AP-MALDI-Q-IMS-TOF MS as a liquid sample mass profiling platform for food authenticity tests.

Oliver J. Hale^a | Michael Morris^b | Rainer Cramer^a

Overview

- Development of workflows suitable for the analysis of crude liquid samples using a custom atmospheric pressure matrix-assisted laser desorption/ionisation (AP-MALDI) source.¹
- High-throughput, real-time classification of samples with ion mobility-time-of-flight mass spectrometry (IMS-TOF MS) profiles is demonstrated.
- Further investigation and identification of individual compounds possible through offline data mining.

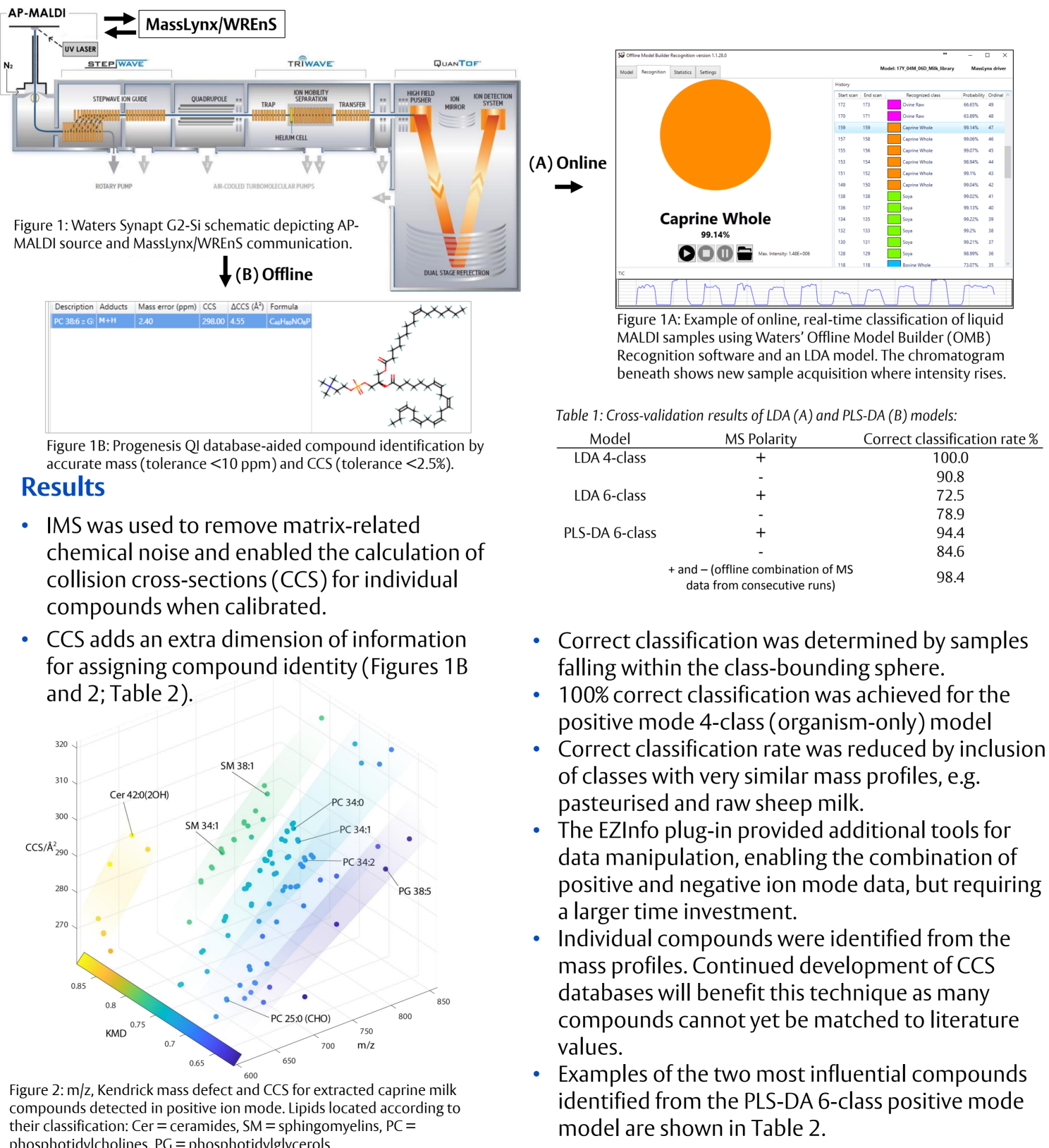
Introduction

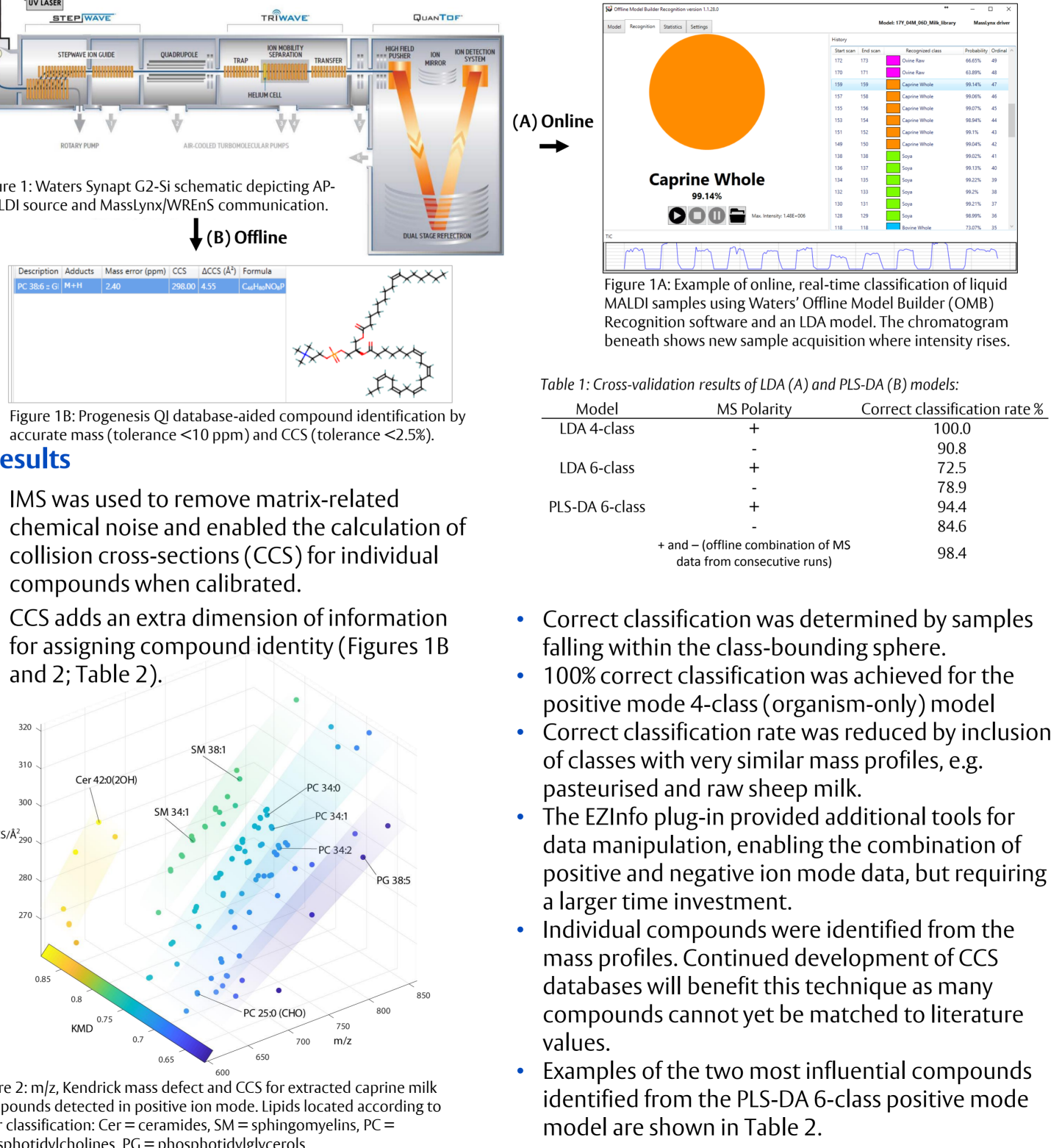
- MALDI is an established, sensitive ionisation technique, but requires crystalline samples for compatibility with its traditionally high-vacuum conditions. The persistent and stable ion beam generated from a glycerol-based liquid support matrix (LSM) at atmospheric pressure is more suitable for rapid, automated acquisition, as well as MS tuning and calibration. Sample longevity is high, enabling multiple analyses of the same droplet, with on-target storage at 4 °C previously demonstrated for many weeks.²
- Presented here are example workflows that take advantage of the AP-MALDI-Q-IMS-TOF MS platform (Figure 1) applied to the analysis of lipids extracted from milk.

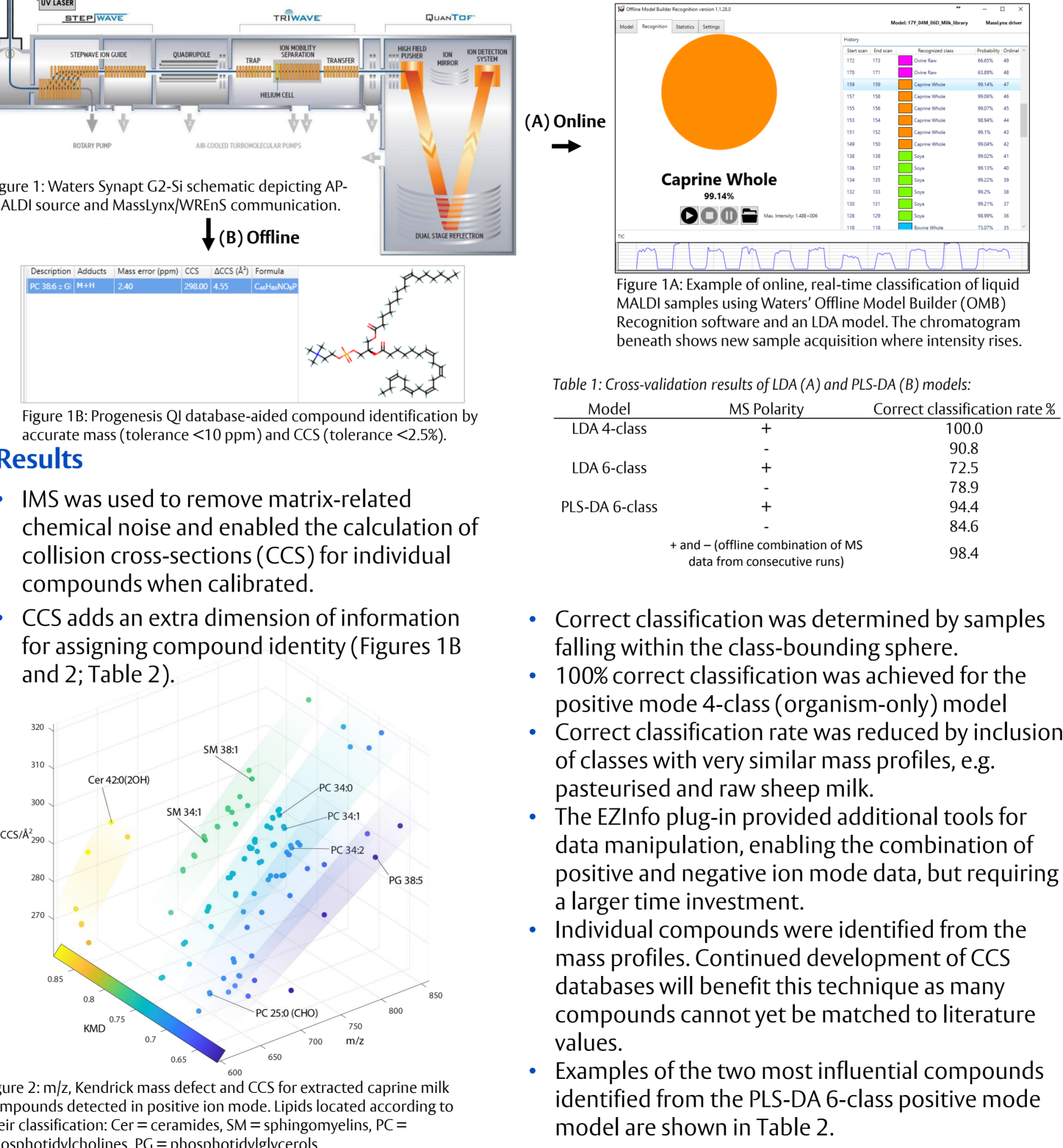
Methods

- Lipids from samples of 6 milk types (bovine whole, bovine skimmed, caprine whole, ovine raw, ovine pasteurised and soya) were extracted with hexane/isopropanol (HIP; 3:2, v/v).
- Each liquid MALDI sample droplet consisted of 0.5 μ L of HIP lipid extract (9:1, v/v) and 0.5 μ L of LSM (2, 5-dihydroxybenzoic acid 25 mg/mL dissolved in $H_2O/MeCN/glycerol$, 3:7:6, v/v/v).
- 10-second analysis (30 Hz laser pulse repetition rate, 28 µJ/pulse) per sample generated IMS-TOF MS profiles in both positive and negative ion modes. The same droplets were used for both ion modes. TOF MS was run in 'sensitivity' mode.

Progenesis QI and the EZInfo plugin (Figure 1B).







phosphotidylcholines, PG = phosphotidylglycerols.

a. Department of Chemistry, University of Reading, Whiteknights, Reading RG6 6AD, UK. b. Waters Corporation, Stamford Avenue, Wilmslow SK9 4AX, UK

• Online, real-time classification of milk types was achieved with Waters' Offline Model Builder (OMB) Recognition tool. TOF MS spectra from 45 samples per class, with an m/z range of 600-900, were used to build linear discriminant analysis (LDA) models (Figure 1A). In-depth analysis was performed with

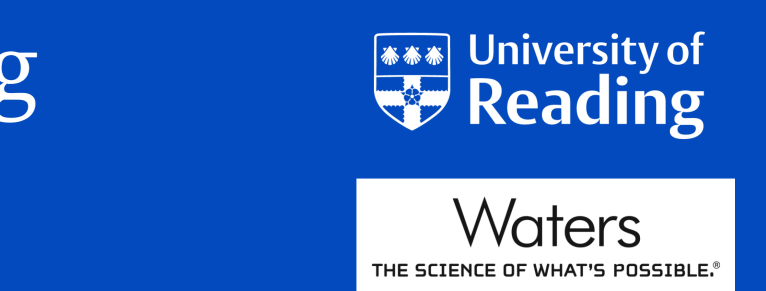


Table 2: The two most influential compounds from PLS-DA 6-class positive mode. Suggested ID m/z (accuracy/ppm) CCS/Å² (% difference from literature³)

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SM 34:1	703.5719(-4.1)	294 (+1%)
PC 34:1	760.5830 (-2.8)	298 (+1%)

The content of binary milk mixtures was determined from orthogonal partial least squares discriminant analysis (OPLS-DA) in EZInfo by relating intergroup separation and %mass (Figure 3).

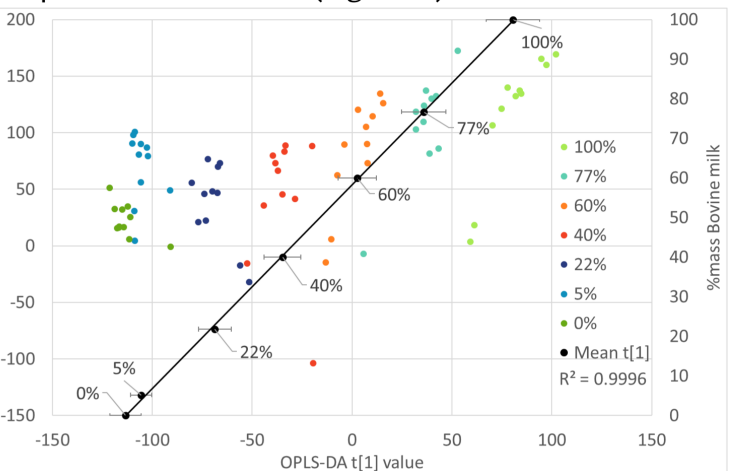


Figure 3: OPLS-DA plot of mixtures of bovine and caprine milk (left y-axis, coloured points) and the linear relationship between mean t[1] and %mass bovine milk (right y-axis, black points). Error bars show the standard deviation of each group.

Conclusions

- Liquid AP-MALDI-Q-IMS-TOF MS provides a good platform for high-throughput, real-time sample classification of milk samples.
- Offline processing enables deeper levels of analysis with existing software tools.
- Combination of positive and negative ion mode profiles has the potential to increase classification accuracy but is not possible online with TOF MS.

References

- 1. Ryumin, P., Brown, J., Morris, M., Cramer, R.: Investigation and optimization of parameters affecting the multiply charged ion yield in AP-MALDI MS. Methods. 104, 11-20 (2016).
- 2. Ryumin, P., Brown, J., Morris, M. & Cramer, R. Protein identification using a nanoUHPLC-AP-MALDI MS/MS workflow with CID of multiply charged proteolytic peptides. International Journal of Mass Spectrometry (2016)
- 3. Paglia, G. et al. Ion mobility-derived collision cross section as an additional measure for lipid fingerprinting and identification. Analytical Chemistry 87, 1137-1144 (2015).

Acknowledgements

- This research is supported by Waters Corporation and EPSRC DTG grant no.1498422
- Martin Wells at Non-Linear Dynamics for access to and continued support with Progenesis QI.
- The British Mass Spectrometry Society and the Association of British Spectroscopists are thanked for their support through travel grants.
- Ovine milk was kindly provided by The Sheep Milk Company (Lancashire, UK).

Contact information

- Department of Chemistry, University of Reading, Whiteknights, RG6 6AD Reading, UK
- Email: o.j.hale@pgr.reading.ac.uk; r.k.cramer@reading.ac.uk
- http://www.reading.ac.uk/chemistry/about/staff/RKCGroupMembers.aspx