Laser-Induced Breakdown Spectroscopy for the Detection and Diagnosis of Bacterial Pathogens in Blood, Urine, and Cerebrospinal Fluid

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66th International Conference on Analytical Sciences and Spectroscopy

Canadian Society for Analytical Sciences and Spectroscopy



Historical Notes

LIBS has been used to detect/identify bacteria since 2003.

1207

Detection of bacteria by time-resolved laser-induced breakdown spectroscopy

Stéphane Morel, Nicolas Leone, Philippe Adam, and Jacques Amouroux

6184 APPLIED OPTICS / Vol. 42, No. 30 / 20 October 2003

Laser-Induced Breakdown Spectroscopy Detection and

Aerospace Engineering, University of California, San Diego, California 92093 (G.A.L., S.G.B.)

JOHN D. HYBL, GREGG A. LITHGOW, and STEVEN G. BUCKLEY

MIT Lincoln Laboratory, 244 Wood Street, Lexington, Massachusetts 02420-9108 (J.D.H.); and Department of Mechanical and

APPLIED SPECTROSCOPY

Classification of Biological Aerosols*

Volume 57, Number 10, 2003

Bacillus globigii BG-1





(sting) (st

Bacillus subtilis var. niger



Laser-induced breakdown spectroscopy of bacterial spores, molds, pollens, and protein: initial studies of discrimination potential

Alan C. Samuels, Frank C. DeLucia, Jr., Kevin L. McNesby, and Andrzej W. Miziolek

20 October 2003 / Vol. 42, No. 30 / APPLIED OPTICS 6205

Bacillus thuringiensis subsp. kurstaki, Bacillus subtilis (also known as Bacillus globigii) subsp. niger, and Bacillus cereus 6E1

800



Historical Notes

Significant achievements followed quickly.

Anal. Chem. 2005, 77, 631-638

Feasibility of Detection and Identification of Individual Bioaerosols Using Laser-Induced Breakdown Spectroscopy

P. B. Dixon and D. W. Hahn*

Department of Mechanical and Aerospace Engineering, University of Florida, Gainesville, Florida 32611-6300

JOURNAL OF APPLIED PHYSICS 99, 084701 (2006)

Femtosecond time-resolved laser-induced breakdown spectroscopy for detection and identification of bacteria: A comparison to the nanosecond regime

Matthieu Baudelet, Laurent Guyon, Jin Yu,^{a)} and Jean-Pierre Wolf Laboratoire de Spectrométrie Ionique et Moléculaire, UMR CNRS 5579, Université Claude Bernard-Lyon 1, 43, Boulevard du 11 Novembre 1918, F-69622 Villeurbanne Cedex, France

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APPLIED PHYSICS LETTERS 90, 163901 (2007)

Escherichia coli identification and strain discrimination using nanosecond laser-induced breakdown spectroscopy

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(Received 26 January 2007; accepted 15 March 2007; published online 16 April 2007)



Detection of single *Bacillus* spores

Escherichia coli and Bacillus subtilis with fs-LIBS



Discrimination of three strains of *Escherichia coli*, mold, and *Candida albicans* yeast



The Reasons?

There did not exist at the time, <u>and</u> <u>there still is not now</u>, a diagnostic test to detect and diagnose bacterial pathogens that can be:

Bioweapons / bioterrorism

Drinking water

- highly-flexible (multiple pathogens)
- very rapid (under 5 minutes)
- robust (field portable)
- relatively simple (minimal prep, performed by a non-expert)
- few to no consumables (biodegradation, cost)
- sensitive
- specific

Environmental surveillance

Food / beverage preparation

Medical diagnosis



To the present

Since then, progress has been extensive on:

- the number / variety of species
- sample preparation methods
- hardware setups
- chemometric algorithms
- number of bacteria detectable
- machine learning



To the present

Table 2

A summary of all the bacterial species/strains tested with LIBS.

Micro-organism	Reference	Form	Chemometric utilized	Laser wavelength ^a
Acinetobacter baumannii ATCC BAA-1789	[66]	Colony on blood agar	PCA/PLS1	1064
Acinetobacter baylyi	[48]	Pellet, freeze-dried powder	Hyperspace projection of	810 (fs)
			trace elements	
Acinetobacter calcoaceticus [FJ816073] ^b	[71]	Colony on glass slide	PCA/PLS-RA	800 (fs)
Arhodomonas sp. [EU308280]	[71]	Colony on glass slide	PCA/PLS-RA	800 (fs)
Bacillus anthracis var. Sterne	[44]	Thin lawn ^c on nylon filter	None	1064
Bacillus anthracis var. Sterne	[38]	Thin lawn on agar, glass slide	PCA/PLS1	1064
Bacillus atrophaeous	[41]	Spore, aerosol stream	None	1064
Bacillus abrophaeous	[52,37]	Dried film on Al disk, steel disk,	NN, MLSRA, PLS-DA	1064
Destillers attended and	(57)	polycarbonate disk	CUM	1064
Bacillus atrophaeous	[57]	Pellet, freeze-dried powder	S VIVI None	1064
Bacillus aureus	[30]	This laws on silver membrane filter	DCA linear correlation and	355
bucillus cereus 6E1	[29,32]	film fawn on silver memorane inter	SIMCA	1064
Bacillus cereus ATCC 14603	[57]	Pellet, freeze-dried powder	SVM	1064
Bacillus globigii ^d BG-1	[39,40]	Pellet, freeze-dried powder	None	1064
Bacillus globigii BG-1	[40]	Spore, aerosol stream	None	1064
Bacillus globigii BG-2	[39,40]	Pellet, freeze-dried powder	None	1064
Bacillus globigii BG-2	[40]	Spore, aerosol stream	None	1064
Bacillus globigii var. niger	[29,42,32]	Thin lawn on silver membrane filter	PCA, linear correlation, and SIMCA	1064
Bacillus globigii var. niger	[30]	Continually refreshed dense aerosol cloud (from powder) and aerosol stream	PCA	1064
Bacillus globigii var. niger	[35,68]	Powder on double-sided sticky tape	No, linear correlation, PCA, PLS-DA	1064×2 (DP)
Bacillus globigii 168	[43]	Colony (wet) on LB medium	None	532
Bacillus globigii	[47]	Thin film lawn on cellulose nitrate	None	810 (fs), 1064
0 0		membrane filter		
Bacillus globigii	[48]	Pellet, freeze-dried powder	Hyperspace projection of	810 (fs)
			trace elements	
Bacillus globigii	[31]	Dried powder on solid substrate	PCA, HCA, PCA + LDA	1064
Bacillus globigii ATCC 23857	[66]	Colony on blood agar	PCA/PLS1	1064
Bacillus megaterium QM B1551	[43]	Colony (wet) on LB medium	None	532
Bacillus stegrathermanhilus ATCC 12070	[43]	Dollat, tracero driad, pourdar	SVM	1064
Bacillus stearoinermophilus ATCC 12979 Bacillus thurangansis	[30,40]	Pellet, freeze-dried powder	None	1064
Bacillus thurengensis var kurstaki	[29,32]	Thin lawn on silver membrane filter	PCA linear correlation and	1064
	[25,02]		SIMCA	1004
Bacillus thurengensis var. kurstaki	[44]	Thin lawn on nylon filter	None	1064
Bacillus thurengensis T34	[43]	Colony (wet) on LB medium	None	532
Bacillus thuringiensis ATCC 51912	[57]	Pellet, freeze-dried powder	SVM	1064
Bacillus sp. [GQ392044]	[71]	Colony on glass side	PUA/PLS-KA	800 (IS)
Bacillus sp. [GQ226038]	[/1]	Colony on glass slide	PGA/PLS-KA	800 (IS)
Enterobacter cloacae [E1104527]	[/1]	Colony on glass slide	PCA/PLS-KA DCA/DISDA	800 (IS) 800 (fe)
Enterobacter cloacae ATCC 12047	[/1]	This lown on putrient free ager	DEA DIS DA	1064
Enterobacter on [CD000652]	[04,07]	Colony on glass slide	DEA /DIS DA	1004 800 (fe)
Enterobacter sp. [CII586319]	(71)	Colony on glass slide	PCA/PLS-RA	800 (IS)
Interoduce ap. [0000017]	[/ 1]	- · · · · · · · · · · · · · · · · · · ·	i ca y i horiori	000 (13)

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Specimen presentation

In my research, I am almost completely focused on developing a realistic clinical testing methodology.

Current methods of LIBS bacterial analysis are completely inadequate for this purpose.

Most samples are obtained as fluid specimens, need to perform LIBS on <u>that</u> to be truly rapid.





Outline for rest of talk

- 1. Explain how we perform LIBS on bacteria in my lab in general
- 2. Three case studies

2a. Using the same sample preparation and data analysis- Blood

- Urine

2b. Using a modified sample preparation and data analysis- Cerebrospinal fluid





LTB Aryelle Echelle Spectrometer ICCD 195-800 nm, 50 pm resolution <1 ns timing control

Spectra-Physics LAB-150 355 nm, 10 ns, 10 mJ/pulse (?)

Argon purge chamber



ESA3000 Echelle Spectrometer ICCD 200-840 nm, 12 pm resolution <1 ns timing control

1111

Spectra-Physics LAB-150 1064 nm, 10 ns, 8 mJ/pulse

LANB 4









Bacteria cultured on agar plate

9.5 mm diameter

disposable filter

Harvested bacteria stored in MΩH₂O



from hospital

Bacteria-negative

specimens collected

A second se

100 μL fluid specimen + 100 μL bacterial suspension pipetted into reassembled apparatus and sealed in centrifuge tube







Disassembled 3D printed centrifuge insert

Centrifuge insert assembled with filter & cone

Aluminum cone with 1 mm diameter aperture at apex





Figure 7.1: (a) Flocked swab used in this work. (b) Flocked swab zoomed-in on the tip.



Figure 7.5: (a) 100 μ L of E. coli pipetted onto surface of metal plate. (b) Metal plate after heated on hot-plate for 2 minutes 20 seconds at 200 °C. Water has evaporated and film of bacteria is observed.

Flocked swab was pre-wet with 10 μL of deionized and swabbed over surface

The swab was placed in 1 mL of deionized water and vortexed for 15 seconds. The entire 1 mL was deposited on a filter paper using the insert and metal cone.







be more)

Samples centrifuged at 5000 RPM for five minutes Insert disassembled, filter mounted on steel with double-sided sticky tape

Sample ablated in Ar environment



Images of filter after data acquisition





be more)





Spectrum From Bacteria





Detecting Species in Fluids with Chemometrics

"Ratio Model" consists of:

- 15 emission line intensities from P, C, Mg, Ca, and Na and 92 simple ratios
- 107 total independent variables





Diagnosing Species in Fluids with Machine Learning





Diagnosing Species in Water with Machine Learning



Late Search Source Bun Debug Convoles Projects Jook Yew Help						
PCA-ANN With Full Spectrum Data						
00 pllssotter(princips 00 pll	E. coli	S. aureus	E. cloacae			
Interference of the second sec	98.04 %	93.27 %	91.23 %			
pt:.short pt:.short	97.71 %	97.22 %	96.12 %			
pt: figure() logiting ("NS") star Classification Error array - defines to array (2.13 %	4.28 %	6.33 %			
<pre>ampliped at 1 sample to zero, how third 'insertion' a submark insertion to the submark submark the submark in a submark insertion to the submark submark the submark insertion to the submark submark insertion for the submark submark insertion to the submark for the submark submark insertion to the submark insertion to the submark submark insertion to the submark submark insert of the submark submark insertion to the submark submark submark submark insertion to the submark insert submark submark submark submark insertion to the submark submark submark submark submark submark insertion to the submark submark submark submark submark submark submark submark submark submark submark submark submark submark submark submark submark submark submark subm</pre>		New Section 0 Section Section 1 0 Section Section 1 0 Section Section 1 0 Section Section 1 0 Section Section Section 1 Section Section Section Se	الا العلم المراكع الله المراكع الله المراكع الله المراكع الله المراكع الله المراكع الله المراكع المراكع الله الله الله الله الله الله الله الل			

- PCA-ANN custom written in Python
- > 42,000 independent variables ⇒ 10 principal components (new independent variables)
- Currently only one hidden layer
- ANN algorithm implemented to optimize the patience and # of hidden nodes
- Models are trained on 80% of single shot data, 20% reserved for testing.
- After PCA, spectra were randomized to demonstrate the ANN was not fitting noise (random classification of cases)



Blood (sepsis)

Sepsis (bacterial blood infection) is a highly dangerous infection and is a common nosocomial (hospital acquired) infection.

Sepsis is not always caused by bacterial infection, but many cases are bacterial in origin.

A blood culture is required for the diagnosis of sepsis, but <u>not all blood cultures will</u> <u>produce a positive test result</u>.

Antibiotics are the preferred treatment for sepsis, but even with the best treatment the <u>mortality for patients</u> who have reached septic shock is <u>no better than 50%</u>.

The time it takes to initiate effective antimicrobial therapy is the single strongest predictor of patient outcome, with <u>every hour of delay</u> increasing patient mortality.

E. Whitnack, Sepsis, in: N.C. Engleberg, V.J. DiRita, T. Dermody, M. Schaechter (Eds.), Mechanisms of Microbial Disease, 3rd ed, Lippincott Williams & Wilkins, 2007, pp. 564–572.

A. Kumar, D. Roberts, K.E. Wood, B. Light, J.E. Parrillo, et al., Duration of hypotension before initiation of effective antimicrobial therapy is the critical determinant of survival in human septic shock, Crit. Care Med. 34 (2006) 1589–1596.



Blood (sepsis)

The standard diagnostic blood cultures are slow and labor intensive, usually requiring a two-step procedure:

1) an initial culture taking up to 120 h

2) a 24-48 h subculture to identify the pathogens detected in a positive initial culture.



Automated blood culture system



Culturing and diagnosis done manually

J.C. Lagier, S. Edouard, I. Pagnier, O. Mediannikov, M. Drancourt, D. Raoult, Current and past strategies for bacterial culture in clinical microbiology, Clin. Microbiol. Rev. 28 (2015) 208–236, https://doi.org/10.1128/cmr.00110-14

D.J. Shin, N. Andini, K. Hsieh, S. Yang, T.-H. Wang, Emerging analytical techniques for rapid pathogen identification and susceptibility testing, Ann. Rev. Anal. Chem. 12 (2019) 41-67, https://doi.org/10.1146/annurev-anchem-061318-115529



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Detecting Infection in Blood with a PLS-DA



Bacterial-negative specimens obtained from Windsor Regional Hospital with the anticoagulant sodium polyanetholesulfonate (SPS) present.

Five different blood samples from different patients were provided to account for the difference between patient physiology.

Specimens were vortex-mixed prior to handling to insure the redistribution of any suspended particles.



DETECTION IN BLOOD (individual spectra)

96.3% sensitivity – 600 bacterial spectra 98.6% specificity – 206 blood spectra 11,000 cells / spectrum

DETECTION IN BLOOD (spectra added)

19 of 19 filters positive, 100% sensitivity7 of 7 filters negative, 100% specificity



Prepared Samples:

- 5 filters each of *S. aureus*, *E. coli*, *E. cloacae*
- 4 filters of *P. aeruginosa*
- 7 filters of sterile blood
- 30 LIBS laser shots from each deposition



Diagnosing Infection in Blood with PCA-ANN



Bacterial-negative specimens obtained from Windsor Regional Hospital with the anticoagulant sodium polyanetholesulfonate (SPS) present.

Five different blood samples from different patients were provided to account for the difference between patient physiology.

Specimens were vortex-mixed prior to handling to insure the redistribution of any suspended particles.



	PCA-ANN With Full Spectrum Data*					
	S. aureus E. coli E. cloacae P. aeruginosa					
Sensitivity	100 %	100 %	100 %	100 %		
Specificity	100 % 100 % 100 % 100 %					
Classification Error	0.00 %	0.00 %	0.00 %	0.00 %		

*classification using 80:20 split

Prepared Samples:

- 5 filters each of *S. aureus*, *E. coli*, *E. cloacae*
- 4 filters of *P. aeruginosa*
- 7 filters of sterile blood
- 30 LIBS laser shots from each deposition



Urine (UTI)

UTI (urinary tract infections) are one of the most common infections in adult women, with up to 50% of women having experienced at least one UTI in their lifetime and upwards of 10% of women experiencing at least one UTI annually.

In 2007, UTI resulted in approximately 8.6 million health care visits in the United States with an estimated cost of 1.6 billion US dollars.

In addition to the prevalence in the population, UTI are also <u>the most common</u> <u>nosocomial infection</u>, with up to 80% of hospital-acquired UTI associated with the use of a bladder catheter.

Even <u>a single catheterization can lead to UTI</u> due to contamination of the catheter tip.

T.K. Price, E.E. Hilt, T.J. Dune, E.R. Mueller, A.J. Wolfe, L. Brubaker, Urine trouble: should we think differently about UTI?, Int. Urogynecol. J. 29 (2018) 205-210, S.M. Schappert, E.A. Rechtsteiner, Ambulatory medical care utilization estimates for 2007, Vital Health Stat. 13 169 (2011) 1-38, PMID: 21614897.

J.M.T. Barfor, K. Anson, Y.Hu, A.R.M. Coates, A model of catheter-associated urinary tract infection initiated by bacterial contamination of the catheter tip, BJU Int. 102 (2008) 67-74.



Urine (UTI)







Detecting Infection in Urine with a PLS-DA



Bacterial-negative urine specimens obtained from Windsor Regional Hospital.

Urine specimens from four different patients were characterized to account for differences in patient physiology.

Specimens were vortex-mixed prior to handling to insure the redistribution of any suspended particles.

DETECTION IN URINE (individual spectra)

98.9% sensitivity – 360 bacterial spectra 100% specificity – 240 urine spectra 11,000 cells per spectrum

DETECTION IN URINE (spectra added)

12 of 12 filters positive, 100% sensitivity 8 of 8 filters negative, 100% specificity





Diagnosing Infection in Urine with PCA-ANN



Bacterial-negative urine specimens obtained from Windsor Regional Hospital.

Urine specimens from four different patients were characterized to account for differences in patient physiology.

Specimens were vortex-mixed prior to handling to insure the redistribution of any suspended particles.

PCA-ANN With Full Spectrum Data*						
S. aureus E. coli E. cloacae						
Sensitivity	100 %	100 %	91.67 %			
Specificity 100 % 95.83 % 100 %						
Classification Error	0.00 %	2.09 %	4.17 %			

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*classification using 80:20 split



External Validation of PCA-ANN in Blood and Urine

- External validation done in both urine and blood (whole filters withheld)
- ANN model run 10 times per excluded filter

Improvements need to be made on quality of spectra, not ML •

										Urine
					Sensitivity of each	E. coli <i>filter remove</i>	d from the model to	be externally valida	ted. ANN run 10 time	s per filter.
					E. coli		P1	redicted		
	а	iverage ser	nsitivity of 7	4.6%	Sample #	S. aureus	Е. со	li	E. cloacae	Sensitivity
	- "				1	30	250		20	0.833
Urine	E. coli	S. aureus	E. cloacae		<2	1	138		161	0.460
					≤ 3	11	153		136	0.510
Average	63.58 %	86.92 %	62.33 %		4	72	222		6	0.74
Sonsitivity					Sum	114	763		323	0.636
Dlaad	F	-								
Average	E. COII	<i>S. aureus</i>	<i>E. cloacae</i>	P. aeruginosa	Sensitivity result:	s for S. aureus filters	removed from the 1	nodel to be externall	v validated. ANN run	Blood
Average	93.73 %	<i>S. aureus</i> 61.73 %	<i>E. cloacae</i> 91.60 %	<i>P. aeruginosa</i> 95.0 %	Sensitivity result: S. aureus	s for S. aureus filters	removed from the r Pr	nodel to be externally edicted	v validated. ANN run	Blood 10 times per filt
Average Sensitivity	93.73 %	5. aureus	<i>E. cloacae</i> 91.60 %	<i>P. aeruginosa</i> 95.0 %	Sensitivity results S. aureus Sample #	5 for S. aureus filters	removed from the r Pr <i>E. coli</i>	nodel to be externally edicted <i>E. cloacae</i>	v validated. ANN run P. aeruginosa	Blood 10 times per filt Sensitivit
Average Sensitivity	93.73 %	<i>S. aureus</i> 61.73 %	<i>E. cloacae</i> 91.60 %	<i>P. aeruginosa</i> 95.0 %	Sensitivity result. S. aureus Sample #	s for S. aureus filters S. aureus 50	removed from the I Pr <u>E. coli</u> 54	nodel to be externally edicted <u>F. cloacae</u> 86	v validated. ANN run <u>P. aeruginosa</u> 110	Blood 10 times per filt Sensitivit 0.167
Average Sensitivity	93.73 %	61.73 %	<i>E. cloacae</i> 91.60 % nsitivity of 8	<i>P. aeruginosa</i> 95.0 % 5.5%	Sensitivity result: S. aureus Sample # 1 2	<i>s for</i> S. aureus <i>filters S. aureus</i> 50 300	removed from the I Pr <u>E. coli</u> 54 0	nodel to be externally edicted <u>F. cloacae</u> 86 0	v validated. ANN run <u>P. aeruginosa</u> 110 0	Blood 10 times per file Sensitivit 0.167 1.000
Average Sensitivity	93.73 %	<i>S. aureus</i> 61.73 % overage ser	91.60 %	<i>P. aeruginosa</i> 95.0 % 5.5%	Sensitivity result: S. aureus Sample # 1 2 3	<i>s for</i> S. aureus <i>filters S. aureus</i> 50 300 1	removed from the 1 Pr <u>E. coli</u> 54 0 27	nodel to be externally edicted <u>F. cloacae</u> 86 0 266	v validated. ANN run <u>P. aeruginosa</u> 110 0 6	Blood 10 times per filt Sensitivit 0.167 1.000 0.033
Average Sensitivity	93.73 %	S. aureus	91.60 %	<i>P. aeruginosa</i> 95.0 % 5.5%	Sensitivity result: Sample # 1 2 3 4	<i>s for</i> S. aureus <i>filters S. aureus</i> 50 300 1 290	removed from the 1 Pr <u>E. coli</u> 54 0 27 10	nodel to be externally edicted <u>F. cloacae</u> 86 0 266 0	v validated. ANN run <u>P. aeruginosa</u> 110 0 6 0	Blood 10 times per filt Sensitivit 0.167 1.000 0.033 0.967
Average Sensitivity	93.73 %	<i>S. aureus</i> 61.73 % werage ser	<i>E. cloacae</i> 91.60 %	<i>P. aeruginosa</i> 95.0 % 5.5%	Sensitivity result: Sample # 1 2 3 4 5	s for S. aureus filters S. aureus 50 300 1 290 285	removed from the r Pr <u>E. coli</u> 54 0 27 10 13	nodel to be externally edicted <u>F cloacae</u> 86 0 266 0 0 0	v validated. ANN run <u>P. aeruginosa</u> 110 0 6 0 2	Blood 10 times per file Sensitivit 0.167 1.000 0.033 0.967 0.950



Cerebrospinal Fluid (meningitis)

Cerebrospinal fluid (CSF) is the fluid the brain, cranium, and spinal cord are bathed in.

Meningitis is an infection of the brain and spinal cord's protective layers (cells in CSF).

Meningitis may be viral or bacterial in nature.

Delays and inappropriate antibiotic use can lead to severe complications, including <u>irreversible brain damage or death</u>.

The gold standard for diagnosis is a spinal tap (lumbar puncture) to remove **cerebrospinal fluid (CSF)** followed by culture, serology, and analysis.



Slane, Valori H. and Chandrashekhar G. Unakal. "Tuberculous Meningitis." StatPearls, StatPearls Publishing, 18 November 2022. Telano, Lauren N. and Stephen Baker. "Physiology, Cerebral Spinal Fluid." StatPearls, StatPearls Publishing, 4 July 2023.



Cerebrospinal Fluid (meningitis)

Spinal taps are rarely performed in Windsor.

Artificial cerebrospinal fluid was used in this study (aCSF).

Artificial cerebrospinal spinal fluid (aCSF) closely matches the electrolyte concentrations of actual CSF and is prepared from analytical grade reagents and high purity water.

lon	Concentrations (mM)
Na⁺	150
K+	3.0
Ca ²⁺	1.4
Mg ²⁺	0.8
P ³⁻	1.0
Cl-	155





New Apparatus Design

Two components, with a built-in concentration cone.

- ✓ Concentrates bacteria in a 1 mm diameter circle
- The ridge on the aperture of β piece prevents bacterial escape, allowing only water and ions to diffuse
- ✓ Indent created on filter is more easily identifiable by the LIBS apparatus and seals bacteria

Stronger bacterial signal and more reproducible data









New Apparatus Data

nitrocellulose filter (0.45 μm pore size)





New Apparatus Data





Detecting Infection in aCSF with a PLS-DA



Bacterial-negative aCSF specimens procured from a biochem supply vendor

Prepared Samples:

- 11 filters each of S. aureus,
 E. coli, M. smegmatis
- 13 filters of sterile aCSF
- 25 LIBS laser shots from each deposition

DETECTION IN URINE (individual spectra)

99.8% sensitivity – 825 bacterial spectra 100% specificity – 325 urine spectra 11,000 cells per spectrum DETECTION IN URINE (spectra added)

33 of 33 filters positive, 100% sensitivity 13 of 13 filters negative, 100% specificity





Diagnosing Infection in aCSF with PCA-ANN



Bacterial-negative aCSF specimens procured from a biochem vendor

Prepared Samples:

- 11 filters each of S. aureus,
 E. coli, M. smegmatis
- 13 filters of sterile aCSF
- 25 LIBS laser shots from each deposition

Internal cross-validation (80:20 split) randomly selects 20% of the dataset to be tested against remaining 80% of spectra.

80:20 Cross-Validation Test Results						
E. coli S. aureus M. smegmatis Average						
Sensitivity	95.8 %	96.0 %	100.0 %	97.2 %		
Specificity	98.2 %	98.0 %	99.7 %	98.6 %		
Classification Accuracy	97.4 %	97.3 %	99.8 %	98.2 %		

External validation tests individually compare one filter (25 spectra) against left over spectra.

External Validation Test Results						
E. coli S. aureus M. smegmatis Average						
Sensitivity	74.6 %	71.1 %	99.5 %	81.7 %		
Specificity	87.6 %	87.9 %	97.0 %	90.9 %		
Classification Accuracy	83.3 %	82.3 %	97.9 %	87.8 %		



The first three PC scores were plotted to visualize the variance in the data



Conclusions

• Bacteria can be **detected** (PLSDA) and **diagnosed** (PCA-ANN) in blood, urine, and CSF.

PCA-ANN With Full Spectrum Data (Urine)						
S. aureus E. coli E. cloacae						
Sensitivity	100 %	100 %	91.67%			
Specificity	100 %	95.83 %	100 %			
Classification Error	0.00 %	2.09 %	4.17 %			

PCA-ANN With Full Spectrum Data (Blood)						
	S. aureus E. coli E. cloacae P. aeruginosa					
Sensitivity	100 %	100 %	100 %	100 %		
Specificity	100 %	100 %	100 %	100 %		
Classification Error	0.00 %	0.00 %	0.00 %	0.00 %		

PCA-ANN With Full Specrum Data (aCSF)							
E. coli S. aureus M. smegmatis Average							
Sensitivity	95.8 %	96.0 %	100.0 %	97.2 %			
Specificity	98.2 %	98.0 %	99.7 %	98.6 %			
Classification Accuracy	97.4 %	97.3 %	99.8 %	98.2 %			



Class 1: aCSF



Conclusions



 Using PCA-ANN on full spectrum data provides the best results for discrimination between bacterial species (using 80:20 split)

E. coli S. aureus E. cloacae Average sensitivity = 94 % Sensitivity 98.04 % 93.27 % 91.23 % Average specificity = 96 % Specificity 97.71% 97.22 % 96.12 % Classification Error 4.28 % 2.13 % 6.33 %

Approximate increase from DFA \approx 30 % (sensitivity), 16% (specificity)



Future Work

Improve external validation using PCA-ANN for bacterial species

- New 3D-printed centrifuge insert with integrated concentration cone help to improve signal-to-noise – revisit blood/urine.
- Work to further optimize the PCA-ANN algorithm (adding hidden layers?)
- Determine what is causing the "between-filter" variance in the measurements

Discrimination of lower concentrations of cells to find limit of identification (LOI)

Engage microbiology students to investigate the behavior of cells in clinical specimens

Extend analysis to other bacterial strains

Mycobacterium tuberculosis and Streptococcus mitis



If you want to learn more:

E.J. Blanchette. *Detection and Diagnosis of Bacterial Pathogens in Blood and Urine Using Laser-Induced Breakdown Spectroscopy*. Master's thesis, University of Windsor, 2022.

E.J. Blanchette et al., "Detection and Classification of Bacterial Cells After Centrifugation and Filtration of Liquid Specimens Using Laser-Induced Breakdown Spectroscopy," *Applied Spectroscopy* **76**, 2022, pp. 894-904. <u>https://doi.org/10.1177/000370282210927</u>

E.J. Blanchette, E.A. Tracey, A. Baughan, G.E. Johnson, H. Malik, C.N. Alionte et al. "Detection and diagnosis of bacterial pathogens in blood using laser-induced breakdown spectroscopy," *Spectrochim. Acta B* **215** (2024) 106911. <u>https://doi.org/10.1016/j.sab.2024.106911</u>

E.J. Blanchette, E.A. Tracey, A. Baughan, G.E. Johnson, H. Malik, C.N. Alionte, et al. "Detection and diagnosis of bacterial pathogens in urine using laser-induced breakdown spectroscopy," *Spectrochim. Acta B* **216** (2024) 106944. <u>https://doi.org/10.1016/j.sab.2024.106944</u>







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Windsor, Ontario, Canada

Thank you!



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LIBS primer

2) removal of samples mass (ablation)









M. smegmatis



E. coli



S. aureus





PCA-ANN Results – Identifying Bacteria

GOAL: Classify the species of bacteria in urine

Input: Entire LIBS spectrum from 200 nm to 590 nm (42,000 variables)



PCA: Spectrum is reduced to 10 principal components (PC) scores, through the use of a Python code developed from several libraries

9 | 10

 urine
 Python libraries used:

 30
 sklearn.decomposition, Pandas,

 ilter
 Numpy, and mpl_toolkits

ANN: Neural network developed on Python code relates 10 PC scores to bacterial species



Python libraries used: Pandas, Numpy, Tensorflow, and Scikit-Learn



LIBS primer

4) expansion and element specific emission (atomic or ionic)

spontaneous emission as atoms/ions decay to ground state

crater debris

The dynamic evolution of the plasma plume is then characterized by a fast expansion and subsequent cooling.

Approximately 1 microsecond after the ablation pulse, spectroscopically narrow atomic/ionic emissions may be identified in the spectrum.





The Centrifuge Insert



Concentrating Bacteria With a Cone

19 mm long Al cone

Centrifuge tube cap presses cone into filter





1 mm hole at apex

Holds 1 mL of fluid

Cone vertex press fit into filter

Shooting Bacteria Concentrated With Cone





Limits of Detection



A calibration curves constructed from forty spectra obtained from each of nine different concentrations.

LIBS bacterial limit of detection of $10,865 \pm 3,712$ CFU per laser ablation event for bacteria deposited on filters using the metal cone.

LOD's calculated using only certain elements observed in the LIBS spectra and present in very low concentrations in the filter were even lower:

1,070 \pm 272CFU for magnesium 1,784 \pm 657 CFU for calcium.

LOD on filter better, but number of cells required in fluid specimen is **WAY** lower!

When performed with no background

- We can identify a bacterial species, certainly its genus, with high sensitivity and specificity (confirmed by others).
- ✓ We can differentiate strains of *E. coli* (demonstrated by others in MRSA).
- Multiple multivariate techniques effective at discriminating spectra.

PLSDA			DFA			
E. COLI	True	False	E. COLI	True	False	
Positive	95.65%	9.17%	Positive	89.63%	15.95%	
Negative	90.83%	4.35%	Negative	84.05%	10.37%	
STAPHYLOCOCCUS	True	False	STAPHYLOCOCCUS	True	False	
Positive	54.05%	0.51%	Positive	86.49%	5.85%	
Negative	99.49%	45.95%	Negative	94.15%	13.51%	
STREPTOCOCCUS	True	False	STREPTOCOCCUS	True	False	
Positive	95.59%	1.02%	Positive	99.26%	13.32%	
Negative	98.98%	4.41%	Negative	88.68%	0.74%	
MYCOBACTERIUM	True	False	MYCOBACTERIUM	True	False	
Positive	88.31%	1.06%	Positive	96.10%	4.08%	
Negative	98.94%	11.69%	Negative	95.92%	3.90%	
DFA:	DEA: Sensitivity: 91 37 + 16 39 % Specificity: 97 46 + 9 35 %					

Specificity: 90.60 ± 21.33 %

PLSDA: Sensitivity: 93.13 ± 10.25 %





Results: We have already demonstrated...

LIBS spectral fingerprint is a sensitive and specific (high rates of true positives, low rates of false positives) test to identify an unknown bacterial specimen or to differentiate between possible identifications

This spectral fingerprint is robust and reliable, and exists through time (multiple tests spanning years on same strains of bacteria)

≻In addition...

9 publications in Applied Physics Letters, Journal of Applied Physics, Applied Optics, Applied Spectroscopy, Spectrochimica Acta B, and others – confirmed by multiple other groups

Results: We have already demonstrated...

LIBS spectral fingerprint is:

- growth-medium independent
- independent of state of growth (how "old" the bacteria are)
- independent of whether the bacteria are live or dead (or inactivated by UV light)
- obtainable even when other types of bacteria or contaminants are present (mixed samples)
- obtainable from urine specimens
- capable of strain discrimination
- obtainable from about 500 bacteria

9 publications in Applied Physics Letters, Journal of Applied Physics, Applied Optics, Applied Spectroscopy, Spectrochimica Acta B, and others – confirmed by multiple other groups

(Poster 344) Bacterial Mounting and Concentration Techniques to Translate Laser-Induced Breakdown Spectroscopy into a Clinical Setting; Alexandra Paulick





