Native Desorption Electrospray Ionization Liberates Soluble and Membrane Protein Complexes from Surfaces

Stephen Ambrose, Nicholas G. Housden, Kallol Gupta, Jieyuan Fan, Paul White, Hsing-Yung Yen, Julien Marcoux, Colin Kleanthous, Jonathan T. S. Hopper,* and Carol V. Robinson*

Abstract: Mass spectrometry (MS) applications for intact protein complexes typically require electrospray (ES) ionization and have not been achieved via direct desorption from surfaces. Desorption ES ionization (DESI) MS has however transformed the study of tissue surfaces through release and characterisation of small molecules. Motivated by the desire to screen for ligand binding to intact protein complexes we report the development of a native DESI platform. By establishing conditions that preserve non-covalent interactions we exploit the surface to capture a rapid turnover enzyme-substrate complex and to optimise detergents for membrane protein study. We demonstrate binding of lipids and drugs to membrane proteins deposited on surfaces and selectivity from a mix of related agonists for specific binding to a GPCR. Overall therefore we introduce this native DESI platform with the potential for high-throughput ligand screening of some of the most challenging drug targets including GPCRs.

A range of applications, including 2D imaging of small molecules and metabolites released from tissue cross-sections, has become possible with the introduction of powerful DESI approaches when coupled with MS.[1,2] The primary goal of DESI applications has been to focus on the small molecules released, for example in the real-time detection of tumour tissue during surgical procedures.[3,4] While DESI has also been adapted to study large biomolecules, via the mixing of ES droplets and solution in a process known as liquid DESI,[5,6] it has not yet been applied to proteins deposited on surfaces and desorbed in solutions that retain their native state interactions. Despite considerable progress in applications of non-denaturing or native MS (nMS) of soluble[7-9] and membrane embedded proteins[10] the possibility of effectively “lifting” intact complexes from surfaces is desirable since many high throughput technologies then become accessible.[11] Moreover the lipid distribution in natural membranes is essentially planar and asymmetric with varying spatial and temporal arrangements in the vicinity of embedded protein complexes.[12] The lipid distribution and the desire for a surface technology that is also able to analyse membrane proteins motivated us to develop a modified DESI approach capable of releasing folded protein molecules from planar surfaces and to construct an interface that we could couple to a high-resolution Orbitrap MS optimised for high mass transmission[13] of membrane proteins.[14] We demonstrate the potential of this methodology in three ways 1) by capturing transient protein substrate products 2) by screening for optimal solution/purification conditions using picoMoles of membrane and soluble proteins and 3) by carrying out ligand binding experiments on planar surfaces.

We modified the design of the original DESI set-up[1] and with our custom-built ion source, ES device and sample stage coupled our interface to an Orbitrap Q-Exactive (Figure 1a and Supporting Information Figure S1). We found that signal intensity was significantly improved if the length of the sample transfer tube, used in conventional DESI set-ups,[15] was minimised and the stage was located directly under the inlet of our ion source. We optimised signal intensity using hen egg-white lysozyme and found that the spectra recorded under these native DESI conditions are similar to those from typical nanoflow ES capillaries, implying that the folded state of the protein is maintained (Figure 1b). If the native fold is maintained then deposited lysozyme should be able to carry out enzymatic functions. Accordingly with N-acetyl-glucosamine (NAG) substrate added to the desorption spray and directed at lysozyme deposited on the stage we observed additional peaks assigned to binding of intact NAG-5 to lysozyme (Figure 1c). The rapid turnover of this substrate precludes its observation in solution-based ES[16] but since substrate binding takes place during rapid desorption, analogous to reactive DESI experiments reported for small molecules,[17] the transient bound state can be captured using this native DESI approach.

To investigate application to larger protein assemblies we chose complexes with a range of oligomeric states: bovine serum albumin, tetrameric alcohol dehydrogenase (ADH) and the GroEL14mer. We were able to record native DESI mass spectra for all three with masses of 66, 148 and 800 kDa,
respectively and with established subunit stoichiometries (Figure 1d,e,f). These results effectively transform DESI from a small molecule approach to a method capable of detecting intact protein assemblies from surfaces. In this regard an obvious next target is membrane proteins since their natural environment is in lipid bilayers.

For this study we selected the outer-membrane protein F (OmpF), a trimer of transmembrane beta-barrels. We deposited 0.4 nmol of OmpF in ammonium acetate (200 mM) containing octyl glucoside (OG) micelles and directed the desorption plume at the deposited protein. Initially a relatively low intensity signal was observed (Figure 2a) followed by rapid deterioration and loss of signal—indicating a dilution effect at the surface leading to disruption of the micelle.[18] Adding OG to the desorption solution (at twice the critical micelle concentration (cmc) (1% w/v OG) we observed recovery of the OmpF trimer signal (Figure 2b) which remained stable for 30 mins (Figure S2).[18] Substituting a different detergent (Lauryldimethylamine N-oxide (LDAO) 0.05% w/v) into the desorption plume induces a shift to higher charge states (Figure 2c), observed previously with LDAO[19] implying that detergent exchange has occurred on the stage. This highlights an important capability of the native DESI approach for rapid detergent screening.

Screening for optimal detergents, as part of membrane protein purification protocols, is time consuming and uses valuable protein resources.[20] We investigated further the possibility of detergent screening on the DESI stage using the sugar transporter semiSWEET from *Vibrio splendidus*[21] since it is extremely sensitive to its detergent environment. Previous MS experiments established that this transporter exists in an entirely monomeric form in DDM and exhibits a monomer–dimer equilibrium in the detergent (C8E4).[22] Depositing semiSWEET on the DESI stage in DDM (Figure 2d) we then added C8E4 (0.5% w/v) directly to the desorption buffer. After 1.8 min of desorption with the C8E4-containing buffer the total ion chromatogram for the (7+) ion (indicative of the dimer) was observed consistent with detergent exchange on the DESI stage from the initial DDM conditions to the C8E4 micelles (Figure 2e). Detergent screening on the DESI stage, with minimal membrane protein consumption, highlights a powerful feature of this approach.

Turing to ligand screening an important criterion to establish is the extent to which protein complexes dissociate during native DESI as opposed to conventional nanoflow ES. Selecting an outer membrane protein receptor FpvA from *Pseudomonas aeruginosa*, which translocates ferric-pyoverdine (Pvd) across outer mem-

![Figure 1. Schematic of the native DESI setup showing deposition of protein on the stage, followed by desorption and analysis in the mass spectrometer with representative spectra for a series of soluble protein their substrates and complexes. a) First, protein is deposited on the native DESI stage (red) from aqueous buffer and second, the ES plume is charged with a voltage of 2.5–3.5 V and directed at the stage. Transfer is effected by positioning the stage close to the orifice of the mass spectrometer. b) Apo lysozyme is deposited on the stage (25 µl, 10 µm) in aqueous ammonium acetate (200 mM, pH 6.8) and the same buffer is used to desorb the protein. c) NAG-S is added to the ES plume directed at the lysozyme deposit, additional peaks reveal binding of the substrate NAG-S prior to its cleavage. Native DESI mass spectra of d) monomeric bovine serum albumin, e) tetrameric alcohol dehydrogenase and f) the GroEL 14-mer.](image-url)
branes,[23] we formed the complex in solution and compared the percentage of complex FpV:A:FvD using the different ionization methods. We found 59% and 61% in the native DESI and nanoES approaches, respectively (Figure 3a and b). We conclude that our native DESI approach, which involves both deposition and desorption within protective micelles, reproduces the nano-ES result in which complexes are electrosprayed directly from solution.

Exploring further the quantitative aspects of this native DESI platform we selected OBS1 a 17-residue peptide known to bind within the pores of OmpF with binding constants determined previously by both ITC and nanoES MS.[24,25] We deposited OmpF on the stage in OG detergent and incubated OmpF with increasing concentrations of OBS1 (0–75 μM) and deposited these protein-peptide complexes onto the native DESI stage. The relative intensities of bound to unbound protein were extracted and plotted as a function of OBS1 concentration (Figure S3). The $K_d$ determined (0.7 ± 0.34 μM) for this membrane protein complex is in agreement with values reported (1.0 ± 0.1 μM).[26] That the solution state...
equilibria is maintained is surprising given that DESI involves ejection of the membrane protein complex, deposited on the surface, by a desorption spray. However detailed kinetics of OmpF–OBS1 peptide complexes are not known so it is unclear how this reflects the kinetics of DESI.

Beyond peptide binding established in solution we wanted to perform binding experiments on membrane proteins deposited on the stage. Delipidated apo OmpF (20 μm) was deposited and OBS1 added to the desorption buffer. Up to three peptides bound per trimer were detected with the predominant peak corresponding to two (Figure S3a). This experiment confirms that the peptide can access binding sites within the short time frame during desorption, even in the presence of detergent. Similarly we added phosphatidylglycerol lipid (POPG) to the desorption buffer, at a concentration of 5 μm, and observed up to three lipids bound to OmpF (Figure S3b). OmpF is also reported to bind to a range of antibiotics within the extracellular and periplasmic pore vestibule. Addition of kanamycin (50 μm) to the desorption buffer reveals binding of one molecule of the antibiotic per OmpF trimer (Figure S3c). Since we anticipate that the peptide binds within the pore, while the lipids bind to the outer surface and kanamycin to the top of the pore, we have highlighted our capability to bind directly to the membrane protein via three different mechanisms. In each of these three scenarios we assume that the small molecule has not reached solution phase equilibrium, since desorption is rapid, but rather has penetrated to some extent the protective micelle that surrounds the membrane protein while deposited on the surface.

Figure 3. The extent of complex formation is comparable to that observed in nano-ES and enables determination of $K_d$ values and competitive binding experiments. Comparison of mass spectra for the complex FvpA:FvD recorded by a) DESI and b) nESI. c) Titration of the peptide OBS1 to OmpF at 0, 25 μm and 50 μm for determination of $K_d$. d) Deposition of the GPCR P2Y1 and desorption in a mixed micelle and e) competitive binding of the specific ligand from a cocktail of six ligands.
Building on this ability to screen for binding to a membrane protein target deposited on a surface we reasoned that it would be possible to add multiple ligands simultaneously. To explore this possibility we selected a Class A G-protein coupled receptor (GPCRs) depositing on the stage 0.4 nmole of P2Y1, responsible for platelet aggregation and a key target for anti-thrombotic therapy.[27] After recording a native DESI spectrum in its apo form we added a cocktail of antagonists/agonists designed to target related GPCRs (Figure 3d,e and Table S1). The native DESI spectrum reveals a discrete mass increase (560.03 Da) in exact agreement with the mass of MR52500 (1′R,2′S,4′S,5′S)-4-(2-Iodo-6-methyl amino-purine-9-yl)-1-{[(phospho)methyl]-}2-(phospho)hexane bound to P2Y1. No other adducts were observed and control experiments, where the specific inhibitor was excluded from the drug cocktail, revealed no ligand binding to P2Y1 (Figure S6). These results reveal that this native DESI platform is capable of detecting selective binding of a specific antagonist to a GPCR from a multicomponent mixture.

In summary we have developed and applied a native DESI platform and shown that it is capable of preserving the native structure of both soluble and membrane proteins and their complexes. Comparing our approach with ambient ionization methods described previously we note the addition of chemicals in the spray solution in reactive DESI applications for small molecule analyses.[17,18] Protein ligand binding experiments have also been achieved with reactions taking place by mixing in solution in a liquid sample DESI approach[19] rather than by interaction following protein deposition on the planar target as shown here. Moreover kinetic approaches have been developed using mixing experiments to monitor the small molecules released during enzymatic cleavage by means of liquid DESI and have increased the range of buffers that can be used.[20,21] Our native DESI approach is largely restricted to volatile buffers and detergents that have been optimised for native MS. A further limitation is imposed by the fact that ligands are observed directly bound to proteins desorbed from a planar surface, high-resolution MS is therefore critical.

The most exciting aspect of this native DESI approach however, is the potential of our method to study intact membrane proteins and their complexes. The ability to place membrane protein targets on planar surfaces in different lipidic environments, without tethering the proteins, and to carry out selective binding from a cocktail of drugs offers possibilities for high throughput screening. Many downstream applications become accessible including the ability to carry out multiple experiments on the same target; for example detergent optimisation, the screening of multiple lipids and ligands that bind to a drug target or the trapping of fast turnover products in enzyme catalysed reactions. Analogous to the powerful native MS methods, now widely accepted as a key component in structural biology, native DESI enables further possibilities for development of spatial, temporal and even directional analyses within artificial bilayers or membrane mimetics.

Acknowledgements

C.V.R. acknowledges funding from an ERC Advanced Grant ENABLE (641317), an MRC Programme Grant (G1000819) and a Wellcome Trust Investigator Award (104633/Z/14/Z). K.G. is a Junior Research Fellow at St Catherine’s College, Oxford and a recipient of the Royal commission for the Exhibition of 1851 fellowship. We thank Dr Renata Kamin ska (Department of Biochemistry, Oxford) for provision of purified OmpF. C.K. acknowledges support from the Wellcome Trust (201505/Z/16/Z) and P.W. acknowledges the Wellcome Trust for studentship funding.

Conflict of interest

The authors declare no conflict of interest.

Keywords: desorption · electrospray ionisation · G-protein coupled receptors · mass spectrometry · membrane protein complexes

How to cite: Angew. Chem. Int. Ed. 2017, 56, 14463–14468

Angew. Chem. Angew. Chem. Int. Ed. 2017, 56, 14463–14468 © 2017 The Authors. Published by Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim www.angewandte.de 14467

[1] Z. Takáts, J. M. Wiseman, R. G. Cooks, J. Mass Spectrom. 2005, 40, 1261–1275.
[2] R. G. Cooks, Z. Ouyang, Z. Takáts, J. M. Wiseman, Science 2006, 311, 1566–1570.
[3] L. S. Eberlin, C. R. Ferreira, A. L. Dill, D. R. Ila, R. G. Cooks, Biophys. Acta Mol. Cell Biol. Lipids 2011, 1811, 946–960.
[4] N. Abbassi-Ghadi, et al., Cancer Res. 2016, 76, 5647–5656.
[5] C. N. Ferguson, S. A. Benchaar, Z. Miao, J. A. Loo, H. Chen, Anal. Chem. 2011, 83, 6468–6473.
[6] P. Liu, J. Zhang, C. N. Ferguson, H. Chen, J. A. Loo, Anal. Chem. 2013, 85, 11966–11972.
[7] A. G. Ngounou Wétie, et al., Proteomics 2013, 13, 538–557.
[8] M. A. Oshina, M. Sharon, Q. Rev. Biophys. 2016, DOI: https://doi.org/10.1017/s0035858516000160.
[9] J. V. van de Waterbeemd, K. L. Fort, D. Boll, M. Reinhardt-Szyba, Nat. Methods 2017, 14, 283–286.
[10] J. A. Martfeld, V. Rajagopalan, D. V. Greathouse, R. E. Koeppe 2nd, Biochim. Biophys. Acta Biomembr. 2015, 1848, 1849–1859.
[11] J. Gault, et al., Nat. Methods 2012, 9, 1084–1086.
[12] J. H. Gault, et al., Nat. Methods 2016, DOI: https://doi.org/10.1038/nmeth.3771.
[13] Z. Takáts, J. M. Wiseman, B. Gologan, R. G. Cooks, Anal. Chem. 2004, 76, 4050–4058.
[14] J. H. T. Hopper, N. J. Oldham, J. Am. Soc. Mass Spectrom. 2009, 20, 1851–1858.
[15] M. A. Oshina, M. Sharon, Q. Rev. Biophys. 2016, DOI: https://doi.org/10.1017/s0035858516000160.
[16] J. V. van de Waterbeemd, K. L. Fort, D. Boll, M. Reinhardt-Szyba, Nat. Methods 2017, 14, 283–286.
[17] J. Gault, et al., Nat. Methods 2016, DOI: https://doi.org/10.1038/nmeth.3771.
[18] Z. Takáts, J. M. Wiseman, B. Gologan, R. G. Cooks, Anal. Chem. 2004, 76, 4050–4058.
[19] J. T. Hopper, N. J. Oldham, J. Am. Soc. Mass Spectrom. 2009, 20, 1851–1858.
[20] I. Cotte-Rodriguez, Z. Takáts, N. Talaty, H. Chen, R. G. Cooks, Anal. Chem. 2005, 77, 6755–6764.
[21] E. Reading, et al., Angew. Chem. Int. Ed. 2015, 54, 4577–4581; Angew. Chem. 2015, 127, 4660–4664.
[22] M. Orwick-Rydmark, T. Arnold, D. Linke, Curr. Protoc. Protein Sci. 2016, 84, 4.8.1–4.8.35.
[23] Y. Xu, et al., Nature 2014, 515, 448–452.
[24] K. Gupta, et al., Nature 2017, 541, 421–424.
[23] D. Cobessi, et al., J. Mol. Biol. 2005, 347, 121–134.
[24] N. G. Housden, et al., Proc. Natl. Acad. Sci. USA 2010, 107, 21412–21417.
[25] N. G. Housden, et al., Science 2013, 340, 1570–1574.
[26] B. K. Ziervogel, B. Roux, Structure 2013, 21, 76–87.
[27] D. Zhang, et al., Nature 2015, 520, 317–321.
[28] G. Huang, H. Chen, X. Zhang, R. G. Cooks, Z. Ouyang, Anal. Chem. 2007, 79, 8327–8332.
[29] S. Cheng, J. Wang, Y. Cai, J. A. Loo, H. Chen, Int. J. Mass Spectrom. 2015, 392, 73–79.
[30] S. Cheng, Q. Wu, H. Xiao, H. Chen, Anal. Chem. 2017, 89, 2338–2344.
Manuscript received: May 11, 2017
Revised manuscript received: August 7, 2017
Accepted manuscript online: September 8, 2017
Version of record online: September 18, 2017