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Nanolithography with metastable neon atoms: Enhanced rate of contamination resist formation for nanostructure fabrication

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A membrane basis for bacterial identification and discrimination using laser-induced breakdown spectroscopy

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Nanosecond single-pulse laser-induced breakdown spectroscopy (LIBS) has been used to discriminate between two different genera of Gram-negative bacteria and between several strains of the Escherichia coli bacterium based on the relative concentration of trace inorganic elements in the bacteria. Of particular importance in all such studies to date has been the role of divalent cations, specifically Ca2+ and Mg2+, which are present in the membranes of Gram-negative bacteria and act to aggregate the highly polar lipopolysaccharide molecules. We have demonstrated that the source of emission from Ca and Mg atoms observed in LIBS plasmas from bacteria is at least partially located at the outer membrane by intentionally altering membrane biochemistry and correlating these changes with the observed changes in the LIBS spectra. The definitive assignment of some fraction of the LIBS emission to the outer membrane composition establishes a potential serological, or surface-antigen, basis for the laser-based identification. E. coli and Pseudomonas aeruginosa were cultured in three nutrient media: trypticase soy agar as a control, a MacConkey agar with a 0.01% concentration of bile salts including sodium deoxycholate, and a trypticase soy agar with a 0.4% deoxycholate concentration. The higher concentration of deoxycholate is known to disrupt bacterial outer membrane integrity and was expected to induce changes in the observed LIBS spectra. Altered LIBS emission was observed for bacteria cultured in this 0.4% medium and laser ablated in an all-argon environment. These spectra evidenced a reduced calcium emission and in the case of one species, a reduced magnesium emission. Culturing on the lower (0.01%) concentration of bile salts altered the LIBS spectra for both the P. aeruginosa and two strains of E. coli in a highly reproducible way, although not nearly as significantly as culturing in the higher concentration of deoxycholate did. This was possibly due to the accumulation of divalent cations around the bacteria by the formation of an extracellular polysaccharide capsule. Lastly, a discriminant function analysis demonstrated that in spite of alterations in the LIBS spectrum induced by growth in the three different media, the analysis could correctly identify all samples better than 90% of the time. This encouraging result illustrates the potential utility of LIBS as a rapid bacteriological identification technology. © 2009 American Institute of Physics [DOI: 10.1063/1.3116141]

I. INTRODUCTION

A. Laser-induced breakdown spectroscopy on biological samples

Laser-induced breakdown spectroscopy (LIBS) is an all-optical analytical technique that utilizes a pulsed laser and optical spectroscopy to detect and identify the constituent elements present in a target material by creating a high-temperature microplasma within the focal region of the laser. The experimental procedures utilized in this technique and the physics used to model and describe the breakdown process and the resulting plasmas have been discussed previously in great detail (see, for example, the books by Cremers and Radziemska1 and Miziolek et al.2 or review articles by Tognoni et al.3 and Pasquini et al.4). While the LIBS technique has been utilized extensively over the past 20 years in a wide variety of spectroscopic and industrial applications,5 only since 2003 has LIBS been used to identify and discriminate a wide variety of microorganisms such as bacteria, fungal/mold spores, pollens, and proteins based on the emission intensity of trace inorganic elements contained in the microorganism.6–15 The success of these early experiments was a result of the positive identification of many elemental lines within the emission spectrum from these biological targets which provided an immediate and unique spectral “fingerprint” which could be used to positively identify the microorganism in the sample.

The ability to rapidly (within seconds or minutes) detect harmful pathogens using LIBS—including those pathogens that cannot be cultured within a reasonable amount of time or cannot be cultured at all—offers an alternative paradigm to the health sciences for the detection, identification, and control of infectious diseases. While the current methods of identifying bacteria can yield accurate results, they also require removal of the sample to a laboratory setting, direct contact (handling) of the potentially pathogenic sample, and above all are often too slow to provide results which could direct responses or treatments which may be time sensitive.
It is also known that the microbiological expertise and cost required to perform these identifications preclude their common use as a screening mechanism to prevent human infection. This lack of convenient testing has led to ever-increasing rates of food-borne outbreaks and secondary infections in hospitals.

Our recent work has focused on two Gram-negative bacteria of concern to the fields of clinical sciences and public health: *Escherichia coli* and *Pseudomonas aeruginosa*. In particular, we have demonstrated the ability to discriminate several nonpathogenic strains of *E. coli* from each other and from the pathogenic strain of enterohemorrhagic *E. coli* (EHEC) O157:H7, which causes kidney failure and deaths in children. A series of recent results by Baudelet and coworkers showed with great success the utility of using both nanosecond and femtosecond LIBS to identify the *E. coli* bacterium. Also, *E. coli* has been completely characterized genetically, has many nonpathogenic strains, and is easy to grow, making it an ideal test bacterium for LIBS studies.

All the experiments described in this paper were conducted on three strains of *E. coli*: an environmental strain, *E. coli* C (Nino) which we will refer to hereafter as Nino C; a derivative of the laboratory K-12 strain, HF4714; and the pathogenic *E. coli* C, EHEC.

*Pseudomonas aeruginosa* is an opportunistic pathogen ubiquitous in hospital environments which complicates patient treatments due to increasing rates of nosocomial infections. While *E. coli* is very much so an intestinal bacteria, *P. aeruginosa* is not. *P. aeruginosa* is best known for its ability to infect burns and to cause complications in patients with cystic fibrosis (CF). CF patients typically develop persistent *P. aeruginosa* lung infections that lead to lung failure, followed by transplants or early death. The treatment of *P. aeruginosa* infections is complicated by its propensity to develop antibiotic resistance.

LIBS experiments were performed on both *E. coli* and *P. aeruginosa* as a demonstration of the technique’s efficacy at discriminating between these two very common yet medically important pathogens.

While the proof of concept use of LIBS to discriminate bacterial targets in idealized laboratory situations has been demonstrated, one of the most significant issues that must be addressed is the inherent biochemical adaptability of the bacteria to their environment. Specifically, it must be proven that the ability to identify a given target organism is not eliminated by the bacteria’s exposure to a wide variety of growth conditions and/or nutrient environments. Early work by Levine et al. concluded that discrimination of two strains of *Bacillus globigii* cultured in different kinds of nutritive media was indeed possible. We have also demonstrated that *P. aeruginosa* cultured in a trypticase soy agar (TSA) medium (a surrogate for a spumt sample) and on a blood agar plate (BAP) (a surrogate for a blood sample) showed no substantial difference in our discrimination. However, we also demonstrated that when the bacterial membrane was purposefully altered via culturing in a nutrient medium containing bile salts or its deoxycholate component, a substantial alteration of the optical LIBS emission was observed. This last experiment was particularly relevant for clinical diagnoses because food-borne and water-borne pathogens are exposed to similar bile salts in the digestive system immediately after ingestion.

The purpose of this article is twofold: to further investigate the extent to which the identification and discrimination of Gram-negative bacteria is disrupted by the intentional alteration of membrane biochemistry due to growth in known environments and to begin to establish a biochemical connection between the current serological (or surface-antigen) basis for bacterial identification and the LIBS identification described herein.

### B. Gram-negative bacteria physiology and its relation to LIBS identification/serological classification

In general, bacteria can be divided into two major groups, called Gram-positive and Gram-negative. The original distinction between Gram-positive and Gram-negative was based on a special staining procedure, the Gram stain, but differences in cell wall structure are at the base of these differences in the Gram staining reaction. Gram-negative bacteria surround themselves with a double membrane and a thin cell wall of peptidoglycans. Gram-positive bacteria possess a cytoplasmic membrane overlaid by a thick cell wall of peptidoglycans. This cytoplasmic membrane is common to both bacteria. By altering membrane biochemistry and observing changes in the observed LIBS spectra, we can prove that inorganic composition of the membrane is at least partially responsible for LIBS-based bacterial identification.

The outer membrane of the Gram-negative bacteria is first and foremost a permeability barrier, but primarily due to its polysaccharide content, it possesses many of the interesting and important characteristics of Gram-negative bacteria.

The inner face of the outer membrane is composed of phospholipids, while the outer face is mainly formed by a different type of amphilic molecule, which is composed of lipopolysaccharide (LPS). The LPS molecule is composed of an inner hydrophobic region, called lipid A, which is attached to a hydrophilic linear polysaccharide region, consisting of the core polysaccharide and the O-specific polysaccharide. The LPS molecule is shown schematically in Fig. 1.

Variations in LPS composition even within a species allow it to be used to identify the bacterium in what is known as a “serological classification.” Based on the reactivity of the outer membrane to introduced antigens, bacteria have been classified for a long time on the basis of their LPS O-antigen which has been a crucial step for diagnostic and epidemiological identification. This serological classification is very frequently used today in clinical laboratories and hospitals and is more commonly used for bacterial identification than the variety of molecular biology techniques that have recently arisen (oligonucleotide microarrays, 16S rRNA probes, polymerase chain reaction (PCR), etc.). However because of antigenic variation, which can be induced by bacteriophage infection or by transposable DNA elements, such classification needs independent confirmation by the development of a diagnostic technology at a molecular or atomic level.

It is known that at least two specific divalent cations, Ca\(^{2+}\) and Mg\(^{2+}\), play a crucial role in stabilizing the mem-
brane by binding the electronegative LPS molecules.\textsuperscript{39} It has been found that these cations can dramatically change immunological and biophysical parameters (such as acyl chain order and aggregate packing structure) which can have profound effects on LPS physiology and bacteria bioactivity\textsuperscript{40,41} although the exact mechanisms of this are not known.\textsuperscript{42–45} Changes in the cation concentration may in some way be linked to epitopes of the bacteria, which are antigen binding sites required for serological identification of the bacteria and it is these metallic cations to which LIBS is particularly sensitive. This is suggestive that the LIBS identification is in some way related to the serological classification of the organism.

II. EXPERIMENTAL SETUP

A. Microbiological sample preparation

Three strains of \textit{E. coli} and one strain of \textit{P. aeruginosa} were cultured in a variety of nutrient media, each chosen to induce a specific reaction within the membrane of these Gram-negative bacteria for 24 h at 37 °C. Such growth conditions yielded a single colony of roughly \((1-5) \times 10^6\) bacteria. One of the substrates was a TSA nutrient medium. TSA is a rich bacteriological growth medium containing pancreatic digest of casein, soybean meal, NaCl, dextrose, and dipotassium phosphate. A second substrate was a BAP. A BAP is a TSA plus blood medium substrate, which is often used to phenologically identify hemolytic activity from the bacteria and was chosen to simulate a sample obtained diagnostically from a blood infection. The TSA substrate more effectively simulated a sample obtained environmentally or from sputum.

A third substrate was a MacConkey plus lactose (MAC) nutrient medium with bile salts (including sodium deoxycholate) at a concentration of 0.01%. It is generally accepted that the hydrophobic bile salts can cause a disruption of the membrane integrity after solubilization of the membrane lipids and that one of the functions of the outer membrane is to prevent the detergent action of such bile salts.\textsuperscript{30,46} These salts are also related to the virulence of certain organisms\textsuperscript{47} and have also been shown to make the bacteria more resistant to lipophilic antibiotics.\textsuperscript{48} At low concentrations of bile salts used in this initial experiment, membrane disruption is not yet expected. Instead, bacteria such as \textit{P. aeruginosa} may form a biofilm during growth in this medium.\textsuperscript{49} In Gram-negative bacteria, a biofilm is an excessive production of exopolysaccharide which forms a mucoid capsule. This biofilm helps to protect the pathogen against the inflammatory defense mechanism of a host.\textsuperscript{50,51} It is possible that bacteria grown in the presence of a low concentration of bile may accumulate the divalent cations is such a polysaccharide capsule which could consequently affect the identification by LIBS. In contrast, in the presence of 0.2% or higher concentration of bile salts (or deoxycholate), it has been shown that the LPS is displaced and dispersed, reducing the exopolysaccharide formation.\textsuperscript{52} To test this, samples were grown in a TSA medium containing 0.4% sodium deoxycholate. Cultures grown in this medium were noticeably smaller due to the inhibitory effect of the deoxycholate and possessed different tactile properties than the other samples. The purity of the cultures of all bacterial strains used in this study was routinely confirmed by streaking on L-agar plates, MAC-lactose plates, and by bacteriophage sensitivity assays.

B. LIBS apparatus

After culturing, the bacteria were transferred to the surface of a 0.7% agar plate with a very thin smear. The choice of agar as an ablation substrate provided a flat, large area substrate with a high-breakdown threshold. The specific advantage of utilizing this highly watery agar was that it did not contribute directly or indirectly to the LIBS spectra of the bacteria. The optical emission from this agar was an order of magnitude smaller than that from the bacteria and lacked many of the elements present in the bacteria. Therefore it did not contribute significantly to a background signal.

Our LIBS apparatus has been described in detail elsewhere.\textsuperscript{19} 10 ns laser pulses from an Nd:YAG (yttrium aluminum garnet) laser (Spectra Physics, LAB-150-10) operating at its fundamental wavelength were focused by a microscope objective to ablate the bacteria and the optical emission from the LIBS microplasma was dispersed in an Échelle spectrometer equipped with a 1024×1024 (24
The ablation pulse, with an ICCD intensifier gate width of 20 μs duration. This delay time was chosen to maximize emission from the strongest, most obvious emission lines while minimizing the contribution from the broadband background emission. Spectra from ten laser pulses were accumulated on the CCD chip prior to readout. The sample was then manually translated 250 μm and another set of ten laser pulses was averaged. Five accumulations were averaged in this way, resulting in a spectrum comprised of 100 laser pulses that took approximately 40 s to obtain. Typically 20–30 such measurements could be made from one colony 24 h after initial streaking.

Initial experiments on P. aeruginosa were conducted in an air atmosphere. However the vast majority of experiments were conducted in an argon environment at atmospheric pressure. The enhancement in optical emission due to the Ar environment is well known and has been detailed in previous works. A comparison of a LIBS spectrum from bacteria ablated in air and in argon is shown in Fig. 2. Both spectra were acquired at a delay time of 1 μs after the ablation pulse, with an ICCD intensifier gate width of 20 μs duration. The enhancement in optical emission due to the Ar environment was consistently strong (+), average sized (○), or weak (−). Weak lines were frequently unobserved in less intense plasmas.

TABLE I. Resolved spectral emission lines observed in LIBS spectra acquired from Escherichia coli (strain Nino C) in an argon environment. The atomic and ionic lines are identified and categorized by whether emission from them was consistently strong (+), average sized (○), or weak (−). Weak lines were frequently unobserved in less intense plasmas.

<table>
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<tr>
<th>Wavelength (nm)</th>
<th>Line identification</th>
<th>Strength</th>
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<td>P I</td>
<td>−</td>
</tr>
<tr>
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<td>P I</td>
<td>o</td>
</tr>
<tr>
<td>213.618</td>
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<tr>
<td>215.294</td>
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<td></td>
</tr>
<tr>
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<td>P I</td>
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<tr>
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<td>C I</td>
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<tr>
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</tr>
<tr>
<td>769.896</td>
<td>K I</td>
<td>−</td>
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areas were utilized as independent predictor variables in a multivariate analysis called discriminant function analysis (DFA). The computerized DFA performed by the SPSS software was described previously. The result of the analysis is to reduce the dimensionality of the 31 independent predictor variables to a few scalar quantities known as discriminant function scores. For discrimination amount \( N \) different groups of spectra, \( N-1 \) discriminant function scores were obtained per spectrum. These scores were then plotted, usually in a two dimensional graph showing each spectrum’s discriminant function one (DF1) score and discriminant function two (DF2) score. Differences in DF1 scores are always more significant than differences in DF2 scores since the first discriminant function is constructed on the basis of the most significant difference among spectra. Higher order discriminant functions always account for less of the variation among spectra.

III. EXPERIMENTAL RESULTS AND DISCUSSIONS

A. Alteration of \( P. aeruginosa \) spectrum by bile salts

The spectra from two strains of \( E. coli \) (Nino C and EHEC) ablated in air were compared to the spectra from \( P. aeruginosa \) cultured in TSA, a BAP, and a MacConkey MAC containing 0.01% bile salts in a DFA. The results are shown in Fig. 3. The \( E. coli \) discriminated well from the \( P. aeruginosa \) and were readily identified. The spectra from \( P. aeruginosa \) cultured in TSA and BAP were identical, as expected, but the spectra from bacteria cultured in MacConkey with 0.01% bile salts were significantly altered. In this analysis if the identity of any given spectrum was assumed to be unknown and the analysis was asked to identify it based on functions constructed from the other spectra (a technique known as a “leave one out” or “cross-validation” analysis), Nino C was correctly identified 97.8% of the time. In such an analysis, EHEC was correctly identified 100% of the time. \( P. aeruginosa \) cultured in TSA 82.6% of the time (the rest of the time identified as being grown in BAP), \( P. aeruginosa \) cultured in BAP 87.5% of the time, (the rest of the time identified as grown in TSA), and the \( P. aeruginosa \) cultured in the bile salt medium correctly identified 100% of the time.

The similarity in DF1 score of the \( Pseudomonas \) samples cultured in the bile salt medium and the two \( E. coli \) samples indicated that these spectra appeared to look more like \( E. coli \). If these \( P. aeruginosa \) spectra were left unclassified altogether, and the analysis was asked to identify the spectra, 72% of the time these spectra were identified as Nino C, 24% of time as EHEC, and only 4% of the time as \( Pseudomonas \). The presence of the bile salts changed the bacterial spectrum in such a way as to make them completely distinct, and the ability to identify them as \( P. aeruginosa \) disappeared. If this spectral alteration was due to the formation of a biofilm capsule, this result is suggestive that the presence of a biofilm may disrupt LIBS-based bacterial identification performed in air. It has previously been observed in non-LIBS experiments that growth of \( P. aeruginosa \) in a biofilm induced phenotypic variation.

The structure matrix of this DFA showed that emission lines from calcium and magnesium were the dominant variables in constructing DF2, which clearly separated the bile salt-cultured bacteria from the others. An analysis of the total spectral power associated with the emission lines in these spectra allowed us to qualitatively describe the changes in the relative concentration of the six elements used in the analysis. This is shown in Fig. 4. The fraction of the total spectral power due to calcium emission increased, while the fraction of the total spectral power due to magnesium emission decreased. The emission from the other four elements was unchanged. We also obtained LIBS spectra from the three substrates themselves. When analyzed by DFA with the bacteria grown on them, no significant difference of any kind was observed among the three and yet the bacteria showed no similarities to the media, proving it was definitely not trace residues of nutrient medium responsible for the discrimination. This can be seen in Figs. 3 and 4 of Ref. 20 as well as in the chemical compositions shown in Fig. 4. We thus attribute the alteration of the bile salt-cultured \( Pseudomonas \) spectra to an increase in calcium and a decrease in magnesium in the LIBS plasma, and thus in the ablated sample. It is not yet possible to discern whether this is due to a biochemical change in the membrane or accumu-
lation in a secreted exopolysaccharide capsule and this will be the subject of future experiments.

**B. Alteration of \( P. \text{ aeruginosa} \) spectrum by 0.4% deoxycholate**

We next investigated whether \( P. \text{ aeruginosa} \) cultured in a TSA medium with 0.4% sodium deoxycholate would yield a spectrum similar to samples cultured in TSA alone or in MacConkey agar with bile salts. While the formation of a biofilm is anticipated in the presence of low concentrations of bile salts, it is unlikely in the presence of higher concentrations of deoxycholate such as the 0.4% used in this experiment. Indeed we expected this concentration to begin to disperse the LPS layer due to the detergent action of the deoxycholate. Samples were cultured in the manner described earlier in TSA, MAC with 0.01% bile salts, and TSA with 0.4% deoxycholate. Cultures were transferred to the 0.7% agar medium, LIBS was preformed, and the results analyzed by a DFA. This experiment and all subsequent experiments were performed in an argon environment at atmospheric pressure. The results of this analysis are shown in Fig. 5, which shows the first two discriminant function scores for spectra from the three bacteria samples and also from the nutrient media themselves.

The spectra from bacteria cultured in the deoxycholate showed a substantial difference from the bacteria cultured in...
TSA and MAC, as the DF2 scores show. DF1 was again able to easily discriminate the media from the bacteria based on their high sodium concentration. The clustering of the media with one DF1 score and the bacteria with a second DF1 score again indicates that it is not possible that it is trace residual quantities of media on the bacteria caused the discrimination. The structure matrix results showed that DF2 was constructed based solely on the calcium concentration. An analysis of the total spectral power associated with the 31 emission lines in these spectra revealed that the overall emission from calcium lines decreased by 20% ± 3%. This does not imply that the concentration of calcium decreased by 20%, but is suggestive of a decrease in the total bacterial calcium concentration. The total emission intensity of the magnesium decreased by 4% ± 3%, which is consistent with no significant change. The MacConkey-cultured bacteria were still distinct from the TSA-cultured samples, as was seen in air, but the largest variation between bacteria was between the deoxycholate-cultured samples and the other two. In a test performed with only the three bacteria, 79% of the total variance in the analysis was accounted for by DF1, which was the variation between the deoxycholate-cultured samples and the other two.

This alteration of *P. aeruginosa*’s spectral fingerprint by a deliberately induced chemical change in the bacterial outer membrane is evidence that membrane biochemistry, the basis of serological identification, is a significant contributor to LIBS-based identification.

C. Alteration of *E. coli* spectrum by 0.4% deoxycholate

*E. coli* and *P. aeruginosa* are both Gram-negative bacteria and thus possess outer membranes with LPS. Therefore we expected the detergent effect of the deoxycholate to be similar (but not necessarily identical) in the two microorganisms. Samples of *E. coli* strain Nino C were grown in the same three nutrient media as described in Sec. III B. A DFA was performed on the LIBS spectra obtained from these samples ablated in argon, as well as the nutrient media, as was performed on *P. aeruginosa*. The results of this analysis are shown in Fig. 6. This analysis showed marked similarities with the analysis of *P. aeruginosa* shown in Fig. 5. Again, all three bacteria groups were easily differentiated from their growth media, which possessed highly similar spectra, by DF1. The *E. coli* cultured in TSA and MAC were highly similar, while the deoxycholate-cultured samples were
highly distinct. However, for the *E. coli*, these last samples were even more distinct from the other two than in the *P. aeruginosa* analysis. This is reflected by the difference between DF2 scores for the two clusters of bacteria being approximately 70% greater than it was in the *P. aeruginosa* analysis.

The cause of this larger difference was a decrease in the total emission from both calcium and also magnesium in these spectra. The total emission intensity from calcium atoms and ions decreased by 12% ± 4%, mirroring the alteration of the *P. aeruginosa* spectrum, but in the *E. coli* samples the magnesium emission intensity also decreased by a significant amount, 25% ± 4%.

In all cases of Gram-negative bacteria tested, reproducible differences in the LIBS emission spectra were induced by exposing the outer bacterial membrane to a known detergent agent during growth. The DFA revealed a significant difference in the spectra from these bacteria due to decreased relative emission from calcium and magnesium. We attribute this to a loss of divalent Mg and Ca cations from the outer membrane, a known effect of exposure to deoxycholate.

D. Decrease in discrimination ability due to growth in bile salts

We have previously reported in the ability to discriminate one strain of *E. coli* (an environmental strain known as Nino C) from a laboratory strain (a K12 derivative known as HF4714) after 24 h of culturing in a standard TSA medium. We next repeated this discrimination in argon and with samples cultured in the MacConkey agar medium with 0.01% deoxycholate bile salts. Samples were cultured in both TSA and MAC as described previously, ablated in an argon environment, and the spectra were analyzed via a DFA. The results are shown in Fig. 7. Figure 7(a) shows the DFA scores of the four groups when analyzed together. The TSA-cultured bacteria were again well-discriminated and the cross-validated discrimination identified the strains of Nino C and HF4714 with 75.9% and 82.1% accuracy, respectively. The MAC-cultured bacteria, however, displayed almost no difference between the two strains with only 59.4% and 63.6% of the Nino C and HF4714 samples correctly identified in the cross-validation test.

While before, spectra from Nino C and HF4714 could be correctly identified with high efficiency, this was no longer true. To test this, the MAC-
cultured Nino C data were removed from the analysis, and the MAC-cultured HF4714 samples were unclassified, allowing the DFA to assign the spectra to either the Nino C or HF4714 category. 76% of the HF4714 spectra were classified as belonging to HF4714 and 24% were classified as belonging to the Nino C category. This test was repeated by eliminating all MAC-cultured HF4714 samples from the analysis and unclassifying the MAC-cultured Nino C samples, allowing the DFA to assign the spectra to either the Nino C or the HF4714 category. Only 47% of the tested Nino C samples were correctly classified as Nino C, the other 53% being incorrectly classified as HF4714 samples. The analysis could do no better than a roughly 50-50 assignment of the ungrouped cases, which is consistent with no discrimination.

We attribute the difference in the drop in discrimination efficiency between HF4714 and Nino C, as well as the relative disparity between the behaviors of the two groups, to their response to the bile salt environment. The inherent difference in the LPS composition of these two strains is most likely the cause of this. The environmental strain Nino C quite possibly forms a polysaccharide capsule in the presence of the low concentration of bile salts, but the HF4714, which has its O-antigen partially missing, may show a reduced proclivity to such formation. Indeed, the severe degradation in the ability to identify Nino C correctly is entirely consistent with a more extreme alteration of the LIBS spectrum due to exopolysaccharide accumulation.

Figure 7(b) shows the same analysis with spectra from the two nutrient media included. It is apparent that the changes induced in the membrane by the presence of the low concentration of bile salts in the MacConkey agar were fairly consistent as the two MAC-cultured samples appeared highly similar in this analysis. Both bacteria were altered in a nearly identical fashion, and this spectral alteration removed the basis for discrimination—based on Ca and Mg content—that was demonstrated previously between the TSA-cultured bacterial samples.

Lasty, we included the previously obtained spectra from Nino C samples cultured in the 0.4% deoxycholate substrate with the TSA/MAC analysis shown in Fig. 7. The results, shown in Fig. 8, again clearly show that the detergent action of the deoxycholate on the bacteria altered the LIBS spectrum in an extreme manner, much more significantly that the low concentrations of bile salts present in the MacConkey agar were able to do. This was attributed to a true biochemical dilution of the outer membrane in the presence of high concentrations of deoxycholate and the possible formation of an extracellular polysaccharide capsule in the presence of the low concentrations of the bile salts. Strain discrimination ability has been compromised, as evidenced by the lack of grouping of HF and Nino samples.

E. Genus discrimination as a function of nutrient medium

Although the alteration of membrane biochemistry and the possible formation of a biofilm reduced the ability to use LIBS to perform strain identification, it did not eliminate the ability to differentiate between two different genera of Gram-negative bacteria, which always exhibited more significant variations between groups than different strains of the same species did.

To demonstrate this, we analyzed the LIBS spectra from two strains of E. coli (Nino C and HF4714) cultured both in TSA and MAC with 0.01% bile salts. We performed a DFA with these four data sets as well as the samples of P. aeruginosa cultured in both TSA and MAC. The results are shown in Fig. 9. This was a significantly more complicated analysis, and the ability to correctly classify spectra was correspondingly less accurate. However, the DFA still was able to accurately discriminate between these two bacteria as evidenced by the DF1 scores for a given type of bacteria. P. aeruginosa samples possessed a positive DF1 score (typically greater than about 1.2), while most of the E. coli samples (regardless of nutrient medium) possessed a DF1 score less than 1.2. Only 72.4% of the E. coli Nino C cultured in TSA and 82.1% of the HF4714 cultured in TSA were correctly classified. When cultured in MAC, that accuracy decreased further to only 62.5% for the Nino C and 69.7% for the HF4714. The P. aeruginosa cultured in TSA and MAC were correctly classified only 65.4% and 57.9% of the time, respectively. However, this was only the ability to iden-
P. aeruginosa. The modification of the LIBS spectrum due to the low concentration of bile salts complicated the proper identification of these bacteria, but nonetheless, 90.4% of all samples were assigned to the correct group in this analysis.

The alteration of the outer membrane composition due to the deoxycholate did not inhibit the ability to correctly identify the bacterial species any more significantly than culturing in the low concentration of bile salts did. E. coli Nino C samples cultured in TSA, MAC plus 0.01% bile salts, and TSA with 0.4% deoxycholate were analyzed in a DFA with P. aeruginosa samples grown in the identical media. This is shown in Fig. 10. Again the bacterial samples were correctly discriminated by genus/species largely on the basis of their DF1 score, the P. aeruginosa possessing negative DF1 scores, and the E. coli possessing positive DF1 scores. While only 76.3% of all cases were correctly classified in this analysis, 91.4% of all cases were correctly assigned to the

![Image](image_url)

**TABLE II.** Ability to correctly identify bacteria on the basis of its LIBS spectrum as a function of nutrient medium (TSA and MacConkey agar with 0.01% bile salts) for two strains of E. coli and one strain of P. aeruginosa. 90.4% of all bacterial samples were assigned the correct identification.

<table>
<thead>
<tr>
<th>Category</th>
<th>predicted group membership (%)</th>
<th>E. coli Nino C (TSA)</th>
<th>E. coli Nino C (MAC)</th>
<th>E. coli HF4714 (TSA)</th>
<th>E. coli HF4714 (MAC)</th>
<th>P. aeruginosa (TSA)</th>
<th>P. aeruginosa (MAC)</th>
<th>Correct genus ID’d Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>E. coli</td>
<td></td>
<td>72.4</td>
<td>13.8</td>
<td>0.0</td>
<td>0.0</td>
<td>13.8</td>
<td>86.2</td>
<td>100.0</td>
</tr>
<tr>
<td>Nino C (TSA)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E. coli</td>
<td></td>
<td>0.0</td>
<td>62.5</td>
<td>3.1</td>
<td>25.0</td>
<td>6.3</td>
<td>3.1</td>
<td>90.6</td>
</tr>
<tr>
<td>Nino C (MAC)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E. coli</td>
<td></td>
<td>2.6</td>
<td>7.7</td>
<td>82.1</td>
<td>2.6</td>
<td>0.0</td>
<td>5.1</td>
<td>94.9</td>
</tr>
<tr>
<td>HF4714 (TSA)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E. coli</td>
<td></td>
<td>3.0</td>
<td>24.2</td>
<td>3.0</td>
<td>69.7</td>
<td>0.0</td>
<td>0.0</td>
<td>100</td>
</tr>
<tr>
<td>HF4714 (MAC)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P. aeruginosa</td>
<td></td>
<td>3.8</td>
<td>0.0</td>
<td>11.5</td>
<td>0.0</td>
<td>65.4</td>
<td>19.2</td>
<td>84.7</td>
</tr>
<tr>
<td>(TSA)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P. aeruginosa</td>
<td></td>
<td>5.3</td>
<td>5.3</td>
<td>10.5</td>
<td>0.0</td>
<td>21.1</td>
<td>57.9</td>
<td>78.9</td>
</tr>
<tr>
<td>(MAC)</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

FIG. 9. (Color) DFA scores for two strains of E. coli (Nino C and HF4714) cultured in TSA (1 and 3); MacConkey agar containing 0.01% bile salts (2 and 4), and one strain of P. aeruginosa cultured in TSA (5) and MacConkey agar with bile salts (6). All MacConkey-cultured bacteria have filled in symbols. Bacterial discrimination by genus is accomplished by DF1 in spite of the spectral changes induced by the low concentration of bile salts in the MacConkey agar. The ability to differentiate one bacterium from another, regardless of nutrient medium, is 90.4%. DF2 discriminates the two strains of E. coli cultured in the TSA medium, but the MacConkey-cultured E. coli samples have compromised strain discrimination.
correct bacterial genus/species. This result is summarized in Table III. The two results shown in Tables II and III, although based on two types of bacteria only, are highly suggestive that the identification of one genus of bacteria relative to another is possible with the LIBS technique regardless of the environmental conditions present during the growth and life cycle of the bacteria.

**IV. CONCLUSIONS**

Nanosecond single-pulse LIBs has been used to identify and discriminate between two bacterial species highly important to the clinical sciences: *E. coli* and *P. aeruginosa*. A DFA was used to analyze the LIBS spectra obtained from colonies of bacteria cultured in three different nutrient media: a standard TS agar, a MacConkey agar containing 0.01% bile salts (MAC) (2), and TSA containing 0.4% deoxycholate (3) when analyzed together with *E. coli* Nino C samples also cultured in TSA (4), MAC with bile salts (5), and TSA containing 0.4% deoxycholate (6). The ability to differentiate one bacterium from another, regardless of nutrient medium, is 91.4%. DF1 performs most of the discrimination between bacterial genus in this analysis.

**TABLE III.** Ability to correctly identify bacteria on the basis of its LIBS spectrum as a function of nutrient medium (TS, MacConkey agar with 0.01% bile salts, and TSA with 0.4% deoxycholate) for a single strain of *E. coli* and a single strain of *P. aeruginosa*. 91.4% of all samples were assigned the correct identification.

<table>
<thead>
<tr>
<th>Category</th>
<th>Predicted group membership (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P. aeruginosa (TSA)</td>
<td>P. aeruginosa (MAC)</td>
</tr>
<tr>
<td>P. aeruginosa (TSA)</td>
<td>70.4</td>
</tr>
<tr>
<td>P. aeruginosa (MAC)</td>
<td>15.0</td>
</tr>
<tr>
<td>P. aeruginosa (0.4% deoxy)</td>
<td>3.4</td>
</tr>
<tr>
<td>E. coli Nino C (TSA)</td>
<td>0.0</td>
</tr>
<tr>
<td>E. coli Nino C (MAC)</td>
<td>6.3</td>
</tr>
<tr>
<td>E. coli Nino C (0.4% deoxy)</td>
<td>0.0</td>
</tr>
</tbody>
</table>
A DFA of the LIBS spectra cultured in these various media clearly revealed reproducible changes to the spectra, confirming that LIBS can be used to probe and quantify Gram-negative outer membrane biochemistry. This is a first step toward establishing an atomic basis for the classification of bacteria, which may correlate with the epitope-based serological grouping based on O-antigen and H-antigen membrane chemistry.

We have demonstrated that the culturing of *P. aeruginosa* in a MacConkey agar plus bile salt medium significantly altered the LIBS spectrum when ablated in air in such a way that it no longer resembled the typical spectrum seen from these bacteria when cultured in TSA or blood agar. This was attributed to an increase in emission from calcium and a decrease in magnesium in the ablated sample possibly due to cation accumulation in a capsule around the bacteria.

The membrane basis for this change in LIBS signal was confirmed by culturing the *P. aeruginosa* in a high-concentration sodium deoxycholate TSA medium, followed by ablation in an all argon environment. The LIBS spectra from these bacteria were significantly altered from both the samples grown in a bile-free environment and the samples grown in the presence of a low concentration of bile salts. This was attributed to the disruption of the Gram-negative outer membrane by the detergent action of the deoxycholate on the LPS molecule. Decreased emission was observed from calcium, which is present in a divalent form in a region of the LPS molecule containing an eight-carbon sugar called KDO (see Fig. 1). No significant change was measured in the emission from magnesium.

This experiment was repeated with an environmental strain of *E. coli* and similar alteration of the LIBS spectrum was observed, again assigned to the disruption of the outer membrane LPS molecules. The total emission intensity from calcium atoms and ions was decreased and the magnesium emission intensity also decreased by a significant amount. This is consistent with both ions being present in the membrane in different concentrations in the two bacteria, making species discrimination possible, as was observed in our earlier works.

Loss of LIBS-based strain discrimination due to culturing in the 0.01% bile salt medium was observed when two different strains of *E. coli* were analyzed after being cultured in this medium and the TSA medium. Although efficient discrimination was possible on the unaltered bacteria, the alteration of cation concentration due to growth in the bile salt containing medium precluded any discrimination of the two strains. When compared to the spectra from bacteria cultured in a deoxycholate-rich medium (0.4%) the low-concentration bile salt-cultured samples appeared almost identical to each other, yet were significantly different from the higher-concentration samples. We again attributed this to the loss of divalent cations from the outer membrane which is known to occur at high concentrations.

We lastly observed that in spite of serious alteration in LIBS emission due to nutrient medium-induced biochemical modification, a DFA still showed accuracy in differentiating spectra on the basis of bacterial genus and species, if not strain. The significant differences that existed between spectra from different genera of bacteria were not completely erased by the alteration of the membrane. This was perhaps the most significant conclusion of this work: the ability to discriminate one genus of bacteria from another with the LIBS technique persisted with better than 90% accuracy regardless of the nutrient medium in which the samples were cultured. By exposing these bacteria to an environmental stressor that could be experienced *in vivo* and subsequently testing them, we have demonstrated that the changes in LIBS signal observed did not inhibit the proper identification of the bacteria. Of course, a larger variety of bacteria will need to be analyzed to generalize this conclusion further. However, these early results are highly encouraging for researchers working to develop an all-optical technique to rapidly identify bacteria in clinical samples.

44S. Snyder, D. Kim, and T. J. McIntosh, Biochemistry 38, 10758 (1999).