

# Investigating UPLC Ion Mobility Mass Spectrometry: A New Approach to Authentication and Routine Screening of Ginsenoside Isomers in Functional Food Products

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## **APPLICATION BENEFITS**

- Leverages the unparalleled selectivity of ion mobility in a unique non-targeted screening workflow for the determination of the presence of ginsenoside markers, including pairs of ginsenoside isomers.
- An additional identification point can be generated using collision cross section (CCS)values enabling characteristic assignment for ginsenoside isomers.
- UPLC® ion mobility CCS screening can be used as a viable approach to perform authentication profiling.
- Processing of complex ion mobility data is made routine with easy-to-use UNIFI® and MassLynx® Software.

# WATERS SOLUTIONS

ACQUITY UPLC® I-Class System

ACQUITY UPLC HSS T3 Column

Waters® Ion Mobility
Mass Spectrometry Systems

**UNIFI Scientific Information System** 

## **KEY WORDS**

Ginsenoside isomers, collision cross section, CCS, ion mobility, spectral clean up

## INTRODUCTION

The most abundant forms of ginseng: Panax (Korean ginseng), *P. japanoicus*, and *P. quinquefolium* (American ginseng) grow in North America. The Korean and American ginseng species are most frequently used for medicinal purposes. For thousands of years, the roots of ginseng plants have been used for therapeutic purposes. However the species are believed to have different therapeutic properties. Ligor et. al. discussed CNS stimulant activity, hypogylcemic properties, and the sedative effects of American ginseng. For each species it is believed that the ginsenoside and polysaccharide content are responsible for the biological activity of products produced from the roots and leaves of ginseng species. Figure 1 shows the structures of the ginsenosides screened in this assay, which are a part of a diverse group of steroidal saponins with four ring structures similar to steroids.

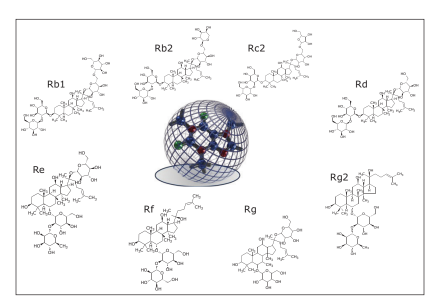


Figure 1. Illustration of rotating three-dimensional conformation of an ion and average collision cross section (shadow), with structures of ginsenosides profiled using UPLC ion mobility mass spectrometry and collision cross section (CCS) screening.

#### **EXPERIMENTAL**

## LC conditions

LC system: ACQUITY UPLC I-Class

Column: ACQUITY UPLC HSS T3,

100 mm x 2.1 mm, 1.8 μm

Column temp.: 40 °C

Flow rate: 0.6 mL/min

Mobile phase A:  $H_2O$  (0.1% formic acid)

Mobile phase B: Acetonitrile (0.1% formic acid)

Gradient:

Time (min) Flow rate %A %В 0.00 0.6 95.0 5.0 3.00 0.6 70.0 30.0 4.00 0.6 70.0 30.0 12.00 0.6 55.0 45.0 18.00 0.6 5.0 95.0 20.00 0.6 95.0 5.0

Injection volume: 1 μL

MS conditions

MS system: SYNAPT® G2-Si

Ionization mode: ESI- at 2.7 kV

Sample cone voltage: 20 V
Desolvation temp.: 600 °C

Lockmass and LockCCS: Leucine enkephalin, [M-H] = 554.2620

Acquisition range: 50 to 1200 *m/z*Acquisition rate: 10 spectra/sec

Collision energy ramp: 35 to 75 eV

Resolution = 20,000 FWHM (Res mode)

# Default IMS parameters

IMS T-Wave™

velocity ramp: Start: 1000 m/s

End: 300 m/s

IMS T-Wave pulse

height: 40V IMS gas flow: 90 mL

# Sample description

Korean ginseng tea (extract in 20 mL of  $\rm H_2O$ ), gingko biloba, red panax (undiluted), red panax (undiluted), extracts and ginsenoside standards (100 pg/ $\mu$ L).

The profiles of ginsenosides in American and Korean ginseng are believed to be different. Ginsenosides are comprised of two main groups: the panaxadiol, or Rb1 group that includes Rb1, Rb2, Rc, Rd, Rg3, Rh2, and Rh3; and the panaxatriol (Rg1 group) that includes Rg1, Re, Rf, Rg2, and Rh1. American ginseng is understood to be richer in the Rb1 group, while Korean ginseng is believed to be richer in the Rg1 group. In addition to the natural differences in the phytochemical profile of the species, the phytochemical profile can also be impacted by the time of harvest, storage conditions, and production processes.

Globally the popularity of nutraceutical and functional food products continues to increase. Functional food/natural product remedies are found in foods, roots, and herbs. Legislative focus has prompted the development of more methods to analyze active compounds in such products. For example in Europe, Directive 2004/24/EC came into full effect on 30 April 2011. Hundreds of traditional herbal remedies were banned, as the EU law aims to protect consumers from possible damaging side-effects of over-the-counter herbal products. More recent regulations allow only long-established and quality-controlled medicines to be sold. Products that have been assessed by the Medicine and Healthcare Products Regulatory Agency (MHRA) may also be sold. Manufacturers have to prove that their products have been made to strict standards and contain a consistent and clearly marked dose.

The ACQUITY UPLC I-Class System and ion mobility mass spectrometry (IM-MS) is a combination of accurate separations with high resolution mass spectrometry (HRMS) and high efficiency ion mobility based measurements that offer some unique advantages for profiling complex mixtures. IM-MS is a rapid orthogonal gas phase separation phase technique that allows another dimension of separation to be obtained within an LC timeframe. Compounds can be differentiated based on size, shape, and charge.<sup>2</sup> The screening assay presented here explores the use of UPLC separations with ion mobility mass spectrometry and UNIFI data processing software as an analytical approach for the characterization of the distribution and content of monodi- and tetra-glycosides in the raw material or processed products. In this case, the aim is to illustrate how the quality and potency of ginseng products can be determined.

## RESULTS AND DISCUSSION

A collision cross section (CCS) value is a robust and precise physicochemical property of an ion. It is an important distinguishing characteristic that is related to its chemical structure and three-dimensional conformation. An illustration is presented in Figure 1, where the shadow of a rotating three dimensional ion is shown. This represents the average collision cross section. Non targeted UPLC-IM-MS has been used to generate travelling wave collision cross sections using a nitrogen buffer gas (TWCCSN<sub>2</sub>), accurate mass precursor/fragment ions, and retention times to profile ginsenoside standards Rb1, (Rb2, Rc), (Rd, Re), (Rf, Rg1), and Rg2. Using TWCCSN<sub>2</sub> measurements can increase non targeted screening specificity. CCS measurements generated have been entered into a scientific library within UNIFI. This allows the expected and determined TWCCSN2 values to be used to screen and confirm the presence of ginsenoside isomer markers. Three extracts, gingko biloba+red panax, red panaz, and Korean ginseng tea were analyzed. These were screened against the created ginsenoside TWCCSN<sub>2</sub> library in UNIFI to determine the presence/ unequivocal identification of ginsenoside isomers.

Here we present TWCCSN<sub>2</sub> values (derived from ion mobility drift times) as a new identification parameter that can distinguish ginsenoside isomers, and also be used to profile unknowns. From Figure 2, the UPLC-IM-MS electrospray negative mode conventional base peak ion chromatogram obtained for the analysis of undiluted Korean ginseng extract is shown, and the complexity of the sample profiled is revealed. However in Figure 3, the UPLC-IM-MS electrospray negative mode plot of drift time (ion mobility resolution) versus retention time for the Korean ginseng tea extract is presented. Using the unique functionality of UNIFI Software, Figure 3 visually illustrates how ion mobility separation orthogonal to chromatographic separation, can increase peak capacity.

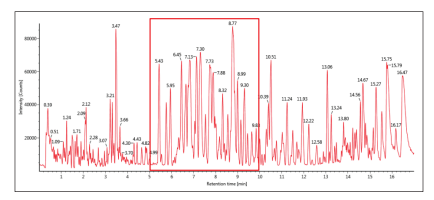


Figure 2. UPLC-IM-MS electrospray negative mode conventional base peak ion chromatogram obtained for analysis of undiluted Korean ginseng tea extract. The complexity of the sample profiled is illustrated.

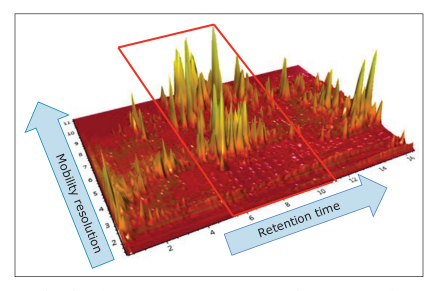


Figure 3. UPLC-IM-MS electrospray negative mode plot of drift time (ion mobility resolution) versus retention time for Korean ginseng tea extract. The true complexity of the sample profiled is illustrated, where separation orthogonal to UPLC separation is obtained when using ion mobility.

# [APPLICATION NOTE]

The retention time region between 6 and 10 minutes shows that there are a large number of compounds that are now resolved compared to the same region on the conventional base peak ion extracted mass chromatogram of Figure 2. The true complexity of the sample profiled is illustrated, when ion mobility resolution and UPLC chromatographic resolution are combined. The ion mobility Data Viewer within UNIFI enables investigative interaction with the acquired ion mobility data. UNIFI includes many easy-to-use functions, such as Zoom to Component and Bookmark, that allow the same investigative interrogation of data across many acquisitions. It is possible to select any one of these components and generate the drift plot, mass spectrum, and extracted mass chromatogram.

For marker ginsenoside isomer pairs Rb2 and Rc (Figure 4), the extracted mass chromatogram (m/z 1123) shows that the ginsenoside marker isomer Rc has a measured <sup>TW</sup>CCSN<sub>2</sub> =350.58.77Å<sup>2</sup> and Rb2 has a measured <sup>TW</sup>CCSN<sub>2</sub> =361.77Å<sup>2</sup>. CCS errors <0.5% were obtained and compared to the expected CCS values that were previously generated from the ginsenoside standards and uploaded into the UNIFI scientific library. The extracted mass chromatogram shows the presence of a series of minor isomeric components between 3.0 and 4.0 mins, 8.0 and 9.0 mins, as well as an unknown isomer at retention time 7.88 min, from which a <sup>TW</sup>CCSN<sub>2</sub> value of 358.80.77Å<sup>2</sup> has been determined. Regardless of the chromatographic methodology used, it it now possible to distinguish the ginsenoside isomer and the unknown isomer using <sup>TW</sup>CCSN<sub>2</sub> measurements. This could not be acheived with accurate mass measurement alone, regardless of the mass resolution capability. It is also worth noting that these acquisitions are non targeted; hence the <sup>TW</sup>CCSN<sub>2</sub> values are generated for all knowns and unknowns. This could not be achieved with other available ion mobility technologies, such as Field Asymmetric Ion Mobility Spectrometry (FAIMS), where a knowledge of the targeted analytes is required, and it is limited to just a few target compounds, compared to unlimited number of possible <sup>TW</sup>CCSN<sub>2</sub> measurements.

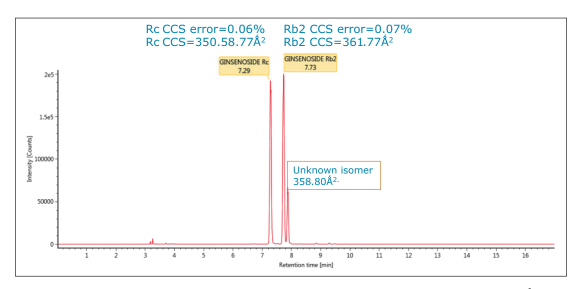


Figure 4. m/z 1123 exact mass extracted mass chromatogram, showing ginsenoside marker isomers Rc ( $^{TW}CCSN_2=350.58.77\text{Å}^2$ )/Rb2 ( $^{TW}CCSN_2=361.77\text{Å}^2$ ), as well as an unknown isomer at retention time 7.88 mins. CCS errors <0.5% are presented. The extracted mass chromatogram shows the presence of a series of minor isomeric components between 3.0 and 4.0 mins, and also 8.0 and 9.0 mins.

In addition to CCS measurements serving as an additional identification point, the combined peak capacity of UPLC and ion mobility brings additional benefits. This can be seen in Figure 5 where the retention time (7.73 mins) and drift time (9.93 ms), aligned precursor, and fragmentation spectra for Rc ginsenoside marker isomer is shown. The spectra presented result from only the Rc ginsenoside, because chromatographically coeluting compounds are resolved using ion mobility. The ion mobility spectral cleanup makes it clear that the unknown isomer at 7.88 mins also has the same characteristic fragment ions as ginsenoside Rc, but it can be differentiated using ion mobility. The characteristic information acquired and processed for the unknown isomer is shown in the UNIFI Component Summary in Figure 6. Illustrated are the candidate Component Summary, mobility trace, and precursor/mobility product ion spectra for the unknown isomer with  $^{\text{TW}}\text{CCSN}_2$ =358.80Å $^2$  at retention time 7.88 mins. The  $^{\text{TW}}\text{CCSN}_2$  for the unknown isomer can be entered into the UNIFI scientific library, along with the  $^{\text{TW}}\text{CCSN}_2$  values generated for all of the compounds detected during the chromatographic run.

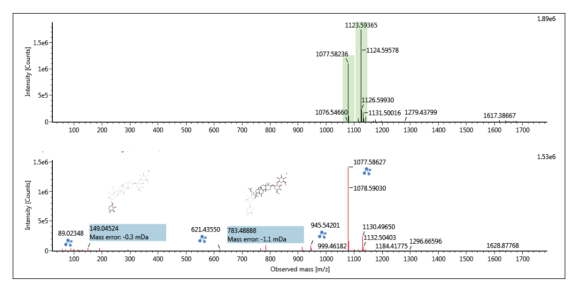


Figure 5. Retention time (7.73 mins)/drift time aligned (9.93 ms) precursor and ion mobility product ion spectra for Rc ginsenoside marker isomers, where their characteristic fragmentation pattern and proposed fragmentation pathways are shown.

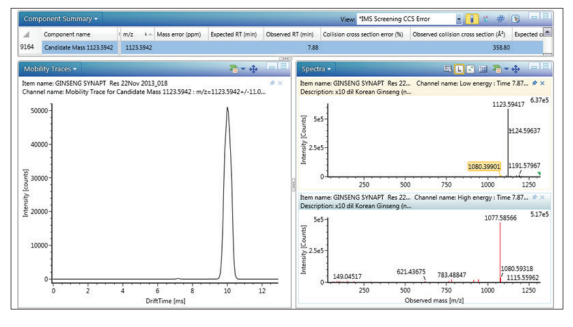


Figure 6. UNIFI Component Summary illustrates the candidate component summary, mobility trace, and precursor/mobility product ion spectra for an unknown isomer TWCCSN<sub>2</sub>=358.80Å<sup>2</sup> at retention time 7.88 mins, as shown in Figure 5.

# [APPLICATION NOTE]

CCS measurements can increase confidence in identification. The results are summarized in Figure 7, where the results obtained for profiling of Korean ginseng over two consecutive weeks is presented. For the marker ginsenoside isomer pairs (Rb2, Rc),  $^{\text{TW}}$ CCSN<sub>2</sub> measurements of 361.77 Å<sup>2</sup>/350.58Å<sup>2</sup> have been determined. For (Rd, Re), 328.89 Å<sup>2</sup>/333.11 Å<sup>2</sup> were determined. For (Rf, Rg1), 304.7 Å<sup>2</sup>/295.83 Å<sup>2</sup> were obtained in week 1. In week two, comparative results were obtained. When comparing the expected against the measured  $^{\text{TW}}$ CCSN<sub>2</sub> results determined (for the eight ginsenosides profiled in the extracts), the measurement errors were typically <0.5%. It is therefore possible to distinguish the marker isomer pairs of ginsenosides in the extracts of the specified products analysed with confidence using  $^{\text{TW}}$ CCSN<sub>2</sub> measurements.

| 4                                    | Component name   | m/z  | Mass error (ppm)                               | Expected RT (min)               | Observed RT (min) 4-  | Collision cross section error (%)                               | Observed collision cross section (Å <sup>2</sup> )   | Expected collision cross section (Å <sup>2</sup> )   | Adducts                                     |
|--------------------------------------|--|--|--|---------------------------------|---|---|--|--|---|
| 1                                    | GINSENOSIDE Re   | 991.5493   | 0.98   | 3.49                            | 3.47  | -0.07   | 328.89   | 329.11   | •HCOO, -H                                   |
| 2                                    | GINSENOSIDE Rg1  | 845.4924   | 2.36   | 3.51                            | 3.49  | -0.38   | 295.83   | 296.95   | -HC00, -H                                   |
| 3                                    | GINSENOSIDE RF   | 845.4902   | -0.21  | 5.46                            | 5.43  | -0.47   | 304.70   | 306.13   | •HCOO, -H                                   |
| 4                                    | GINSENOSIDE Rg2  | 829.4979   | 2.91   | 6.48                            | 6.44  | -0.01   | 300.51   | 300.54   | +HCOO, -H                                   |
| 5                                    | GINSENOSIDE Rb1  | 1153.6052  | 3.48   | 6.84                            | 6.80  | 0.25  | 358.07   | 357.18   | -HCOO, -H                                   |
| 6                                    | GINSENOSIDE Rc   | 1123.5938  | 2.85   | 7.32                            | 7.29  | 0.06  | 350.58   | 350.36   | •HCOO, -H                                   |
| 7                                    | GINSENOSIDE Rb2  | 1123.5937  | 2.74   | 7.77                            | 7.73  | 0.07  | 361.77   | 361.50   | -HCOO, -H                                   |
| 8                                    | GINSENOSIDE Rd   | 991.5495   | 0.02   |                                 | 0.00  |   |  |  |   |
| ٧                                    |  |  | an gins  | seng te                         | 8.76<br>Ta  | 0.00  | 333.11   | 333.12   | +HC00, -H                                   |
| ۷                                    | Veek 2:  | Kore   | an gins  | seng te                         | a   |   |  |  |   |
| V                                    | Veek 2:  | Kore   | an gins  | seng te                         | Observed RT (min)   | Collision cross section error (%)                               | Observed collision cross section $(\hat{A}^k)$   | Expected collision cross section $\langle A^2 \rangle$   | Adducts                                     |
| V                                    | Veek 2:  | Kore   | ean gins                                       | seng te                         | Cobserved RT (min)  | Collision cross section error (%) -0.28                         | Observed collision cross section $(\hat{A}^k)$   | Expected collision cross section $(\mathbb{A}^k)$ 329.11   | Adducts<br>+MCOO                            |
| V                                    | Veek 2:<br>Component name<br>GINSENOSIDE RE  | Kore<br>m/z<br>991.5489  | ean gins                                       | seng te                         | Observed RT (min)  9 3.47 51 3.49                                     | Collision cross section error (%) -0.28 -0.55                   | Observed collision cross section (Å <sup>2</sup> )<br>328.18<br>295.33                     | Expected collision cross section (Å <sup>2</sup> ) 329.11 296.95   | Adducts                                     |
| 1<br>2<br>3                          | Veek 2:<br>Component name<br>GINSENOSIDE Re<br>GINSENOSIDE Rg1   | M/z<br>991.5489<br>845.4898                                      | Mass error (ppm) 0.64 -0.71                    | Seng te                         | Observed RT (min)  99 3.47  51 3.49  66 5.42                          | Collision cross section error (%) -0.28 -0.55 -0.30             | Observed collision cross section (Å <sup>2</sup> ) 328.18 295.33 305.24                    | Expected collision cross section (Å <sup>2</sup> ) 329.11 296.95 306.15  | Adducts +HCOO +HCOO                         |
| 1<br>1<br>2<br>3<br>4<br>5           | Veek 2:<br>Component name<br>GINSENOSIDE Re<br>GINSENOSIDE Rg1<br>GINSENOSIDE Rf                                   | M/z<br>991.5489<br>845.4898<br>845.4935                          | Mass error (ppm) 0.64 -0.71 3.70               | Seng te                         | Observed RT (min)  49 3.47  51 3.49  66 5.42  88 6.43                 | Collision cross section error (%) -0.28 -0.55 -0.30 0.04        | Observed collision cross section (Å <sup>2</sup> )<br>328.18<br>295.33<br>305.24<br>300.65 | Expected collision cross section (A <sup>2</sup> )<br>379.11<br>296.95<br>306.15<br>300.54                     | Adducts<br>+HCOO<br>+HCOO<br>+HCOO          |
| 1<br>2<br>3<br>4<br>5                | Veek 2: Component name GINSENOSIDE RE GINSENOSIDE Rg1 GINSENOSIDE Rf GINSENOSIDE Rg2                               | M/z<br>991.5489<br>845.4898<br>845.4935<br>829.4972              | Mass error (ppm)  0.64 -0.71 3.70 2.02         | Seng teo                        | Observed RT (min)  9 3.47  51 3.49  66 5.42  88 6.43  94 6.78         | Colision cross section error (%) -0.29 -0.55 -0.30 -0.30        | Observed collision cross section (Å*) 388.18 295.33 300.65 358.53                          | Expected collision cross section (A <sup>2</sup> )<br>329.11<br>296.95<br>306.15<br>300.54<br>357.18           | Adducts<br>+HC00<br>+HC00<br>+HC00<br>+HC00 |
| V<br>1<br>2<br>3<br>4<br>5<br>6<br>7 | Veek 2: Component name GINSENOSIDE Re GINSENOSIDE Rf GINSENOSIDE Rf GINSENOSIDE Rf GINSENOSIDE Rg2 GINSENOSIDE Rb1 | M/z<br>991.5489<br>845.4898<br>845.4935<br>829.4972<br>1153.6031 | ean gins Mass error (ppm) 0.64 -0.71 3.70 2.02 | Expected RT (min) 1 3.3 5.6 6.6 | Cobserved RT (min) 189 3.47 51 3.49 166 5.42 18 6.43 54 6.78 192 7.27 | Collision cross section error (%) -0.28 -0.35 -0.30 -0.34 -0.38 | Observed collision cross section (Å*) 308.18 295.33 305.24 300.65 3985.35 3950.70          | Expected collision cross section (A <sup>2</sup> )<br>329-11<br>296-95<br>306-15<br>307-18<br>357-18<br>350-36 | Adducts ++COO ++COO ++COO ++COO ++COO       |

Figure 7. Consecutive ginsenoside  $^{TW}CCSN_2$  screening results obtained for the analysis of Korean ginseng tea, illustrating the  $^{TW}CCSN_2$  reproducibility obtained.

This approach offers a unique selectivity for profiling complex mixtures. The results obtained clearly show the benefits of using CCS measurements and the combined peak capacity of UPLC with ion mobility. Coeluting analytes and isomers have been resolved as well as unequivocally identified in the three extracts profiled. In addition it is possible to acquire the cleaned up mobility specific product ion spectra, which are ion mobility resolved from coeluting components. In this case the reason for such a screening approach is to generate a new analysis to enable the characterization/distribution and content of mono- di- and tetra-glycosides in the raw material or processed functional food/nutraceutical products. The new assay approach would add confidence when assessing quality, potency, and consistency of a final product, incorporating ingredients such as gingko biloba, Korean ginseng and red panax. This new analytical approach changes the scope of authentication profiling.

# [APPLICATION NOTE]

While gaining confidence in the identifications made from the use of accurate mass measurement and CCS measurements, there is also a clear potential to reduce the amount of high purity standards required, where confirmation relies on retention time and accurate mass measurement. The costs of the standards for this assay, shown in Table 1, totalled £2483.00, a significant expense. The use of a CCS screening approach has the potential to provide significant cost savings across many application areas. For further studies to profile ginsenosides, Rb1, (Rb2, Rc), (Rd, Re), (Rf, Rg1), and Rg2, the consumption and costs of high purity standards has been significantly reduced within the laboratory where the study was performed.

| rn | M    | ^ I | IIС | 10 | M C  |
|----|------|-----|-----|----|------|
|    | IN I |     | 117 |    | IN 2 |

- The ACQUITY UPLC I-Class System with IM-MS can be used to routinely screen and authenticate the true ginsenoside phytochemical makeup in Korean ginseng, gingko biloba, and red panax extracts.
- Ginsenoside Rb1, (Rb2, Rc), (Rd, Re), (Rf, Rg1), and Rg2 have been profiled in gingko biloba, korean ginseng and red panax.
- CCS measurements of <0.5% have been obtained routinely, as well as excellent reproducibility.
- Isomeric ginsenosides have been differentiated using IM-MS and CCS measurements.
- TWCCSN<sub>2</sub> screening can be used to profile sample makeup and uniquely differentiate ginsenoside isomer composition, increasing confidence that ginsenosides are not missed due to chromatographic coelution.
- TWCCSN<sub>2</sub> authentication profiling has the potential to reveal the true phytochemical profile across many species.
- The data processing functionality within UNIFI provides an easy-to-use platform that allows analysts to routinely and confidently profile complex samples.

| Ginsenoside     | Quanity | Price    |
|-----------------|---------|----------|
| Ginsenoside Re  | 10 mg   | £260.50  |
| Ginsenoside Rg1 | 10 mg   | £259.50  |
| Ginsenoside Rf  | 10 mg   | £372.00  |
| Ginsenoside Rg2 | 10 mg   | £143.50  |
| Ginsenoside Rb1 | 10 mg   | £259.50  |
| Ginsenoside Rc  | 10 mg   | £800.00  |
| Ginsenoside Rb2 | 10 mg   | £244.50  |
| Ginsenoside Rd  | 10 mg   | £143.50  |
|                 |         | £2483.00 |

Table 1. Cost of ginsenoside standards purchased for ion mobility study.

#### References

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