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### Article

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1       **Enrichment and analysis of quaternary alkaloids from *Zanthoxylum simulans* using weak**  
2       **cation exchange solid-phase extraction coupled with LC-MS**

3  
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22  
23      **ABSTRACT**

24      **Introduction:** Quaternary alkaloids (QAs) are the major alkaloids in several traditional Chinese  
25      medicines, especially in *Zanthoxylum simulans* (*Z. simulans*). However, few studies on  
26      enrichment of QAs from *Z. simulans* were conducted due to their high polarity and low content.

27      **Objective:** To develop a weak cation exchange solid-phase extraction coupled with LC-MS  
28      method to enrich and identify QAs from *Z. simulans*. Meanwhile, the qualitative and quantitative  
29      analyses of QAs were carried out based on the optimum conditions of the method.

30      **Methods:** Fresh stem bark of *Z. simulans* was extracted with 70% aqueous methanol and enriched  
31      by weak cation exchange (WCX) solid-phase extraction (SPE). A high performance liquid  
32      chromatography coupled with tandem mass spectrometry (HPLC-MS/MS) with an electrospray  
33      ionisation (ESI) source was used for the qualitative and quantitative analyses of QAs.

34      **Results:** Significant improvements were observed in resolution and abundance of the peaks with  
35      WCX-SPE. The linearity, limit of detection (LOD) and limit of quantification (LOQ) were  
36      determined for this analytical method. The linear relationship ( $A = 338.85C - 187.72$ ,  $R^2 = 0.99$ )  
37      was explored in the range of 0.5-312.5  $\mu\text{g/mL}$  for chelerythrine. The LOD and LOQ for

38 chelerythrine standard solutions were 0.0539 µg/mL and 0.1798 µg/mL, respectively. In addition,  
39 twenty two peaks were detected successfully with WCX-SPE and nine of them are undetectable  
40 without the processing of WCX-SPE.

41 **Conclusion:** A highly selective and efficient method for simultaneous enrichment and  
42 identification of QAs from crude extract of *Z. simulans* was developed for the first time by  
43 combining WCX-SPE with LC-MS.

44

#### 45 **KEYWORDS**

46 *Zanthoxylum simulans*, quaternary alkaloids, weak cation exchange, solid-phase extraction

47

#### 48 **1 | INTRODUCTION**

49 The genus *Zanthoxylum* (Rutaceae) consists of about 250 species of deciduous shrubs, and 39  
50 species with 14 varieties have been found in China.<sup>1</sup> *Zanthoxylum simulans* (*Z. simulans*), a  
51 common prickly shrub, has the effects of relieving swelling and pain, detoxification, and  
52 diminishing inflammation.<sup>2</sup> Previous phytochemical studies on this plant have led to the  
53 identification of various compounds, such as volatile oils,<sup>3</sup> alkaloids, coumarins, lignans,  
54 terpenoids, and sterols.<sup>4,5</sup> Among them, lignans and neolignans mainly distribute in the stem wood  
55 when compared with any other parts of this plant.<sup>4</sup> Alkaloids, mainly found in the root or the stem  
56 bark <sup>6,7</sup> of *Z. simulans* have been of immense interest due to their bioactivities,<sup>8-10</sup> and  
57 quaternary alkaloids (QAs) as major alkaloids in this plant possess significant antitumour  
58 activities. For instance, nitidine displayed inhibitory activity on hepatic carcinoma cells by  
59 inhibiting the JAK1/STAT3 signal pathway *in vitro*,<sup>11</sup> chelerythridine and sanguinarine showed  
60 dose-dependent inhibitory activity by damaging the DNA of leukaemia carcinoma cells (L1210) *in*  
61 *vitro*,<sup>12</sup> and fagaronine acted as angiogenesis inhibitors on leukaemia cancer cells.<sup>13</sup> To explore the  
62 antitumour activities and the subsequent mechanisms of action regarding QAs from *Z. simulans*,  
63 one of the most important steps is to enrich and identify them from complex crude plant extracts.

64 In this context, we set out to develop and optimize the method for the simultaneous enrichment  
65 and analysis of alkaloids of interest. Alkaloids, as an important subgroup of plant secondary  
66 metabolites, are a type of nitrogen-containing organic compounds, and most of which have  
67 complex nitrogen heterocyclic structures. Meanwhile, alkaloids often co-exist with a large number

68 of other compounds, and are in extremely low content. Most types of alkaloids are easy to be  
69 dissolved in organic solvents rather than in water, while quaternary alkaloids can be dissolved  
70 both in water and alcohol. QAs, as soluble alkaloids, cannot be extracted directly from aqueous  
71 solution by organic solvents due to their specific polarity with very low contents in the plant, thus  
72 the pre-concentration of QAs from crude plant extract is extremely difficult. Thus it is urgent to  
73 develop an efficient method to enrich QAs. Therefore, effective enrichment of QAs from *Z.*  
74 *simulans* is the first important step prior to their qualitative and quantitative analyses.<sup>14</sup> In the  
75 previous study, repeated column chromatography and further purification combined with nuclear  
76 magnetic resonance (NMR) techniques were commonly employed to isolate and identify pure  
77 QAs from *Z. simulans*.<sup>15</sup> Although these methods could offer efficient and precise results, it was  
78 relatively time-consuming, labour-intensive, and expensive for precious samples. In an effort to  
79 improve the extraction efficiency as much as possible, Reinecke salt based colorimetric method  
80 was reported.<sup>16</sup> Although precipitation, filtration and further purification were seemed to be  
81 effective in this method, the cations, such as Cr<sup>4+</sup> and Ag<sup>+</sup>, are not environment-friendly, and also  
82 the labile factors. For instance, the selected reagents to precipitate, cation varieties, and time to  
83 precipitate could cause troubles leading to the failure to enrich alkaloids of interest. In addition,  
84 with the fast development of separation and analytical technologies, some instruments with high  
85 separation efficiency and sensitivity have been applied to the alkaloids analysis from the  
86 *Zanthoxylum* genus, including but not limited to high-speed counter current chromatography  
87 (HSCCC), capillary electrophoresis (CE), reversed phase high-performance liquid  
88 chromatography (RP-HPLC), and high performance liquid chromatography combined with  
89 electrospray tandem mass spectrometry and nuclear magnetic resonance (LC-ESI/MS/NMR).<sup>17-21</sup>  
90 However, the aforementioned instruments could not improve the analytical results significantly  
91 without proper pre-purification due to the complexity and high polarity of QAs. Therefore, we  
92 aimed to develop a cost-effective and environment-friendly method to enrich and purify QAs from  
93 *Z. simulans*.

94 Currently, some researches and applications of new technologies have greatly improved the  
95 efficiency of enrichment, meanwhile saved energy and time. For instance, macroporous resin is a  
96 kind of organic polymer adsorbents, which has frequently been reported in the enrichment and  
97 purification of alkaloids because of its good efficiency,<sup>22</sup> but the method has some limitations

98 because the efficiency is restricted by many factors such as pH. To overcome the above limitations,  
99 ion exchange resin has demonstrated the advantages of low cost, high efficiency, good  
100 maneuverability, and low pollution, which has been widely used in the enrichment and  
101 purification of acid and alkaline components in natural products.<sup>23, 24</sup> Fortunately, WCX resin,  
102 with anions such as RCOO<sup>-</sup>, can be unexceptionably combined with QAs in the aqueous solution,  
103 <sup>25</sup> thus, successful enrichment of QAs can be achieved by using this strategy. With the carboxylic  
104 acid functional groups, WCX could combine with positive ion in water in the form of negatively  
105 charged groups because of the dissociation and cation exchange. It is difficult to dissociate in  
106 accompany with cation exchange in low pH and prefers to alkaline, neutral, and slightly acidic  
107 solutions. At this point, the WCX material, with the advantage of easy to be regenerated, might be  
108 applied to enrich and obtain higher content of QAs. WCX chromatography has played an  
109 important role in the field of separation and purification of samples, such as drug, urine and  
110 plasma samples.<sup>21, 26-30</sup> With WCX, the effectiveness of separation and purification was markedly  
111 improved. However, WCX has not been reported to enrich and purify QAs from *Z. simulans*.  
112 Taken all the points above into consideration, the application conditions of WCX were optimized  
113 and applied to the enrichment and analysis of QAs from *Z. simulans* in this study. As a result, a  
114 simple WCX based method for the simultaneous enrichment and analysis of QAs was firstly  
115 developed and proved to be an efficient method for fast analysis and quality control of QAs from  
116 plants.

## 117 **2 | EXPERIMENTAL**

### 118 **2.1 | Materials, chemicals and reagents**

119 Fresh stem bark of *Z. simulans* was collected from Wuhan Botanical Garden of the Chinese  
120 Academy Sciences (Wuhan, China) in April 2015. After drying below at 30 °C, the stem bark  
121 materials were crushed and stored in drying apparatus before use.

122 The reference standards, including magnoflorine, laurifoline, magnocurarine, fagaronine,  
123 chelerythrine, were purchased from the national standard substance center and stored at room  
124 temperature. Acetonitrile of HPLC grade was purchased from Thermo Fischer Scientific Inc.  
125 (USA). Other organic solvents were of analytical grade and purchased from Sino-pharm Chemical  
126 Reagent Co. Ltd. (Shanghai, China). Three standards, nuciferine, lycorine and bhelerythrine, were

127 obtained from Tauto Bio-tech Co. Ltd. (Shanghai, China). Water for LC and LC-MS were  
128 prepared from EPED (Nanjing Yeap Esselte Technology Development Co., Ltd. Nanjing, China).  
129 Mix-mode Weak Cation Exchange Solid Phase Extraction columns (WCX) were bought from  
130 ANPEL Scientific Instrument (Shanghai, China) Co., Ltd.

## 131 **2.2 | Instruments and conditions**

132 The flow rate of SPE was 1 mL/min and the samples drying were recorded on nitrogen drier  
133 (Organomation N-EV AP) with 45 °C and 12 L/min. The analysis was performed on a Thermo  
134 Accela 1250 HPLC (Thermo Fisher Scientific, USA) combined with an auto-sampler and a VWD  
135 detector. A 10 µL aliquot of sample solution was injected and separated on a Phenomenex ODS (2)  
136 column (5 µm, 2 mm × 150 mm) at 25 °C. The chromatograms were observed at a wavelength of  
137 280 nm, and the flow rate was set at 0.2 mL/min. Mobile phase A and B were 5 mM ammonium  
138 acetate solution, and acetonitrile, respectively. The gradient was set as follows: 0-5 min, 5% (B);  
139 5-50 min, 5-60% (B). For ESI-MS/MS analysis, a Thermo Accela 600 LC system with a VWD  
140 detector and a TSQ Quantum Access MAX mass spectrometer (Thermo Fisher Scientific, San Jose,  
141 CA, USA) were applied to detect alkaloids in the positive ion mode. The MS conditions were  
142 optimized as follows: mass range from 200-1000 Da; spray Voltage, 3.0 kV; capillary temperature,  
143 250 °C; sheath gas pressure, 40 psi; aux gas pressure, 10 psi.

## 144 **2.3 | WCX extraction procedures and sample preparation**

145 Immersing in 70% aqueous methanol for 12 h, crushed stem bark of *Z. simulans* was extracted  
146 ultrasonically for 30 min in triplicates. After centrifugation, the supernatants were combined and  
147 condensed to afford residues, which were then dispersed in methanol (equal to 1 g plant materials  
148 in 1 mL methanol). The WCX Solid Phase Extraction conditions were optimized and used to  
149 enrich crude QAs from *Z. simulans* as follows: 1) WCX cartridges were preconditioned with  
150 methanol (3 mL) and 10% methanol (3 mL); 2) An aliquot of 200 µL crude methanol extract of *Z.*  
151 *simulans* was diluted in 10% methanol and further dispersed to 10 mL for three times. After  
152 centrifuged at 10000 rpm for 5 min, the samples were loaded to the preconditioned cartridges; 3)  
153 Water (1 mL) and methanol (5 mL) were used to wash out the unbounded components; 4) Crude  
154 QAs were eluted by methanol with 5% formic acid (5 mL). Then, the final eluents were collected

155 and dried by nitrogen drier, and the residues were dissolved with 10% methanol and filtered with  
156 0.22  $\mu\text{m}$  micro-filter membrane before the LC-MS analysis.

## 157 **2.4 | Method validation**

158 To validate the analysis method for the determination of QAs, a series of experiments were  
159 carried out. The regression equation and correlation coefficient were determined by integrating  
160 peak area of different concentrations of standard solutions accompany with a linear regression  
161 analysis. To assess the precision and accuracy of the method by calculating the relative standard  
162 deviation (RSD) and the recovery rate of inter-day, reference solution of chelerythrine with three  
163 different concentrations (0.94  $\mu\text{g/g}$ , 0.63  $\mu\text{g/g}$ , 0.31  $\mu\text{g/g}$ ) was added to the sample of *Z. simulans*,  
164 then the sample was extracted though WCX solid-phase extraction column and injected for HPLC  
165 analysis. For intra-day stability test, the chelerythrine standard solution was analyzed 5 times  
166 within one day ( $n = 5$ ), the results were measured by relative standard deviations (RSD). The  
167 recovery of method was evaluated by adding spiked samples. The limit of detection (LOD) and  
168 limit of quantification (LOQ) for chelerythrine standard solutions were calculated based on  
169 signal-to-noise ratio (S/N) of 3 and 10, respectively.

## 170 **3 | RESULTS AND DISCUSSION**

### 171 **3.1. Effectiveness of WCX SPE on the enrichment of quaternary alkaloids (QAs)**

172 As expected, crude extracts of *Z. simulans* were then processed by WCX-SPE, and subjected to  
173 HPLC analysis. Fig.1 shows the chromatograms in the HPLC (280 nm) analysis of crude extracts  
174 without (A) and with (B) WCX-SPE. It was observed that the peak shape of the sample with  
175 WCX-SPE was remarkably better than that without WCX-SPE. Meanwhile, the resolutions of  
176 most peaks observed were significantly improved, and 22 peaks were clearly resolved in Fig.1B,  
177 which clearly indicated that WCX-SPE can greatly enhance the enrichment and analysis of QAs  
178 from *Z. simulans*.

### 179 **3.2 | Optimization of WCX procedures**

180 In order to achieve the best enrichment efficiency for QAs from plant extracts, some relevant  
181 key parameters, including pH of samples, desorption reagents, specificity, and amount of rinse

182 reagents, sample sizes were systematically investigated. For instance, proper acidic samples might  
183 be good for QAs to dissolve, but improper acidic levels could cause low recoveries of QAs. As for  
184 alkalized samples, significant higher pH might lead to poor extraction efficiencies for QAs  
185 because of the competitive binding to carboxyl group between QAs and cation in solution. Thus,  
186 the effects of pH on the enrichment efficiency were evaluated in the range of 3.0-11.0 to obtain the  
187 best binding capacity. As shown in Fig. 2A, after subjected to WCX, the highest recovery for QAs  
188 was achieved at around pH 7.0. Thus, pH 7.0 was selected in the following studies. Based on  
189 another application by Qiu *et al.*,<sup>30</sup> 5 mL of desorption reagents were used to elute QAs from  
190 WCX in the following experiments. To select the suitable desorption reagents to improve the  
191 efficiency, the most commonly used desorption reagents, such as 5% formic acid in methanol, 5%  
192 formic acid in acetonitrile, and 5% formic acid in acetone were investigated.<sup>26, 29, 35</sup> Fig. 2B shows  
193 that 5% formic acid in methanol was the best with 96.0% recovery rate of chelerythrine, which  
194 was eventually selected as the desorption reagents for the complex samples from *Z. simulans*. To  
195 evaluate the specificity of WCX to QAs, the recovery of two pure compounds, chelerythrine and  
196 nuciferine, were compared, since distinctive polarity discrepancy of compounds could make a  
197 great contribution to the specificity of cation exchange solid-phase extraction.<sup>29</sup>

198 In general, a variety of alkaloids, like different types of tertiary alkaloids and quaternary  
199 alkaloids from *Z. simulans*, could possess different polarities. Most of the alkalogenic compounds  
200 present alkalinity in 10% methanol solution, and the alkalinity of the type of tertiary alkaloids is  
201 slightly weaker than that of QAs. Accordingly, the mixed standard solution, containing nuciferine  
202 (tertiary alkaloid) and chelerythrine (quaternary alkaloid), was extracted by WCX solid-phase  
203 extraction column, in order to investigate the special properties of the mixed weak cationic solid  
204 phase extraction column to quaternary alkaloid. Based on evidences above, WCX-SPE had  
205 obvious enrichment for chelerythrine (quaternary alkaloid) and no obvious enrichment effect to  
206 nuciferine (tertiary alkaloid). To investigate the volumes of the rinse reagents for WCX-SPE, the  
207 mixed standard solution with lycorine (tertiary alkaloid) and chelerythrine (QA), was extracted by  
208 WCX-SPE, and then eluted with 3 mL, 5 mL and 7 mL methanol, respectively. Finally, the  
209 optimal volume of the rinse reagent (methanol) was confirmed by calculating the recovery rate of  
210 the two standard alkaloids. As shown in Fig. 2C, the recovery rate of chelerythrine slightly  
211 increased with the volume of methanol from 3 mL to 5 mL, while the recovery rate of lycorine



212 decreased significantly in the meantime. The recovery rates of lycorine and chelerythrine showed  
213 no significant change, with the volumes of methanol increase from 5 mL to 7 mL. Thus, the  
214 volume of rinse reagent (methanol) was defined as 5 mL in this study, in order to reduce the  
215 non-QAs content in the sample with WCX.

216 Since the mixed mode weak cationic solid-phase extraction column used in this study was  
217 packed with 500 mg of material in a 3 mL column, the ion exchange capacity of each column was  
218 limited. In this way, appropriate sample size was extremely vital for the successful enrichment of  
219 QAs, and underloading or overloading the sample would cause some alkaloids from the *Z.*  
220 *simulans* undetectable or with poor resolution, and even damage the column. Chelerythrine, a  
221 representative type of QAs, while not presented in *Z. simulans*, was then selected to optimize the  
222 loading amount of samples. To get the recovery rate of the chelerythrine, 0.2 g and 0.4 g samples  
223 of *Z. simulans* were prepared with the addition of certain amount of reference substance  
224 chelerythrine, respectively. Then, the samples were subjected to WCX solid-phase extraction  
225 column. The recovery rates of added chelerythrine were 93.9% and 106.0%, respectively, when  
226 the samples loaded are 0.2 g and 0.4 g. By comparison of the chromatography analyzed under 280  
227 nm, same peaks were detected for 0.2 g and 0.4 g crude material. Therefore, the sample amount  
228 for the analysis of quaternary alkaloid in final could be economically set as 0.2 g.

### 229 **3.3 | Validation of the proposed method**

230 Table 1 shows the results of linear ranges (LR), limits of detection (LOD), limits of  
231 quantitation (LOQ), and relative standard deviations (RSD) for a representative QA  
232 (chelerythrine). The linearity ( $A = 338.85C - 187.72$ ; where A is absorbance, and C refers to the  
233 concentration of chelerythrine) was good with a correlation coefficient ( $R^2$ ) greater than 0.9999,  
234 when the concentrations of chelerythrine ranged from 0.5  $\mu\text{g/mL}$  to 312.5  $\mu\text{g/mL}$ . LOD and LOQ  
235 for chelerythrine were 0.0539  $\mu\text{g/mL}$  and 0.1798  $\mu\text{g/mL}$ , respectively. To assess the precision and  
236 accuracy of the method by calculating the RSD and the recovery rate of inter-day, the reference  
237 solutions of chelerythrine with three different concentrations (0.94  $\mu\text{g/g}$ , 0.63  $\mu\text{g/g}$ , and 0.31  $\mu\text{g/g}$ )  
238 was added to the sample of *Z. simulans*, then the sample was subjected to WCX solid-phase  
239 extraction column prior to HPLC analysis. For intra-day stability test, the chelerythrine standard  
240 solution was analyzed 5 times within one day ( $n = 5$ ), and the RSD was 0.67%, which indicated

241 good stability of the method. The recovery rates (RR) of three difference concentrations of  
242 chelerythrine were 107.8%, 105.8%, and 93.4%, respectively; and the average recovery rate (ARR)  
243 was 102.3%. The results clearly indicated that the proposed method had good accuracy, and could  
244 be used to detect the content of other QAs.

### 245 **3.4 | Applications of WCX-SPE for the enrichment and fingerprinting analysis of quaternary** 246 **alkaloids**

247 Under the optimized conditions, 0.2 g of *Z. simulans* was used, and the total QAs of *Z.*  
248 *simulans* were enriched by WCX, and eluted with 5% formic acid in methanol solution, and then  
249 subjected to HPLC and LC-MS in order to further confirm the selectivity of WCX to QAs, and  
250 identify quaternary alkaloids in *Z. simulans*.

#### 251 **3.4.1 | Fingerprinting analysis of total QA in *Z. simulans***

252 Chelerythrine, the crude extract of *Z. simulans*, and total quaternary alkaloids enriched from *Z.*  
253 *simulans* by WCX were eluted under the chromatographic conditions shown in section 2.2, the  
254 chromatograms of three samples were presented in Fig. 3 (0-40 min). With the application of the  
255 WCX-SPE, QAs from the crude extract of *Z. simulans* were successfully enriched and a majority  
256 of non-QAs were removed because of the high selectivity of WCX for quaternary amines. There  
257 were 22 compounds detected under 280 nm, and peak 18 was identified according to the retention  
258 time of chelerythrine and the corresponding fragmentation patterns in LC-MS/MS.

#### 259 **3.4.2 | Qualitative and quantitative analysis of QAs in *Z. simulans***

260 The samples with WCX-SPE were analyzed by LC coupled with a TSQ Quantum Access  
261 MAX mass spectrometer system, and the LC chromatogram is presented in Fig. 3. The  
262 information of detected compounds is tabulated in Table 2, which lists contents, retention times ( $t_R$ )  
263 and MS fragment ions. Based on MS spectra of detected peaks, 15 alkaloids exhibited their  
264 quasi-molecular ions  $[M]^+$  or  $[M+H]^+$ , and were identified from *Z. simulans* and shown in Fig. 4,  
265 which included four benzophenidine alkaloids (fagaronine 3-glucoside, fagaronine,  
266 8-*O*-demethylchelerythrine, chelerythrine), three *N*-methyl-tetrahydrocorberine alkaloids  
267 (*N*-methyltetrahydrocolumbamine, *N*-methylcanadine or its isomer), three aporphine alkaloids

268 (magnoflorine, laurifoline, 10-demethyl-magnoflorine), one protoberberine alkaloid (palmatine),  
269 and four benzyloquinoline alkaloids (magnocurarine or its isomers, 8-methoxy-isotembetarine,  
270 and isotembetarine). Their MS/MS data and fragmentation patterns were in good agreement with  
271 those reference compounds or literatures. In more details, the interpretations of MS/MS spectra for  
272 different types of alkaloids would be discussed by taking some representative alkaloids as  
273 examples. For benzophenidine type of alkaloids, peak 18 with the parent ion at  $m/z$  348 was  
274 further discussed; the product ions at  $m/z$  332, 318, 304 and 290 indicated different neutral losses  
275 of 16, 30, 44 and 58 Da, which were corresponding to  $\text{CH}_4$ ,  $\text{CH}_2\text{O}$ ,  $\text{C}_2\text{H}_4\text{O}$  and  $\text{C}_2\text{H}_5\text{CHNH}$ ,  
276 respectively. Based on the fragment ions and the analysis above as well as the data in the literature,  
277 peak 18 could be proposed as chelerythrine.<sup>36</sup> As shown in the MS/MS spectrum,  $m/z$  190 and 165  
278 were the most intensive abundance fragments, indicating the RDA cleavage of the mother ion. The  
279 further losing of  $\cdot\text{CH}_3$  from  $m/z$  190 produce the fragment at  $m/z$  175. Based on the reported  
280 literature, peak 14/16 with the molecular ion ( $[\text{M} + \text{H}]^+$  at  $m/z$  354 were tentatively identified as  
281 *N*-methylcanadine or its isomer.<sup>37</sup> In the MS/MS spectrum of peak 3 and 8, same molecular ion  
282  $[\text{M} + \text{H}]^+$  at  $m/z$  342 and common fragment ions  $[\text{M} - (\text{CH}_3)_2\text{NH}]^+$  at  $m/z$  297 and  
283  $[\text{M} - (\text{CH}_3)_2\text{NH} - \text{CH}_3]^+$  at  $m/z$  282 were observed because of the cleavage of atom adjacent to N  
284 atom. Due to the drop of  $\text{CH}_3\text{OH}$  from  $m/z$  at 297 led to  $[\text{M} - (\text{CH}_3)_2\text{NH} - \text{CH}_3\text{OH}]^+$  at  $m/z$  265. The  
285 distinguishing fragment  $m/z$  at 237 was observed in the MS/MS spectrum of compound. Based on  
286 the reported literature, peak 3 and 8 were identified as magnoflorine and laurifoline,<sup>38</sup> respectively.  
287 Peak 19 had  $[\text{M} + \text{H}]^+$  at  $m/z$  352. The MS/MS fragments of peak 19 at  $m/z$  337, 334, 322, 320, 308  
288 and 294 were consistent with palmatine reported previously. As a result, peak 19 was definitely  
289 identified as palmatine.<sup>39</sup> For peak 4 and 9, the characteristic and intensive fragment at  $m/z$  209  
290 and 107 were observed due to the  $\beta$  cleavage of  $[\text{M} + \text{H}]^+$  at  $m/z$  314. The further cleavage of bond  
291 adjacent to *N* atom from  $m/z$  209 led to fragments at  $m/z$  194 and 166. Due to different retention  
292 time at 19.91 and 24.54 min but same MS/MS fragments, peak 4 and 9 with same molecular ion  
293  $[\text{M} + \text{H}]^+$  at 314  $m/z$  were identified as magnocurarine or its isomers.<sup>40</sup> As shown in the Fig.3, the  
294 contents of QAs were determined using the method with external standard, and the contents of  
295 magnoflorine (peak 3), laurifoline (peak 8), magnocurarine or its isomer (peak 9), fagaronine  
296 (peak 15) and chelerythrine (peak 18) are higher than the other alkaloids, which are 234.2  $\mu\text{g/g}$ ,  
297 68.8  $\mu\text{g/g}$ , 87.4  $\mu\text{g/g}$ , 371.7  $\mu\text{g/g}$  and 193.0  $\mu\text{g/g}$ , respectively. Meanwhile, the contents of these

298 five compounds account for more than 5% of the detected alkaloids in *Z. simulans*. In addition,  
299 nine alkaloids were detected from the sample enriched by WCX-SPE, but were not detected  
300 without enrichment.

301

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412 with the isoquinoline alkaloids biosynthetic pathway. *J Pham Biomed Anal.* 2015;103:26-34.  
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Analytical	LR ( $\mu\text{g/mL}$ )	$R^2$	LOD ( $\mu\text{g/mL}$ )	LOQ ( $\mu\text{g/mL}$ )	RSD (%) / Intra-day	ARR (%)
Chelerythrine	0.5-312.5	0.9999	0.0539	0.1798	0.67	102.3

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415

416



417 **Table 2** Alkaloids detected in *Z. simulans* by LC-MS/MS.

418

Peak NO.	Rt / min	M <sup>+</sup> / [M+H] <sup>+</sup>	MS/MS spectrum	Identification	Contents (µg/g)	Relative contents (%)
1	4.50	314	314, 213, 140, 121, 103, 97	Unidentified*	35.9	2.98
2	10.18	222	222, 207, 191, 179, 164, 58	Unidentified*	3.0	0.25
3	18.10	342	342, 297, 282, 265, 247, 237, 219, 207, 191	Magnoflorine	234.2	19.46
4	19.91	314	314, 269, 237, 219, 209, 194, 192, 166, 137, 119, 115, 107	Magnocurarine or its isomer	9.4	0.78
5	21.60	358	358,313,298,283,267,206,189,174,163,158,151,137	8-Methoxy-isotembetarine	8.7	0.73
6	22.28	344	344, 314, 301, 269, 239, 207, 175, 143, 137	Isotembetarine	9.5	0.79
7	22.90	328	328, 313, 283, 268, 189, 151, 121, 107	Unidentified	4.8	0.40
8	23.51	342	342, 297, 282, 265, 250, 233, 222, 205	Laurifoline	68.8	5.72
9	24.54	314	314, 269, 237, 219, 209, 194, 192, 166, 137, 119, 115, 107	Magnocurarine or its isomer	87.4	7.26
10	25.75	512	512, 350, 335, 307	Fagaronine 3-glucoside	21.7	1.80
11	26.26	328	328, 283, 251, 223, 208, 175, 143, 121	10-Demethyl-magnoflorine	2.5	0.20
12	27.61	454	454, 340, 322, 226, 209, 114, 96	Unidentified*	5.8	0.48
13	28.31	356	356, 192, 177, 149	<i>N</i> -Methyltetrahydrocolumbamine	1.9	0.16
14	28.82	354	354, 190, 175, 165	<i>N</i> -Methylcanadine or its isomer*	4.4	0.36
15	30.58	350	350, 335, 320, 307, 292, 264	Fagaronine*	371.7	30.88
16	32.21	354	354,190,175,165	<i>N</i> -Methylcanadine or its isomer	43.0	3.57
17	33.57	334	334, 319, 304, 291, 276, 262	8- <i>O</i> -Demethylchelerythrine	2.5	0.21
18	33.98	348	348, 332, 318, 304, 290	Chelerythrine	193.0	16.04
19	38.48	352	337, 334, 322, 320, 308, 294	Palmatine*	21.5	1.79
20	39.19	378	378, 363, 334, 319	Unidentified*	19.2	1.59
21	39.97	274	274, 256, 106, 88	Unidentified*	22.76	1.89
22	41.29	594	594, 533, 385, 348, 193, 149	Unidentified*	8.0	0.67

419 \*compounds were not detected before WCX-SPE, but were successfully detected after WCX-SPE

420

421 **Figure captions:**

422

423 **FIGURE 1** The chromatograms of total extracts of *Z. simulans* without (A) and with (B) WCX.

424 Twenty two peaks were detected with WCX-SPE and nine of them are undetectable in absence of

425 WCX-SPE.

426

427 **FIGURE 2** Effects of pH of sample loading solvents (A), desorption reagents (B), and volumes of

428 rinse reagents (C) on the extraction efficiency. A): The highest recovery for QAs was achieved at

429 around pH 7.0 after subjected to WCX; B): 5% formic acid in methanol was eventually selected as

430 the desorption reagents; C): The volume of rinse reagent (methanol) was defined as 5 mL.

431

432 **FIGURE 3** The chromatograms of chelerythrine (a), crude extract of *Z. simulans* (b) and total

433 QAs enriched from *Z. simulans* by WCX (c). The sample with WCX-SPE has twenty-two QAs in

434 c, while the resolution and abundance of the peaks in b is not so satisfactory without WCX.

435

436 **FIGURE 4** The structures of compounds detected from *Z. simulans*.

437