# JMS-S3000 SpiralTOF

## Biomolecules Applications Notebook

Edition July 2016

### Table of Contents

**Introduction and Fundamentals**

<table>
<thead>
<tr>
<th>Topic</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Development of JMS-S3000: MALDI-TOF/TOF Utilizing a Spiral Ion Trajectory (Takaya Satoh, JEOL News, 45, 34-37, 2010)</td>
<td>1</td>
</tr>
<tr>
<td>The Relationship between Crystal Condition and Mass Resolving Power, Mass Accuracy</td>
<td>5</td>
</tr>
</tbody>
</table>

**Lipids**

<table>
<thead>
<tr>
<th>Topic</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Structural Analysis of Tristearin</td>
<td>17</td>
</tr>
<tr>
<td>Structural Analysis of Triolein</td>
<td>19</td>
</tr>
<tr>
<td>Structural Analysis of Oxidized Triolein</td>
<td>21</td>
</tr>
</tbody>
</table>

**See also in the JMS-S3000 SpiralTOF Imaging Applications Notebook:**

- MALDI-Imaging MS of Lipids on Mouse Brain Tissue Sections Using Negative Ion Mode
- Fingerprint Analyses Using MALDI Imaging and SEM Imaging

**Peptides and Proteins**

<table>
<thead>
<tr>
<th>Topic</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Analysis of Bovine Serum Albumin</td>
<td>23</td>
</tr>
<tr>
<td>High-Energy CID Analysis of Bovine Serum Albumin</td>
<td>25</td>
</tr>
<tr>
<td>Structural Analysis of a High Molecular Weight Peptide</td>
<td>27</td>
</tr>
<tr>
<td>Analysis of Phosphopeptide Using TOF-TOF</td>
<td>29</td>
</tr>
<tr>
<td>Distinguishing Lysine and Glutamine in a Peptide</td>
<td>31</td>
</tr>
<tr>
<td>Comparison of the JMS-S3000 SpiralTOF-TOF and a 4-Sector Tandem Double-Focusing Mass Spectrometer</td>
<td>35</td>
</tr>
<tr>
<td>High Sensitivity Peptide Measurement with the New Matrix α-Cyano-4-Chlorocinnamic Acid</td>
<td>37</td>
</tr>
<tr>
<td>High Sensitivity Analysis of Intact Proteins Using Linear TOF</td>
<td>41</td>
</tr>
<tr>
<td>MALDI-ISD Measurements Using Both the SpiralTOF Mode and the Linear Mode</td>
<td>43</td>
</tr>
<tr>
<td>High Resolution and High Mass Accuracy MALDI-ISD Measurements</td>
<td>45</td>
</tr>
</tbody>
</table>
Other biomolecules (natural products, oligosaccharides) .................................................................. 49
Analysis of the Natural Organic Compound SAAF by Using the TOF-TOF Option ............... 49
Analysis of the Natural Organic Compound YTX by Using the TOF-TOF Option ............. 51
High-energy CID Mass Spectrometry of Oligosaccharides ....................................................... 53

JMS-S3000 SpiralTOF bibliography ................................................................................................. 57

- Microsoft and Excel are either registered trademarks or trademarks of Microsoft Corporation in the United States and/or other countries.
- MASCOT is either a registered trademark or a trademark of Matrix Science, Ltd. in the United States and/or other countries.
Development of JMS-S3000: MALDI-TOF/TOF Utilizing a Spiral Ion Trajectory

Takaya Satoh
MS Business Unit, JEOL Ltd.

Introduction

The time-of-flight mass spectrometer (TOFMS) is one of mass spectrometry techniques, which include the quadrupole mass spectrometer, the magnetic sector mass spectrometer, the ion trap mass spectrometer and the Fourier transform ion cyclotron resonance mass spectrometer. In the case of TOFMS, ions of various m/z values, which are generated in the ion source, are accelerated to the detection plane by a pulse voltage applied from a starting time of data acquisition. Since the time-of-flight of ions at the detection plane are proportional to the square root of their m/z values, the ions generated in the ion source can be separated. One of the TOFMS feature is fast measurement, which is due to the unnecesity of scan for any physical parameters such as electric or magnetic fields. Recently, not only a single type mass spectrometer, but also a tandem type mass spectrometer connected with the quadrupole mass spectrometer (Q/TOF) or tandemly connected two TOFMSs (TOF/TOF) are available.

The mass resolution of TOFMS is expressed by $T/2\Delta T$, where $\Delta T$ is the time-of-flight distribution of the ion group with the same m/z value (ion packet) at the detection plane (that is, spatial distribution of the ion packet in the flight direction at the detection plane) and, $T$ is centroid of the time-of-flight distribution. Since TOFMS was invented in 1964 [1], its mass resolution has been improved by increasing $T$ and decreasing $\Delta T$. In 1955, a unique acceleration technique was developed, which focuses the initial space and energy distributions at the detector surface in the flight direction. Applying this technique, the mass resolution was increased by decreasing $\Delta T$ [2]. Furthermore, in the early 1970s, a new technique was developed. In this technique, the focus position defined by the above-mentioned acceleration technique is chosen as the start point, and an ion optical system that is composed of ion mirror [3] or electrostatic sectors [4] is placed at the post stage. This innovation made it possible to increase the time-of-flight $T$ without increasing $\Delta T$, and led to a dramatic improvement of the mass resolution. Recently, most of commercially available TOFMS instruments use ion mirrors, and their flight paths are 1 to 3 m. For further improvement in the mass resolution of TOFMS, another types of ion optical systems have been proposed. They are the multi-reflecting type [5] and the multi-turn type [6-7] ion optical system where ions fly multiple times on the certain trajectory. These two ion optical systems theoretically achieve an infinitely long flight path in a compact space, and improved the mass resolution. However, they have the limitation of the mass range because ions with large speed (ions with small m/z) lap the ions with small speed (ions with large m/z) when the ions flying on the same trajectory multiple times.

We have developed an original ion optical system that utilizes a spiral ion trajectory. This ion optical system can overcome the “lap” problem present in multi-reflecting and multi-turn type ion optical systems. In addition, it is possible to achieve mass resolution and mass accuracy higher than those of widely used reflectron ion optical systems. In this paper, we describe the design of the spiral trajectory.
type ion optical system, and basic performance of a MALDI-TOF/TOF system applying it. The system consisted of the spiral trajectory type ion optical system and reflectron type ion optical system using offset parabolic reflector for the first and second TOFMSs, respectively. The instrument achieves higher mass resolution, mass accuracy and precursor ion selectivity due to utilizing a spiral ion optical system for the first TOFMS, thus enabling more precise analysis.

Design of the spiral trajectory ion optical system

Multi-turn type ion optical system technique was applied for development of the spiral trajectory ion optical system. Especially, a combination of the “perfect focusing” and “multi-turn” [12] techniques developed at Osaka University, which achieved highest mass resolution in the world, was considered the most suitable for development of the spiral trajectory ion optical system. For conversion of a multi-turn type ion optical system for a spiral trajectory ion optical system, it is necessary to move ion trajectory perpendicular to the orbit plane. In order to achieve this, we have designed the system so that ion injection is slightly tilted to the orbit plane. The advantage of the design is that there is no need for the mechanism to transfer the ions to the next layer. There are concerns about degradation of mass resolution due to the trajectory deviation from a multi-turn type ion optical system. However, the effect should be negligible by keeping the injection angle to several degrees.

Practically, we have designed the spiral trajectory ion optical system based on MULTUM II [7] construction, which consists of four toroidal electrostatic sectors (cylindrical electrodes with two Matsuda plates). The schematic of the ion optical system is shown in Fig. 1. To achieve a spiral trajectory, we have constructed a layered toroidal electric field (TES) by placing (number of cycles +1) Matsuda plates into the cylindrical electrostaticic sectors. The Matsuda plates are arranged within certain equal distances \( L_e \) in the space \( L_e \), between the external and internal electrodes. The three types of voltages applied on TESs 1 to 4 is that of the internal electrode, external electrodes and Matsuda plates. Corresponding voltages are supplied to every Matsuda plates, internal and external electrodes of TESs 1 to 4.

Also, four TESs were placed so that they correspond to MULTUM II when looked from the orbit plane. Y direction was set perpendicular to the periodic orbit plane. In development of the MALDI-TOF/TOF, we have made Y direction to horizontal. The TES1 in the Fig. 1 shows the external electrode is removed so that it can be seen the Matsuda plates are equally spaced. Ions fly through the center of the space, formed by \( L_o \) and \( L_e \). Ion passes the same layer of TESs 1 to 4, and after passing the TES 4, it enters to the next layer of TES 1. The process is repeated for several cycles; the ion thus draws a spiral trajectory and reaches the detector (DET1) (Green line in the Figure 1 represents the ion trajectory). The injection angle \( \theta \) into the layered toroidal electric field can be expressed as follows,

\[
\tan \theta = (L_o + L_m) / L_e
\]

where, \( L_m \) is the thickness of a Matsuda plate and \( L_e \) is the one cycle length.

As mentioned above, owing to the usage of four TESs of the same structure in it’s construction, the ion optical system can achieve a complicated trajectory within a simple structure.

Production of MALDI-TOF/TOF utilizing spiral trajectory ion optical system

We have developed MALDI-TOF/TOF utilizing the spiral trajectory ion optical system. It consisted of the first TOFMS using the spiral trajectory ion optical system and the second TOFMS using the reflectron ion optical system. The mass spectrum measurement in the first TOFMS is referred as spiral mode, and the product ion spectrum measurement in the second TOFMS as TOF/TOF mode.

An schematic of the system is shown in Fig. 2 (ion source and the detector DET1 of the first TOFMSs are omitted). Spiral trajectory is set to eight cycles of 2.093 m per each. A distance between central trajectories of the adjacent layers is 58 mm, an injection angle is 1.6 degree according to equation (1). Y direction is set as horizontal, so the injection angle is achieved by tilting the extraction direction of the ion source 1.6 degrees from a horizontal plane.

In the spiral mode, ions fly a spiral trajectory and are detected with the spiral mode detector (though not specified in Fig. 2, it is located similarly to DET1 in Fig. 1). Ion gate is placed in the 7th cycle. It allows eliminating high-intensity matrix ions, which are outside of the data acquisition m/z range.

In TOF/TOF mode, selection width of the ion gate is made narrower and monoisotopic ions of precursor ions are selected out of all isotopic ions of them. It is possible to mechanically move the spiral mode detector out of the trajectory so that precursor ions can be introduced into the collision cell. Ions, that entered a collision cell, collide with rare gas inside of the cell with a kinetic energy of approximately
20 keV, and generate fragment ions. Precursor ions and fragment ions are mass-separated in a reflectron ion optical system that combines an offset parabolic reflectron (OPR) [13] and a reacceleration mechanism. OPR is a reflectron connecting a linear and parabolic electric fields. It allows simultaneous observation of ions, from low \( m/z \) fragment ions up to precursor ions. In addition, in order to increase transmission of ions, fine adjustment of the ion trajectory is enabled by installing two deflectors (DEF1 and DEF2) on both sides of the collision cell.

**Evaluation of MALDI-TOF/TOF with spiral trajectory ion optical system utilized**

Figure 3 shows mass spectrum of six types of peptide mixtures (in order of \( m/z \) increase: Bradykinin fragment 1-7, Angiotensin II, Angiotensin I, P14R, ACTH fragment 1-17, ACTH fragment 18-39). The mass spectrum of Angiotensin II and ACTH fragment 1-17 are also displayed as an enlarged image. Mass resolution is 58000 (FWHM) and 73000 (FWHM) respectively. The mass error of ACTH fragment 1-17 is 0.16 ppm, when internal calibration is performed among five peptides except ACTH fragment 1-17. It became clear from the above mentioned facts that distance of flight for spiral trajectory ion optical system is 17 m, which is 5 times longer than that of the conventional reflectron type ion optical systems. This allows enhancement of mass resolution and mass accuracy.

Figure 4 shows the relation between \( m/z \) value and mass resolution when mass resolution is adjusted with ACTH fragment 1-17. Figure 4 shows that it is possible to achieve high mass resolution simultaneously in a wide \( m/z \) range. This overcomes the problem of MALDI-TOFMS utilizing conventional reflectron type ion optical system that could achieve high mass resolution only in a narrow \( m/z \) range.

Figure 5a shows a product ion spectrum diagram of Poly (oxypropylene), acquired in TOF/TOF mode. Selected precursor ions are monoisotopic ions from [M+Na]\(^+\) series with \( m/z \) 1027. A numbers of fragmentation channels from sodium ions as fragment ion to precursor ion are observed. The enlarged spectrum around \( m/z \) 780 is shown in Fig. 5b. The system is able to select only monoisotopic ions of precursor ions, therefore each fragment channels can be observed as one peak without any isotopic peaks. Two peaks in Fig. 5b indicate different fragmentation channels. It indicates that 2u different fragmentation channels can be clearly separated. Figure 5c displays an image of the same \( m/z \) range as in Fig. 5b when measured with conventional MALDI-TOF/TOF. Pecursor ion selectivity of traditional TOF/TOF is insufficient so that the fragment ions from all isotopic ions of precursor ions are analyzed in the second TOFMS. Thus every fragmentation channels of product ion spectrum include isotopic peaks. As a result, when \( m/z \) values of monoisotopic ions of two fragment channels are close, such as 2 u, their isotopic peaks are overlapped and are impossible to be clearly identified. The high precursor ion selectivity originated from the spiral trajectory ion optical system used in this system makes the structural analysis of chemical compounds much easier.

**Conclusion**

This paper reports on the development of the spiral trajectory ion optical system. Also, the paper describes the development of MALDI-TOF/TOF, which combines a spiral trajectory ion optical system and reflectron type ion optical system utilizing offset parabolic ion mirrors. Innovative ion optical system introduced to the JMS-S3000 has overcome preexisting problems related to conventional MALDI-TOF and MALDI-TOF/TOF. Thus, the JMS-S3000 is expected to play a significant role in various areas.

**References**


---

Fig. 3 Mass spectrum of peptide mixtures.
Fig. 4 Relation between m/z value and mass resolution.

Fig. 5 Product ion spectrum of Poly(oxypropylene).

SpiralTOF™
The Relationship between Crystal Condition and Mass Resolving Power, Mass Accuracy

Introduction
The JMS-S3000 SpiralTOF™ has a unique 17 m flight path that offers the highest resolution MALDI-TOF MS system currently available. With an extended flight distance, the SpiralTOF reduces topographic effect of matrix crystal to a minimum and achieves highly reproducible mass resolving power and high mass accuracy with external mass calibration.

In this work, we demonstrate the measurement of a polymer standard with 4 types of matrices that are typically used for MALDI polymer measurement by using the JEOL SpiralTOF system. Additionally, we looked at the crystal condition using the JEOL JSM-7600F thermal field emission scanning electron microscope (FE-SEM).

Experimental
Sample information and preparation conditions are shown in Table 1. PEG1500 was dissolved in water at a concentration of 10 mg/mL. Each matrix was dissolved in THF at a concentration of 10 mg/mL. NaI used as the cationization agent was dissolved in THF at a concentration of 1 mg/mL. Next, the PEG1500, NaI and matrix solutions were mixed together 1:1:2 (1:1:4 for DIT) by volume. Afterwards, 0.75 µL of this mixture was placed on the hairline finish stainless steel plate (MTP format, 384 spots for samples and 96 spot for calibrant). Finally, the dried sample was measured using the JMS-S3000 SpiralTOF MS system. We also obtained SEM images for each crystal condition with the JSM-7600F.

<table>
<thead>
<tr>
<th>Polymer standard</th>
<th>Conc.</th>
<th>Solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyethylene glycol (PEG) 1500</td>
<td>10 mg/mL</td>
<td>Water</td>
</tr>
</tbody>
</table>

Cationization agent

<table>
<thead>
<tr>
<th>Matrix</th>
<th>Conc.</th>
<th>Solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaI</td>
<td>1 mg/mL</td>
<td>Water</td>
</tr>
<tr>
<td>α-Cyan-4-hydroxyphenylacetic acid (CHCA)</td>
<td>10 mg/mL</td>
<td>Tetrahydrofuran (THF)</td>
</tr>
<tr>
<td>2,5-Dihydroxybenzoic acid (DHB)</td>
<td>10 mg/mL</td>
<td>THF</td>
</tr>
<tr>
<td>Dithranol (DIT)*</td>
<td>10 mg/mL</td>
<td>THF</td>
</tr>
<tr>
<td>trans-3-Indoleacrylic acid (IAA)</td>
<td>10 mg/mL</td>
<td>THF</td>
</tr>
</tbody>
</table>

Sample

| PEG1500: NaI/Matrix = 1/1/2 (v/v) | 0.75 µL of this sample solution mixture was placed on the MALDI target plate |

JSM-7600F conditions

<table>
<thead>
<tr>
<th>Sample preparation</th>
<th>Uncoated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acceleration voltage</td>
<td>1 kV</td>
</tr>
<tr>
<td>Magnification</td>
<td>x500 and x2,000</td>
</tr>
</tbody>
</table>

Figure 1. Reduced topographic effect of matrix crystal.

Figure 2. A difference of flight time arises due to the topographic effect of the matrix crystal.

Figure 3. JMS-3000 SpiralTOF.

Figure 4. JSM-7600F Thermal FE-SEM.

Table 1. Sample information and preparation conditions.
Results & discussion:
The MALDI mass spectra of PEG1500 are shown in Figure 4 for each matrix. We set the delay time to achieve the maximum mass resolving power at m/z 1537.9. Therefore, the resolving power was approximately 70,000 for the [HO(C,H,O),H + Na]+ peaks, well in excess of that needed to separate isotope peaks. Additionally, we observed excellent mass distributions.

We determined the average mass resolving power (n=10) and external mass accuracy (n=8) for m/z 1097.6, m/z 1537.9 and m/z 1978.2 for each matrix. The results are shown in Figure 5 and 6, respectively. We achieved high mass resolving power for the three selected ions with each matrix. In addition, we obtained excellent mass error (less than 10 ppm) with external calibration for each matrix.

Figure 4. MALDI mass spectra of PEG1500.
Figure 5. Averaged mass resolving power (n=10) for m/z 1097.6, m/z 1537.9 and m/z 1978.2.

Figure 6. Mass error with external calibration (n=8) for m/z 1097.6, m/z 1537.9 and m/z 1978.2.
We examined the crystal condition with the JEOL JSM-7600F thermal field emission scanning electron microscope. The SEM images are shown Figures 7-10. The crystal shape, size and dispersion were quite different in each matrix crystal. However, SpiralTOF performance was not affected by the topographic effects because these spatial differences were a negligible fraction of the 17 m flight path.

**Conclusions**
SpiralTOF achieved highly reproducible mass resolving power and high mass accuracy with external mass calibration for all samples. These values were not significantly influenced by the different crystal morphologies for the different matrices. This is attributed to the SpiralTOF’s very long (17 meter) flight path.

![Figure 7. SEM images of CHCA crystal with PEG1500: left: x500, right: x2,000.](image)

![Figure 8. SEM images of DHB crystal with PEG1500: left: x500, right: x2,000.](image)
Figure 9. SEM images of DIT crystal with PEG1500: left: x500, right: x2,000.

Figure 10. SEM images of IAA crystal with PEG1500: left: x500, right: x2,000.
This page intentionally left blank for layout purpose.
Structural Analysis of Triacylglycerols by Using a MALDI-TOF/TOF System with Monoisotopic Precursor Selection

Ayumi Kubo,1 Takaya Satoh,1 Yoshiyuki Itoh,1 Masahiro Hashimoto,1 Jun Tamura,1 Robert B. Cody2

1JEOL Ltd., Tokyo, Japan
2JEOL USA, Inc., Peabody, MA 01960, USA

Abstract. A new MALDI-TOF/TOF system with monoisotopic precursor selection was applied to the analysis of triacylglycerols in an olive oil sample. Monoisotopic precursor selection made it possible to obtain product-ion mass spectra without interference from species that differed by a single double bond. Complete structure determination of all triacylglycerols, including structural isomers, was made possible by interpreting the charge-remote fragmentation resulting from high-energy collision-induced dissociation (CID) of the sodiated triacylglycerols.

Key words: MALDI, TOF/TOF, Charge-remote fragmentation, High-energy CID, Triacylglycerols, MS/MS, Lipids, Collision-induced dissociation

Introduction

Triacylglycerols (TAGs or triglycerides) are comprised of three fatty acids esterified with glycerol. Because TAGs are the major components in animal fats and vegetable oils, the analysis of TAGs is biologically important and crucial for quality control of food products.

Recent mass spectrometric approaches to the analysis of TAGs have made use of atmospheric pressure chemical ionization (APCI) [1–5] or electrospray ionization (ESI) [6–10] and tandem mass spectrometry (MS/MS).

Using ESI and a triple quadrupole mass spectrometer, Hsu and Turk [6] reported that collisional activation of lithium adducts of TAGs can provide structural information about the acyl groups. Low-energy collision-induced dissociation (CID) of cationized TAGs does not provide information about the position of the double bonds. However, the CID fragments of unusual dilithiated species was shown to be dependent on double bond location. Byrdwell and Neff [7] reported a method based on dual parallel ESI and APCI combined with tandem mass spectrometry for the analysis of TAGs and their oxidation products. McAnoy et al. [8] used ESI with a linear ion trap to characterize TAG components within a complex mixture of neutral lipids from cell extracts.

High-energy CID is an especially attractive approach for TAG analysis because charge-remote fragmentation [11–22] provides a great deal of information about lipid structure. The complete structural characterization of TAGs was reported in 1998 by Cheng et al. using fast atom bombardment (FAB) and tandem magnetic sector mass spectrometry with high-energy CID fragmentation of the [M + Na]+ species [21]. All TAG structural features could be determined except stereochemistry.

However, large tandem magnetic sector mass spectrometers have fallen out of favor in recent years and high-energy CID appeared destined to become a “lost art” until the introduction of tandem time-of-flight (TOF/TOF) mass spectrometers by Cotter and Cornish in 1993 [22]. Recently, Pittenauer and Allmaier showed that TOF/TOF mass spectrometers have the potential to provide the same complete structural information as a tandem magnetic sector mass spectrometer [23]. The principal limitation of this method was found to be the poor MS-I selectivity (a 4 to 6 u window) of the TOF/TOF system, making it impractical to select precursor ions for TAGs with compositions that differ by two hydrogens. The authors concluded that a LC/MALDI-MS/MS approach might be required to make use of charge-remote fragmentations to characterize TAGs in complex mixtures.

We have developed a tandem time-of-flight mass spectrometer featuring high precursor ion selectivity that resolves the problem of poor MS-I selectivity [24]. The mass
spectrometer uses multi-turn and “perfect focusing” ion optics [25] to fit a very long (17-m) flight path into a compact space [26]. In TOF/TOF mode, an ion gate positioned at the 15 m point in the spiral ion flight path is used to isolate and guide the precursor ion into a gas-filled collision chamber. The long flight path provides ample time separation prior to precursor ion selection, resulting in unit precursor selectivity. The precursor ions undergo 20 kV collisions with a target gas and are subjected to a 9 kV post-acceleration into an offset parabolic reflector with wide energy acceptance.

Monoisotopic precursor selection combined with high-energy CID is the key to using TOF/TOF for structural analysis of triacylglycerols in complex mixtures. This paper describes the method for structural analysis with this system and reports the complete structural analysis of TAGs, including isomers, in a commercial olive oil sample.

**Experimental**

**Materials and Chemicals**

A triacylglycerol standard (1-palmitoyl-2-oleoyl-3-linoleoyl-rac-glycerol), matrix (2,5-dihydroxybenzoic acid or DHB), and cationizing agent (sodium trifluoroacetate), were purchased from Sigma-Aldrich (St. Louis, MO). A trioleoylglycerol (triolein) standard was purchased from TCI. Tetrahydrofuran (THF) was purchased from Wako (Osaka, Japan) and olive oil was purchased from local stores. The triacylglycerol standard, including triolein, and the olive oil were dissolved in THF at respective concentrations of 100 pmol/μL and 10 μg/μL. A solution of sodium trifluoroacetate and DHB was dissolved in THF at respective concentrations of 1 μg/μL and 20 μg/μL, and added to the samples at a volume ratio of 1:1:2. The resulting mixture was loaded onto an MTP 96-hole hairline plate (JEOL Ltd., Akishima Japan) at a volume of 1 μL per spot.

**MALDI Mass Spectrometry**

A JMS-S3000 Spiral TOF (JEOL Ltd., Akishima, Japan) equipped with the TOF/TOF option was used for all measurements. The laser was a Nd-YLF laser operated at a wavelength of 349 nm. The laser intensity and the detector voltage were set to prevent triacylglycerol peaks from saturating. The extraction delay was optimized to 400 ns to provide a resolving power (FWHM) of approximately 50,000 for the TAG peaks in MS-I mode. For product-ion mass spectrum acquisition, helium collision gas was introduced to attenuate the precursor ion abundance to approximately 50 % of the initial value. The laser was operated at a repetition rate of 1000 Hz. Spectra were acquired at a rate of two spectra per s and 500 spectra were accumulated for each product-ion mass spectrum shown here.

**Results and Discussion**

Figure 1 shows the structure of 1-palmitoyl-2-oleoyl-3-linoleoyl-rac-glycerol. The structure shows (18:1) oleic acid, (16:0) palmitic acid, and (18:2) linoleic acid substituents at positions sn-2 (the site that determines the stereochemistry) and positions sn-1 and sn-3, respectively. In this article, we have labeled the fatty acid substituents at positions sn-1 and sn-3 as “sn-1/sn-3.” The substituents at sn-1 and sn-3 are indistinguishable by mass spectrometry because the steric structure of triacylglycerol cannot be identified by mass spectrometry. Each fragmentation path is assigned as shown in Figure 1, and is labeled alphabetically. Each letter represents the initial letter of the fatty acid, and the accompanying number represents the bonding position in each fatty acid. The labeling for TAGs such as “TAG(54:3)” follows the convention where the numeral on the left in parentheses represents the total number of acyl carbon chains and the numeral on the right represents the total number of unsaturated bonds at fatty acid moieties.

The major species observed for 1-palmitoyl-2-oleoyl-3-linoleoyl-rac-glycerol was the sodiated molecule [M + Na]+. Figure 2 shows the product ion spectrum acquired by selecting the monoisotopic ion of this species. The resulting fragment ions are solely monoisotopic ions as well because a monoisotopic precursor ion was selected. Thus, each fragmentation path is observed as a single peak on the product-ion mass spectrum. Figure 2a shows the entire mass range of the product-ion spectrum. The Na+ peak detected at m/z 23.0 confirms that the precursor ion is indeed [M + Na]+. Peaks characteristic of fatty acid fragmentation are predicted as A-, B-, C-, D-, E-, F-, G-, H-, I-, J-, K-, L-, M-, N-, O-, P-, Q-, R-, S-, T-, U-, V-, W-, X-, Y-, Z-.
C-, E-, G-, and J-type ions using the nomenclature defined in reference [21]. Figure 2a demonstrates that all of A-, B-, C-, E-, G-, and J-type ions predicted in reference [21] were observed for this example and "G+2" ions (mentioned in reference [21]) were observed. The structure of "G+2" ions and their fragmentation pathway are not clear, but "G+2" ions were also observed in the product ion spectrum of the triolein standard (shown in Figure 5b) at a relatively lower intensity than that of the G-type ions. Figure 2b also shows that signals resulting from charge-remote fragmentation were detected in the high mass range above m/z 650. When the fragment ion at each bonding position is defined as in Figure 1, the peaks can be assigned as shown in Figure 2b. The intensities of fragment ions corresponding to unsaturated bonding positions, such as LΔ9, LΔ12, and OΔ9, are relatively weak or are not observed, resulting in a peak pattern that reflects the structure of 3 fatty acids.

In the analysis of triacylglycerols in the olive oil sample, particular attention was focused on the G- and J-type ions. These ions help determine the numbers of carbon chains and unsaturated bonds in each fatty acid. In the G-type ion, fatty acids remain at sn-1/sn-3, while the J-type ion, where a fatty acid remains at sn-2, has one less CH₂ at the end. This makes it possible to estimate the bonding positions of three fatty acids because fatty acids having an odd acyl carbon number rarely exist in the natural world.

Figure 3 shows the mass spectrum of the olive oil. Sodiated triacylglycerols [M + Na]+ were observed for this sample that included TAG (52:3) (m/z 879.7), TAG (52:2) (m/z 881.7), TAG (54:4) (m/z 905.8), and TAG (54:3) (m/z 907.8). The monoisotopic ions of these four TAGs were selected as the precursor ions, and their product-ion mass spectra were acquired. Figure 4 shows the spectra of ions at m/z 905.8 acquired before and after the precursor ion selection. The figure demonstrates that only the ions at m/z 905.8 were selected, completely eliminating ions at other mass values.

Figure 5 shows comparison between the product-ion mass spectra of TAG (54:3) [M + Na]+ at m/z 907.8 from olive oil and from triolein standard. Given that olive oil is...
rich in oleic acid, the ion at \( m/z \) 907.8 is expected to contain three oleic acids (18:1). Both of the product-ion mass spectra show a \( J_2 \)-type ion at \( m/z \) 331.3, indicating that an oleic acid is bonded at the \( sn-2 \) position, and a \( G \)-type ion at \( m/z \) 345.3, indicating that an oleic acid is bonded at the \( sn-1/sn-3 \) positions. The spectra show only one peak that is considered an \( A \)-, \( B \)-, and \( C \)-type ion, suggesting that TAG (54:3) is trioleoylglycerol, which contains three oleic acid molecules. The signals in high mass region resulting from charge-remote fragmentation were identical between the sample and standard, and the spectral patterns were consistent with structure of oleic acid.

![Figure 4](www.jeol.com)

**Figure 4.** Precursor-ion selection for ions at \( m/z \) 905.8 (a) before selection, (b) after selection

![Figure 5](www.jeol.com)

**Figure 5.** Comparison of product-ion mass spectra for the precursor at \( m/z \) 907.8 with that of sodiated triolein standard, (a) the ions at \( m/z \) 907.8 from olive oil, (b) triolein standard
Next, the ion at \( m/z \) 905.8 was selected as the precursor ion. The \( m/z \) value of this ion suggests that it is a monoisotopic \([M + Na]^+\) ion of TAG (54:4). Figure 6 shows the product-ion mass spectrum. It is expected that this triacylglycerol is also composed of two (18:1) oleic acids and one (18:2) linoleic acid, given that the major component of olive oil is oleic acid. The product-ion mass spectrum shows fragment ions assigned as G-type ions, at \( m/z \) 343.4 and \( m/z \) 345.3. If the ion at \( m/z \) 343.4 is a G-type ion, the fatty acid molecule at \( sn-1/sn-3 \) is linoleic acid, and if the ion at \( m/z \) 345.4 is a G-type ion, the fatty acid molecules at \( sn-1/sn-3 \) are oleic acid. The ion at \( m/z \) 347.3 is assigned as a “G+2” ion because in the product-ion mass spectrum “G+2” ions were observed at lower intensity than G-type ions as discussed above, and the intensity of the ion at \( m/z \) 347.3 is relatively lower than that of the ion at \( m/z \) 345.3. This is consistent with the assignment of G+2 ions by Cheng et al. in reference [21]. Since the G-type ion suggests that both oleic acid and linoleic acid are bonded, the remaining fatty acid is (18:1) oleic acid. Next, the product-ion spectrum shows J-type ions: a J-type ion containing oleic acid (J2O) and a J-type ion containing linoleic acid (J2L) at \( m/z \) 331.3 and \( m/z \) 329.3, respectively. In the high-mass region, the signals resulting from charge-remote fragmentation were consistent with the structures of oleic acid and linoleic acid. This demonstrates that \( m/z \) 905.8 is triacylglycerol that contains two molecules of oleic acid and one molecule of linoleic acid and is a mixture of the structural isomers 1,3-dioleoyl-2-linoleoyl-glycerol and 1,2-dioleoyl-3-linoleoyl-glycerol. Table 1 summarizes the structures of triacylglycerols determined for the olive oil samples from the product-ion mass spectra. The ions associated with the peak at \( m/z \) 879.7 are a mixture of the structural isomers 1-palmitoyl-2-oleooyl-3-linoleoyl-glycerol and 1-palmitoyl-2-linoleoyl-3-oleoyl-glycerol.

### Table 1. Summary of TAGs Found in the Olive Oil Sample

<table>
<thead>
<tr>
<th>( m/z )</th>
<th>Acyl carbon number and number of double bond</th>
<th>Composition of each fatty acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>879.7</td>
<td>52:3</td>
<td>(16:0,18:1,18:2) (16:0,18:2,18:1)</td>
</tr>
<tr>
<td>881.7</td>
<td>52:2</td>
<td>(16:0,18:1,18:1)</td>
</tr>
<tr>
<td>905.8</td>
<td>54:4</td>
<td>(18:1,18:1,18:2) (18:1,18:2,18:1)</td>
</tr>
<tr>
<td>907.8</td>
<td>54:3</td>
<td>(18:1,18:1,18:1)</td>
</tr>
</tbody>
</table>

**Figure 6.** Product-ion mass spectrum for the precursor at \( m/z \) 905.8

**Conclusion**

Monoisotopic precursor selection was demonstrated for TOF/TOF analysis of a standard TAG and TAGs in an olive oil sample. This selectivity made it possible to use charge-remote fragmentation to determine the complete structure (except stereochemistry) for all of the TAGs, including structural isomers, present in the sample. Multiple structural isomers in the precursor ion were identified through the observation of G- and J-type ions. These results demonstrate that the MALDI-TOF-TOF system with high precursor ion selectivity can fully analyze the structure of triacylglycerols without prior chromatographic separation, and that the method is effective for the analysis of complex fat composites in food.

**Open Access**

This article is distributed under the terms of the Creative Commons Attribution License which permits any use, distribution, and reproduction in any medium, provided the original author(s) and the source are credited.

**References**


SpiralTOF-TOF
Structural Analysis of Tristearin

Introduction:
The JMS-S3000 SpiralTOF™ is a MALDI-TOF MS that uses an innovative spiral ion optical system to achieve the highest resolution currently available for a MALDI instrument. Additionally, this system can be equipped with a TOF-TOF option that can acquire high-energy collision-induced dissociation (HE-CID) product ion spectra for monoisotopically selected precursor ions. The resulting HE-CID product ion spectra provide detailed structural information about compounds like triglycerides by means of charge-remote fragmentation (CRF).

Tristearin is a triglyceride found in fats that consists of three stearic acid moieties (Fig. 1). In this work, we report the structural analysis of tristearin by using the HE-CID capabilities of the SpiralTOF-TOF.

Results and Discussion:
The sample was dissolved with NaI in methanol to promote Na⁺ adduct formation. PEG 1000 was used as an external calibration standard. Initially, the sample was measured in Spiral mode and showed a signal at m/z 913.8231 (Fig. 2). This ion was consistent with the expected [M+Na]⁺ calculated mass of 913.8194. Next, the product ion spectrum of this monoisotopically selected m/z was obtained using the TOF-TOF mode (Fig. 3). The resulting mass spectrum showed a fragmentation pattern consistent with the occurrence of CRF. Fig. 4 shows the enlarged m/z 600-920 mass range in which the labeled peaks are clearly reflected in the structure shown in Fig. 5. It should be noted that these fragment ions were detected when CRF occurred to one of the three stearin branches present in the molecule.

Figure 1. Structure of tristearin.

Figure 2. Mass spectrum of tristearin.
Conclusions:
Using the TOF-TOF method, the JMS-S3000 “Spiral-TOF” can acquire HE-CID product ion spectra that show clear CRF mass spectral patterns. Therefore, the structural analyses of compounds like tristearin are easily performed using this technique.

References:
SpiralTOF-TOF
Structural Analysis of Triolein

Introduction:
The JMS-S3000 “SpiralTOF™” is a MALDI-TOF MS that uses an innovative spiral ion optical system to achieve the highest resolution currently available for a MALDI instrument. Additionally, this system can be equipped with a TOF-TOF option that can acquire high-energy collision-induced dissociation (HE-CID) product ion spectra for monoisotopically selected precursor ions. The resulting HE-CID product ion spectra provide detailed structural information about compounds like triglycerides by means of charge-remote fragmentation (CRF). In this work, we report the structural analysis of triolein, a triglyceride that contains 3 oleic acid moieties (Fig. 1), by using HE-CID capabilities of the SpiralTOF-TOF.

\[
\begin{align*}
&\text{CH}_2-O-\text{C}=	ext{C}-\text{C}=	ext{C} \quad \text{CH}_2-O-\text{C}=	ext{C}-\text{C} \quad \text{CH}_2-O-\text{C}=	ext{C}-\text{C} \\
&\text{CH}-O-\text{C}=	ext{C}-\text{C} \quad \text{CH}-O-\text{C}=	ext{C}-\text{C} \quad \text{CH}-O-\text{C}=	ext{C}-\text{C}
\end{align*}
\]

Figure 1. Structure of triolein.

Results and Discussion:
The sample was dissolved with NaCl in methanol to promote Na\(^{+}\) adduct formation. PEG 1000 was used as an external calibration standard. Initially, the sample was measured in Spiral mode and showed a signal at m/z 907.7782 (Fig. 2). This ion was consistent with the expected [M+Na]\(^+\) calculated mass of 907.7725. Next, the product ion spectrum for this peak was acquired using the TOF-TOF mode (Fig. 3). The resulting mass spectrum showed a fragmentation pattern consistent with the occurrence of CRF. Fig. 4 shows the enlarged m/z 600-920 mass range in which the labeled peaks are reflected in the structure shown in Fig. 5. Each of the major peaks between m/z 684-753 and between m/z 807-821 (except m/z 808) shows a difference of 14Da which is consistent with successive losses of CH\(_2\) along the oleic acid backbone. However, the m/z 808, which is 1Da from m/z 807, is a peak that specifically appears when unsaturated bonds are present in the structure. Additionally, the low intensity peaks between m/z 753-807 also confirm the presence of the unsaturated bonds at this position. As noted previously, all of these product ion peaks are consistent with CRF occurring along an oleic acid branch within triolein.

\[
\begin{align*}
\text{Intensity} \\
0 & \quad 200 \quad 400 \quad 600 \quad 800 \quad 1000
\end{align*}
\]

Figure 2. Mass spectrum for the sodium adduct of triolein.
Conclusions:
The data for triolein demonstrates that the JMS-S3000 "SpiralTOF" can acquire HE-CID product ion spectra with clear CRF mass spectral patterns. Additionally, this method allows for easy determination of unsaturated bond locations in the carbon chains.

References:
**SpiralTOF-TOF**

**Structural Analysis of Oxidized Triolein**

**Introduction:**
In previous work, we showed that the JEOL SpiralTOF-TOF system’s high-energy collision-induced dissociation (HE-CID) is useful for the structural analysis of triglycerides.\(^1\) The resulting HE-CID mass spectra provided detailed information about the fatty acid moieties such as the positions of double bonds, branching, hydroxylation, and oxidation by means of charge-remote fragmentation (CRF).\(^2\)

In this work, we report the structural analysis of oxidized triglycerides by HE-CID using the SpiralTOF-TOF.

**Experimental:**
Triolein, a triglyceride with 3 oleic acid moieties (Fig.1), was used in this experiment. The triolein sample was kept at 160°C for 60min so that thermal oxidation would occur. The resulting sample was then dissolved in tetrahydrofuran (THF) at a concentration of 10 mg/mL. 2′,4′,6′-Trihydroxyacetophenone monohydrate (THAP) was used as the matrix and sodium trifluoroacetate (NaTFA) as the cationizing reagent. The THAP and NaTFA were dissolved in THF at a concentration of 10mg/mL and 1mg/mL, respectively. These two solutions and the sample solution were then mixed 1:1:1 by volume. Afterwards, 0.5 μL of this mixture solution was deposited and dried on the MALDI target plate.

**Results:**
The MALDI mass spectrum of oxidized triolein is shown in Fig. 2. PEG10000 was used as an external calibrant. A series of sodiated peaks were observed for non-oxidized (m/z 907), mono-oxidized (m/z 923), di-oxidized (m/z 939), and tri-oxidized (m/z 955) triolein. These results were supported by elemental composition calculations that showed a mass accuracy of < 2 ppm for each ion.

The product-ion spectra for m/z 907 and m/z 923 are shown in Fig. 3. Many product-ions caused by CRF were observed which simplified the analysis of each structure. The A, B, J2, and G ions in the product-ion spectrum for m/z 923 were shifted by 16 Da when compared to the product-ion spectrum for m/z 907. An enlarged product-ion spectrum for m/z 923 is shown in Fig. 4. The correlating structures for these ions are based on the structural fragmentation assignments shown in Fig. 5. These results clearly show that the high-energy CID product ions collected by SpiralTOF-TOF provide a very straightforward pattern for identifying the sample structure and the position of oxidation.

**Conclusion:**
In this study we demonstrated the structural analysis of an oxidized triglyceride using the high-energy CID available on the MALDI-SpiralTOF-TOF. Additionally, CRF using high-energy CID is a very useful technique for assigning the position of oxidation.

**Reference:**
Figure 3. Product ion mass spectrum from m/z 907 (top) and m/z 923.

Figure 4. Product ion spectrum from m/z 923 (enlarged between m/z 680 and m/z 920).

Figure 5. Assignment of product ions from m/z 923.
SpiralTOF™
Introduction of the JMS-S3000 SpiralTOF™
Analysis of Bovine Serum Albumin

Introduction:
The JMS-S3000 “SpiralTOF™” is a MALDI-TOF MS incorporating an innovative SpiralTOF™ ion optics system (Fig. 1). JEOL’s patented technology\(^1\) achieves a spiral ion trajectory of 17m within a compact 1m space. The ions are sent through four sets of layered toroidal electric fields, which are implemented by four pairs of cylindrical electrodes and nine Matsuda plates that are incorporated within every pair of cylindrical electrodes. Ions are accelerated to 20 kV in the ion source and fly sequentially through the layered toroidal electric fields to reach the detector (Fig. 2). With the extended flight distance, the JMS-S3000 SpiralTOF™ achieves high mass resolving power and high mass accuracy over a wide mass range. However, ions with a very short lifetime or that undergo spontaneous dissociation during their flight (e.g., protonated molecules of high-mass proteins, multiply-phosphorylated peptides, etc.) cannot be detected by either the SpiralTOF™ or a conventional reflectron TOF. Considering the wide application of the MALDI technique, a mass analyzer that can detect such short-lived ions is necessary in addition to the SpiralTOF™. A Linear TOF option is available for the JMS-S3000 in order to satisfy this requirement (Fig. 2). In this application note, the analysis of the tryptic digest of bovine serum albumin (BSA) is shown as a Spiral mode example while the analysis of intact BSA is shown as a Linear mode example.

Figure 1. JMS-S3000 SpiralTOF™.

Figure 2. Ion trajectories of the JMS-S3000 SpiralTOF™.
Methods:
The tryptic digest of BSA was analyzed in SpiralTOF mode. An external mass calibration was used for this data. The intact BSA was analyzed using the Linear TOF mode. The data was acquired automatically by using the msTornado™ Control, which is the instrument control and data acquisition software for the JMS-S3000.

Results and discussion:
The mass spectrum of the tryptic digest equivalent to 25 fmol of BSA is shown in Fig. 3. The peak list from this mass spectrum was submitted to the MASCOT peptide mass fingerprint (PMF) search, and the protein was identified as BSA. The mass error plot of all matched peptides is shown in Fig. 4. The RMS error was 3 ppm. The mass spectrum of the intact BSA acquired in Linear mode is shown in Fig. 5. Single- and double-charge protonated molecular ion peaks were observed at the m/z values expected for the primary structure of this protein.

Conclusions:
A mass spectrum of the tryptic digest of BSA with very high mass accuracy was obtained using Spiral mode. Using the MASCOT PMF search method, a peptide mass tolerance set as narrow as 10 ppm can lead to highly reliable protein identification with very few false positives. Additionally, the molecular weight of an intact protein was readily obtained by using the JMS-S3000 Linear mode.

References:

SpiralTOF-TOF
High-Energy CID Analysis of Bovine Serum Albumin

Introduction
The JMS-S3000 “SpiralTOF™” is a MALDI-TOFMS that uses an innovative SpiralTOF ion optics system to achieve the highest resolution currently available for a MALDI instrument (Fig. 1 and 2). Additionally, this system can be equipped with a TOF-TOF option that can acquire high-energy collision-induced dissociation (CID) product ion spectra for monoisotopically selected precursor ions. The distance to the ion gate is 15 m, more than one order of magnitude longer than that of conventional MALDI TOF-TOF instruments, thus allowing the monoisotopic selection of the precursor ion. The second TOFMS incorporates a re-acceleration mechanism and an offset parabolic reflector, another innovative ion optical system developed by JEOL. This unique design enables the seamless observation of product ions ranging from very low m/z up to that of the precursor ion.

In a previous application note1, we reported the analysis of bovine serum albumin (BSA) by Spiral and Linear modes. In this work, we show the analytical result for BSA by using the JMS-S3000 “SpiralTOF™” with the TOF-TOF option.

Results and Discussion
As a starting point, the tryptic digest of BSA (tBSA) was analyzed using just the Spiral mode. For the calibration step, a peptide mixture was used as an external mass calibrant, which resulted in an RMS error of 4 ppm. Subsequently, the mass spectrum peak list for this sample was submitted to the MASCOT (Marix Science, Ltd.)

Figure 1. JMS-S3000 SpiralTOF with TOF-TOF attachment.

Figure 2. Ion trajectory of Spiral-TOF and TOF-TOF attachment.
peptide mass fingerprint search, and the protein was identified as BSA. These results are in very good agreement with the results obtained previously for Spiral mode.\(^1\)

Next, the 10 highest intensity ions in the tBSA mass spectrum were selected as the precursor ions for automatic measurement using the TOF-TOF option. The product ion spectra for \(m/z\) 927.5, 1439.8, and 1567.7 shown in Fig.3 had an immonium ion along with the \(a\)-ion, \(d\)-ion, and \(w\)-ion series. These spectra along with the other 7 product ion spectra were then submitted to the MASCOT MS/MS Ion Search, and the protein was identified as BSA (see Fig.4).

Conclusions
The high resolution tBSA mass spectrum measured using Spiral mode was easily identified using the MASCOT peptide mass fingerprint method, even with a peptide mass tolerance set as narrow as 10 ppm (not shown in this note). Additionally, the high energy CID of the 10 highest intensity monoisotopically selected peaks measured using the SpiralTOF-TOF mode produced product ion spectra that readily identified BSA as the protein through the MASCOT MS/MS Ion Search.

Reference

Figure 3. Product ion spectra for \(m/z\) (a) 927.5, (b) 1439.8, and (c) 1567.7.

Figure 4. MASCOT MS/MS Ion search result of tBSA by TOF-TOF mode.
Introduction:
The JMS-S3000 SpiralTOF™ is a MALDI-TOF MS that uses an innovative spiral ion optics system to achieve the highest resolution currently available for a MALDI instrument. Additionally, the JMS-S3000 is available with a TOF-TOF option that acquires high-energy collision-induced dissociation (CID) product-ion spectra for monoisotopically selected precursor ions. In this work, we analyzed a high molecular weight peptide by using the JMS-S3000 SpiralTOF with the TOF-TOF option.

Experimental:
Sample: ACTH18-39 (Arg-Pro-Val-Lys-Val-Tyr-Pro-Asn-Gly-Ala-Glu-Asp-Glu-Ser-Ala-Glu-Ala-Phe-Pro-Leu-Glu-Phe)
Matrix: α-Cyano-4-hydroxycinnamic acid (CHCA)

Results and Discussion:
The high-energy CID product-ion spectrum for protonated ACTH18-39 [M+H]+ (m/z 2456.2) is shown in Fig. 1. Each product ion was labeled using the Biemann convention (Fig. 2) in which the a-, b-, and d-ion series are fragments generated from the N terminus of the ACTH18-39 molecule. The SpiralTOF MS/MS data in Fig. 1 show that the sequence information is clearly represented by the a-ion series from a2 to a21. The immonium ions (Fig. 3) and the d-ion series (produced by fragmentation of the a-ion series) were also observed in the mass spectrum.

Conclusions:
The JMS-S3000 SpiralTOF with the TOF-TOF mode produced high energy CID product ion spectra that clearly identified the sequence for the high molecular weight peptide ACTH18-39.
SpiralTOF-TOF
Analysis of Phosphopeptide Using TOF-TOF

Introduction:
Phosphorylation is a type of post-translational modification of proteins that is used for the intracellular signal transduction in a wide range of living species. For this reason, it is very important to determine where the protein (amino acid) is phosphorylated. In this work, we measure a monophosphopeptide (FQ pS EEQQQTEDELQDK) that was obtained from the tryptic digestion of β-casein (Bovine) using the TOF-TOF option that is available for the JEOL SpiralTOF™ system.

Experimental:
The monophosphopeptide sample was dissolved in water containing 0.1% trifluoroacetic acid at a concentration of 10 pmol/μL. The matrix for this analysis was made by mixing 150 μL of CHCA (Methanol) at a concentration 30 mg/mL with 35 mg of 3-Aminoquinoline. Next, the monophosphopeptide sample solution and matrix solution were mixed together 1:1 by volume. Afterwards, 0.5 μL of this mixture was placed on the MALDI target plate (2.5 pmol/spot). Finally, the sample spot was measured using the TOF-TOF option available on the JMS-S3000 SpiralTOF MS system.

Results & discussion:
The MALDI-TOF mass spectrum of the monophosphopeptide is shown in Figure 1. The peak corresponding to the protonated molecule for this compound was observed at m/z 2061.8. The monoisotopic peak was then selected for high-energy CID TOF-TOF analysis. The resulting product-ion mass spectrum for this protonated molecule is shown in Figure 2. Because Lysine is present at the C-terminal for basic amino acids, we were able to observe the y-ion series generated by cleavage of the main-chain peptide bonds. Additionally, the w-ion series generated by elimination of the side-chains from the corresponding y-ions were observed.

Generally, the identification of the phosphorylation position using low-energy CID is considered very difficult because the phosphate group is lost before the cleavage of the main-chain peptide bond [1]. In contrast, high-energy CID does provide information about the location of phosphorylated amino acids because the y- and w-ion series were observed without the loss of the phosphate group (Figure 2) [2]. In Figure 2, the mass difference of the y13-ion and the y14-ion is 167u, which shows the position of the phosphorylated serine.

![Figure 1. Mass spectrum of the monophosphopeptide.](image-url)
Figure 2. High-energy CID product ion spectrum of the monophosphopeptide.

Conclusions:
As this work shows, we can identify the location of phosphorylated amino acids in a peptide by using the high-energy CID TOF-TOF option available for the JMS-S3000 SpiralTOF system. Furthermore, this data shows the possibility of using this instrument for the primary structural analysis of peptides, particularly for de novo sequencing.

References:
**SpiralTOF-TOF**  
Distinguishing Lysine and Glutamine in a Peptide

**Introduction:**
Lysine and glutamine are not easily distinguished by the most common approaches to peptide sequencing which involve mass spectrometers with low to moderate resolving power and low-energy collision-induced dissociation (CID). Lysine (C6H14N2O2 with a mass of 146.1055 u) and glutamine (C5H10N2O3 with a mass of 146.0691) differ by only 0.036 u. In this study, we demonstrate the measurement of a mixture of Substance P and a synthesized peptide (3-Gln ) with glutamine substituted for lysine in the Substance P sequence. Because the mass difference between Substance P and 3-Gln is 0.036 u, a resolving power of greater than 37,000 is required to separate each peptide. Additionally, we show that the TOF-TOF mode can be used to distinguish lysine and glutamine in these peptides by comparing the peak area ratio between a ions and d ions in the high-energy CID mass spectra.

**Experimental:**
Substance P standard was obtained from Sigma-Aldrich. The 3-Gln sample was synthesized and then provided by Hayashi Kasei.

1. Substance P, Arg-Pro-Lys-Pro-Gln-Gln-Phe-Phe-Leu-Met (Sigma-Aldrich)
2. 3-Gln, Arg-Pro-Gln-Pro-Gln-Gln-Phe-Phe-Leu-Met (Hayashi Kasei)

The peptide standard samples were dissolved in water containing 0.1% trifluoroacetic acid (TFA) at a concentration of 10 pmol/μL. α-Cyano-4-hydroxycinnamic acid (CHCA) was used as the matrix and was dissolved in 1:1 water/acetoneitrile containing 0.1% TFA. Next, the peptide standard solution and CHCA solution were mixed together 1:1 by volume. Afterwards, 0.5 μL of this mixture was placed on the MALDI target plate (2.5 pmol/spot). Finally, the dried sample was measured using the SpiralTOF and TOF-TOF option available on the JMS-S3000 Spiral-TOF MS system.

**Results:**

1. **SpiralTOF measurement**
The MALDI mass spectra are shown in Figure 1, and expanded mass spectra are shown in Figure 2 for the monoisotopic ion of the protonated molecules [M + H]+. The mass resolving power was approximately 60,000 for each of the [M + H]+ peaks in both substance P and 3-Gln mass spectra. Excellent mass accuracy (less than 10 ppm) was obtained with external calibration in all mass spectra. Additionally, we demonstrated that the high resolving power of the SpiralTOF could separate both components in the mixture.

2. **TOF-TOF measurement**
The product-ion mass spectra are shown in Figure 3. In both product-ion spectra, the characteristic a ions and d ions were observed in the high-energy CID. We cannot distinguish lysine and glutamine by mass accuracy and mass resolution in the product-ion mass spectra measured in TOF-TOF mode. However, we can identify lysine and glutamine by examining the a3/d3 ion intensities ratio [1]. The d3 ion intensity is higher than a3 ion for glutamine due to the stable ejected radical (CH2=CH2CH2NH2). Conversely, the d3 ion intensity is lower than a3 ion for lysine due to the unstable ejected radical (CH3CH=CHNH2) [2]. This information clearly identifies the presence of lysine in Substance P and glutamine in 3-Gln, respectively, as the third amino acid in each peptide sequence.
Figure 1. MALDI mass spectra of peptides: (a) Substance P, (b) 3-Gln, (c) Mixture

Figure 2. Expanded MALDI mass spectra showing the [M + H]+ monoisotopic ions for (a) Substance P, (b) 3-Gln, and the (c) mixture.

(a) Substance P
C$_{63}$H$_{99}$N$_{18}$O$_{13}$S
Exact m/z: 1347.73542
Mass error: -3.6 ppm

(b) 3-Gln
C$_{62}$H$_{95}$N$_{18}$O$_{14}$S
Exact m/z: 1347.69904
Mass error: 4.3 ppm

(c) Mixture
3-Gln
Mass error: -6.9 ppm
Substance P
Mass error: -8.8 ppm
Conclusion:
In this study we demonstrated that the JEOL SpiralTOF MALDI mass spectrometer can distinguish lysine and glutamine in a peptide.

SpiralTOF mode provides
- Ultra high resolving power sufficient to separate isobaric ions (0.036 u in this study)
- Excellent mass accuracy with external calibration (less than 10 ppm)

TOF-TOF mode provides
- High-energy (20 kV) CID
- Lysine and glutamine can be distinguished by examining the a3/d3 peak area ratio differences

Acknowledgement:
These data were obtained through collaborative research between Mass Spectrometry Group, Department of Physics, Graduate School of Science, Osaka University and JEOL Ltd.

Reference:
This page intentionally left blank for layout purpose.
**SpiralTOF-TOF**

Comparison of the JMS-S3000 SpiralTOF-TOF and a 4-Sector Tandem Double-Focusing Mass Spectrometer

**Introduction**

The JMS-3000 SpiralTOF™ has an optional TOF-TOF mode that features monoisotopic precursor ion selectivity, elimination of post source decay (PSD) ions, and high energy collision induced dissociation (CID). The JMS-700T MStation four-sector tandem double focusing mass spectrometer, although featuring a different analyzer and ionization techniques, has similar capabilities that have been previously used for a wide variety of applications including the structural analysis of complex biological molecules.

In this work, we compare the SpiralTOF-TOF with the MStation four sector tandem double focusing mass spectrometer using the same sample for MS/MS analysis.

**Experimental**

The sample analyzed by both systems was renin substrate tetradecapeptide (porcine), a peptide having an amino-acid sequence of DRVYHFPFILLVYS. Table 1 shows the measurement conditions and sample sizes used for both systems.

<table>
<thead>
<tr>
<th></th>
<th>JMS-S3000</th>
<th>JMS-700T</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ionization method</td>
<td>MALDI</td>
<td>FAB (Xe 6 kV)</td>
</tr>
<tr>
<td>Sample quantity</td>
<td>2.5 pmol</td>
<td>1 nmol</td>
</tr>
<tr>
<td>Precursor ion</td>
<td>mono isotopic</td>
<td>mono isotopic</td>
</tr>
<tr>
<td>Target gas</td>
<td>He</td>
<td>Ar</td>
</tr>
<tr>
<td>Matrix</td>
<td>CHCA</td>
<td>Glycerin, NBA</td>
</tr>
<tr>
<td>Collision energy</td>
<td>20000 eV</td>
<td>5000 eV</td>
</tr>
</tbody>
</table>

Table 1. Experimental conditions and specifications.

**Results and Discussion**

The SpiralTOF ion optics with the TOF-TOF option are shown in Fig. 1a. This design allows for the acquisition of high-energy CID product ion spectra for monoisotopically selected precursor ions. This precursor selection is made possible by the fact that the distance to the ion gate is 15 m. Additionally, the second TOF incorporates a re-acceleration mechanism into an offset parabolic reflectron (OPR) which enables the seamless observation of product ions ranging from very low m/z up to that of the precursor ion.

![Diagram of ion trajectory](image)

Figure 1. Ion trajectory - Left: JMS-S3000 SpiralTOF, Right: JMS-700T tandem MStation.
The MStation tandem has the ion trajectory shown in Fig. 1b, which also allows the selection of ions that have a specific \( m/z \) but in this case using the first magnetic and electric fields as the MS1 analyzer. The selected ions then pass into the collision chamber which is floated at a given voltage level to reduce their overall speed before colliding with the collision gas. Subsequently, the resulting product ions are re-accelerated to give them sufficient kinetic energy to be analyzed efficiently by the second magnetic and electric fields.

For comparison, the renin substrate tetradecapeptide (porcine) was analyzed using both systems. Fig. 2 shows the product ion spectra acquired for each system using the [M+H]+ \( (m/z \ 1758.93) \) as the precursor ion. Because of the difference between the FAB (Fast Atom Bombardment) and MALDI (Matrix Assisted Laser Desorption/Ionization) ionization techniques, the SpiralTOF-TOF acquired the product ion spectrum using 1/100 less volume than was used to get a comparable spectrum for the MStation. The resulting spectra show the immonium ions and the same pattern of a- and d-series ions which are the characteristic product ions produced by high energy CID. For the product ion spectrum acquired with the MStation in Fig. 2, the floating voltage was set to a slightly high level of 5000 V in order to observe low mass ions efficiently. However, this actually reduced the resolution and transmission rate of the high mass ions. In contrast, the SpiralTOF was able to analyze the full mass range efficiently, as a result of the second accelerator and the OPR not requiring the process of voltage selection for the target mass range.

**Conclusions**

These results clearly demonstrate that the SpiralTOF-TOF produced the same structural information as a traditional four sector tandem double focusing mass spectrometer. Additionally, the SpiralTOF-TOF required a smaller sample volume and a simpler analysis procedure to produce comparable product ion spectra.

---

Figure 2. Product ion spectra of Renin-substrate tetradecapeptide by a) JMS-S3000 SpiralTOF-TOF and b) JMS-700T tandem MStation.
SpirealTOF™

High Sensitivity Peptide Measurement with the New Matrix α-Cyano-4-Chlorocinnamic Acid

Introduction:
Matrix assisted laser desorption ionization (MALDI) is a powerful and useful ionization technique that is commonly used for the analysis of biomolecules such as peptides and proteins. Typically, α-Cyano-4-hydroxycinnamic acid (CHCA) is the matrix used for MALDI peptide measurement. Recently, a new matrix “α-Cyano-4-chlorocinnamic acid (CCICA)” was investigated for peptide analysis [1]. In this study, we demonstrate the measurement of a BSA digest to evaluate the improvement in peptide sensitivity with CCICA in comparison with CHCA by using the JMS-S3000 SpirealTOF MS system.

Experimental:
Sample information and preparation conditions are shown in Table 1. The BSA digest was obtained from Michrom. The CHCA matrix and peptide standard were obtained from Sigma-Aldrich, and CCICA matrix was obtained from Cayman.

The BSA digest standard samples were dissolved in water containing 0.1% trifluoroacetic acid (TFA). CHCA was dissolved in 1:1 water/acetoniitrile (ACN) containing 0.1% TFA. And CCICA was dissolved in 1:4 water/ACN containing 0.1% TFA. Next, the BSA digest standard solution and matrix solution were mixed together 1:1 by volume. Afterwards, 0.5 μL of this mixture was placed on the MALDI target plate. Finally, the dried sample was measured using the JMS-S3000 SpirealTOF MS system.

Results:
(1) Mass resolving power and mass accuracy
The comparison of mass resolving power and isotopic pattern for three of the peptide standards (Bradykinin1-7, Angiotensin II and ACTH18-39) with CHCA and CCICA are shown in Figure 1. The data for each matrix showed similar isotopic patterns and high mass-resolving power. The mass resolving power was approximately 23,000 at m/z 757.4 for Bradykinin1-7, over 30,000 at m/z 1046.5 for Angiotensin II and 73,000 at m/z 2466.2 for ACTH18-39. A residual plot for the calibration curves are shown in Figure 2. We used six peptides (Bradykinin1-7, Angiotensin II, Angiotensin I, P14R, ACTH1-17 and ACTH18-39) for the calibration. The Root Mean Squares (RMS) mass error of the actual measured m/z value against the calibration curve was just less than 0.4 ppm with both matrices. We achieved high mass resolving power and good mass calibration using the new matrix CCICA with SpirealTOF.

(2) BSA digest measurements
Next, we examined peptide sensitivity using the tryptic digest of BSA with both matrices. MALDI mass spectra of BSA digests with CCICA are shown in Figure 3. And MALDI mass spectra of BSA digests with CHCA are shown in Figure 4. There were a number of peptide peaks observed in the CCICA 250 amol/spot that had better signal-to-noise ratio than the peptide peaks observed in the CHCA 250 amol/spot (Figure 5). Additionally, the peptide mass fingerprint analysis using the peak list from CCICA 250 amol/spot mass spectrum resulted in a higher score and better coverage for the MASCOT (Matrix Science, Ltd.) search results (Figure 6). The protein was identified as BSA for this sample.

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Conc.</th>
<th>Solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>BSA digest</td>
<td>100 fmol/μL</td>
<td>0.1% TFA</td>
</tr>
<tr>
<td></td>
<td>10 fmol/μL</td>
<td>0.1% TFA</td>
</tr>
<tr>
<td></td>
<td>1 fmol/μL</td>
<td>0.1% TFA</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Matrix</th>
<th>Conc.</th>
<th>Solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHCA</td>
<td>10 mg/mL</td>
<td>ACN/0.1% TFA = 1/1 (v/v)</td>
</tr>
<tr>
<td>CCICA</td>
<td>10 mg/mL</td>
<td>ACN/0.1% TFA = 4/1 (v/v)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sample</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>BSA digest/Matrix = 1/1 (v/v)</td>
<td></td>
</tr>
<tr>
<td>0.5 μL of this sample solution mixture was placed on the MALDI target plate</td>
<td></td>
</tr>
</tbody>
</table>

Table 1. Sample information and preparation conditions.
Figure 1. The comparison of mass resolving power and isotopic pattern:
(a1) Bradykinin1-7 with CCICA, (a2) Angiotensin II with CCICA and (a3) ACTH18-39 with CCICA,
(b1) Bradykinin1-7 with CHCA, (b2) Angiotensin II with CHCA and (b3) ACTH18-39 with CHCA.

Figure 2. The residual plot for the calibration curves obtained by using six-peptide mixture:
(a) CCICA and (b) CHCA.
Figure 3. MALDI mass spectra of BSA digests with CCICA: (a) 25 fmol/spot, (b) 2.5 fmol/spot, (c) 250 amol/spot.

Figure 4. MALDI mass spectra of BSA digests with CHCA: (a) 25 fmol/spot, (b) 2.5 fmol/spot, (c) 250 amol/spot.
Figure 5. Comparison of BSA digests mass spectra each 250 amol/spot: (a) CCICA, (b) CHCA.

Conclusion:
We have done an initial study in which the new matrix “α-cyano-4-chlorocinnamic acid (CCICA)” was coupled with the SpiralTOF. The CCICA matrix provides high mass resolving power and excellent mass calibration curves that are comparable to the CHCA matrix. Additionally, the CCICA matrix improves the sensitivity of peptides. This new matrix will be a powerful and useful tool for the analysis of peptides and proteins.

Reference:

Figure 6. MASCOT search result using 250 amol/spot of CCICA.
SpiralTOF™
High Sensitivity Analysis of Intact Proteins Using Linear TOF

Introduction:
The JMS-S3000 SpiralTOF has a unique 17m flight path that offers the highest resolution MALDI-TOF MS system currently available. However, ions with a very short lifetime or that undergo spontaneous dissociation during their flight cannot be detected by the SpiralTOF (or a conventional reflectron TOF). To address this situation, the SpiralTOF with Linear TOF option can be used for the high sensitivity analysis of intact proteins.

In this work, we demonstrate the measurement of intact proteins by using the Linear TOF option for the JEOL SpiralTOF system.

Experimental:
Protein standards were obtained from Sigma-Aldrich.

a. Pepsin (porcine gastric mucosa), P-6887
b. Albumin (bovine serum), A8471
c. Conalbumin (chicken egg white), C0755
d. IgG (bovine serum), 15506

The protein standard samples were dissolved in water at a concentration of 1 pmol/μL. Sinapinic acid (SA) was used as the matrix and was dissolved in 1:1 water/acetonitrile containing 0.1% trifluoroacetic acid. Next, the protein standard solution and SA solution were mixed together 1:1 by volume. Afterwards, 0.5μL of this mixture was placed on the MALDI target plate (250 fmol/spot). Finally, the dried sample was measured using the Linear TOF option available on the JMS-S3000 SpiralTOF MS system.

Figure 1. SpiralTOF with Linear TOF option.
Results & discussion:

The MALDI mass spectra are shown in Figure 2 for each protein sample. Peaks corresponding to single- and double-charge protonated molecules were observed at the expected m/z values for the primary structure of these proteins.

Figure 2. MALDI mass spectrum of protein standard, (a) Pepsin (250 fmol/spot), (b) Albumin (250 fmol/spot) (c) Conalbumin (250 fmol/spot), (d) IgG (250 fmol/spot).

The SpiralTOF with Linear TOF option provided:

- Good peak shape in the high m/z region
- Excellent signal-to-noise ratio for these samples at this concentration (250 fmol)

High-quality mass spectra were achieved for intact proteins by using the JEOL JMS-S3000 system. Also worth noting, the Linear TOF can measure intact molecules up to m/z 500,000 (data not shown in these examples).
SpiralTOF™

MALDI-ISD Measurements Using Both the SpiralTOF Mode and the LinearTOF Mode

Introduction
Matrix assisted laser desorption/ionization (MALDI) combined with in-source decay (ISD) is a useful tool for doing top-down sequencing of intact proteins. In this work, we measured and compared the ISD fragment ions generated for several proteins by using both the high resolution MALDI-Spiral mode and the high sensitivity MALDI-Linear mode available on the JEOL SpiralTOF MALDI-MS system.

Experimental
Myoglobin and Bovine serum albumin (BSA) protein samples were separately dissolved into 0.1% trifluoroacetic acid aqueous with the concentration fixed at 10 pmol/μL. 1,5-diaminonaphthalene (DAN), which can provide good S/N for ISD fragment ions [1], was used as the MALDI matrix. The DAN matrix was dissolved to 0.1% trifluoroacetic acid aqueous/ 50% acetonitrile with the matrix concentration fixed at 10 mg/mL. Subsequently, the matrix and the sample solutions were mixed 1/1 (v/v), and then 1 μL of each solution was deposited and dried on the MALDI target plate. Afterwards, each sample was analyzed in triplicate on the JEOL JMS-3000 SpiralTOF by using both the SpiralTOF mode and the LinearTOF mode.

Results
The Myoglobin and BSA ISD mass spectra for both the LinearTOF and SpiralTOF measurements are shown in Figures 1 and 2, respectively. Both sets of ISD spectra were dominated by the c-ion series. The Linear mode mass spectra showed higher sensitivity overall, especially for ions over m/z 5,000 (shown in the m/z 4000-7000 zoomed regions). However, as expected, the Linear TOF did not provide full isotopic separation of these c-ions (see insets for the myoglobin c35 and BSA c33 ions). Conversely, the Spiral mode measurements fully resolved these ions and their associated isotopes, as shown in the Figures 1b and 2b insets, respectively, but had less sensitivity at higher m/z values.

![ISD Spectra of myoglobin (MW: 16,952Da) using (a) LinearTOF mode, (b) SpiralTOF mode.](image-url)
A MASCOT (Matrix Science, Ltd.) MS/MS Database Search using the LinearTOF data identified each sample as myoglobin and BSA, respectively. As an example, Figure 3 shows the Mascot database search result for the myoglobin sample. These results showed that, despite the fact that the ISD Linear data does not contain exact masses and does not provide isotopic separation of the ions, the data can still be used with a database search function like MASCOT to identify proteins.

**Conclusion**

In this work, we showed a brief study in which the ISD measurements for standard proteins were measured by using the SpiralTOF mode and the LinearTOF mode. The SpiralTOF mode provides high mass accuracy and fully separated isotopic ions while the LinearTOF mode provides higher sensitivity, particularly for ions in the higher m/z region. Additionally, the LinearTOF data can be used in conjunction with a database search analysis to identify proteins.

**Reference**

SpiralTOF™
High Resolution and High Mass Accuracy MALDI-ISD Measurements

Introduction
Matrix assisted laser desorption/ionization (MALDI) combined with in-source decay (ISD) is a useful tool for doing top-down sequencing of intact proteins. This technique can provide enough information to determine both N- and C-terminal sequences. In this work, we measured the ISD fragment ions generated for several peptides using the JEOL SpiralTOF MALDI-MS system.

Experimental
ACTH18-39 and oxidized Insulin B chain peptide samples were separately dissolved in 0.1% trifluoroacetic acid aqueous with the concentration fixed at 10 pmol/μL. 1,5-diaminonaphthalene (DAN), which can provide good S/N for ISD fragment ions [1], was used as the MALDI matrix. The DAN matrix was dissolved to 0.1% trifluoroacetic acid aqueous/50% acetonitrile with the matrix concentration fixed at 10 mg/mL. Subsequently, the matrix and sample solutions were mixed 1/1 (v/v), and then 1 μL of each solution was deposited and dried on the MALDI target plate. Afterwards, each sample was analyzed in triplicate on the JEOL JMS-3000 SpiralTOF MALDI-MS system.

Results
As a starting point, MALDI imaging was used to visualize the location of the sample ion distributions within the DAN crystal matrix. A comparison between an optical microscope image and the MALDI image for the oxidized Insulin B chain y10 fragment ion (m/z 1215.6) are shown in Figure 1. These results clearly showed that the ISD fragment ions were concentrated along the edges of the DAN matrix crystals. As a result, the MALDI laser irradiation for each sample spot was focused on the crystal edges in order to obtain sufficient sensitivity for the ISD measurements.

Afterwards, the ISD mass spectra were measured for each peptide, and the results are shown in Figure 2. The ACTH18-39 peptide produced ISD spectra that mostly consisted of the c-ion series. However, for the oxidized Insulin B chain, y-ion series dominated the ISD spectra. It should be noted here that when the basic amino acids are located on the N-terminal side, c-ion series are the dominant ions generated during ISD. However, when the basic amino acids are located on the C-terminal side, the y-ion series are the dominant ions generated during ISD [2]. These results highlight the difference in location for the basic amino acids for each of the peptide chains tested.

Next, each ISD fragment ion series was evaluated for mass accuracy using an external calibration. These analyses showed an average RMS of 4.0 ppm for the ACTH18-39 c ion series and an average RMS of 4.7 ppm for the y-ion series produced from the oxidized Insulin B chain, as shown in Tables 1 and 2, respectively.

Figure 1. (a) Optical microscope image of DAN crystal, (b) MALDI imaging of m/z 1215.6 (y10 ISD fragment ion of oxidized Insulin B chain (width 3.0mm x height 3.0mm, interval 0.05mm).
Figure 2. ISD mass spectra for (a) ACTH18-39, (b) oxidized Insulin B chain.

Table 1. Accurate mass measurement results for c-ion series of ACTH18-39.

<table>
<thead>
<tr>
<th>Ion species</th>
<th>Calc. m/z</th>
<th>Data 1 (ppm)</th>
<th>Data 2 (ppm)</th>
<th>Data 3 (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>c7</td>
<td>857.5356</td>
<td>3.73</td>
<td>13.18</td>
<td>2.92</td>
</tr>
<tr>
<td>c8</td>
<td>971.5785</td>
<td>1.96</td>
<td>12.87</td>
<td>3.50</td>
</tr>
<tr>
<td>c9</td>
<td>1028.6000</td>
<td>0.49</td>
<td>11.76</td>
<td>3.01</td>
</tr>
<tr>
<td>c10</td>
<td>1099.6371</td>
<td>0.36</td>
<td>11.55</td>
<td>2.46</td>
</tr>
<tr>
<td>c11</td>
<td>1228.6797</td>
<td>1.06</td>
<td>10.09</td>
<td>2.28</td>
</tr>
<tr>
<td>c12</td>
<td>1343.7066</td>
<td>0.52</td>
<td>7.59</td>
<td>0.52</td>
</tr>
<tr>
<td>c13</td>
<td>1472.7492</td>
<td>0.95</td>
<td>6.52</td>
<td>0.88</td>
</tr>
<tr>
<td>c14</td>
<td>1559.7812</td>
<td>0.26</td>
<td>4.62</td>
<td>1.15</td>
</tr>
<tr>
<td>c15</td>
<td>1630.8184</td>
<td>0.06</td>
<td>3.07</td>
<td>1.59</td>
</tr>
<tr>
<td>c16</td>
<td>1759.8610</td>
<td>1.88</td>
<td>1.99</td>
<td>2.84</td>
</tr>
<tr>
<td>c17</td>
<td>1830.8981</td>
<td>0.66</td>
<td>0.44</td>
<td>0.64</td>
</tr>
<tr>
<td>c18</td>
<td>1977.9665</td>
<td>6.77</td>
<td>3.08</td>
<td>6.32</td>
</tr>
<tr>
<td>c19</td>
<td>2075.0192</td>
<td>1.11</td>
<td>0.82</td>
<td>5.73</td>
</tr>
<tr>
<td>c20</td>
<td>2188.1033</td>
<td>5.12</td>
<td>4.30</td>
<td>8.91</td>
</tr>
<tr>
<td>c21</td>
<td>2317.1459</td>
<td>4.19</td>
<td>3.88</td>
<td>8.03</td>
</tr>
<tr>
<td>Abs. Average</td>
<td>13.44</td>
<td>6.38</td>
<td>3.65</td>
<td></td>
</tr>
</tbody>
</table>

Table 2. Accurate mass measurement results for y-ion series of oxidized Insulin B chain.

<table>
<thead>
<tr>
<th>Ion species</th>
<th>Calc. m/z</th>
<th>Data 1 (ppm)</th>
<th>Data 2 (ppm)</th>
<th>Data 3 (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>y9</td>
<td>1086.7351</td>
<td>1.76</td>
<td>0.10</td>
<td>0.08</td>
</tr>
<tr>
<td>y10</td>
<td>1215.7651</td>
<td>0.76</td>
<td>0.48</td>
<td>1.22</td>
</tr>
<tr>
<td>y11</td>
<td>1272.3872</td>
<td>0.98</td>
<td>0.04</td>
<td>1.06</td>
</tr>
<tr>
<td>y12</td>
<td>1423.8311</td>
<td>11.51</td>
<td>11.51</td>
<td>9.12</td>
</tr>
<tr>
<td>y13</td>
<td>1522.6995</td>
<td>9.25</td>
<td>4.98</td>
<td>7.81</td>
</tr>
<tr>
<td>y14</td>
<td>1635.7836</td>
<td>4.19</td>
<td>5.41</td>
<td>3.82</td>
</tr>
<tr>
<td>y15</td>
<td>1788.8469</td>
<td>4.55</td>
<td>6.27</td>
<td>7.49</td>
</tr>
<tr>
<td>y16</td>
<td>1911.9310</td>
<td>8.29</td>
<td>4.89</td>
<td>9.81</td>
</tr>
<tr>
<td>y17</td>
<td>1982.9681</td>
<td>3.56</td>
<td>6.33</td>
<td>5.33</td>
</tr>
<tr>
<td>y18</td>
<td>2112.0107</td>
<td>2.30</td>
<td>6.37</td>
<td>3.29</td>
</tr>
<tr>
<td>y19</td>
<td>2211.0791</td>
<td>4.42</td>
<td>8.94</td>
<td>5.69</td>
</tr>
<tr>
<td>y20</td>
<td>2324.1631</td>
<td>3.97</td>
<td>5.31</td>
<td>8.92</td>
</tr>
<tr>
<td>y21</td>
<td>2461.2220</td>
<td>1.15</td>
<td>1.15</td>
<td>0.96</td>
</tr>
<tr>
<td>y22</td>
<td>2548.2541</td>
<td>2.77</td>
<td>4.42</td>
<td>3.44</td>
</tr>
<tr>
<td>y23</td>
<td>2605.2755</td>
<td>3.82</td>
<td>4.81</td>
<td>2.28</td>
</tr>
<tr>
<td>y24</td>
<td>2756.2695</td>
<td>5.68</td>
<td>5.43</td>
<td>2.96</td>
</tr>
<tr>
<td>y25</td>
<td>2869.3535</td>
<td>4.47</td>
<td>8.76</td>
<td>2.87</td>
</tr>
<tr>
<td>y26</td>
<td>3006.4124</td>
<td>5.47</td>
<td>0.55</td>
<td>4.40</td>
</tr>
<tr>
<td>y27</td>
<td>3134.4710</td>
<td>6.13</td>
<td>13.47</td>
<td>16.65</td>
</tr>
<tr>
<td>y28</td>
<td>3248.5139</td>
<td>1.40</td>
<td>16.76</td>
<td>13.22</td>
</tr>
<tr>
<td>y29</td>
<td>3347.5824</td>
<td>3.33</td>
<td>6.83</td>
<td>4.04</td>
</tr>
<tr>
<td>Abs. Average</td>
<td>4.37</td>
<td>5.10</td>
<td>4.55</td>
<td></td>
</tr>
</tbody>
</table>
Conclusion

In this work, we showed a brief study in which the ISD measurements for standard peptides were measured by using the SpiralTOF. It should be noted that ISD measurements generally require a relatively high laser power to generate ISD fragment ions which in turn detrimentally impacts the mass resolving power of the instrument. As a result, it is generally very difficult to achieve high mass resolving power using a high laser power. However, the Spiral trajectory ion optics provides a 17m flight path that overcame this problem to produce resolving powers of 20,000-35,000 for each of the measured ions. Furthermore, as shown with these peptide sequences, the system provided excellent mass accuracy even when an external calibration was used for the data.

Reference

This page intentionally left blank for layout purpose.
SpiralTOF-TOF
Analysis of the Natural Organic Compound SAAF by Using the TOF-TOF Option

Introduction
Sperm activating and attracting factor (SAAF) is a naturally occurring substance produced by ascidians (sea squirts) to promote the fertilization process. Previously, the structure of this compound was characterized by high-energy collision induced dissociation (CID) using a traditional four sector tandem double focusing mass spectrometer. In this work, we analyzed the SAAF from a ciona intestinalis by using a JMS-S3000 SpiralTOF™ equipped with the optional TOF-TOF analyzer to generate a high energy CID product ion spectrum for this compound.

Results and Discussion
Fig. 1 shows the SAAF mass spectrum acquired using PEG sulfate 600 as an external calibrant. The spectrum shows a peak at m/z 515.301, which is suspected to be the monoisotopic ion [M-2Na-SO₄₊H⁺] (calculated value 515.305). In addition, peaks from SAAF are observed at two other masses: [M-2Na+H⁺] at m/z 595.255 (calculated value 595.261) and [M-Na] at m/z 617.237 (calculated value 617.242). Among these peaks, the product ion spectrum was acquired for the [M-2Na-SO₄₊H⁺] ion, which had been previously analyzed using a traditional four sector tandem double focusing mass spectrometer. Fig. 2 shows the high energy CID results generated by the SpiralTOF-TOF. The negative charge was fixed to the sulfate ester end of the molecule, causing charge remote fragmentation (CRF) to occur. As a result, the spectrum showed peaks that systematically reflected the structure of SAAF, as

![Figure 1. Mass spectrum of SAAF.](image1)

![Figure 2. Product ion spectrum of [M-2Na-SO₄₊H⁺].](image2)
shown in Fig. 3. This data closely resembles the results obtained using a traditional four sector MS/MS system.1

Conclusions
As this data demonstrates, high energy CID analysis by using SpiralTOF-TOF clearly detected the peaks resulting from CRF, which enabled the analysis of samples that are complex in structure, such as natural organic compounds like SAAF.

Acknowledgment
The analysis above was conducted in collaboration with the Mass Analysis Group, Department of Physics, Graduate School of Science, Osaka University. We wish to express our thanks to Prof. Michio Murata of the Department of Biomolecular Sciences, Department of Physics, Graduate School of Science, Osaka University, for providing the SAAF samples.

Reference

Figure 3. Structure of SAAF and fragmentation pattern. (Asterisks show dehydrated ions.)
SpiralTOF-TOF
Analysis of the Natural Organic Compound YTX by Using TOF-TOF Option

Introduction
Yessotoxin (YTX) is one of the substances that cause food poisoning when humans consume toxic shellfish. The structure of this compound has been previously analyzed by high-energy collision induced dissociation (CID) using a traditional four sector tandem double focusing mass spectrometer.[1]

In this work, we analyzed YTX by using a JMS-S3000 SpiralTOF™ equipped with the optional TOF-TOF analyzer to generate a high energy CID product ion spectrum.

Results and Discussion
Fig. 1 shows the YTX mass spectrum acquired using polyalanine as an external calibrant. The spectrum shows a peak at m/z 1061.513, which is suspected to be the monoisotopic ion [M-2Na-SO3+H]+ (calculated value 1061.609). There was also a [M-Na+] peak observed for YTX at m/z 1164.450 (calculated value 1163.548). Among these peaks, the product ion spectrum was acquired from [M-2Na-SO3+H]+, which had been previously analyzed using a traditional four sector tandem double focusing mass spectrometer.[1] Fig. 2 shows the high-energy CID results generated by the SpiralTOF-TOF. The negative charge was fixed to the sulfate ester end of the molecule, causing charge remote fragmentation (CRF) to occur. As a result, the spectrum showed peaks that systematically reflected the structure of YTX, as shown in Fig. 3. This data closely resembles the results obtained using a traditional four sector MS/MS system.[1]
Conclusions
As this data demonstrates, high energy CID analysis using SpiralTOF-TOF clearly detected the peaks resulting from CRF, Which enabled analysis of samples that are complex in structure such, as natural organic compounds.

Acknowledgement
We wish to express our thanks to Prof. Michio Murata of the Department of Biomolecular Sciences, Department of Physics, Graduate School of Science, Osaka University, for providing the YTX samples.

Reference
**SpiralTOF-TOF**

**High-energy CID Mass Spectrometry of Oligosaccharides**

**Introduction:**
Matrix assisted laser desorption ionization (MALDI) is a powerful and useful ionization technique that is commonly used for the analysis of biomolecules such as oligosaccharides. There are many applications of oligosaccharides in which various ionization techniques and mass spectrometers were used for their analysis [1]. In particular, tandem mass spectrometry techniques are often used to sequence these molecules.

Recently, JEOL developed a new tandem TOF-TOF instrument coupled with MALDI that is called the SpiralTOF. The 1st TOF consists of 4 toroidal electric sectors that fold a 17 meter flight path into a one meter box. This design provides several unique advantages for TOF-TOF analysis. The 2nd TOF has (a) 20 kV high-energy CID, (b) monoisotopic precursor ion selection, and (c) no PSD ions in the product ion mass spectrum.

In this study, we analyzed several oligosaccharides by using the JMS-S3000 SpiralTOF-TOF tandem mass spectrometer system.

**Experimental:**
All oligosaccharides (Laminaritetrose, Stachyose, α-Cyclodextrin, β-Cyclodextrin, γ-Cyclodextrin) were commercially available items that were used without further purification. Each oligosaccharide standard solution was dissolved in water. 2,5-Dihydroxybenzoic acid (DHB) was dissolved in 40% ethanol at a concentration of 10 mg/mL. Next, the oligosaccharides standard solution and matrix solution were mixed together 1:1 by volume. Afterwards, 0.5 µL of this mixture was placed on the MALDI target plate. Finally, the dried sample was measured using the JMS-S3000 SpiralTOF-TOF.

**Results:**
The A, B and C fragment ions for oligosaccharides are labeled as the non-reducing terminal ends while the X, Y and Z fragment ions are labeled as the reducing terminal ends. The nomenclature fragmentation pathway of oligosaccharides by tandem MS is shown in Figure 1 [modified from Reference 2].

In this study, we did not do further purification for all oligosaccharide standards, matrix and solvents. As a result, the sodiated molecules were the most intense peak in each positive-ion MALDI mass spectrum (see Figure 2). Therefore, these sodiated molecules were selected as the precursor ions for the SpiralTOF-TOF analysis. MALDI TOF-TOF spectra of Laminaritetrose and Stachyose are shown in Figure 3. Laminaritetrose is a tetrascacharide that consists of four β-D-glucose units that are linked as Glc(β1→3)Glc(β1→3)Glc(β1→3)Glc. Stachyose is also a tetrascacharide that consists of two α-D-galactose units, one α-D-glucose unit, and one β-D-fructose unit that are linked as Gal(α1→6)Gal(α1→6)Glc(α1→2β) Fru. These isomers showed significantly different mass spectral patterns. B and Y fragment ions were generated from the glycosidic bond cleavage were observed as the dominant components in each MALDI TOF-TOF spectrum. The fragmentation scheme for Laminaritetrose is shown in Figure 4. High-energy CID fragmentation occurs within a shorter time-scale than low-energy CID, which means that it provides a cross-ring cleavage that gives useful structural information about the oligosaccharides and their glycoconjugates. This cross-ring cleavage was also observed for both oligosaccharides, as indicated by the presence of the fragment ions m/z 599, 569, 555, etc. (see Figure 3). All of this information is essential for determining the structure of each oligosaccharide and differentiating structural isomers from each other.

Cyclodextrins are cyclic oligosaccharides consisting of five or more α-D-glucose units that are linked as Glc(α1→4)Glc. The MALDI TOF-TOF spectrum for each Cyclodextrin is shown in Figure 5. In each case, the B ion series from glycosidic bond cleavage were the dominant ions observed in the TOF-TOF spectra. Additionally, a number of fragment ions were observed that were the result of cross-ring cleavage.

![Figure 1. Fragmentation pathways.](www.jeol.com)
Figure 2. MALDI TOF spectra. A) Stachyose, B) \( \gamma \)-Cyclodextrin

Figure 3. MALDI TOF-TOF spectra. A) Laminaritetrose, B) Stachyose
Figure 4. The glycosidic bond cleavage to generate B and Y ion series for Laminaritetaose

![Diagram showing glycosidic bond cleavage]

Figure 5. MALDI TOF-TOF spectra. A) α-Cyclodextrin, B) β-Cyclodextrin, C) γ-Cyclodextrin

![Graphs showing MALDI TOF-TOF spectra]

Conclusion:
In this work, we showed a brief study in which the high-energy CID measurements for several oligosaccharides were measured by using the SpiralTOF-TOF. The B and Y ion series were observed as the main components in each TOF-TOF spectrum. Furthermore, we could also see a number of fragment ions that were generated from cross-ring cleavage, which was helpful in differentiating structural isomers. These results show that high-energy CID coupled with MALDI provides a good platform for determining oligosaccharide structural information that will not generally be available when analyzing these compounds by other mass spectrometry techniques.

Reference:
This page intentionally left blank for layout purpose.


佐藤貴弥. らせん状のイオン軌道を利用した飛行時間質量分析計の開発. ぶんせき, **688-691** (2009).


佐藤貴弥. らせん状のイオン軌道をもつタンデム飛行時間質量分析装置の開発と応用例. ぶんせき, **532-536** (2011).


Abe, T., Matsuzaka, M., Isoda, K. & Tadokoro, M. Liquid-Crystalline and Electronic Properties


This page intentionally left blank for layout purpose.