

AFRL-AFOSR-JP-TR-2016-0074

Electrokinetic enrichment and detection of neuropeptide for performance monitor

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06/14/2016 Final Report

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	REPOR	Form Approved OMB No. 0704-0188				
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1. REPORT DA	TE (DD-MM-YYY	Y) 2. R	EPORT TYPE			3. DATES COVERED (From - To)
14-06-2016 4. TITLE AND S	UBTITLE	H H	nal		5a.	
Electrokinetic monitoring	enrichment an	d detection of r	neuropeptide for pe	rformance		
					5b.	GRANT NUMBER FA2386-14-1-4070
					5c.	PROGRAM ELEMENT NUMBER 61102F
6. AUTHOR(S) Nathan Swan	ni				5d.	PROJECT NUMBER
						TASK NUMBER
					5f.	WORK UNIT NUMBER
7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES) VIRGINIA UNIV CHARLOTTESVILLE						8. PERFORMING ORGANIZATION REPORT NUMBER
CHARLOTTESV	/ILLE, VA 22904-	4160 US				
9. SPONSORING/MONITORING AGENCY NAME(S) AND ADDRESS(ES) AOARD						10. SPONSOR/MONITOR'S ACRONYM(S) AFRL/AFOSR IOA
APO AP 96338	3-5002					11. SPONSOR/MONITOR'S REPORT NUMBER(S) AFRL-AFOSR-JP-TR-2016-0074
12. DISTRIBUTIO	ION/AVAILABILI N UNLIMITED: PE	TY STATEMENT 3 Public Release				
13. SUPPLEME	NTARY NOTES					
14. ABSTRACT The real-time missions requi present at dilu devices, so th conditions of minimal user i conditions for biomarker pre application to 15. SUBJECT T biomarkers, n	r monitoring of h res devices tha ute levels, there at each releva these force field ntervention. In t key neurologic econcentration wards unraveli ERMS anofluidics, pre	uman performa t enable facile o is a need for bio nt target may b ds can then be r his project, we o al biomarkers o was coupled to ng the signaling -concentration	nce biomarkers for f detection of multiple omarker enrichment e detected versus h routinely applied wit developed nano-slit f interest, by using na various detection p pathways for assess devices	atigue, vigilan targets with n under electric igh levels of int hin field setting devices and c anoparticles an paradigms to a sing and mitigo	ce, and stre ninimal user al and mag erfering mo gs for facile optimized th nd aptamer chieve high ating stress.	ess among DoD personnel during key intervention. Since the biomarkers are gnetic force fields within microfluidic olecules within biofluids. The optimized electrical and optical detection, with e electrokinetic preconcentration rs to enhance specificity. Additionally, n-sensitivity biomarker profiles for future
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						Standard Form 298 (Rev. 8/98) Prescribed by ANSI Std. Z39.18

Report for AOARD Grant # FA2386-14-1-4070: "Electrokinetic enrichment and detection of neuropeptides for human performance monitoring"

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Period of Performance: 12/01/2014 - 12/02/2015

Abstract: The real-time monitoring of human performance biomarkers for fatigue, vigilance, and stress among DoD personnel during key missions requires devices that enable facile detection of multiple targets with minimal user intervention. Since the biomarkers are present at dilute levels, there is a need for biomarker enrichment under electrical and magnetic force fields within microfluidic devices, so that each relevant target may be detected versus high levels of interfering molecules within biofluids. The optimized conditions of these force fields can then be routinely applied within field settings for facile electrical and optical detection, with minimal user intervention. In this project, we developed nano-slit devices and optimized the electrokinetic preconcentration conditions for key neurological biomarkers of interest, by using nanoparticles and aptamers to enhance specificity. Additionally, biomarker preconcentration was coupled to various detection paradigms to achieve high-sensitivity biomarker profiles for future application towards unraveling the signaling pathways for assessing and mitigating stress.

Introduction: Assessment and enhancement of the capabilities and alertness of the largest asset of the Air Force, namely their field personnel, is a key vision within the AFOSR and the Human Performance Wing of AFRL¹. Implementation of this vision requires the development of technologies for real-time monitoring of human performance biomarkers for fatigue, vigilance, and stress through micro-sampling from biofluids such as saliva, sweat, blood, and urine. However, characterizing and regulating stress conditions through biomarker expression analysis is a particularly challenging task due to the involvement of multiple inter-related targets with complex inter-regulating circuitry. Additionally, the biomarkers are typically present over a wide concentration range (mg/mL - pg/mL) within biofluids², thereby requiring the application of selective pre-concentration approaches for analyte enrichment over interfering proteins and small molecules. Herein, we seek to develop biomarker preconcentration methodologies based on electrical or magnetic force fields within micro/nanofluidic devices^{3,4} to achieve rapid localized biomarker enrichment due to the ensuing volume reduction (Fig. 1). As part of this initiative, this particular collaboration between Nathan S. Swami (Virginia) and Taiwan group led by Chia-Fu Chou (Academia Sinica) seeks to develop preconcentration and detection methodologies based on biomarkers from AFRL's 711th Human Performance Wing (Nancy Kelley-Loughnane). In this current year of the grant, we focused on coupling biomarker enrichment with nanoparticle immunoassays and aptamer-based approaches for enhancing detection specificity.

Experiment: Towards addressing AFRL's grand challenge of enabling Human Performance Monitoring, the experimental program was based on biomarkers, nanoparticles and assay chemistries from AFRL's 711th Human Performance Wing (Jorge Chavez & Nancy Kelley-Loughnane), detection systems from N. Swami's group (Virginia) and device technologies enabled by the group of C-F. Chou (Academia Sinica). Table 1 describes the experimental organization and supported researchers on this collaborative work.

Project	Supported Personnel	(Location) Outcome ^{citation}
Device for AC electrokinetic preconcentration of biomarkers and nanoparticles	Student: W. Varhue (50%); Postdoc: K.T. Liao (50%) (Year 1)	(Virginia & Taiwan) Nano-slit device with constrictions to frequency-selective enrichment ⁵ , ⁶
Modifying device fabrication for coupling electrochemical detection to preconcentration	Walter Varhue (50%) (Year 2 & 3)	(Virginia) Microfabricated cover-slip with nano-device for electrochemistry ⁷ , ⁸
Applying prior devices for preconcentration and detection of neuropeptides	Bankim Sanghavi (30%); AFRL collaborator: Jorge B. Chavez (Year 3)	(Virginia & Taiwan) Detection of NPY and Orexin A in various biological matrices ⁹
Electrochemical (EC) assay for aptamer-based detection of cortisol and NPY	Bankim Sanghavi (50%); AFRL collaborator: Jorge B. Chavez (Year 3)	(Virginia) Aptamer-based EC assay for cortisol and NPY detection (in progress)
Functionalization & mapping transcription factor sites	Yih-Li, Lin (50%) – Year 1 & 2	(Taiwan) Highly parallel analysis for resolving protein-binding locations on DNA probes ¹⁰ , ¹¹ , ¹²
Dielectrophoretic enrichment coupled to Raman spectroscopy	L. Lesser-Rojas (50%) – Year 3	(Taiwan) High sensitivity biomarker detection ¹³

Results and Discussion: The outcomes of the collaborative project are briefly described below: **1. "Nano-constriction device for rapid protein pre-concentration in physiological media** by electrokinetic force balance", K.T. Liao, M. Tsegaye, V. Chaurey, C.F. Chou, N. Swami*. *Electrophoresis* (2012), 33, 1958-1966. Impact Factor =3.3; DOI: 10.1002/elps.201100707



Fig. 1: Biomarker enrichment in physiological media by electrokinetic force fields at nano-constrictions.

Herein, we developed a methodology to steeply enhance biomarker pre-concentration within physiological media over that achieved through negative dielectrophoresis at nanoscale constriction gap devices, by utilizing an additional DC field offset to exponentially enhance the extent of protein depletion across the device. These protein pre-concentration methodologies may be applied towards biomarker discovery, protein crystallization, and rare target sensing for early disease diagnostics.

2. "Scaling down constriction-based dielectrophoresis for trapping nanoscale biomolecules in high conductivity media", V. Chaurey, A. Rohani, Y.-H. Su, W. Varhue, K.T. Liao, C. F. Chou, N. S. Swami*, *Electrophoresis* (2013), 34, 1097-1104. Journal Impact Factor =3.3



Fig. 2: Temperature rise at nano-Constrictions limits DEP trapping

Selective trapping of nanoscale biomarkers is significant for the separation and high-sensitivity detection of biomarkers. Dielectrophoresis is capable of highly selective trapping of bio-particles based on their characteristic frequency response. However, the trapping forces fall steeply with particle size, especially within physiological media of high-conductivity where the trapping can be dissipated by electrothermal flow due to localized Joule heating. Herein, we investigate the influence of device scaling within the electrodeless insulator dielectrophoresis geometry through the application of highly constricted channels of successively smaller channel depth, on the net balance of dielectrophoretic trapping force versus electrothermal drag force on bio-particles.

3. Real-time electrochemical monitoring of ATP at graphene-modified electrodes", B. Sanghavi, S. Sitaula, M. Griep, S. Karna, M. Ali, N. S. Swami* Anal. Chem. (2013), 85, 8158–8165 Journal Impact Factor =5.8: We report on a competitive electrochemical detection system that is free of wash-steps and enables the real-time monitoring of adenosine triphosphate (ATP) over a five-log concentration range, with the ability to speed-up target binding kinetics by increasing capture probe concentration. This displacement based assay enables biomarker detection by using nanoparticle-immobilized receptors, thereby obviating the need for functionalization of microfluidic devices to enable biomarker recognition.



Fig. 3: Example sensing paradigm based on competitive displacement of pre-bound electroactive FAD From aptamer receptors for enabling monitoring of ATP through electrochemical detection.

4. "Electrokinetic preconcentration and detection of neuropeptides at patterned graphene-modified electrodes in a nanochannel", Sanghavi, B. J., Varhue, W., Chávez, J. L., Chou, C. F., & Swami, N. S.*; Analytical chemistry, 86(9), 4120-4125. Impact Factor = 5.83;



Neuropeptides with electrochemical detection.

Neuropeptides are vital to the transmission and modulation of neurological signals, with Neuropeptide Y (NPY) and Orexin A (OXA) diagnostic information on offering stress. depression, and neurotrauma. NPY is an especially significant biomarker, since it can be noninvasively collected from sweat, but its detection has been limited by poor sensitivity, long assay times, and the inability to scale-down

sample volumes. Herein, we apply electrokinetic preconcentration of the neuropeptide onto patterned graphene-modified electrodes in a nanochannel by frequency-selective dielectrophoresis for 10 s or by electrochemical adsorptive accumulation for 300 s, to enable the electrochemical detection of NPY and OXA at picomolar levels from subnanoliter samples, with sufficient signal sensitivity to avoid interferences from high levels of dopamine and ascorbic acid within biological matrices. Given the high sensitivity of the methodology within small volume samples, we envision its utility toward off-line detection from droplets collected by microdialysis for the eventual measurement of neuropeptides at high spatial and temporal resolutions.

5. "Quantifying spatio-temporal dynamics of biomarker pre-concentration and depletion in microfluidic systems by intensity threshold analysis", <u>A. Rohani, W. Varhue, Y.-H. Su</u>, N. S. Swami; *Biomicrofluidics* (2014) 8, 052009. <u>Journal Impact Factor</u> =3.8

Microfluidic systems are commonly applied towards pre-concentration of biomarkers for enhancing detection sensitivity. Quantitative information on the spatial and temporal dynamics of pre-concentration, such as its position, extent and time evolution are essential towards sensor design for coupling pre-concentration to detection. Current quantification methodologies are based on the time evolution of fluorescence signals from biomarkers within a statically defined region of interest, which does not offer information on the spatial dynamics



Fig. 5: Biomarker preconcentration under force fields (a) over varying spatial (b) and temporal spreads (c & d) is quantified for alignment to sensing region (e).

of pre-concentration and leads to significant errors when the pre-concentration zone is delocalized or exhibits wide variations in size, shape and position over time under the force field. We present a dynamic methodology for quantifying the region of interest by using a statistical description of particle distribution across the device geometry to determine the intensity thresholds for particle pre-concentration. This method is applied to study the delocalized pre-concentration dynamics under an electrokinetic force balance driven by negative dielectrophoresis, for aligning the pre-concentration and detection regions of neuropeptide Y, and for quantifying the polarizability dispersion of silica nano-colloids with frequency of the force field. We envision the application of this automated methodology on data from 2D images and 3D Z-stacks for quantifying pre-concentration dynamics over delocalized regions as a function of the force field.

6. "DNA Combing on Low-Pressure Oxygen Plasma Modified Polysilsesquioxane Substrates For Single-Molecule Studies", K. K. Sriram, C.L. Chang, U. R. Kumar, C.F. Chou* (2014). *Biomicrofluidics* 2014, 8, 052102; DOI: 10.1063/1.4892515. (IF: 3.771): Molecular combing and flow-induced stretching are the most commonly used methods to immobilize and stretch DNA molecules. While both approaches require functionalization steps for the substrate surface and the molecules, conventionally the former does not take advantage of, as the latter, the of microfluidics regarding robustness. buffer exchange capability. versatility and molecule manipulation using external forces for single molecule studies. Here, we demonstrate a simple one-step combing process involving only low-pressure oxygen (O2) plasma modified polysilsesquioxane (PSO) polymer layer to facilitate both room temperature microfluidic device bonding and immobilization of stretched single DNA molecules without molecular functionalization step. Atomic force microscopy and Kelvin probe force microscopy experiments revealed a significant increase in surface roughness and surface potential on low-pressure O2 plasma treated PSQ, in contrast to that with high-pressure O2 plasma treatment, which are proposed to be responsible for enabling effective DNA immobilization. We further demonstrate the use of our platform to observe DNA-RNA polymerase complexes and cancer drug cisplatin induced DNA condensation using wide-field fluorescence imaging.

7. "Direct Optical Mapping of Transcription Factor Binding Sites on Field-stretched λ -DNA in Nanofluidic Devices", K.K. Sriram, J.W. Yeh, Y.L. Lin, Y.R. Chang, C.F. Chou* (2014). Nucleic Acids Research 2014, 42, e85. DOI: 10.1093/nar/gku254 (IF: 8.808): Mapping transcription factor (TF) binding sites along a DNA backbone is crucial in understanding the

regulatory circuits that control cellular processes. Here, we deployed a method adopting bioconjugation, nanofluidic confinement and fluorescence single molecule imaging for direct mapping of TF (RNA polymerase) binding sites on field-stretched single DNA molecules. Using this method, we have mapped out five of the TF binding sites of E. coli RNA polymerase to bacteriophage -DNA, where two promoter sites and three pseudo-promoter sites are identified with the corresponding binding frequency of 45% and 30%, respectively. Our method is quick, robust and capable of resolving protein-binding locations with high accuracy (\sim 300 bp), making our system a complementary platform to the methods currently practiced. It is advantageous in parallel analysis and less prone to false positive results over other single molecule mapping techniques such as optical tweezers, atomic force microscopy and molecular combing, and could potentially be extended to general mapping of protein–DNA interaction sites.

8. "Low-Copy Number Protein Detection by Electrode Nanogap-Enabled Dielectrophoretic Trapping for Surface-enhanced Raman Spectroscopy and Electronic Measurements", L. Lesser-Rojas, P. Ebbinghaus, G. Vasan, M.L. Chu, A. Erbe*, C.F. Chou* (2014). Nano Letters 2014, 14(5), 2242–2250. http://dx.doi.org/10.1021/nl4046685. (IF=13.025):



We report a versatile analysis platform, based on a set of nanogap electrodes, for the manipulation and sensing of biomolecules, as demonstrated here for low-copy number protein detection. An array of Ti nanogap electrode with sub-10 nm gap size function as templates for alternating current dielectrophoresis-based molecular trapping, hot spots for surface-enhanced Raman spectroscopy as well as electronic measurements, and fluorescence imaging. During molecular trapping, recorded Raman spectra, conductance measurements

Fig. 6: Coupling DEP to detection

across the nanogaps, and fluorescence imaging show unambiguously the presence and characteristics of the trapped proteins. Our platform opens up a simple way for multifunctional low-concentration heterogeneous sample analysis without the need for target preconcentration.

9. **"Tandem array of nanoelectronic readers embedded coplanar to a fluidic nanochannel** for correlated single biopolymer analysis", L. Lesser-Rojas, K. K. Sriram, K.T. Liao, S.C. Lai, P.C. Kuo, M.L. Chu, C.F. Chou* (2014). Biomicrofluidics 2014, 8, 016501. http://dx.doi.org/10.1063/1.4861435 (IF=3.771): We have developed a two-step electron-beam lithography process to fabricate a tandem array of three pairs of tip-like gold nanoelectronic detectors with electrode gap size as small as 9 nm, embedded in a coplanar fashion to 60 nm deep, 100 nm wide, and up to 150 μ m long nanochannels coupled to a world-micro-nanofluidic interface for easy sample introduction. Experimental tests with a sealed device using DNA-protein complexes demonstrate the coplanarity of the nanoelectrodes to the nanochannel surface. Further, this device could improve transverse current detection by correlated time-of-flight measurements of translocating samples, and serve as an autocalibrated velocimeter and nanoscale tandem Coulter counters for single molecule analysis of heterogeneous samples.

List of Publications and Significant Collaborations that resulted from your AOARD supported project: In standard format showing authors, title, journal, issue, pages, and date, for each category list the following:

a) papers published in peer-reviewed journals:

(i) K.T. Liao, M. Tsegaye, V. Chaurey, C.F. Chou, N. Swami. "Nano-constriction device for rapid protein pre-concentration in physiological media by electrokinetic force balance", *Electrophoresis* (2012), 33, 1958-1966. DOI: 10.1002/elps.201100707

(ii) V. Chaurey, A. Rohani, Y.-H. Su, W. Varhue, K.T. Liao, C. F. Chou, N. S. Swami.
"Scaling down constriction-based dielectrophoresis for trapping nanoscale biomolecules in high conductivity media", *Electrophoresis* (2013), 34, 1097-1104. 10.1002/elps.201200456
(iii) B. J. Sanghavi, S. Sitaula, M. Griep, S. Karna, M. Ali, N. S. Swami. "Real-time electrochemical monitoring of ATP at graphene-modified electrodes", *Anal. Chem.* (2013), 85, 8158–8165. DOI: 10.1021/ac4011205

(iv) A. Rohani, W. Varhue, Y.-H. Su, N. S. Swami. "Quantifying spatio-temporal dynamics of biomarker pre-concentration and depletion in microfluidic systems by intensity threshold analysis", Biomicrofluidics (2014) 8, 052009. http://dx.doi.org/10.1063/1.4897283 (v) B. Sanghavi, W. Varhue, J. Chavez, C.F. Chou, N. S. Swami. "Electrokinetic preconcentration and detection of neuropeptides at patterned graphene-modified electrodes in a nano-channel device", Anal. Chem. (2014), 86, pp 4120-4125. DOI:10.1021/ac500155g (vi) K. K. Sriram, C.L. Chang, U. R. Kumar, C.F. Chou* (2014). "DNA Combing on Low-Pressure Oxygen Plasma Modified Polysilsesquioxane Substrates For Single-Molecule Studies", *Biomicrofluidics* 2014, 8, 052102; DOI: 10.1063/1.4892515. (IF: 3.771) (vii) K.K. Sriram, J.W. Yeh, Y.L. Lin, Y.R. Chang, C.F. Chou* (2014). "Direct Optical Mapping of Transcription Factor Binding Sites on Field-stretched λ -DNA in Nanofluidic Devices", Nucleic Acids Research 2014, 42, e85. DOI: 10.1093/nar/gku254 (IF: 8.808) (viii) L. Lesser-Rojas, K. K. Sriram, K.T. Liao, S.C. Lai, P.C. Kuo, M.L. Chu, C.F. Chou* (2014). "Tandem array of nanoelectronic readers embedded coplanar to a fluidic nanochannel for correlated single biopolymer analysis", *Biomicrofluidics* 2014, 8, 016501. http://dx.doi.org/10.1063/1.4861435 (IF=3.771)

(ix) L. Lesser-Rojas, P. Ebbinghaus, G. Vasan, M.L. Chu, A. Erbe*, <u>C.F. Chou*</u> (2014). "Low-Copy Number Protein Detection by Electrode Nanogap-Enabled Dielectrophoretic Trapping for Surface-enhanced Raman Spectroscopy and Electronic Measurements", *Nano Letters* 2014, 14(5), 2242–2250. <u>http://dx.doi.org/10.1021/nl4046685</u>. (IF=13.025)

b) papers published in non-peer-reviewed journals or in conference proceedings: None c) conference presentations (Selected)

(i) B.J. Sanghavi, W. Varhue, A. Rohani, J. Chavez, C.-F. Chou, N.S. Swami; "Conformation-selective biomarker preconcentration by dielectrophoresis", MicroTAS 2014, San Antonio, USA

(ii) K.-T. Laio, N. S. Swami, C.-F. Chou. "Rapid monitoring of low abundance prostate specific antigen by protein nanoconstriction molecular dam." MicroTAS, Germany (2013). http://www.rsc.org/images/loc/2013/PDFs/Papers/471_0719.pdf

(iii) B.J. Sanghavi, W. Varhue, A. Rohani, J. Chavez, C.-F. Chou, N.S. Swami; "Electrokinetic preconcentration and detection of neuropeptides", AIP Advances in Micro/Nanofluidics, Academia Sinica, Taipei, Taiwan.

(iv) N. S. Swami, "Frequency selective trapping of biomarkers using conformation-specific aptamers", Microscale Bioseparations: MSB 2013, Charlottesville, VA, March 2013.

(v) N.S. Swami, "Frequency-selective polarization of the electrical double-layer of nano-colloids", American Electrophoresis Society Annual Meeting, AIChE, San Francisco, Nov 2013

(vi) N.S. Swami, "Coupling dielectrophoresis to ion concentration polarization for enhanced protein enrichment" Advances in Micro-Nanofluidics, AMN 2013, University of Notre Dame,

May 2013

(vii) N.S. Swami, "Nano-slit device for dielectrophoretic enrichment of proteins", ITP Separations, Baltimore, 2012.

DD882: No inventions disclosures (form submitted).

Important Note: Abstracts of refereed publications have been submitted above as part of "Results & Discussion".

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⁶ V. Chaurey, A. Rohani, Y.-H. Su, W. Varhue, K.T. Liao⁺, C. F. Chou, N. S. Swami. "Scaling down constriction-based dielectrophoresis for trapping nanoscale biomolecules in high conductivity media", *Electrophoresis* (2013), 34, 1097-1104.

⁷ B. J. Sanghavi, S. Sitaula, M. Griep, S. Karna, M. Ali, N. S. Swami. "Real-time electrochemical monitoring of ATP at graphene-modified electrodes", *Anal. Chem.* (2013), 85, 8158–8165.

⁸ A. Rohani, W. Varhue, Y.-H. Su, N. S. Swami. "Quantifying spatio-temporal dynamics of biomarker pre-concentration and depletion in microfluidic systems by intensity threshold analysis", *Biomicrofluidics* (2014) 8, 052009.

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¹⁰ K. K. Sriram, C.L. Chang, U. R. Kumar, <u>C.F. Chou</u>* (2014). "DNA Combing on Low-Pressure Oxygen Plasma Modified Polysilsesquioxane Substrates For Single-Molecule Studies", *Biomicrofluidics* 2014, 8, 052102; <u>DOI: 10.1063/1.4892515</u>. (IF: 3.771)

¹¹ K.K. Sriram, J.W. Yeh, Y.L. Lin, Y.R. Chang, <u>C.F. Chou</u>* (2014). "Direct Optical Mapping of Transcription Factor Binding Sites on Field-stretched λ-DNA in Nanofluidic Devices", *Nucleic Acids Research* 2014, 42, e85. DOI: 10.1093/nar/gku254 (IF: 8.808)

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¹³ L. Lesser-Rojas, P. Ebbinghaus, G. Vasan, M.L. Chu, A. Erbe*, <u>C.F. Chou*</u> (2014). "Low-Copy Number Protein Detection by Electrode Nanogap-Enabled Dielectrophoretic Trapping for Surface-enhanced Raman Spectroscopy and Electronic Measurements", *Nano Letters* 2014, 14(5), 2242–2250. <u>http://dx.doi.org/10.1021/nl4046685</u>. (IF=13.025)