suspension, tungsten powder (10401, Alfa Aesar) with an average particle size of 12 μ m was used to simulate a contaminant. Tungsten powder was chosen for its inertness, cost and availability, safety of use, ease of preparation, biologically relevant size, and elemental purity. LIBS performed on the filters yielded easily measured and identified tungsten emission lines in the LIBS spectra which were not ever observed in the bacterial spectra. This allowed for the easy identification of the presence or absence of tungsten powder on a filter as well as a quantification of the amount of tungsten deposited. Tungsten powder was added to a suspension of *E. coli*, vortexed, and ~ 0.1 mL was pipetted into the top of the insert prior to centrifugation, which was performed with the conditions described in section 2.4.



Fig. 4. Disassembled centrifuge tube insert with the main body of the insert alongside two base pieces (a). The assembled insert indicating the location and pore size of the two filters (b) and the insert in the centrifuge tube (c). After centrifugation filtration, tungsten powder with nominal grain size 12 μ m was readily apparent on the first-stage 5 μ m filter (d) while none was detected using LIBS analysis on the second-stage 0.45 μ m filter.

After centrifugation, the filter papers were removed and images of them were acquired prior to LIBS testing, shown in Fig. 4(d) and 4(e). Tungsten powder was clearly observed on the 5 μ m filter, while none was observed on the 0.45 μ m filter, nor in the filtrate after centrifugation. Each filter paper was then tested with LIBS. Forty-five spectra were acquired across each filter and each of the 45 spectra was an average of three single-shot spectra acquired at different locations. Representative spectra from each filter are shown overlaid in Fig. 5. Readily identifiable tungsten emission lines were observed in the spectrum of the 5 μ m filter. Bacterial emission lines were always observed in the spectrum of the 0.45 μ m filter. These results indicated that the tungsten powder was caught by the first filter while the majority of the bacteria passed through this first filter and were subsequently caught on the second filter.



Fig. 5. LIBS spectrum from 5 μ m filter (black) overlaid with spectrum from 0.45 μ m filter (red) after centrifugation filtration of a bacteria/tungsten powder suspension. Tungsten emission lines in the spectrum from the 5 μ m filter are evident in the 200 – 250 nm range. No tungsten lines were observed in the spectrum from the 0.45 μ m filter, but bacterial emission lines were evident.

To determine what fraction of the bacterial cells were inadvertently caught on the first-stage 5 μ m filter, the average total LIBS intensity from the 45 spectra acquired on the filter papers was used. The total measured LIBS intensity used in this study was calculated as the sum of the

intensities of all bacterial emission lines shown in Table 1 but not including the intensity of the observed carbon 247 nm line. The intensity of this line was not used because a significant fraction of this line is due to ablation of the filter medium, and this can be effected by coverage of the filter with the tungsten material. As well, the intensity of the measured carbon 247 nm line was different for the blank 5 μ m filter compared to a blank 0.45 μ m due to the difference in the amount of carbon ablated, which is a function of the pore size of the filter. The experiment was performed three times: once with the suspension of *E. coli* mixed with tungsten powder and twice with a suspension of just *E. coli*.

The measured LIBS bacterial signal on the 5 μ m filters was not the same within error as a 5 μ m blank control filter, indicating that some bacteria were in fact caught on the 5 μ m filter. In all three cases, the LIBS bacterial signal was significantly greater on the 0.45 μ m filter compared to the 5 μ m filter, indicating that the majority of the bacteria bypassed the first filter and were caught on the second one. An approximation of how much of the bacterial suspension was caught on the 5 μ m filter was done by calculating a ratio of the bacterial signal measured on the 5 μ m first-stage filter (after subtracting off the contribution from a blank control filter) to the total bacterial signal measured on both filters (again after subtracting off any possible contribution from blank control filters). The three experiments indicated that 10%, 12%, and 9% of the bacteria in the suspension were caught on the first filter. These values are consistent within the uncertainty of the measurement. These values are based on the assumption that all of the bacterial cells in the suspension settled on the two filter papers and none of them settled anywhere else. If this assumption was not true, these values are artificially high, thus establishing an upper bound for the percentage of bacterial cells inadvertently lost on the first-stage filter.

4. Conclusions

A unique two-stage centrifuge insert device has been fabricated for rapid and convenient deposition of bacterial cells on disposable and inexpensive nitrocellulose filter media from liquid

suspensions of arbitrary concentration. The bacteria deposited in this manner formed a uniform layer of bacterial cells which could be detected and identified on the basis of their LIBS spectra.

Using a 0.22 µm pore-size filter, 1.5 mL of suspension was centrifuge-filtered in three minutes and over 90% of the bacterial cells in the suspension were caught on the filter for subsequent LIBS testing. Single-shot LIBS spectra were easily acquired from a range of concentrations and a limit of detection with an upper limit of 90,000 cells per ablation event was calculated. Experiments are already underway which further concentrate cells in a much reduced area of the filter and an order of magnitude reduction in this limit of detection is anticipated.

The uniformity of the deposited bacterial lawn was studied by obtaining the total measured LIBS intensity from spectra acquired across the surface of the filter and the intensities were found to vary by approximately 20% from the mean which was also found to be the approximate fractional standard deviation for emission lines in single-shot spectra. Some systematic dependence on position, most likely related to the orientation of the filter relative to the fixed-angle of the centrifuge was observed. Overall, coverage was found to be quite uniform and adequate to obtain up to 45 high single-to-noise spectra from one 1.5 mL specimen, ideal for performing experiments with good statistics.

A two-stage filtration was performed by utilizing a 5 μ m pore-size filter in the first-stage of the insert and a 0.45 μ m pore-size filter in the second stage of the insert. Performance was evaluated by using a bacteria/tungsten powder mixture to quantify the amount of tungsten caught on both filters and the amount of bacteria caught on both filters. It was found that the first-stage filter caught 100% of the 12 μ m grain size tungsten (within the limit of our ability to detect tungsten on the filters and in the filtrate liquid) and an upper limit of 11% of the bacterial cells. This result needs to be confirmed with more biologically-relevant interferents, such as human eukaryotic cells like red or white blood cells.

All experiments were performed with Gram-negative *E. coli*. Because the capture and retention of bacterial cells may be dependent on the cell size and morphology as well as cell envelope biochemistry, these experiments need to be reproduced with other phenotypes, such as Grampositive cocci (*Staphylococcus* or *Streptococcus*), Gram-positive rods (*Bacillus subtilis*), and even Gram-negative cocci (such as a *Neisseria* species.)

Taken together, these results indicate that even with a minor loss in available cells for testing, this mounting method provides a quick and convenient method for sample preparation that makes use of common techniques and apparatus that should be familiar to clinicians or microbiology/pathology laboratory personnel. Because the method efficiently and easily separates the bacterial cells from the liquid filtrate, it could be equally applied to clinical specimens such as blood or urine, environmental specimens such as contaminated tap water, or waters used to wash and prepare food items prior to consumption. The efficiency and ease with which large contaminants of both a biological and abiotic nature can be separated from the bacterial cells suggests that the adoption of this method prior to any laser-based interrogation of the filter medium in a wide variety of situations could be advantageous.

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