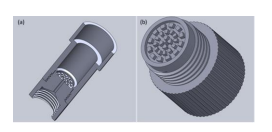
EMSLIBS 2019

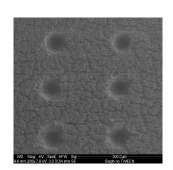
10th Euro-Mediterranean Symposium on Laser-Induced Breakdown Spectroscopy 8 – 13th September 2019 | Brno, Czech Republic

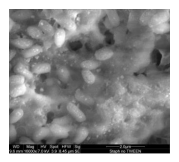
Bacterial Limit of Detection Reduction Utilizing a Novel Sample Preparation Protocol

Steven J. Rehse, Jeremy Marvin, Alexandra E. Paulick, Emma Blanchette, Sydney Sleiman

Department of Physics, University of Windsor, Windsor, Ontario, Canada

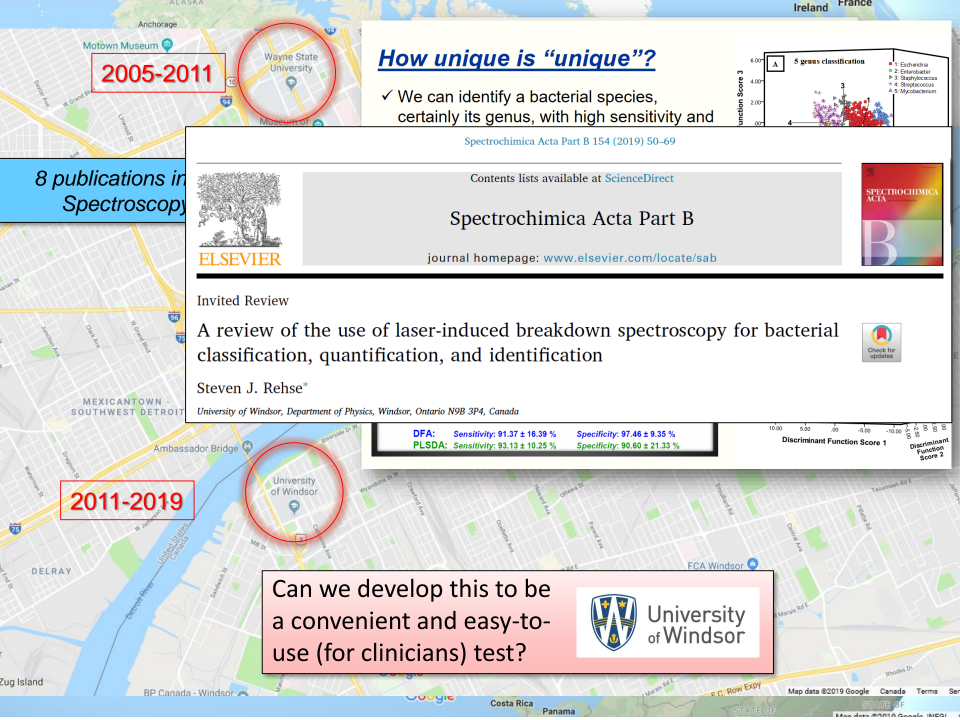












Early days (at Wayne State)

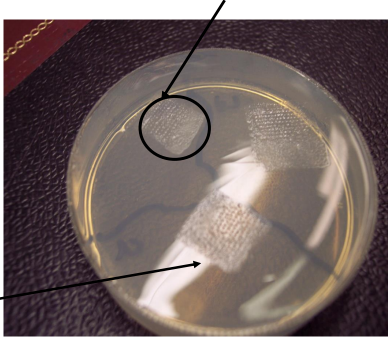
Advantages:

✓ Background free mounting substrate

Disadvantages:

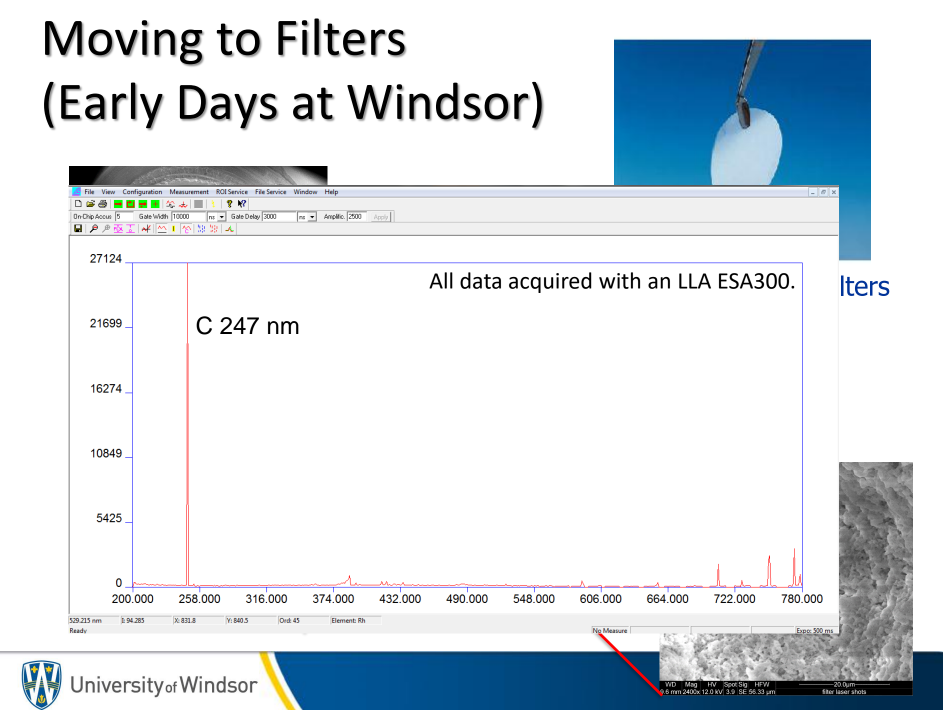
- X Not really flat
- X Degrades
- X Hard to make
- X Watery

University of Windsor



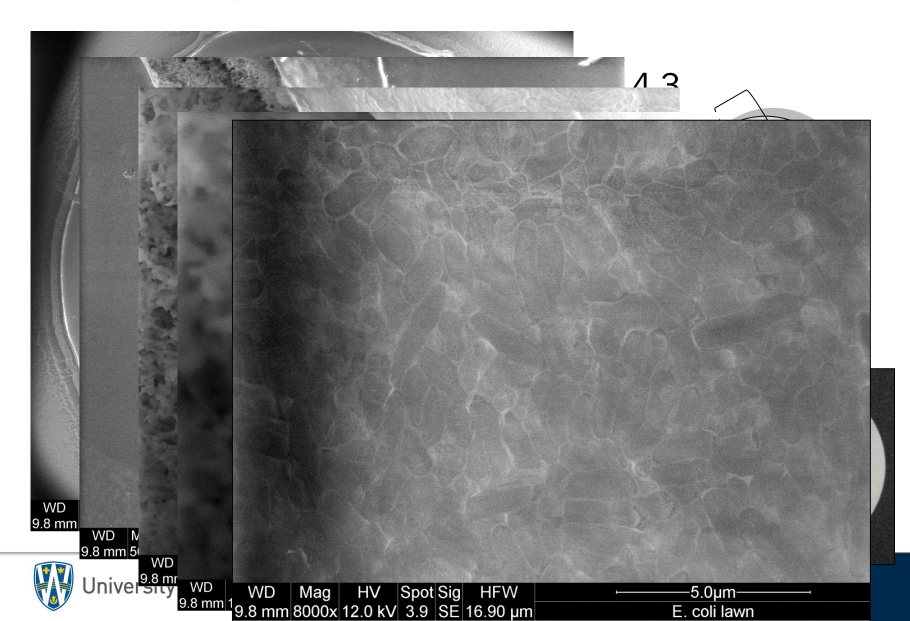
10 microliter

Nutrient-free bacto-agar



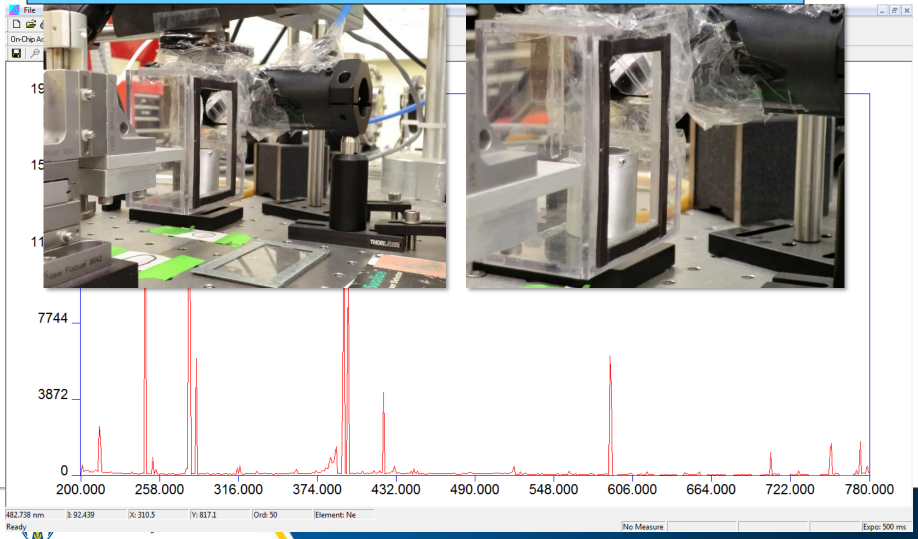
How to get bacteria onto filters?

How to get bacteria onto filters?

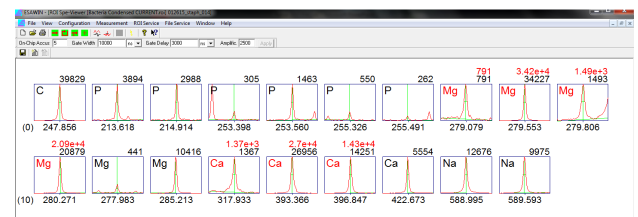


Typical bacterial LIBS spectrum

Currently: using matched parabolic reflectors into fiber for UV



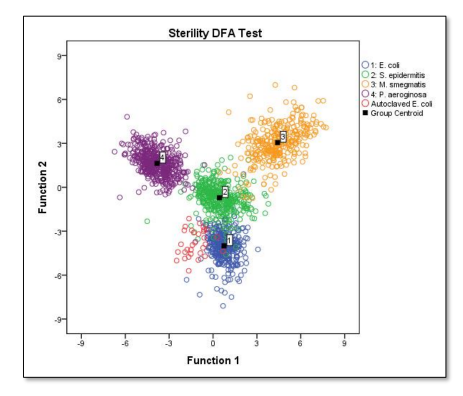
Data analysis with variable down selection



- 164 independent variables
 - 19 line intensities (all divided by sum)
 - 145 ratios of intensities



Performance on filters



DF	A Classif	ication (Grouped by Spe	ecies	
Escherichia	TRUE	FALSE	Staphylococcus	TRUE	FALSE
Positive	98.28%	0.77%	Positive	97.75%	1.44%
Negative	99.23%	1.72%	Negative	98.56%	2.25%
Mycobacterium	TRUE	FALSE	Pseudomonas	TRUE	FALSE
Positive	95.36%	0.33%	Positive	99.57%	0.22%
Negative	99.67%	4.64%	Negative	99.78%	0.43%
Sensitivity: 98 ± 2% Specificity: 99 ± 1%					

	PLS-DA Classification Grouped by Species					
Escl	herichia	TRUE	FALSE	Staphylococcus	TRUE	FALSE
Pc	sitive	96.55%	1.12%	Positive	96.75%	1.53%
Ne	gative	98.88%	3.45%	Negative	98.47%	3.25%
Мусо	bacterium	TRUE	FALSE	Pseudomonas	TRUE	FALSE
Pc	ositive	97.02%	0.41%	Positive	98.92%	0.33%
Ne	gative	99.59%	2.98%	Negative	99.67%	1.08%
	Sensitivity: 97 ± 3%			Specificity: 9	9±2%	

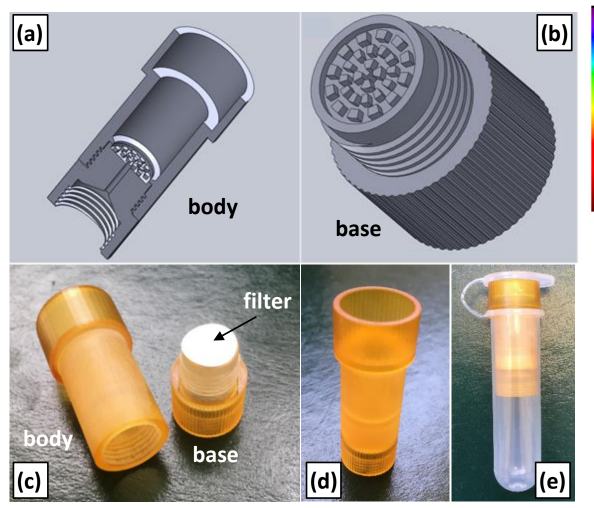
All external validation results

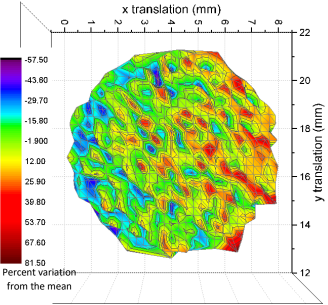
	DFA (by filter)	DFA (by species) above	PLSDA (by species) above
Sensitivity	0.93±0.07	0.98±0.02	0.97±0.03
Specificity	0.98±0.03	0.99±0.01	0.99±0.02

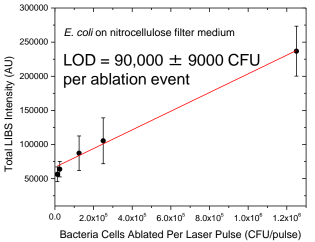


"by filter" means approximately 30 groups in DFA, no relationships between groups assumed

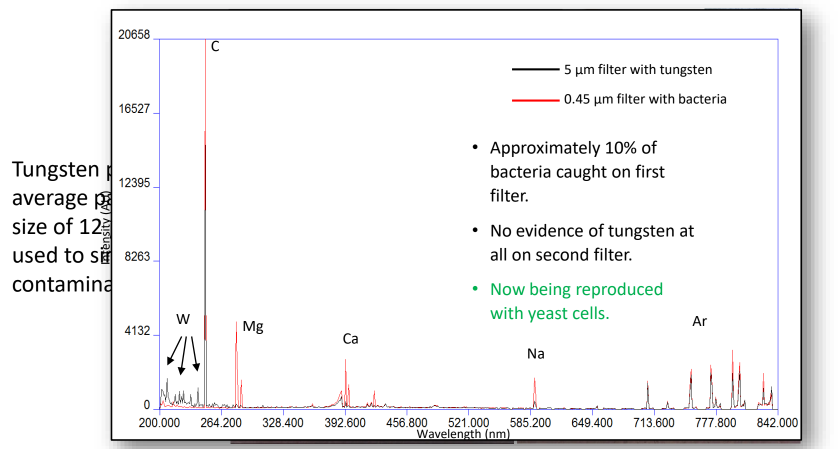
Using filters, a better way: the centrifuge insert







The centrifuge insert for cell sorting



- Tungsten powder was chosen for its inertness, cost and availability, safety of use, ease of preparation, biologically relevant size, and elemental purity.
- 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.

Using filters, a better way: the centrifuge insert

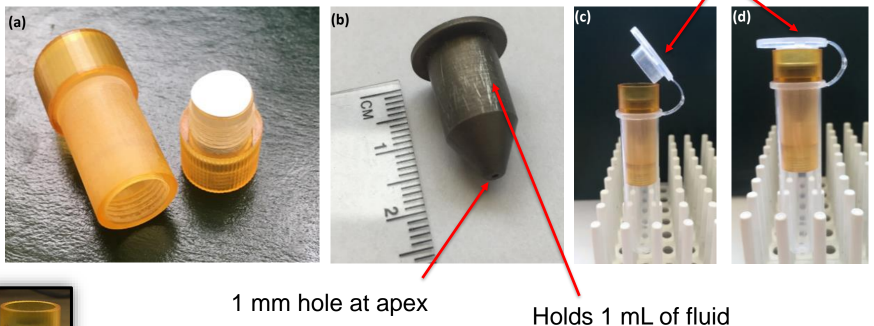




Using filters, an <u>even better</u> way: the centrifuge cone

19 mm long Al cone

Centrifuge tube cap presses cone into filter

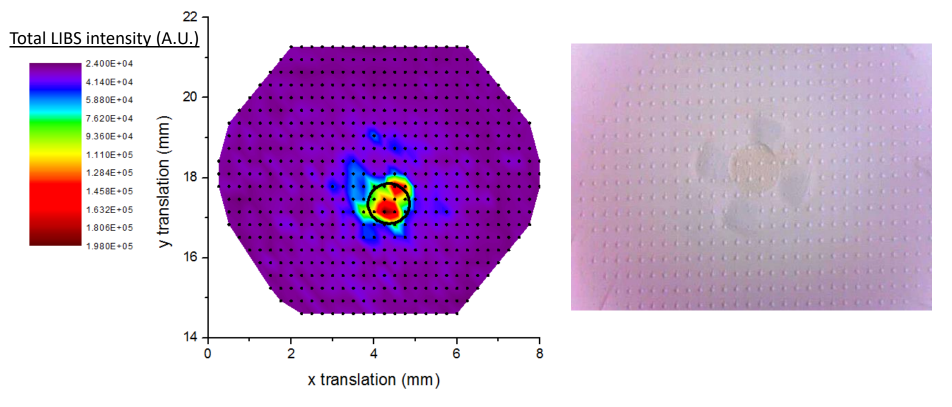


Cone vertex press fit into filter



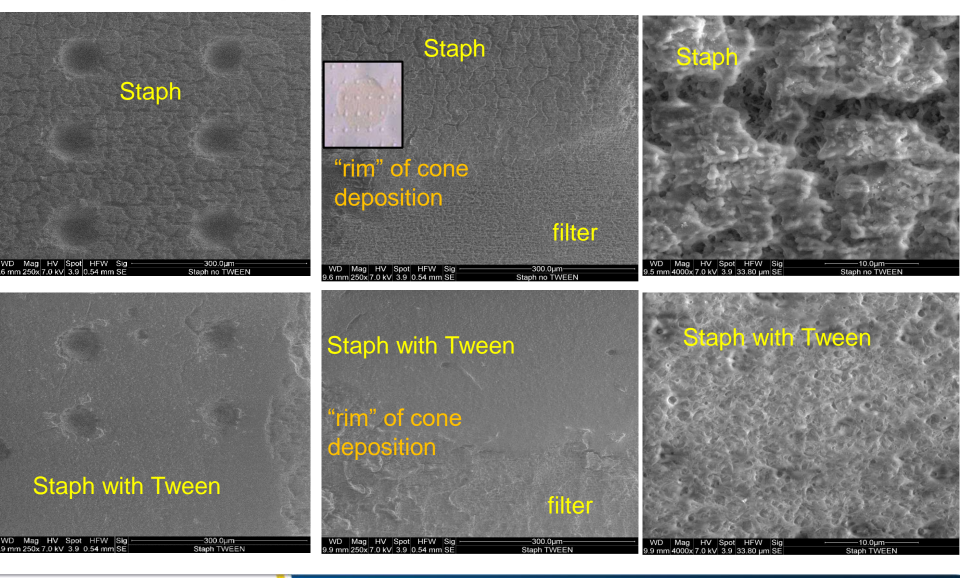
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Using filters, an <u>even better</u> way: the centrifuge cone



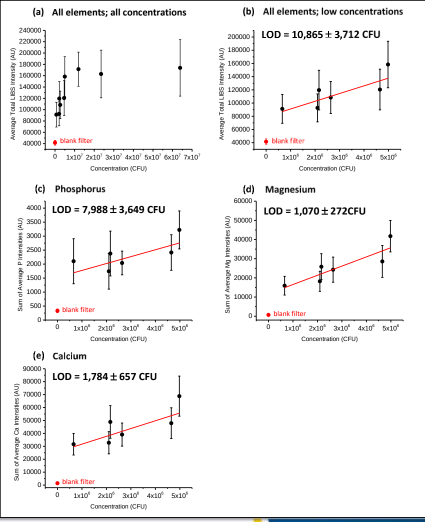


SEM micrographs





Using filters, an <u>even better</u> way: the centrifuge cone



niversity₀fWindsor

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

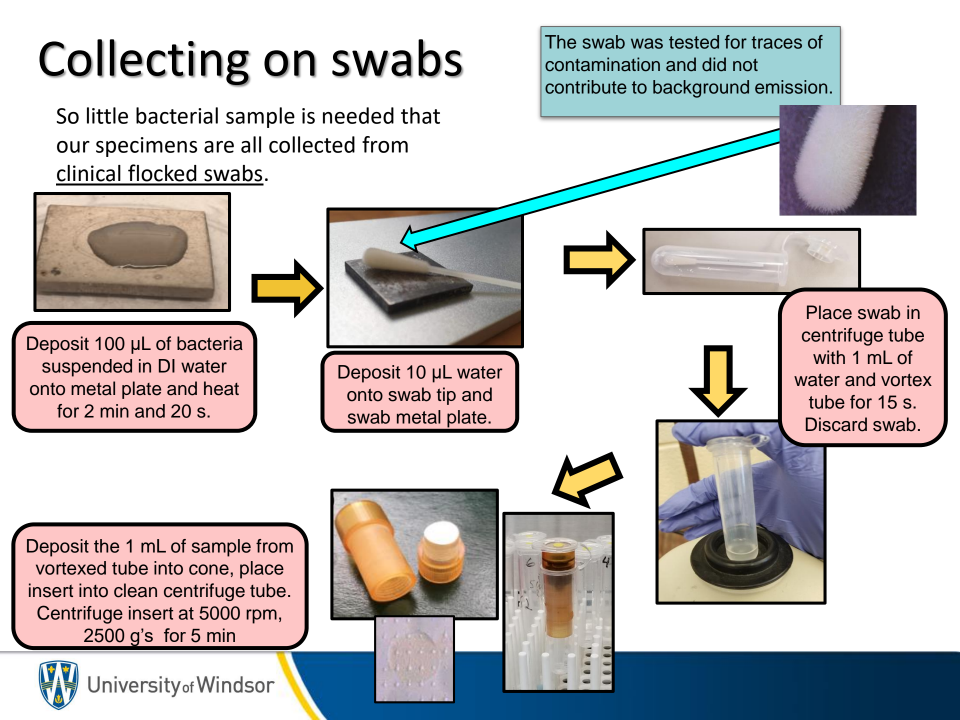
LIBS bacterial limit of detection of 10,865 \pm 3,712 CFU per laser ablation event for bacteria deposited on filters using the metal cone.

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

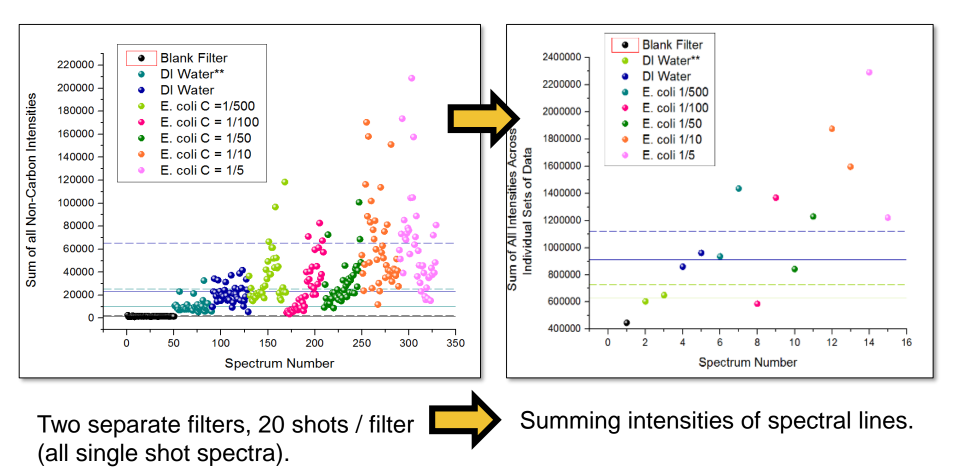
1,070 \pm 272CFU for magnesium 1,784 \pm 657 CFU for calcium.

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





Collecting on swabs





The people who did the work...



NSERC Discovery Grant



Natural Sciences and Engineering Research Council of Canada

Conseil de recherches en sciences naturelles et en génie du Canada

CFI-LOF grant

University of Windsor

University of Windsor



University₀fWindsor

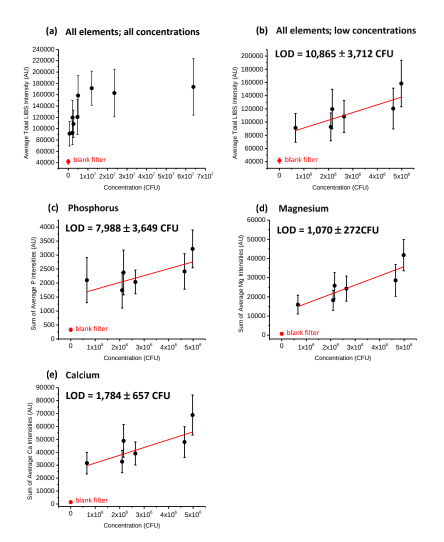
Thank you!



Using filters, a better way: the centrifuge insert







A calibration curve constructed from forty spectra obtained from each of the nine different concentrations returned a 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. Limits of detection 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 and 1,784 \pm 657 CFU for calcium. This represents a factor of 50 reduction in the limit of detection compared to our previously reported value.



Bacterial LOD

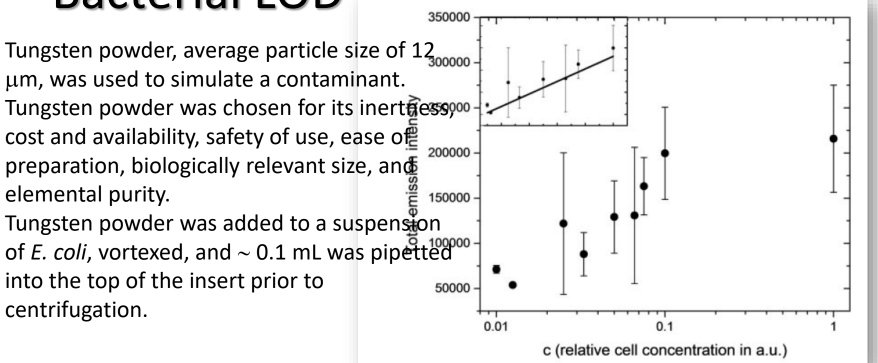
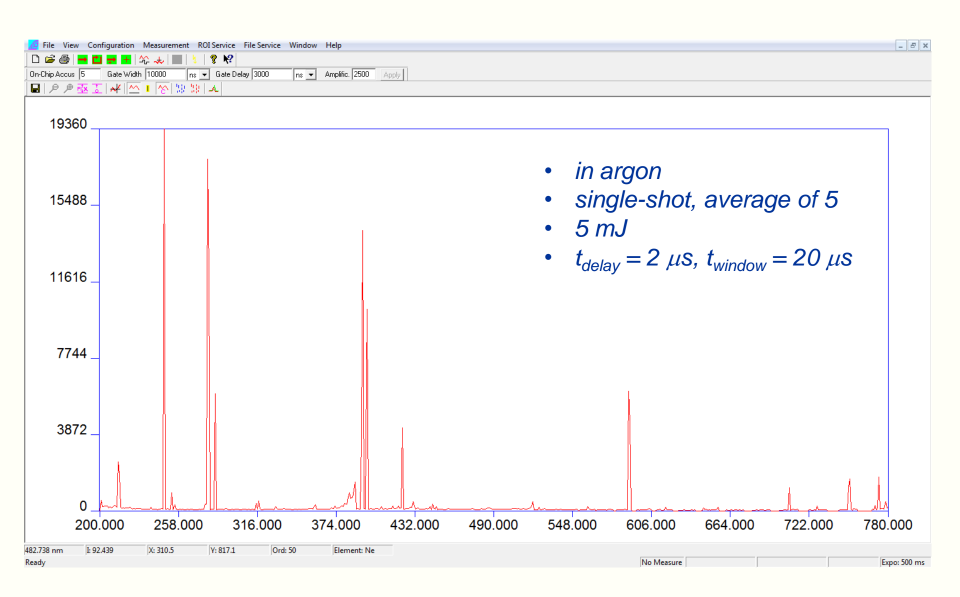


Figure 6. A log-lin calibration curve for bacterial samples. A concentration of c = 1 corresponded to 10^{11} cells/mL as determined by optical densitometry resulting in approximately 10^6 cells per ablation. This was the concentration achieved by transferring 24 h of growth for *E. coli* from a TSA plate to 1.5 mL distilled water. The inset shows a linear fit to the lowest eight concentrations on a lin-lin plot.

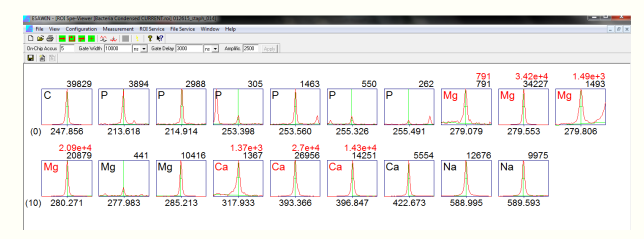




Typical Bacterial Spectrum

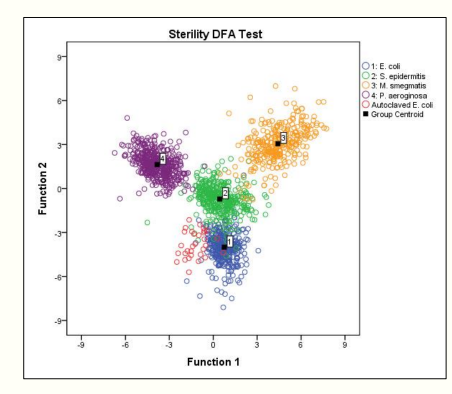


Variable Down-Selection



- New classification model
- 164 independent variable
 - 19 line intensities (all divided by sum)
 - 145 ratios of intensities
- No other metals. Beware?
 - Farooq (2014) sees S, CI, Mn, Fe, AI, Cu, etc.
 - Sivakumar (2015) only sees Ca, Na, Mg, K, O, H, C, P
 - We see Ni, Fe, Ti only when contaminated!

Performance With New Library



	DFA	A Classif	ication (Grouped by Spe	cies	
Escheric	hia	TRUE	FALSE	Staphylococcus	TRUE	FALSE
Positive	е	98.28%	0.77%	Positive	97.75%	1.44%
Negativ	'e	99.23%	1.72%	Negative	98.56%	2.25%
Mycobacte	rium	TRUE	FALSE	Pseudomonas	TRUE	FALSE
Positiv	е	95.36%	0.33%	Positive	99.57%	0.22%
Negativ	'e	99.67%	4.64%	Negative	99.78%	0.43%
ę	Sensi	tivity: 98	± 2%	Specificity: 99	9±1%	

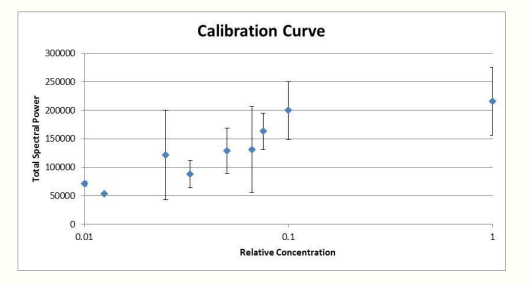
	PLS-DA Classification Grouped by Species					
Esch	herichia	TRUE	FALSE	Staphylococcus	TRUE	FALSE
Po	sitive	96.55%	1.12%	Positive	96.75%	1.53%
Ne	gative	98.88%	3.45%	Negative	98.47%	3.25%
Mycol	bacterium	TRUE	FALSE	Pseudomonas	TRUE	FALSE
Pc	sitive	97.02%	0.41%	Positive	98.92%	0.33%
Ne	gative	99.59%	2.98%	Negative	99.67%	1.08%
	Sensit	ivity: 97	± 3%	Specificity: 9	9±2%	

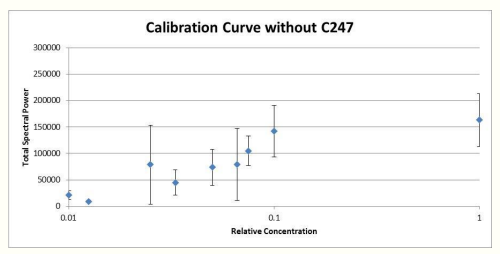
All external validation results

	DFA (by filter)	DFA (by species) above	PLSDA (by species) above
Sensitivity	0.93±0.07	0.98±0.02	0.97±0.03
Specificity	0.98±0.03	0.99±0.01	0.99±0.02

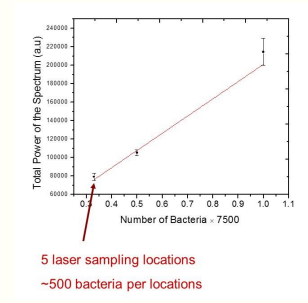
"by filter" means approximately 30 groups in DFA, no relationships between groups assumed

New Concentration Study

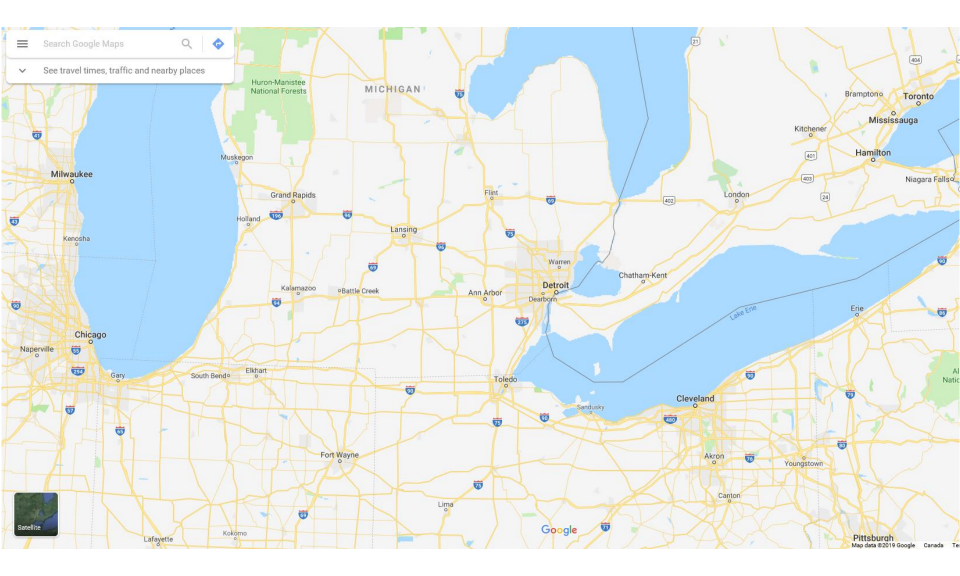




Previous result



- Performed with serial dilutions.
- Concentration 1" → harvest entire plate of colonies off TSA, suspend in 1.5 mL distilled H₂0
- Measure with optical densitometry
- OD=0.1 measured for C=0.001 (from literature OD 0.1=10⁸ cells/mL).
- * C=1 \rightarrow 10¹¹ cells/mL
- Implies for C=1, 10⁶/shot





How unique is "unique"?

- ✓ 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	, in the second s	
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%
	98.94%	11.69%	Negative	95.92%	3.90%

Specificity: 90.60 ± 21.33 %

PLSDA: Sensitivity: 93.13 ± 10.25 %

