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# DART-ultraFAIMS-MS testing

**Results & comments**

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## UltraFAIMS with DART



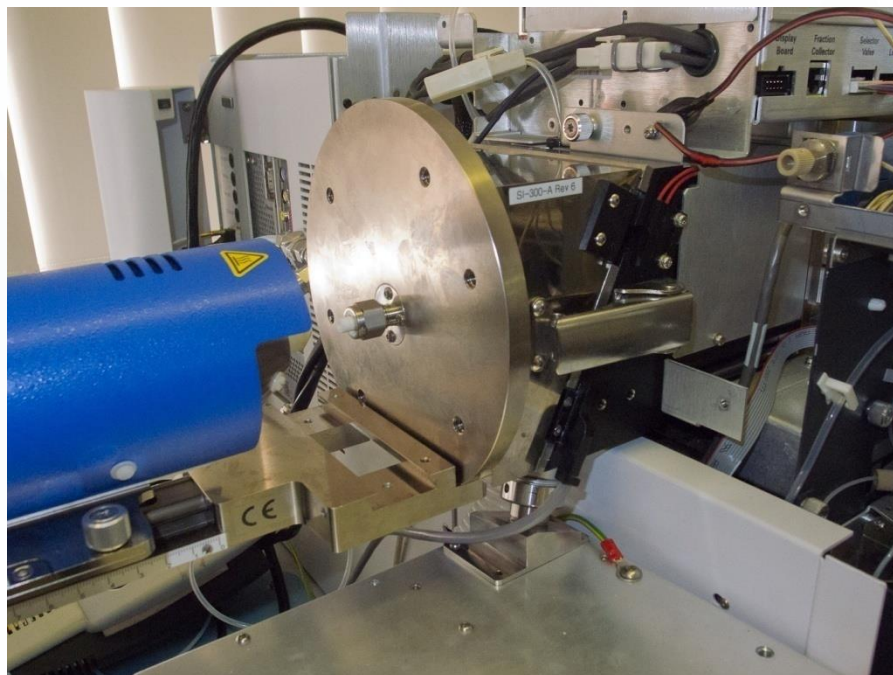
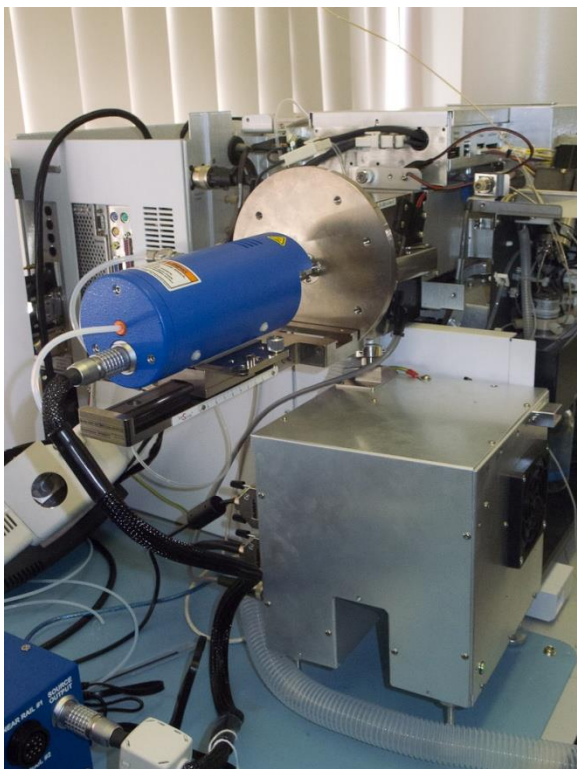
- Many mass spec users have highlighted the potential for using FAIMS pre-separation with direct ionisation sources as a gas-phase analog to the LC pre-separation used with electrospray sources
- The ultraFAIMS pre-separation should reduce general chemical noise, and may also allow selective transmission of isomers and isobars
- We recently coupled our ultraFAIMS-A1 system with the IonSense DART source for proof-of-principle testing, on an Agilent 6230 TOF
- This confirmed that the two systems can be coupled without the need for modifications to either instrument
- Photos of the set-up and results from initial test samples are shown on the following slides.



## UltraFAIMS with DART



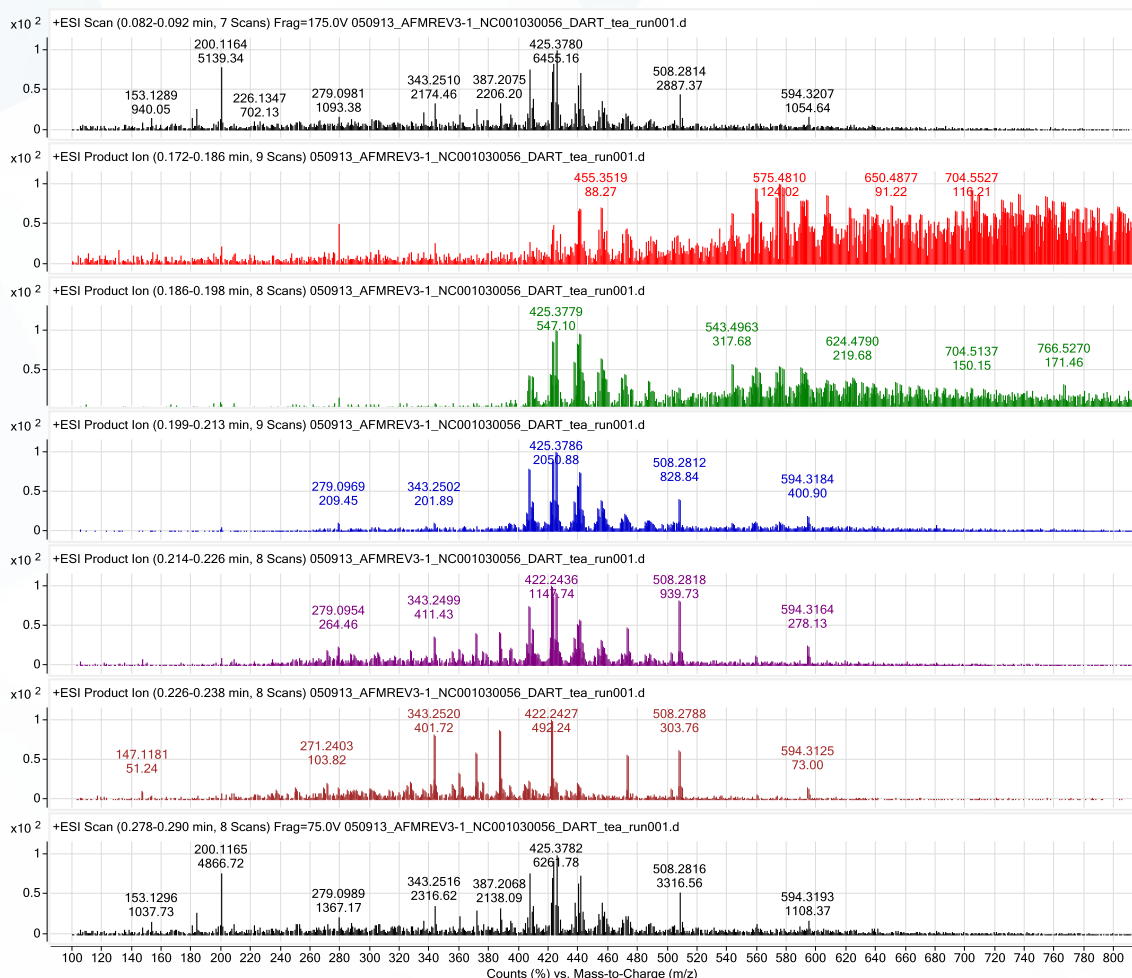
- UltraFAIMS interfaces are designed to avoid the need for modifications to the ionisation source – this means that any source that fits the standard mounting (on Agilent or Thermo mass spectrometers) can be coupled with the UltraFAIMS system – setting up here took a matter of minutes



# Example results 1: Green tea in methanol



The black plots show spectrum before and after FAIMS sweep, the coloured plots show (average) spectrum with FAIMS active at 260Td DF, at a sequence of different CF settings. At each setpoint, different subsets of ions are transmitted. The duration of the sweep was 10 secs.



Immediately before start of sweep

CF 1.0-1.4Td

CF 1.4-1.8Td

CF 1.8-2.2Td

CF 2.2-2.6Td

CF 2.6-3.0Td

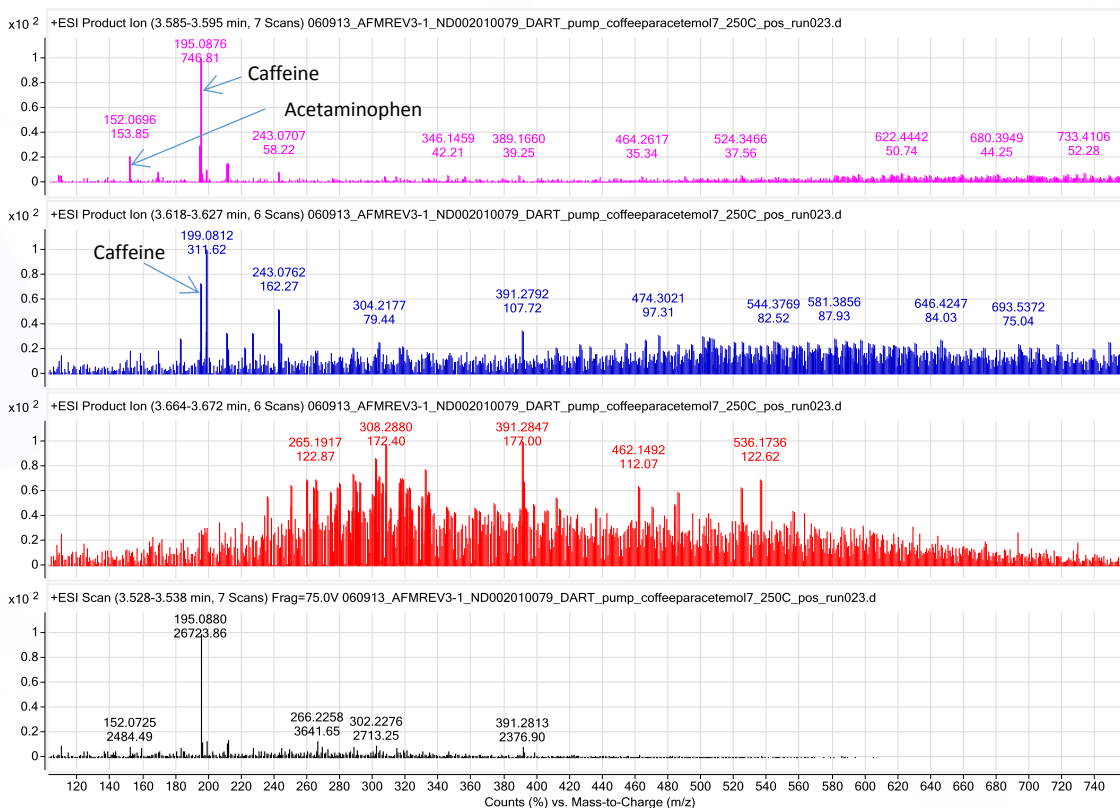
Immediately following end of FAIMS sweep



# Example results 2: Coffee/acetaminophen mix



The black plot shows the spectrum just before the FAIMS sweep, the coloured plots show (average) spectrum with FAIMS active at 260Td DF, at a sequence of different CF settings. At each setpoint, different subsets of ions are transmitted. The duration of the sweep was 10 secs.



CF 1.3-1.4Td

CF 1.6-1.7Td

CF 2.0-2.1Td

Immediately before FAIMS sweep

## Summary



- Coupling the two systems together is straightforward, where both DART and ultraFAIMS interfaces are available for a given mass spectrometer
- ultraFAIMS scan speed is fast enough to provide separation on DART timescales
- The approach shows potential to enable detection of more low abundance analytes through improvements in signal to noise, though more work is needed to quantify the improvement
- Further testing would be needed to develop methods to separate specific problematic isomeric interferences – this is better done initially with standards
- Owlstone is keen to support users interested in exploring DART-ultraFAIMS-MS applications further – please contact us to discuss



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**THANK YOU**  
FOR YOUR ATTENTION!

FEEL FREE TO CONTACT US:

Alasdair Edge

[Alasdair.edge@owlstone.co.uk](mailto:Alasdair.edge@owlstone.co.uk)

[www.ultrafaims.com](http://www.ultrafaims.com)