Detection of oxygen addition peaks for terpendole E and related indole–diterpene alkaloids in a positive-mode ESI-MS

This report describes that a regular positive electrospray ionization mass spectrometry (MS) analysis of terpendoles often causes unexpected oxygen additions to form \([M + H + O]^+\) and \([M + H + 2O]^+\), which might be a troublesome in the characterization of new natural analogues. The intensities of \([M + H + O]^+\) and \([M + H + 2O]^+\) among terpendoles were unpredictable and fluctuated largely. Simple electrochemical oxidation in electrospray ionization was insufficient to explain the phenomenon. So we studied factors to form \([M + H + O]^+\) and \([M + H + 2O]^+\) using terpendole E and natural terpendoles together with some model indole alkaloids. Similar oxygen addition was observed for 1,2,3,4-tetrahydrocyclopent[b]indole, which is corresponding to the substructure of terpendole E. In tandem MS experiments, a major fragment ion at \(m/z\) 130 from protonated terpendole E was assigned to the substructure containing indole. When the \([M + H + O]^+\) was selected as a precursor ion, the ion shifted to \(m/z\) 146. The same 16 Da shift of fragments was also observed for 1,2,3,4-tetrahydrocyclopent[b]indole, indicating that the oxygen addition of terpendole E took place at the indole portion. However, the oxygen addition was absent for some terpendoles, even whose structure resembles terpendole E. The breakdown curves characterized the tandem MS features of terpendoles. Preferential dissociation into \(m/z\) 130 suggested the protonation tendency at the indole site. Terpendoles that are preferentially protonated at indole tend to form oxygen addition peaks, suggesting that the protonation feature contributes to the oxygen additions in some degrees. © 2014 The Authors. Journal of Mass Spectrometry published by John Wiley & Sons, Ltd.

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Keywords: terpendoles; indole–diterpene alkaloids; ESI oxidation; protonation site; oxygen addition for indole

Dear Sir,

Terpendoles, which are indole–diterpene alkaloids discovered from a soil-isolated fungal strain, were initially reported as inhibitors of acyl-CoA:cholesterol acyltransferase.\(^1\)\(^2\) Terpendole E (Compound 1, Fig. 1) was re-discovered as a novel Eg5 inhibitor.\(^3\) Some analogues have been regarded as candidates for lead compounds of anticancer drugs. For that reason, they are becoming attractive targets of chemical and biological research. Elucidation of molecular formulae with accurate mass measurement using mass spectrometry is the first task to distinguish and characterize analogues. Furthermore, hyphenation with liquid chromatography electrospray ionization mass spectrometry (LC-MS) is a sensitive identification technique that is useful for the analysis of terpendoles in a culture broth with far fewer preparation steps than those used for other techniques, such as X-ray diffraction and NMR analysis. However, unexpected \([M + H + 16]^+\) and \([M + H + 32]^+\) were often detected for terpendoles in a regular ESI-MS. Exact mass analyses defined them as one or two oxygen additions (Table 1). Undesirable oxygen additions cause mistakes in characterization and reduce the targeted ion signal intensity. It is also troublesome that the intensities of \([M + H + O]^+\) and \([M + H + 2O]^+\) fluctuated greatly and that they are unpredictable in each analysis. Simple electrochemical oxidation in ESI could not explain the observation as described later. Therefore, we studied factors affecting the formation of \([M + H + O]^+\) and \([M + H + 2O]^+\) using terpendole E and related natural indole–diterpenes together with some model indole alkaloids.

Electrospray ionization mass analysis of terpendole E (1) caused \([M + H]^+\) (\(m/z\) 438) with \([M + H + O]^+\) (\(m/z\) 454), \([M + H + 2O]^+\) (\(m/z\) 470) (Fig. 2(1)). Under the same regular ESI conditions, the oxygen addition peaks were observed at least for emindole SB (2), terpendole I (3) and paspalline (4). All their spectra were shown in Supporting Information 1 with similar results of four model compounds: \(\alpha\)-paxitriol (5), androsterone derivative alcohol (7), 1,2,3,4-tetrahydrocyclopent[b]indole (9) and 1,2,3,4-tetrahydrocarbazole (10). An LC-MS succeeded in a separation of an isobaric impurity of the oxygen addition peak assigned to \([M + 2O] + H]^+\), which increased in analyses for a preserved sample solutions with a time course of a few months (Supporting Information 2).

In this study, terpendoles (1, 2, 3 and 4) were extracted and purified from the culture broth as reported previously.\(^4\) \(\alpha\)-Paxitriol (5) was prepared by chemical reduction of purchased pasxilinie (6) (Sigma-Aldrich Corp.).\(^5\) Two simple steroidal indole alkaloids, alcohol (7) and ketone (8), were synthesized from androsterone in our laboratory. The detailed procedures are presented in Supporting Information 3. Yohimbine (11), reserpine (12), 1,2,3,4-tetrahydrocyclopent[b]indole (9) and 1,2,3,4-tetrahydrocarbazole (10) were purchased from Sigma-Aldrich Corp. and used without purification. For the ESI experiment, LC-MS grade methanol and water (Kanto Chemical Co. Inc) were used. Sodium formate was purchased from Sigma-Aldrich and Wako Pure Chemical Inds. Ltd. Special grade toluene (Wako Pure Chemical Industries. Ltd) was dried using molecular sieve.

All LC-ESI-MS analyses in this study were conducted (Synapt G2 HDMS; Waters Corp., Manchester, UK) with a regular ESI-source

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unit, but the oxygen additions on terpendoles are also observed by another devise (4000 Q Trap LC-MS system; AB Sciex) with a turbo spray ion source. Isocratic LC separation system (Acquity; Waters Corp.) with 20% water and 80% methanol at a flow rate of 300 μl/min. A BEH C18 column (2.1 × 50 mm, 1.7 μm) at 40 °C for the separation. The time-of-flight analyzer was set for resolution-mode with resolving power of 20 000 at m/z 556 (leucine enkephalin). The m/z range of 100–1500 was calibrated using 100 ppm sodium formate solution. The desolvation temperature was fixed at 250 °C. Regular ESI conditions in this study were set to obtain the maximum intensity of [M + H]+ for terpendole E; a stainless steel ESI capillary was applied high voltage of 3.0 kV, and the cone and desolvation gas (N2) flows were set to 200 l/h.

Table 1. Observed accurate masses of terpendoles and analogues

| Compound                  | Formula       | Theoretical mass (m/z) | Observed mass (m/z) | Error (ppm) | R.I.* |
|---------------------------|---------------|------------------------|---------------------|-------------|-------|
| Terpendole E (1)          | C28H39NO3     | 438.3008               | 438.3002            | 0.4          |       |
| Emindole SB (2)           | C28H39NO      | 406.3110               | 406.3113            | 0.7          |       |
| Terpendole I (3)          | C27H35NO5     | 454.2593               | 454.2585            | 0.9          |       |
| Psapaline (4)             | C28H39NO2     | 422.3059               | 422.3058            | 0.2          |       |
| α-Paxitriol (5)           | C27H35NO4     | 438.2644               | 438.2640            | 0.9          |       |
| Paxilline (6)             | C27H33NO4     | 436.2488               | 436.2484            | 0.8          |       |
| Alcohol (7)               | C25H33NO      | 364.2640               | 364.2643            | 0.8          |       |
| Ketone (8)                | C25H31NO      | 362.2484               | 362.2481            | 0.8          |       |
| 1,2,3,4-tetrahydrocyclopent(b)indole (9) | C11H11N | 158.0970               | 158.0972            | 1.3          |       |
| 1,2,3,4-tetrahydrocarbazole (10) | C12H13N | 172.1126               | 172.1125            | 0.6          |       |
| Yohimbine (11)            | C21H26N2O3    | 355.2022               | 355.2019            | 0.8          |       |
| Reserpine (12)            | C33H40N2O9    | 609.2812               | 609.2816            | 0.7          |       |

* R.I., relative intensity normalized to the intensity of [M + H]+.

Figure 1. Structures of 12 compounds including five natural terpendoles and seven model compounds: 1, terpendole E; 2, emindole SB; 3, terpendole I; 4, paspaline; 5, α-paxitriol; 6, paxilline; 7, alcohol; 8, ketone; 9, 1,2,3,4-tetrahydrocyclopent(b)indole; 10, 1,2,3,4-tetrahydrocarbazole; 11, yohimbine; 12, reserpine.
through line, and a new peak was observed at reaction because of the low oxidation potentials (2H2O=O 2 +4e−). Reactive oxygen formation from water is the most favorable additional oxygen originated from water in the solvent system. Isotope labeled water was added into the post-column flow, and 300 l/h, respectively. Samples were set as 5 ppm; 5 μl of the solution was injected into the system. The tandem MS (MS/MS) experiments were conducted with collision-induced dissociation technique with laboratory collision energies at 20 or 25 V. Precursor ions of interest were isolated using a Q mass analyzer with a 1 u mass isolation window (high-resolution setting) for [M + H]+ to separate them from [M]+. Ions were let through the collision-induced dissociation cell where Ar 2.44 × 10−7 e− mbar was filled as the collision gas.

Hydroquinone is known as one of redox buffers that partially participate directly in the neutralization of free radicals and reactive oxygen species.1–4 With the addition of hydroquinone (300 ppm in resultant solution), [M + H + O]+ and [M + H + 2O]+ formation are suppressed in the terpendole E (1) analysis (Fig. 2(2)). It indicated that the reactive oxygen formed by the oxidation process in ESI causes the oxygen addition for terpendole E. Using dry toluene as a solvent, the oxygen addition was diminished, and [M]+ (m/z 437) became prominent (Fig. 2(3)). To confirm the origin of the oxygen, incorporation experiment using 18O-labeled water was performed in the α-paxitriol analysis. Isotope labeled water was added into the post-column flow through line, and a new peak was observed at m/z 474 (obs. 474.2620) assigned to [M + H + 218O]+ (calc. 474.2627) for α-paxitriol (5) (Supporting Information 4). Hence, we concluded that the additional oxygen originated from water in the solvent system. Reactive oxygen formation from water is the most favorable reaction because of the low oxidation potentials (2H2O=O2+4H++4e− (E°(2H+/H2)=1.25 V)). Successive formation of reactive oxygen species (such as H2O2 (2OH−=H2O2+2e−)) can also take place via electrolysis.10,11 Obvious mass shift (2 Da) was absent for [M + H + O]+ by the addition of 18O−water. It might be a result from the insufficient formation of [M + H + O]+ and isobaric interference of the isotope of [M + NH4]+ on [M + H + 18O]+.

The anodic side reaction of analytes in positive-mode ESI has been well documented for alklylation and polymerization of phenyленediamines16,10 gas-phase methanol addition of aromatic aldehydes11 and halogen addition to π-conjugated phosphasilenes.12 Multiple electrochemical parameters, containing material of capillary metal, flow rate, needle voltage and gas flows, contribute to the reactions. Oxygen additions to the compound in positive-mode ESI were also documented for reserpine13 and zotepine14 with some byproducts using potentiostat system incorporated into ESI system. Without potentiostat system, no oxidized ions were observed for yohimbine (11) and reserpine (12) under any ESI settings in this study (Supporting Information 5(1 and 2)). Therefore, the molecular feature of terpendoles partially contributed on the oxygen addition as well as electrochemical formation of reactive oxygen. However, the structural character that causes oxygen additions in ESI has only rarely been understood. Reduction of both cone and desolvation gas flows increased the signal of oxygen additions for terpendole E (Fig. 2(4)), indicating that the balance of gas parameter settings was one of controlling factors. On the other hand, no systematic trend was observed between the needle voltage setting and the relative intensities of oxygen addition peaks for terpendole E (1) (Supporting Information 6). The parameters giving the maximum intensity of [M + H + O]+ or [M + H + 2O]+ varied among compound. To compare the degrees of oxygen additions among structures, we examined the identical regular ESI condition on the indole containing compounds (Table 1, Supporting Information 1 and 6). The smallest compound that gave the oxygen additions under the regular ESI settings was 1,2,3,4-tetrahydrocyclopent[b]indole (9). Also, 1,2,3,4-tetrahydrocarbazole (10) gave the oxygen addition peaks [M + H + O]+ and [M + H + 2O]+ with intense [M + O]+. The oxygen additions for 1,2,3,4-tetrahydrocyclopent[b]indole (9) and 1,2,3,4-tetrahydrocarbazole (10) are favorable to take place at reactive C-2 and C-3 positions of indole.15

To elucidate the oxidation site of terpendole E, MS/MS experiments were conducted on [M + H]+ and the oxygen added peaks. Protonated terpendole E (1) ([M + H]+ at m/z 438) was fragmented into m/z 130 (130.0660) attributed to 3-methylenediindolium or N-protonated quinoline ion (C9H8N+, calc. 130.0657),16 which was
identical fragment ion observed for MS/MS on protonated 1,2,3,4-
tetrahydrocyclopent[b]indole (9) (Fig. 3). When the [M + H + O]⁺
was selected as the precursor ion, m/z 130 of both terpendole E
(1) and 1,2,3,4-tetrahydrocyclopent[b]indole (9) shifted to m/z 146
(146.0595, C₁₉H₂₂NO⁺ calc. 146.0606). The MS/MS on [M + H + 2O]⁺
also gave fragment ions assigned to oxidized 3-methyleneindolinium
or N-protonated quinoline ion for both compounds. The peak at m/z
145 (145.0519) from [M + H + 2O]⁺ of 1,2,3,4-tetrahydrocyclopent[b]
indole (9) was assigned to C₁₀H₁₀NO⁺ (calc. 145.0528), containing
oxidized indole. These observations indicated that the site of the
oxygen additions of terpendole E (1) was indole portion same as
that of 1,2,3,4-tetrahydrocyclopent[b]indole (9).

Terpendole E (1) is composed with some ring units; indole part
with rings A, B and diterpene part from ring C to the
tetrahydropyran ring F, which connects with fused cyclopentene
ring C. (Fig. 1). The ring A–C portion can be recognized as the
substructure of indole fused with cyclopentene ring C at C-2
and C-3 double bond of the indole. Two hydroxyl groups exist
on the tetrahydropyran ring F and a 2-hydroxyisopropyl side
chain of the ring F. The resemblablence of oxygen addition and
MS/MS features of terpendole E to 1,2,3,4-tetrahydrocyclopent[b]
indole (9) suggested that the ring A–C portion of terpendole E
(1) was a significant structure in the oxidation reaction.

It is particularly interesting that the oxygen additions were rarely
detected for some terpendoles, even having an identical ring A–C
structure. Paxilline (6) possesses a conjugated ketone carbonyl
group in dihydroxy group ring F (Supporting Information 6 (3)).
Although small oxygen addition peaks were detected with changing
condition from regular ESI, the intensities of both [M + H + O]⁺ and
[M + H + 2O]⁺ of protonated paxilline (6) were smaller than those of
α-paxitriol (5), which was a reduction product of paxilline (6) having
an α-oriented hydroxyl group at the allylic position in ring F. The
core structure were unchanged; however, fragmentation character-
istics described by means of breakdown curves in MS/MS exper-
iments showed the significant difference between them. Fragment
ion at m/z 130 was characteristic for α-paxitriol (5) (Fig. 4(a)). On the
other hand, a water loss competed to the formation of m/z 130 for
paxilline (6) (Fig. 4(b)). Preferential fragmentation into m/z 130
required protonation at the indole portion for α-paxitriol (5), and
a water loss of protonated paxilline (6) should be triggered by the
proton localization at a hydroxyl group in the ring D–F portion.
For paxilline (6), proton can migrate into the hydroxyl group on
the 2-hydroxyisopropyl group followed by the first protonation at
the carbonyl site at the ring F via 6-membered ring intermediate.
Available data of gas-phase proton affinities predict the favorable
protonation site in the terpendoles. Proton affinities of indole and
alkyl ketone were reported to be 891–986 kJ/mol[16–19] and up to
864 kJ/mol[18], respectively. Their comparable proton affinities in
paxilline (6) could cause the competitive proton localization be-
tween the ring A–C portion and the ring D–F portion with a ketone.
However, if the ring D–F portion contained only hydroxyl group like
α-paxitriol (i-C₅H₅OH: 793 kJ/mol[18]), a proton should localize at
the indole preferentially because of its higher proton affinity.

Therefore, we can assume the additional factor to facilitate the
oxygen additions; preferential localization of a charged proton at
the indole portion as well as 1,2,3,4-tetrahydrocyclopent[b]indole
(9). Although the reaction mechanism to add oxygen to protonated
indole has not been clear yet, the structure containing the ring A–C
portion may be insufficient to the substantial reaction.

In the same way, oxygen additions were observed for alcohol (7),
but absent for ketone (8). Although the protonated (8) fragmented
into m/z 130 predominantly as well as (7), higher energy was nec-
essary to give the fragment than (7) (Fig. 4(c) and 4(d)). If the behav-
ior upon protonation was identical between these two, the energy
requirements to form m/z 130 should be similar. The comparable
proton affinities between indole and ketone group at ring F for
compound (8) could reduce the population of indole protonated
species rather than the case of compound (7).

The artificial oxygen additions to terpendoles in positive ESI-MS
were described here. Multiple factors control the phenomenon:
water contents in the ESI system, oxidation of water to form
reactive oxygen species, and protonation site of molecules. To

Figure 3. Tandem mass spectrometry spectra of terpendole E: (1)[M + H]⁺ (collision energy = 20 V), (2) [M + H + O]⁺ (20 V), (3) [M + H + 2O]⁺ (25 V) and
1,2,3,4-tetrahydrocyclopent[b]indole: (4) [M + H]⁺ (20 V), (5) [M + H + O]⁺ (20 V), (6) [M + H + 2O]⁺ (25 V).
reduce the unexpected oxygen addition as possible, hydroquinone addition after LC separation is expected to be one of effective means. The oxygen additions also depend on the analyte structure. Protonation at indole portion of the terpendole structure seemed to be the key to the oxygen addition. Unfortunately, it would be difficult to control protonation site for compounds in each ESI settings, but source tuning to give dominant \([M]+\) or \([M+Na]+\) formation may also be effective to reduce unexpected ions.

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