Fragmentation of organic ions bearing fixed multiple charges observed in MALDI MS

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Abstract
Matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI TOF MS) was used to analyze a series of synthetic organic ions bearing fixed multiple charges. Despite the multiple intrinsic charges, only singly charged ions were recorded in each case. In addition to the pseudo-molecular ions formed by counterion adduction, deprotonation and electron capture, a number of fragment ions were also observed. Charge splitting by fragmentation was found to be a viable route for charge reduction leading to the formation of the observed singly charged fragment ions. Unlike multivalent metal ions, organic ions can rearrange and/or fragment during charge reduction. This fragmentation process will evidently complicate the interpretation of the MALDI MS spectrum. Because MALDI MS is usually considered as a soft ionization technique, the fragment ion peaks can easily be erroneously interpreted as impurities. Therefore, the awareness and understanding of the underlying MALDI-induced fragmentation pathways is essential for a proper interpretation of the corresponding mass spectra. Due to the fragment ions generated during charge reduction, special care should be taken in the MALDI MS analysis of multiply charged ions. In this work, the possible mechanisms by which the organic ions bearing fixed multiple charges fragment are investigated. With an improved understanding of the fragmentation mechanisms, MALDI TOF MS should still be a useful technique for the characterization of organic ions with fixed multiple charges.

KEYWORDS
artificial impurities, false interpretation, fixed multiple charges, ion fragmentation, matrix-assisted laser desorption/ionization time-of-flight mass spectrometry

1 | INTRODUCTION

Matrix-assisted laser desorption/ionization mass spectrometry (MALDI MS) is a powerful and indispensable technique for the analysis of various biological and synthetic compounds.1-3 An important characteristic of MALDI is that analytes, with the exception of very high molecular weight molecules such as proteins, are predominantly recorded as singly charged ions.4,5 This feature facilitates interpretation of the resulting mass spectra. The uncomplicated spectra together with many other advantages, including relatively straightforward operation, high sensitivity, tolerance to contamination, and rapid analysis to provide absolute molecular mass, have also rendered MALDI MS a popular analytical technique for the characterization of nonvolatile synthetic organic compounds in many chemistry laboratories across the world.

Although singly charged ions are usually the most prominent species recorded for small molecules in MALDI MS, analytes can be multiply charged in a sample solution and this original charge state might be preserved in the sample spot after the sample is dried on a MALDI target plate.6 However, highly charged molecular ions can only be observed under special conditions.7-10 Upon laser irradiation, a series of events or reactions are initiated, which lead to the observed singly charged ions.4,5,11-16 According to Karas et al.4,5 singly charged ions are the “lucky survivors” in MALDI MS. Depending on the properties of the analyte and the matrix, a number of gas phase processes are responsible for the formation of singly charged ions. For analytes such as proteins/peptides that carry more than one basic group, multiply protonated ions accompanied by counter anions will be incorporated in typically acidic matrices. Proton neutralization within matrix clusters
or proton transfer reactions in the MALDI plume can effectively reduce the multiply charged protonated ions to singly charged ions.\textsuperscript{4,5,12,13} Interestingly, even for complexes with multivalent metal ions, singly charged ions are still the dominant species in MALDI MS. When multivalent metal ions are added as cation reagents in the MALDI MS analysis of synthetic apolar polymers like polystyrene, the multivalent ions will be reduced to singly charged ions by the capture of free electrons or by gas phase charge exchange with matrix molecules.\textsuperscript{17,18}

For polar synthetic polymers, on the other hand, the singly charged quasi-molecular ions can also be formed by deprotonation (e.g., [analyte + metal\textsuperscript{\textit{2+}} – H\textsuperscript{+}]\textsuperscript{19,20}). Schäfer and Budzikiewicz reported that deprotonation reactions and one-electron reduction are the predominant paths of transforming preformed doubly charged porphyrins to singly charged ions.\textsuperscript{21} Besides the deprotonation and electron capture reactions, another important route for charge reduction in MALDI is the attachment of counterions and/or deprotonated matrix ions.\textsuperscript{5,22}

In all the examples cited above, it could be argued that the ions recorded can still be considered as (pseudo)-molecular ions because no fragmentation or rearrangement of the molecular structure occurred. The resulted MALDI MS spectra are still easy to interpret. Charge reduction in these examples is fulfilled by deprotonation, electron capture, and/or counterion attachment.

In this contribution, we report that the situation of forming singly charged ions can be more complicated for organic ions bearing fixed multiple charges. For such multiply charged ions, fragmentation into a series of singly charged ions can also be a viable route in addition to the charge reduction reactions mentioned above. Furthermore, organic ions might concurrently fragment and/or rearrange after catching electrons, which is in clear contrast with the reduction of multivalent metal ions. Obviously, these reactions can give extra fragment peaks and thus significantly complicate the interpretation of the final MALDI mass spectra. As MALDI is usually considered as a soft ionization technique, the fragments might easily be falsely interpreted as impurities. With the widespread application of MALDI MS, it can be envisaged that many new types of multiply charged organic ions will be analyzed using this technique. Therefore, it is extremely important for analysts to be aware of the possible fragmentation in the MALDI MS analysis of organic ions with fixed multiple charges.

Four types of ionic organic compounds bearing fixed positive multiple charges were studied in this work, namely, a doubly charged poly(propylene imine) diaminobutane dendrimer,\textsuperscript{22} a doubly and a quadruply charged viologen derivatives, a doubly and a triply charged benzene-1,3,5-tricarboxamide derivatives,\textsuperscript{24,25} and a doubly charged perylene derivative (Figure 1). The results clearly indicate that fragmentation and rearrangement reactions can occur in the charge reduction of multiply charged organic ions in MALDI MS. Possible mechanisms for fragmentation and rearrangement of these compounds were investigated. The aim of this contribution is to stress that an awareness of possible fragmentation and the understanding of the fragmentation mechanisms are essential for proper interpretation of the mass spectra in the MALDI MS analysis of organic ions bearing fixed multiple charges.

## EXPERIMENTAL

The structures of all the analytes studied are given in Figure 1. The syntheses of QDendr1 and the 2 benzene-1,3,5-tricarboxamide (BTA) derivatives have been reported in previous publications by our group.\textsuperscript{23-25} Syntheses and nuclear magnetic resonance characterization of the 2 viologen derivatives and the perylene derivative are provided in the electronic supplement information (ESI).

Three matrices were used in this work. \(\alpha\)-Cyano-4-hydrocinnamic acid (CHCA) and tetrathiafulvalene (TTF) were purchased from Fluka (Zwijndrecht, The Netherlands), and 2-[(2E)-3-(4-tert-butylphenyl)-2-methylprop-2-enylidene] malononitril (DCTB) were purchased from Fluka (Zwijndrecht, The Netherlands).
was synthesized according to Ulmer et al. Tetrahydrofuran (THF, analytical reagent grade stabilized with butylated hydroxytoluene) was obtained from Biosolve (Biosolve BV, Valkenswaard, The Netherlands).

Matrix solutions were freshly prepared in THF at a concentration of approximately 20 mg/mL. All the sample solutions were also freshly prepared. Two percent (v/v) of water in THF was required to dissolve QDendr1. 4% (v/v) of water in THF to dissolve VP(F3)2 and BV(PF3)4, and 2% (v/v) dimethyl sulfoxide (DMSO) in THF to dissolve QPB. BTA2CySCl2 and BTA3CySCl2 were dissolved in methanol. A sample solution and a matrix solution were mixed in an Eppendorf tube. All mass spectra shown in this article were recorded with an analyte/matrix mole ratio of 1:100 since no considerable difference in the MALDI MS spectra concerning the fragmentation patterns was observed with the analyte/matrix mole ratios from 1:20 to 1:1000. A total of 0.5 µL of the mixed solution was pipetted onto a stainless steel MALDI target plate and allowed to dry.

The matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI TOF MS) measurements were performed with an Autoflex Speed (Bruker, Bremen, Germany) instrument equipped with a 355-nm Nd:YAG smartbeam laser with maximum repetition rate of 1000 Hz, capable of executing both linear and reflector modes. The accelerating voltage was held at 19 kV and the delay time at 130 nanoseconds for all experiments. Mass spectra were acquired in the reflector positive ion mode by summing spectra from 500 random laser shots at an acquisition rate of 100 Hz. The MS spectra were calibrated with Csl clusters of known masses.

3 RESULTS AND DISCUSSION

3.1 Quaternized dendrimer

Dendrimers are highly branched globular molecules, which emanate from a central core. Dendrimers have attracted significant scientific interest owing to their distinctive molecular structure. In our laboratory, a large number of poly(propylene imine) dendrimers were synthesized, and MS has been found to be a powerful tool for the analysis of these dendrimers. In this study, a first generation quaternized dendrimer (QDendr1, Figure 1) consisting of a doubly charged ion with 2 quaternized nitrogen atoms at the core and iodide as the counter ion was studied. The excellent matching between the ions recorded in Figure 2 and the expected ions, except H+ and CH3+, which might be too small to be detected in our system, could easily be found in Figure 2.

The excellent matching between the ions recorded in Figure 2 and the expected ions based on Scheme 1 strongly supports the fragmentation routes of Scheme 1. Although we could not detect [QDendr12+ + e+] ions, we do not rule out the possibility of charge splitting or electron capture and subsequent fragmentation. In the analysis of proteins and peptides, multiply charged ions can be fragmented by electron capture dissociation (ECD) leading to z and c ions.

The matrix-assisted laser desorption/ionization time-of-flight mass spectrometry of QDendr12+ using α-cyano-4-hydroxycinnamic acid (CHCA) as the matrix. The molecular structure of the analyte is shown in Figure 1 (QDendr12+ = C36H32N4O4). The assignment of the peaks can be found in Scheme 1 and Table S1. * indicates unidentified peaks. CHCA is dissolved in THF and the analyte in THF with 2% of water. The mole ratio of analyte/CHCA is 1:100 simply catching one electron in the same way as multivalent metal ions, and hence no [QDendr12+ + e+] was recorded either. Although no [QDendr12+ + e+] was observed, it will be shown below that singly charged molecular ions formed on electron capture were recorded for analytes if the molecular ions could be sufficiently stabilized via π-π conjugation which is lacking for QDendr1. Based on the above reasoning and the m/z values of the extra peaks, we propose that the corresponding ions were formed through charge loss via fragmentation.

Considering the molecular structure of QDendr12+, fragmentation is expected to occur around the 2 charged N-atoms in the core. Each charged N-atom forms 4 N–C bonds, 2 of which carry identical substituent groups (Figure 1). Hence, 3 hypothetical cleavage mechanisms are envisaged (see Scheme 1). In routes 1 and 2, a doubly charged ion is split into 2 singly charged ions. In route 3, the cleavage first yields another doubly charged fragment which is further reduced to singly charged ions by charge-charge association with an anion, cleavage of one of the N–C bonds, or by losing either a single H+ or CH3+ fragment. The molecular formula and molecular weights of the possible fragment ions predicted according to the routes shown in Scheme 1 are listed in Table S1 of the ESI. Convincingly, all the expected fragment ions, except H+ and CH3+, which might be too small to be detected in our system, could easily be found in Figure 2.

The excellent matching between the ions recorded in Figure 2 and the expected ions based on Scheme 1 strongly supports the fragmentation routes of Scheme 1. Although we could not detect [QDendr12+ + e+] ions, we do not rule out the possibility of charge reduction by electron capture and subsequent fragmentation. In the analysis of proteins and peptides, multiply charged ions can be fragmented by electron capture dissociation (ECD) leading to z and c ions. It might also be possible that the recombination energy between QDendr12+ and electron would be redistributed throughout the reduced product to induce fragmentation. Therefore, it is very well possible that both ECD and fragmentation via charge splitting are important routes for the formation of the QDendr1 fragment ions.
Based on the discussion above, it is evident that Qdendr12+ could generate a number of fragment ions in MALDI. The fragmentation of Qdendr12+ is not unexpected because of the small molecular size and the lack of easily cleavable protons. The driving force for the fragmentation is the tendency to form singly charged ions in MALDI. Because of the fragmentation, additional care should be taken for the interpretation of the MALDI mass spectra for which a good understanding of the fragmentation pathways as illustrated in Scheme 1 is vital.

3.2 | Viologen derivatives

Viologens are bipyridium derivatives of 4,4′-bipyridyl which are widely used for electrochromic systems because of their ability to reversibly change color under redox conditions. In addition, there are significant recent interests in the synthesis and application of viologen derivatives in supramolecular chemistry and material science.31,32 In this work, 2 viologen derivatives were studied and their structures are shown in Figure 1. V(PF6)2 consists of one dicationic core of viologen, while BV(PF6)4 consists of 2 dicationic cores.

Figure 3 shows a MALDI TOF MS spectrum of V(PF6)2 using DCTB as the matrix. With DCTB, major signals of singly charged ions corresponding to V2+ with and without a single PF6− counter ion were recorded. By contrast, no [V2+ + PF6−]1+ ions were detected using CHCA as matrix (see ESI, Figure S4), probably because CHCA is a “hot matrix” and [V2+ + PF6−]1+ cannot survive in MALDI under these experimental conditions. Although V2+ is doubly charged, it is readily reduced to a singly charged cation radical [V2+ + e]1+ through capture of a single electron owing to the highly electron-deficient core.33 At first glance, it would appear as if V2+ is reduced by simple one-electron reduction, in a similar way to multiply charged metal ions. However, the actual mechanisms for charge reduction of multivalent metal ions and of multivalent organic ions are very much different. For a multivalent metal ion, the charges are confined to one metal atom. By contrast, the 2 charges of a viologen core are shared among several atoms of the 2 pyridine rings via π-conjugation. On electron capture, the viologen core rearranges as shown in Scheme 2.33,34 Depending
on the experimental conditions, the odd-electron radical ion of \([V^{2+} + e]^-\) can further lose one \(H\) forming \([V^{2+} + e - H]^-\). A small but clear peak of the \([V^{2+} + e - H]^-\) fragment was observed with either DCTB (see the red inset of Figure 3) or CHCA matrix (ESI, Figure S4).

The MALDI TOF MS spectrum of \(\text{BV(PF}_6\text{)}_4\) using DCTB as the matrix is shown in Figure 4. \(\text{BV}^{4+}\) has 2 highly electron-deficient dicationic cores of viologen. With DCTB matrix, 4 pseudo-molecular ion peaks with mass difference of 145 Dalton were observed which can easily be assigned to \(\text{BV}^{4+}\) ions with different numbers of \((\text{PF}_6^-)\) counter ions. However, no peaks of \([\text{BV}^{4+} + 3(\text{PF}_6^-)]^+\) and only tiny peaks of \([\text{BV}^{4+} + e + 2(\text{PF}_6^-)]^+\) were observed with CHCA matrix (ESI, Figure S5). Once again, the absence of \([\text{BV}^{4+} + 3(\text{PF}_6^-)]^+\) is probably because of the use of the hot CHCA matrix, which disfavors its survival. In forming the singly charged ions, different numbers of electrons must be involved for charge reduction. After electron capture, the viologen cores might further rearrange and/or fragment. As in Scheme 2 for \(V^{2+}\), fragment ions resulted from the loss of different numbers of \(H\) radical could also be expected. The possible routes of rearrangement and \(H\) radical dissociation for \(\text{BV}^{4+}\) after catching electrons are illustrated in the ESI (Scheme S3). Indeed, fragment ions formed through the loss of \(H\) radicals, such as \([\text{BV}^{4+} + e + 2(\text{PF}_6^-) - H]^+\), \([\text{BV}^{4+} + 2e + (\text{PF}_6^-) - H]^+\), \([\text{BV}^{4+} + 2e + (\text{PF}_6^-) - 2H]^+\), \([\text{BV}^{4+} + 3e - H]^+\) and \([\text{BV}^{4+} + 3e - 2H]^+\), were observed (see insets of Figure 4).

Additional supporting evidence for charge reduction by electron capture instead of by loss of protons can be found by analyzing the cluster of ions at \(m/z\) around 692. If the charge reduction is to be achieved by loss of protons, then the loss of 3 protons is required for \(\text{BV}^{4+}\) to form a singly charged ion. However, we could not detect \([\text{BV}^{4+} - 3H]^+\) ions (see Figures 4 and S5). The lack of \([\text{BV}^{4+} - 3H]^+\) and the observation of \([\text{BV}^{4+} + 3e]^+\), \([\text{BV}^{4+} + 3e - H]^+\) and \([\text{BV}^{4+} + 3e - 2H]^+\) ions strongly support the mechanisms proposed above for the charge reduction and fragmentation for viologen derivatives.

Our results also reveal that mechanisms other than electron capture exist to reduce the multiple charges of \(\text{BV}^{4+}\). In addition to the pseudo-molecular ions and the fragments discussed above, a number of other fragment ions were also clearly observed in Figure 4. The formation of these fragments can be attributed to the cleavage of the linkage between the two doubly charged bipyridinium cores. Considering the quadruple charges and the small molecular size of \(\text{BV}^{4+}\), the charge repulsion within the ion must be significant. Because of the charge repulsion, the most fragile bond within the \(\text{BV}^{4+}\) ion should be the aliphatic chain between the 2 bipyridinium cores. A schematic diagram of possible pathways for the cleavage is given in Scheme 3.
Compellingly, all the predicted fragment ions were observed with both DCTB and CHCA matrices (Figures 4 and S5).

3.3 | Cyanine dye-labelled BTA derivatives

Benzene-1,3,5-tricarboxamide (BTA) derivatives can self-assemble into supramolecular polymers via reversible intermolecular hydrogen bonding and π–π stacking, which together with a variety of secondary interactions, is used to make novel supramolecular materials.35 Recently, 2 cyanine dyes (Cy3 and Cy5) were used as fluorescent labels to study the spatiotemporal control and selectivity in supramolecular polymers.24,25 In this work, 2 BTA derivatives with 2 and 3 Cy5 attached as dye labels, (BTA2Cy5)Cl2 and (BTA3Cy5)Cl3, were measured by MALDI TOF MS. The structures of the derivatives are provided in Figure 1.

The MALDI TOF MS spectra of (BTA2Cy5)Cl2 and (BTA3Cy5)Cl3 using CHCA matrix are shown in Figure 5. Similar results were also obtained using DCTB as matrix. Since one Cy5 bears one positive charge, the number of fixed charges is determined by the number of Cy5 attached in the BTA derivative. Based on the m/z values, the most abundant ion peaks can be assigned as [BTA2Cy52+ – H+] and [BTA3Cy53+ – 2H+] for (BTA2Cy5)Cl2 and (BTA3Cy5)Cl3, respectively. Judging on these data, it would appear that these ions form through loss of protons.

From Figure 5, it can also be seen clearly that deprotonation is not the only mechanism by which these 2 BTA compounds undergo charge reduction. Fragment peaks of [BTA2Cy52+ – CH3]+, [BTA3Cy53+ – H+ – CH3]+, and [BTA3Cy53+ – 2CH3+] were also observed for (BTA2Cy5)Cl2 and (BTA3Cy5)Cl3, respectively. Considering the chemical structure of Cy5+, the methyl groups on the C atoms close to the benzene rings would appear to be easily cleaved as doing...
so would not interfere with the \( \pi \)-conjugation through the dye moiety. Scheme 4 shows the possible pathways for charge reduction of BTA2Cy5\(^{2+} \). Obviously, for BTA3Cy5\(^{3+} \), cleaving off 2 CH\(_3\) groups from 2 different Cy5s is required for the formation of a singly charged ion.

### 3.4 | Perylene bisimide derivative

Perylene bisimide derivatives are an important class of dyes useful for textile applications and as additives in high-grade industrial paint.\(^{36,37} \) As perylene bisimides feature a relatively low reduction potential which enables their use as an n-type semiconductor and as an electron acceptor in photoinduced charge transfer reactions, they have also been widely used in various optical and optoelectronic applications.\(^{38,39} \) In this study, a quaternized ammonium perylene bisimide (QPB) was measured using MALDI TOF MS, of which the structure is given in Figure 1.

Figure 6 shows the MALDI TOF MS spectra of QPB. Like Qdendr1, QPB is also doubly charged, bearing 2 quarternized ammonium centers. The peaks at \( m/z \) of 831.5 and 786.4, which were observed using both CHCA and DCTB as matrices, can be assigned as \([\text{QPB}^{2+} - \text{CH}_3]^{+}\) and \([\text{QPB}^{2+} - \text{N(CH}_3)_3 - \text{H}]^{+}\), respectively. As discussed above for Qdendr1, charge reduction for a quarternized ammonium can be achieved through cleavage across 1 of the 4 N–C bonds. A schematic diagram of fragmentation pathways for QPB is provided in Scheme 5, which is similar to that for Qdendr1 (Scheme 1).

Interestingly, QPB also consists of a highly electron-deficient perylene group. Owing to the electron deficient nature of the perylene group, an electron could easily be captured and stabilized within the \( \pi \)-conjugated system of perylene. It is known for similar electron deficient groups, a singly charged ion such as \([M + 2\text{Na}^+ + e]^+\) can be formed through double cation adduction combined with electron capture.\(^{40} \) Although not detected when using CHCA as matrix (the red spectrum of Figure 6), an intense singly charged molecular ion peak \([\text{QPB}^{2+} + e]^+\) of \( m/z \) at 846.5, formed by one electron reduction, was clearly observed for QPB with DCTB as the matrix (the black spectrum of Figure 6).

### 3.5 | Tetrathiafulvalene as matrix

In the experiments discussed above, 2 popular MALDI matrices of CHCA and DCTB were studied. Asakawa et al reported the use of tetrathiafulvalene (TTF), an electron donating compound, as a matrix for the analysis of some industrial pigments.\(^{41} \) Inspired by their nice results, we also tested TTF for the analytes containing ions bearing fixed multiple charges. Under our conditions, no analyte ions were detected for \((\text{QDendr1})_2\), \((\text{BTA2Cy5})_2\text{Cl}_2\), and \((\text{BTA3Cy5})_2\text{Cl}_3\), indicating that TTF is not a good matrix for these compounds. The MALDI TOF MS spectra for \(\text{V(PF}_6)_2\), \(\text{BV(PF}_6)_4\), and \((\text{QPB})_2\text{Br}_2\) are given in the ESI (Figures S6–S8), which are similar to the corresponding spectra obtained with CHCA and DCTB matrices. Apparently, the strong...
electron donating property of TTF did not yield considerably more reduced analyte molecular ions via electron capture. The reason for this observation is still under investigation.

It is well known that MALDI MS results are strongly matrix dependent. Although noticeable differences were observed in the mass spectra using different matrices, all the ions recorded in the spectra are singly charged because of the strong tendency to form singly charged ions in MALDI. For multiply charged organic ions, the fixed charges are not confined to one atom, which is in clear contrast with multivalent metal ions. Upon electron capture, fragmentation will occur concurrently for the organic ions. In addition, reduction via charge splitting is also an important route for the formation of singly charged fragment ions. Because of the fragmentation reactions, extra care should be taken in the MALDI MS analysis of organic ions bearing fixed multiple charges.

4 | CONCLUSIONS

A series of fragment ion peaks were observed in the MALDI TOF MS analysis of organic ions bearing fixed multiple charges. Although all of the organic ions studied in this work are intrinsically multiply charged, only singly charged ions were observed. In addition to counterion adduction, deprotonation, and electron capture, fragmentation was also found to be an important route for charge reduction. Cleavage of a chemical bond to the charged centers and/or rearrangement of the ions after catching electron(s) are found to be the main possible pathways of fragmentation. Therefore, care should be taken when interpreting MALDI MS data measured on ions bearing fixed multiple charges. With an improved understanding of the possible fragmentation mechanisms, MALDI TOF MS should still be a very useful technique for the characterization of organic ions bearing fixed multiple charges.

ACKNOWLEDGEMENTS

The authors would like to thank Dr J.A.J.M Vekemans and Dr K. Pieterse for valuable discussions and constructive suggestions.

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How to cite this article: Lou X, Li B, de Waal BFM, et al. Fragmentation of organic ions bearing fixed multiple charges observed in MALDI MS. J Mass Spectrom. 2018;39–47. https://doi.org/10.1002/jms.4035