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High-Resolution Ion Cyclotron Mobility Spectrometry

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A novel ion mobility spectrometry instrument incorporating a cyclotron geometry drift tube is presented. The drift tube consists of eight regions, four curved drift tubes and four ion funnels. Packets of ions are propagated around the drift tube by changing the drift field at a frequency that is resonant with the ion’s drift time through each region. The approach trims each packet of ions as it leaves and enters each new region. An electrostatic gate allows ions to be kept in the drift tube for numerous cycles, increasing the ability to resolve specified ions. We demonstrate the approach by isolating the [M + 2H]^{2+} or [M + 3H]^{3+} charge state of substance P as well as individual trisaccharide isomers from a mixture of melezitose and raffinose. Resolving powers in excess of 300 are obtainable with this approach.

We describe the development of the first cyclotron geometry drift tube. In this instrument a packet of ions is released into the cyclotron and transmitted through multiple cycles by oscillating the applied drift fields at an appropriate frequency. As the ions cycle, species with different mobilities are pulled apart in time. After a desired number of cycles, only species with mobilities that are in resonance with the drift field application frequency are trapped, and subsequently these ions are released into a mass analysis region and detected. We demonstrate storage of ions for more than 10 cycles (a total drift distance of 20.75 m) and an experimental resolving power, $R = t/\Delta t > 300$, where $t$ is the drift time of an ion measured at the center of its peak in a drift time distribution and $\Delta t$ is the full width at half-maximum (fwhm) of the peak.

For the past decade, there has been significant interest in developing new ion mobility spectrometry (IMS) technologies that couple with mass spectrometry (MS) analyses. The primary motivation for coupling these techniques is that IMS separates ions based on differences in their shapes and MS provides primary structural information. It is anticipated that combinations of IMS-MS with other condensed-phase separation methods will lead to a more comprehensive understanding of complex molecular systems, such as molecular mixtures that arise in analysis of biological systems.

Although some remarkable separations of isomers and different conformers have been reported, IMS is generally thought of as a low-resolution technique. Several groups have constructed instruments with resolving powers in excess of 100. To date the highest reported resolving power for an IMS separation was obtained for the [M – 4H]^{4+} ion of $\text{CH}_3\text{SO}_2\text{NH}_2\text{O}_3\text{CH}_3\text{SO}_2\text{NH}_2\text{CH}_3$ reported as $t/\Delta t = 240$. Jarrold and co-workers developed a high-resolution instrument for studying cluster ions and reported $R = 172$ for carbon clusters and sodium dichloride ($\text{NaCl}$) ions. Smith and their collaborators have developed high-resolution instruments for separating biomolecular ions and have reported resolving powers ranging from 100–216. In general, the resolving power of traditional IMS methods becomes limited.

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because it increases as the square root of the key experimental values $E$ and $L$ (the drift field and length, respectively). Our current efforts to develop cyclotron geometry instruments are aimed at ultimately increasing the resolving power by allowing ions to experience very long drift regions through multiple cycles. As a final note, the term cyclotron is used here to draw parallels with ion cyclotron resonance MS$^{20}$ with respect to the ability to achieve a high-resolution measurement; no magnetic fields are employed to manipulate ion trajectories.

**EXPERIMENTAL SECTION**

**General.** Detailed descriptions of IMS theory and instrumentation are provided elsewhere; only a short description of the instrument used for these studies is provided. A diagram of the instrument and pulse sequence is shown in Figure 1. Ions generated by electrospray ionization$^{27}$ accumulate in a Smith-geometry ion funnel (F1).$^{28}$ Packets of ions are periodically introduced into the cyclotron drift tube via an electrostatic gate (G1) following the pulse sequence shown in Figure 1. This initial pulse is also used to trigger the data acquisition system that is used to record drift times.

As the packet of ions leaves the source, it drifts around a curve into the main body of the multipass drift tube. The cyclotron region of the drift tube consists of eight separate components: four curved drift regions (D1–D4), each 30.01 cm in length, and four ion funnels (F2–F5), each 15.21 cm in length. Each of these has voltages applied to the first and last lens of the segment by a home-built drift field wavedriver.$^{1}$ A series of resistors ($5 \, \text{M}\Omega$ for each curve, 1.5 M$\Omega$ for each funnel) is used as voltage dividers and establishes the drift field along the length of each section.

Two sections (D1 and D4) have Y-shaped geometries. These regions are used to maintain ions inside the cyclotron or to move ions into (D1) or out of (D4) the cyclotron. Both ion trajectory simulations with the program SIMION$^{29}$ and experimental data (not shown) have demonstrated that the geometry of D1 focuses ions exiting either F1 or F5 into F2, with very little deviation in path length when compared to D2 or D3 under the same field conditions. With respect to D4, the propagation of ions in this region is controlled by electrostatic gates. The lenses in the center of the Y are split (as shown in the inset in Figure 1). By application of the appropriate voltages to the lenses labeled G2a and G2b, both of which are isolated from the resistor chain, it is possible to maintain ions inside the multipass cyclotron region or let ions out of the cyclotron drift tube (bottom two pulse sequences). Again, SIMION simulations and experimentation show that this has a negligible effect on drift times through this region. Once out of the cyclotron drift tube, ions are focused in F6 prior to entering the D5 region, where they exit into the vacuum and are ultimately mass analyzed and detected.

A square wave at a fixed frequency (generated by a Wavetek model 195 universal waveform generator, Wavetek Dalton Inc.) controls the drift field. This is shown as the third pulse sequence in Figure 1. The wavedriver is arranged such that the drift field is applied across the first lens of an ion funnel to the last lens of a curved region for one-half of the period (the A regions in Figure 1). Under these conditions, the constant field regions are F2 through D2, F3 through D3, F4 through D4, and F5 through D1.

![Figure 1](image.png)
For the next half of the period (B), the field is defined by the first lens of a curved region to the last lens of an ion funnel; D1 through F2, D2 through F3, D3 through F4, and D4 through both F5 (and F6). A large positive potential across the 1.27 cm gaps (between adjacent continuous drift field regions) keeps ions from entering the next region.

It is instructive to describe how a packet of ions is propagated around this drift tube. The injection pulse (G1) occurs simultaneously with the A half of the period. During this time, ions enter the D1 region but can travel no further because of the large positive potential between the exit of this region and the entrance to F2. Once the drift field wavedriver switches to the B phase, ions can diffuse from D1 into F2 but only those ions that enter F2 before the next A phase are now stable within the instrument.

Any ions at the leading edge of the distribution are defocused by the large positive potential and diffuse into surrounding electrodes; ions at the tailing edge are cut off by the switching potential and diffuse away as well. This process is repeated twice more to guide ions around 3/4 of the cyclotron; for each subsequent cycle around the drift tube, 4 more periods are required. For simplicity, this multiple of the wavedriver period is used as the delay time on the split lens G2a/G2b assembly in order to keep ions within the cyclotron for the number of cycles desired. As an example, the pulse sequence shown in Figure 1 is for two ion injections at one cycle at a frequency of 200 Hz.

The result of these voltage applications is the creation of four transmission and four elimination regions that are present at any given time within the instrument. In the present configuration the transmission regions are significantly smaller than the elimination regions. Within each application of the drift field (a full A–B–A transition or the period of the wavedriver), ions must travel ~45.22 cm (the length of one funnel and one curved region) to remain stable. In one full cycle, 180.88 cm are traversed.

All drift regions and ions funnels are maintained at 7 V cm⁻¹. The length of the linear region (F6 through F7) is 97.58 cm. The current design will allow this length to be reduced to 30.93 cm; the utility of this will be addressed in future publications. Thus, the distance an ion travels for a 13/4 cycle experiment would be 447.09 cm; each additional cycle adds 180.88 cm in the linear region (F6 through F7) are maintained at 500 kHz, 130 Vp-p. Since each ion funnel (F2–F6) are maintained at 500 kHz, 130 Vp-p. Since each ion funnel F2–F5 is operated at two different dc potentials, their rf generators must be floated by the dc potential applied to each funnel. The entire drift tube is filled with 2.80 atm of 3/4 of the cyclotron and monitoring the 300 K He buffer gas, and the pressure is monitored by a capacitance manometer.

Electrospray Conditions. Substance P (98% purity), melezitose hydrate (99% purity), and raffinose pentahydrate (98% purity) were purchased from Sigma Aldrich and used without purification. A 2.0 × 10⁻⁴ M solution of substance P was prepared in 49:49: 2% water/acetonitrile/acetic acid (by volume). Solutions of 5.0 × 10⁻⁴ M concentration (containing each of the oligosaccharides) were prepared in 50:50% water/acetonitrile, 2 mM NaCl. A mixture of the two was prepared by mixing equal volumes of the two solutions, for a final concentration of 2.5 × 10⁻⁴ M for each sugar.

Figure 2. Intensities of the substance P [M + 2H]²⁺ and [M + 3H]³⁺ charge states measured with the TOF mass analyzer as a function of frequency, for injected ions experiencing from 3/4 of a cycle to 33/4 cycles around the cyclotron drift tube. Frequencies of 128.2 Hz (a period of 7.80 ms) to 250.0 Hz (4.00 ms) were examined, in increments of 0.10 ms periods, for a total of 39 mass spectra for each individual number of cycles. The two intense frequencies observed for the 3/4 cycles data (158.7 Hz for the [M + 2H]²⁺ ion and 208.3 Hz for the [M + 3H]³⁺ ion) were those employed in Figure 3.

RESULTS AND DISCUSSION

Illustration of Multiple Passes of Substance P [M + 2H]²⁺ and [M + 3H]³⁺ Ions. Figure 2 illustrates the multipass capability of the cyclotron drift tube for the [M + 2H]²⁺ and [M + 3H]³⁺ ions of substance P. To obtain these results, we begin by determining the optimal drift field application frequency for transmission of each of the ions. This is done by propagating the ions around 3/4 of the cyclotron and monitoring the intensities of the m/z values corresponding to [M + 2H]²⁺ or [M + 3H]³⁺. The intensities at frequencies spanning 128.2–250.0 Hz (39 different frequencies in total) are plotted at the top of Figure 2. The [M + 2H]²⁺ ion is optimally transmitted using a drift field frequency of 158.7 Hz; whereas the [M + 3H]³⁺ ion is favored at 208.3 Hz. Increasing the number of cycles for which the ions are stored within the cyclotron yields interesting results. While the [M + 2H]²⁺ ion remains centered about 158.7 Hz, two maxima appear for the [M + 3H]³⁺ ion, 196.1 and 232.6 Hz. Increasing from 3/4 cycles to 33/4 cycles leads to a sharpening of these peaks in frequency space, as more of the ions with mobilities not corresponding to the maxima are lost.
Myung et al.\textsuperscript{30} we derive $K_0 ([M + 2H]^{2+}) = 3.41 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. Drift time distributions for the $[M + 3H]^{3+}$ ion shows that there are two resolvable conformers, a compact state (having $K_0 = 5.02 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$),\textsuperscript{30} and a relatively more stable elongated state (with $K_0 = 4.44 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$).\textsuperscript{31} In the data shown in Figure 2, the peak at 158.7 Hz for the $[M + 2H]^{2+}$ ion allows us to determine a value of $K_0 = 3.45 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, in good agreement with the value determined from Myung’s data.\textsuperscript{30} The peak at 208.3 Hz for $[M + 3H]^{3+}$ yields a value of $K_0([M + 3H]^{3+}) = 4.52 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. This value is most consistent with the measured mobility for the elongated state for $[M + 3H]^{3+}$. With the use of the $3V/4$ cycles data, the peak at 232.6 Hz gives us a value of $K_0 = 5.06 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, in agreement with the mobility calculated for the compact state. The peak at 196.1 Hz yields a value of $K_0([M + 3H]^{3+}) = 4.26 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$.

With this understanding of which ions are favored as a function of frequency, consider the drift time distributions in Figure 3. In separate experiments, using transmission frequencies of 158.7 and 208.3 Hz, we have examined ions for multiple cycles around the cyclotron. The results appear very promising. At $1V/4$ cycles (4.47 m total length), we determine drift times of 47.04 and 62.88 ms for the $[M + 3H]^{3+}$ and $[M + 2H]^{2+}$ ions, respectively. The measured fwhm’s of typical peaks under these conditions are 0.72 and 0.96 ms, respectively. This yields an IMS resolving power of 65 for each peak.

Upon increasing the delay of the G2a/G2b gate, we allow the ions to experience $5V/4$ cycles. Under these conditions, each ion travels 11.71 m. Although peak centers increase to 124.08 and 163.68 ms, there is little change in the fwhm of these peaks. The fwhm of the $[M + 3H]^{3+}$ peak is 1.20 ms, and the $[M + 2H]^{2+}$ peak is 0.96 ms. Thus, the peaks have moved apart but do not appear to broaden significantly. This provides an important illustration of the separation process that we are utilizing. The widths of peaks are not solely determined by diffusion broadening. If they were dictated only by diffusion, the fwhm of peaks would always increase upon increasing the number of cycles. Instead, ions with mobilities that are not resonant with the drift field frequency wind up in positions that cannot be transmitted around the instrument. The net result is that the ion packet is periodically trimmed. As the ions move through each drift region, populations that have diffused to the leading and trailing edges are eliminated at each new section. We have recently written about this phenomenon in a multidrift region approach called overtone mobility spectrometry.\textsuperscript{1} This trimming also leads to substantial losses in signal at multiple cycles, and it becomes apparent that only frequencies for which there exist a stable mobility will have signal after many passes.

The net result of this trimming process is that it allows us to access relatively high resolving powers under relatively moderate experimental conditions. This effect becomes more pronounced with an increasing number of cycles. Figure 3 also shows data acquired for $10V/4$ cycles. Under these conditions, the ions have traveled 20.75 m. The peak centers increase to 220.32 and 289.68 ms. In this case, the fwhm([M + 2H]^{2+}) = ~0.84–0.96

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure3.png}
\caption{Isolation and separation of the $[M + 2H]^{2+}$ and $[M + 3H]^{3+}$ charge states of substance P at $1V/4$, $5V/4$, and $10V/4$ cycles with the cyclotron drift tube. Two spectra were collected for each number of cycles; one at a frequency of 158.7 Hz for the isolation of the $[M + 2H]^{2+}$ ion and another at a frequency of 208.3 Hz to isolate the $[M + 3H]^{3+}$ ion. The total distance traveled by the ions for each number of cycles is shown, as well as the resolving power for both charge states after $10V/4$ cycles.}
\end{figure}


\begin{align*}
K_0 &= \frac{L^2 f 73.2 P}{V T} 760
\end{align*}

where $L$ is the length of the drift region, $V$ is the drift voltage, $f$ is the drift field frequency, and $T$ and $P$ are the temperature and pressure at which the experiment was conducted, respectively (for normalization purposes). Although we have not published extensively about substance P, the mobilities of the $[M + 2H]^{2+}$ and $[M + 3H]^{3+}$ species are well characterized. From the data of
ms, and fwhm((M + 3H)3+) = 0.96 ms (see inset). From these values, we determine $t/\Delta t = 230$ for (M + 3H)3+ and $\sim 305–345$ for (M + 2H)2+.

The results for other trapping conditions are shown in Figure 4, which shows a plot of the drift times for the (M + 2H)2+ and (M + 3H)3+ ions of substance P for data obtained for $1\frac{3}{4}$ to $10\frac{3}{4}$ cycles. The line through the data has a slope that corresponds to the time required for a single pass around the cyclotron drift tube. The intercept represents the time to traverse the linear region as well as $3/4$ of the cyclotron region. The bottom plot shows the IMS resolving power as a function of the number of cycles for both the (M + 2H)2+ and (M + 3H)3+ charge states of substance P. Lines projecting IMS resolving powers for both ions at higher numbers of cycles are shown.

Figure 4. Total drift time as a function of the number of cycles for the (M + 2H)2+ and (M + 3H)3+ charge states of substance P through the cyclotron drift tube are shown in the top plot. Ions have experienced from $1\frac{3}{4}$ to $10\frac{3}{4}$ cycles. Times are shown as points, with the lines of best fit drawn through each set of times. The slope of each line represents the time the ions spend in one cycle of the drift tube, while the intercept represents the time to traverse the linear region as well as $3/4$ of the cyclotron region. The bottom plot shows the IMS resolving power as a function of the number of cycles for both the (M + 2H)2+ and (M + 3H)3+ charge states of substance P. Lines projecting IMS resolving powers for both ions at higher numbers of cycles are shown.

It is also interesting to examine the resolving power after additional cycles. Figure 4 also shows a plot of the resolving power for peaks measured using 10 different cyclotron conditions such that ions experience $1\frac{3}{4}$ to $10\frac{3}{4}$ cycles. The resolving power increase with the number of cycles is roughly linear. Note, $t/\Delta t$ does not increase as $L^{1/2}$, as is expected for peak shapes dictated by diffusion. As mentioned above, this is because only ions with a mobility that is resonant with the drift field frequency remain stable in this instrument. Thus, as ions are transmitted from region to region, only species that are stable within the next region will remain. Leading and trailing edges of the ion distribution are eliminated. While this effectively trims the ion packet and limits the fwhm of the peak that is observed, the values of $t/\Delta t$ that are obtained rigorously correspond to values of resolving power; only, in this case, diffusion has a minimal effect on peak shape until ions are released into the final linear drift region. It is interesting to consider projected resolving powers for experiments that utilize even more passes. From Figure 3 we anticipate that the resolving powers associated with the (M + 2H)2+ ion peak of substance P will exceed 500 after 20 cycles. We also note that the peak widths could be minimized by using a shorter linear drift region (as well as a higher drift field in this region) at the exit of the cyclotron, as this limits broadening due to diffusion.

Effects of Curves, Fields, And Frequencies on Separations and Measured Mobilities. In the present experimental design, only an ion that traverses the central axis of the drift tube exactly will experience a constant drift field. Any ion that diffuses toward the side of the tube will travel a shorter distance and experience a higher drift field. Likewise, any ion that diffuses toward the outside of the drift tube will traverse a longer distance and experience a lower drift field. We have attempted to minimize this type of effect by incorporating ion funnels immediately before and after each curve. In this way, all ions are focused to the center of the drift region prior to entering each curve. At this early point in development, we cannot entirely rule out the possibility that the use of curved drift regions introduces some systematic errors in these measurements. However, we note that it is comforting that so far, in the systems we have studied, mobilities appear to be in reasonable agreement with accepted values. Thus, it appears that ions experience a drift field and length that corresponds to the center of the drift region.

With this analysis we expect that accurate mobilities are attainable with this approach. However, such accuracy may be somewhat artificial. As long as an ion’s mobility is in resonance with the drift field frequency, it should be transmitted. The ion need not be at the center of a distribution of ions. The requirement is that the structure is stable. Over multiple cycles, all other mobilities will ultimately drift into regions of the drift tube that do not lead to transmission. Therefore, while it is possible to measure which mobilities are stable at a particular number of cycles around the cyclotron, those mobilities may not correspond to the center point of a peak determined by traditional IMS methods. Rather, these ions may correspond to small stable...
populations that are centered near the leading or trailing sides of large populations.

Example Involving the Separation of Two Isomers: Raffinose and Melezitose. A much more difficult system to separate is the pair of isobaric trisaccharides melezitose and raffinose. Figure 5 shows that such species are readily isolated using the multipass capabilities of the cyclotron drift tube. We begin by recording the ion intensities as a function of drift field frequency around $3/4$ of the cyclotron for the $m/z = 527$ ion which corresponds to the $[M + Na]^+$ ion of each monomer. A total of 24 frequencies were scanned, and this was carried out separately for each trisaccharide. Melezitose possesses a more compact structure and is therefore observed at higher frequencies, with the highest intensity found at 169.5 Hz. The lower mobility raffinose shows a maximum at 149.3 Hz; however, we used a frequency of 144.9 Hz for the isolation of this sugar, since there was overlap at 149.3 Hz. With the use of the experimental conditions, these values can be used to define $K_0 = 4.05$ and 3.47 cm$^2$ V$^{-1}$ s$^{-1}$, respectively, for melezitose and raffinose. This matches well for melezitose, which has $K_0 = 4.08$ cm$^2$ V$^{-1}$ s$^{-1}$; the agreement is not as good for raffinose, which has $K_0 = 3.88$ cm$^2$ V$^{-1}$ s$^{-1}$, for the reasons described in the above section (i.e., we did not transmit the center of the ion distribution).

Example data obtained for the separation of the mixture of melezitose and raffinose after multiple cycles are also shown in Figure 5. At 169.5 Hz for melezitose or 144.9 Hz for raffinose we find that $1^{3/4}$ cycles is sufficient to isolate each ion. Under these conditions, melezitose has a drift time of 58.56 ms and raffinose has a drift time of 68.40 ms. At $3^{3/4}$ cycles, the ions have traveled 8.09 m and are further separated. Finally, at $6^{3/4}$ cycles, the two sugars have now traveled 13.52 m. The drift time increases to 176.64 ms for melezitose, and this ion has a fwhm of 1.20 ms, or $R = 147$. The drift time of raffinose is 206.64 ms; combined with a fwhm of 0.96 ms, we obtain $R = 215$ for this peak.

SUMMARY AND CONCLUSIONS

The first data collected with a multipass cyclotron geometry drift tube were presented. By variation of the field across regions of the drift tube at a fixed frequency, ions of specific mobilities are isolated and this isolation refined over numerous cycles through the instrument. The operation and precision of the instrument was demonstrated by examining the $[M + 2H]^+_{2+}$ and $[M + 3H]^3+$ ions of the peptide substance P for up to $10^{3/4}$ cycles. Additionally, we presented an isolation of a mixture of the isomers melezitose and raffinose. Peak widths do not increase significantly with an increase in the number of cycles. This is a result of the trimming away of ions in the leading or trailing regions of the distribution as species are transmitted to new drift regions. While not the focus of this first result, we note that this approach leads to limitations in sensitivity. We are currently investigating this and working on methods to improve sensitivity. Clearly, many other improvements also remain to be made as well; however, this first prototype appears promising as a means of increasing the resolving power of IMS methods.

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