Characterization of main components in Xiao’er Xiaoji Zhike oral liquid by UPLC-MS and their taste evaluation

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Abstract
This paper provided a method for determining the potential quality markers in Xiao’er Xiaoji Zhike oral liquid (XXZOL) based on the concordant tastes of compounds with their respective originated Chinese medicinal pieces. UPLC-Q-Exactive-Orbitrap-MS technology was used to identify the main chemical constituents in XXZOL. The electronic tongue collected the electronic responses of the sour, bitter, sweet, pungent, and salty sample solutions, and the discriminant factor analysis (DFA) model was established to recognize the taste characteristics of 23 liquid samples. Fifteen high content ingredients in XXZOL were identified by UPLC-MS, and the established DFA model recognized their respective tastes. The accurate recognition rate of the DFA model was 73.33%, the false rate was 6.67%, and the unrecognized rate was 20%. The concordance rate of their authentic tastes with the tastes of their respective originated Chinese medicinal pieces was 78.57%. Trigonelline, malic acid, citric acid, and caffeic acid were the potential sour material bases of XXZOL. Mannitol was the potential sweet material basis of XXZOL. L-phenylalanine, sinapine, forsythoside I, pinoresinol-4-O-β-D-glucopyranoside, naringin, and neohesperidin were the potential bitter material bases of XXZOL. So the above 11 main compounds were the potential quality markers of XXZOL.

Keywords: Xiao’er Xiaoji Zhike oral liquid; UPLC-Q-Exactive-Orbitrap-MS; electronic tongue; potential quality marker.

Practical Application: A method for determining the potential quality markers in Xiao’er Xiaoji Zhike oral liquid based on the concordant tastes of compounds with their respective originated Chinese medicinal pieces.

1 Introduction
The electronic senses electronically reproduce the responses similar to the five senses presenting in human beings, such as electronic eye (Orlandi et al., 2018), electronic ear, electronic skin (Zhang et al., 2020b), electronic tongue (e-tongue) (Toko, 1998) and electronic nose (Mohd Ali et al., 2020). These electronic sensory technologies can be used alone (Gan et al., 2019; Pascual et al., 2018; Nategh et al., 2021) or combined (Orlandi et al., 2019; Banerjee et al., 2019; Xu et al., 2019) to achieve experimental purposes. The e-tongue is a kind of liquid analytical instrument consisting of three parts, including sensor array, signal acquisition system, pattern recognition system (Toko, 1998), and taste sensations of the sensor array can usually be classified into sweet, sour, salty, bitter, and umami. Sensor array imitates the taste cells in taste buds that interact with the flavorful substance and transmit the taste information to nerve fibers and the relevant areas in the central nervous system (CNS) (Pandurangan & Hwang, 2015). The non-specific and poorly selective sensors in the sensory array are immersed into a sample solution and acquire a global characteristic response signal of the substances in the sample (Jiang et al., 2018). Then the corresponding signals are sent to a signal processing system equivalent to CNS, which can analyze the acquired signals using appropriate pattern recognition methods. The e-tongue can realize the qualitative and quantitative analysis of simple or complex liquid samples and obtain the results reflecting the taste characteristics of samples (Leggin et al., 2000). Based on the working principle of the sensor array, the e-tongue can be classified into electrochemical (Wei et al., 2018; Ciosek & Wróblewski, 2011), optical and enzymatic e-tongue. Besides the advantages of low cost and simple operation, sensitivity, reliability, and robustness, e-tongue has its particular advantage to analyze the taste of some toxic substances (Wadehra & Patil, 2016). The e-tongue has been widely used in different fields of food (Wadehra & Patil, 2016; Ghasemi-Varnamkhasti et al., 2018; Peris & Escuder-Gilabert, 2016), medicine (Wasilewski et al., 2019; Guedes et al., 2021), or environmental detection (Shimizu et al., 2019; Magro et al., 2018). The e-tongue system has good applications in quality monitoring of Chinese medicinal pieces (Shi et al., 2018), origins identification (Wu et al., 2018) and harvest period determination of herbal medicines, and the evaluation of the pharmaceutical process of traditional Chinese medicine (TCM) (Bi et al., 2020), such as correcting or masking the taste of the decoction from TCM (Lin et al., 2016; Feng et al., 2019).

Xiao’er Xiaoji Zhike oral liquid (XXZOL), a prescription preparation of TCM, with the effect of clearing heat in lungs, relieving cough due to lung heat, and helping digestion, usually is applied to treat children’s respiratory diseases such as mycoplasma pneumonia (Zhang et al., 2020a), asthma (Zhou et al., 2020) and cough (Liang et al., 2018). Because of its safety and definite effects, the market sales of XXZOL have been increasing steadily. There are 10 Chinese medicinal pieces in the prescription of XXZOL...
according to their taste deployment to reduce the toxicity and bias of a single Chinese medicinal piece and achieve the overall synergy of all Chinese medicinal pieces. This prescription is the compatibility of stir-baked *Crataegi Fructus*, *Arecae Semen*, honeyed *Eriobotryae Folium*, *Trichosanthis Fructus*, *Platycodonis Radix*, *Forsythiae Fructus*, *Aurantii Fructus Immaturus*, stir-baked *Raphani Semen*, stir-baked *Descurainiae Semen Lepidii Semen*, and *Cicadae Periostracum* (China, 2020). The taste of a single herbal decoction or a prescription preparation decoction is stable only when the type and quantity of the chemical compounds in the sample solution do not alter. According to the main tastes of Chinese medicinal pieces from the prescription, to understand the recipe-construction rule of XXZOL, high content components in XXZOL were identified. Their tastes were recognized respectively by the e-tongue system. Comparing with the main tastes of Chinese medicinal pieces from the prescription, components with consistent tastes are more likely to be the material bases of XXZOL.

In our work, the UPLC-Q-Exactive-Orbitrap-MS platform was used to analyze the chemical constituents of XXZOL, and the main components were identified by the high-resolution mass spectrometry (MS) technology and control substances. The e-tongue collected the sour, bitter, sweet, pungent, and salty samples, which were 1.0 mg/mL solutions of the reference components with above tastes, and the discriminant factor analysis (DFA) model to recognize the taste characteristics of liquid samples was established to predict the taste of every main component in XXZOL. Considering the main tastes of Chinese medicinal pieces from XXZOL, we obtained several main components in XXZOL with similar tastes, which were likely to be the important quality markers of the prescription, providing a basis for the determination of the quality markers of XXZOL.

### 2 Materials and methods

#### 2.1 Experimental materials

Acetosulfam potassium, andrographolide, berberine, citric acid, fructose, gallic acid, gentiopicroside, glycine, oxymatrine, piperine, saccharin, salicin, sodium chloride, sucrose, and zingerone were purchased from Shanghai Yuanye Bio-Technology Co., Ltd (Shanghai, China). Liquiritin and potassium iodide were purchased from Shanghai Macklin Biochemical Co., Ltd (Shanghai, China). Citric acid monohydrate was purchased from Xilong Scientific Co., Ltd (Guangdong, China). Potassium chloride was purchased from Beijing Chemical Works (Beijing, China). Nonivamide was purchased from Shanghai Aladdin Biochemical Technology Co., Ltd (Shanghai, China). Stevioside was purchased from Chengdu Push Bio-Technology Co., Ltd (Chengdu, Sichuan province, China). Caffeic acid, forsythoside A, forsythoside E, forsythoside I, L-phenylalanine, malic acid, maltose monohydrate, naringin, neohesperidin, pinosol-4-O-β-D-glucopyranoside, sinapine, synephrine, and trigonelline were purchased from Chengdu DeSiTe Biological Technology Co., Ltd (Chengdu, Sichuan province, China). Mannitol was purchased from Tianjin Fuchen Chemical Reagent Co., Ltd (Tianjin, China). The information of the reference components used for the e-tongue experiments was shown in Table 1, and their contents were greater than or equal to 98%. MS grade acetonitrile and formic acid were purchased from Thermo Fisher Scientific (USA). Analytical grade ethanol was purchased from Shanghai Rhawn Chemical Technology Co., Ltd (Shanghai, China).

XXZOL was purchased from Lunan Hope Pharmaceutical Co., Ltd (Linyi, Shandong province, China). Take the oral liquid 0.25 mL precisely to 10 mL volumetric flask, water was added to the mark. Then, the solution was filtered with a membrane filter (0.22 µm) to collect the successive filtrate as the sample solution.

The powder (25 mg) of each reference component was dissolved in 25 mL of 50% v/v aqueous ethanol with ultrasonication at room temperature for 10 minutes. The obtained solutions were collected for electronic tongue analysis.

#### 2.2 LC-MS/MS analytical conditions

LC-MS/MS was carried out on a UPLC-Q-Exactive-Orbitrap-MS platform (Thermo Fisher Scientific, USA) with a Waters ACQUITY UPLC HSS T3 Column (100 mm×2.1 mm, 1.8 μm). The mobile phases consisted of acetonitrile (A) and 0.1% aqueous formic acid (v/v) (B) using gradient elutions of 0-2%A at 0-10 min, 2-7%A at 10-20 min, 7-14%A at 20-30 min, 14-14.5%A at 30-32 min, 14.5-15%A at 32-37 min, 15-15.5%A at 37-40 min, 15.5-17%A at 40-45 min, 17-24%A at 45-53 min, 24-30%A at 53-60 min.

| Taste | Type of reference component | Reference component (No.) |
|-------|-----------------------------|---------------------------|
| Sour  | Organic acid                | Citric acid (SO-1), citric acid monohydrate (SO-2), gallic acid (SO-3), malic acid (SO-4) |
| Bitter| Alkaloid                    | Berberine (B-1), oxymatrine (B-2) |
|       | Glycoside                   | Salicin (B-3)             |
|       | Terpenoid                   | Andrographolide (B-4), gentiopicroside (B-5) |
| Sweet | Amino acid                  | Glycine (SW-1)            |
|       | Carbohydrate                | Fructose (SW-2), maltose monohydrate (SW-3), sucrose (SW-4) |
|       | Flavonoid                   | Liquiritin (SW-5)         |
|       | Terpenoid                   | Stevioside (SW-6)         |
| Other | Aromatic hydrocarbon        | Acesulfame potassium (SW-7), saccharin (SW-8) |
| Pungent| Alkaloid                    | Nonivamide (P-1), piperine (P-2) |
|       | Aromatic hydrocarbon        | Zingerone (P-3)           |
| Salty | Salt                        | Potassium chloride (SA-1), potassium iodide (SA-2), sodium chloride (SA-3) |
24-31% A at 53-58 min, 31-37% A at 58-59 min, 37-38% A at 59-63 min, 38-55% A at 63-64 min, 55-58% A at 64-69 min, 58-100% A at 69-70 min, 100% A at 70-75 min. The mobile phase flow rate was 0.4 mL/min, and the column temperature was maintained at 30 °C. The injection volume was 5 μL. The ESI-MS was performed in both positive and negative modes with the source settings as follows: spray voltage, 3800 V (+), 3200 V (-); auxiliary gas flow rate, 15 arb; sheath gas flow rate, 35 arb; capillary temperature, 350 °C; scan range, m/z 100-1500; collision dissociation energy, 30 eV, 40 eV, 50 eV. MS data was collected with Thermo Xcalibur software.

2.3 The electrochemical e-tongue device and data collected parameters

The ASTREE e-tongue (Alpha M.O.S., France) equipped with the fifth set of sensor systems was used, and the sensor system had 7 electrochemical sensors (SRS, GPS, STS, UMS, SPS, SWS, BRS) and an Ag/AgCl reference electrode. After 7 electrochemical sensors were activated and calibrated, these sensors were put into a liquid sample to collect the data, including taste characteristics of the liquid sample. The parameters for the e-tongue data collected were as follows, the data acquisition time 120 s, acquisition period 1.0 s, acquisition delay 0 s, stirring rate 1.0 r·s⁻¹. Nine replicates for each sample, only the average of the 100-120 s data of the last 3 replicates were kept as 3 results for each sample to ensure the stability of the sensor response values, indicating that the sensor responses were stable when the RSD value of 3 results for each sample was less than 5%.

2.4 Data processing

The Xcalibur 4.0 software (Thermo Fisher Scientific, USA) was used to calculate the high-resolution accurate mass of the compounds. Based on error less than 5 ppm and MS/MS fragment matching, we identified main compounds in XXZOL, and the identified substances were used to verify the identified results.

Principal component analysis (PCA) and DFA were used to recognize the taste characteristics of liquid samples of reference components and establish the classification model of five kinds of tastes. The taste of every main component in XXZOL was discriminated by the FDA model.

3 Results and discussion

3.1 Identification of main chemical compounds in XXZOL

The negative and positive total ion chromatograms of XXZOL were shown in Figure 1. Due to their high responses in the mass spectrometer and UV spectrometer, high content compounds in XXZOL were focused on, and 15 high content compounds in XXZOL were chosen considering the compounds’ solubility in 50% ethanol. The high-resolution accurate mass and major MS/MS fragments of the above 15 main compounds were presented in Table 2, and the mass relative errors between the theoretical mass and measured mass of 15 main compounds were all smaller than 5 ppm, and the MS/MS fragments measured of each compound existed rationally and accorded with the MS/MS fragments reported in the works of literature.

Table 2. 15 main components of XXZOL identified by LC-MS/MS.

| No. | Retention time | Compound                  | Molecular formula | Ion type | Theoretical m/z | Measured m/z | Error ppm | Major MS/MS fragments |
|-----|----------------|---------------------------|-------------------|----------|-----------------|--------------|-----------|-----------------------|
| 1   | 0.66           | Mannitol                  | C₆H₁₄O₆ [M-H]-    | 181.07066| 181.07085       | 1.05         |           | 163.06062, 119.03394, 101.02320, 89.02316, 85.02821, 73.02821, 71.01254, 59.01259 |
| 2   | 0.80           | Trigonelline              | C₇H₇NO₂ [M+H]+   | 138.05495| 138.05490       | 0.36         |           | 110.06025, 94.06551, 92.04988 |
| 3   | 0.98           | Sucrose                   | C₁₂H₂₂O₁₁ [M-H]- | 341.10784| 341.10858       | 2.17         |           | 119.03384, 113.02325, 101.02320, 89.03313, 71.01256, 59.01260 |
| 4   | 1.03           | Malic acid                | C₆H₈O₅ [M-H]-     | 133.01315| 133.01302       | 0.98         |           | 133.01315, 115.00250, 89.02316, 72.99179, 71.01254 |
| 5   | 1.40           | Synephrine                | C₁₃H₁₅NO₃ [M+H]+ | 168.10191| 168.10173       | 1.07         |           | 150.09122, 135.06776, 119.04924, 91.05463 |
| 6   | 2.12           | Citric acid               | C₆H₈O₇ [M-H]-     | 191.01863| 191.01889       | 1.36         |           | 111.00758, 87.00749, 85.02821 |
| 7   | 7.59           | L-phenylalanine           | C₉H₁₁NO₂ [M+H]+ | 166.08626| 166.08623       | 0.18         |           | 131.04921, 120.08099, 103.05453 |
| 8   | 21.71          | Forsythoside E            | C₂₀H₃₀O₁₂ [M-H]- | 461.16353| 461.16635       | 2.17         |           | 205.07127, 143.03397, 135.04413, 131.03394, 101.02319, 89.03235, 71.02821, 71.01257, 59.01260 |
| 9   | 22.32          | Caffeic acid              | C₉H₈O₄ [M-H]-     | 179.03389| 179.03413       | 1.34         |           | 135.04410 |
| 10  | 25.93          | Sinapine                  | C₁₆H₂₄NO₅ [M]+   | 310.16490| 310.16473       | 0.55         |           | 251.09106, 175.03885 |
| 11  | 34.36          | Forsythoside I            | C₁₉H₂₃O₁₁ [M-H]- | 623.19705| 623.19788       | 1.33         |           | 179.03410, 161.02353, 135.04408 |
| 12  | 38.42          | Forsythoside A            | C₁₉H₂₃O₁₁ [M-H]- | 623.19705| 623.19800       | 1.52         |           | 179.03423, 161.02354, 135.04408 |
| 13  | 39.21          | Pinoresinol-4-O-β-D-      | C₂₆H₃₂O₁₁ [M-H]- | 519.18609| 519.18707       | 1.89         |           | 357.13467, 151.03914, 136.01556 |
| 14  | 42.70          | Naringin                  | C₁₇H₂₀O₉ [M-H]-  | 579.17083| 579.17081       | 0.03         |           | 271.06143, 151.00270, 119.04909 |
| 15  | 44.98          | Neohesperidin             | C₁₈H₂₂O₁₁ [M-H]- | 609.18140| 609.18231       | 1.49         |           | 301.07172 |
Furthermore, 12 main compounds were identified with the corresponding control substances in the DAD spectrometer at the detection wavelength of 254 nm, (A) the sample chromatogram, (B) the mixed control substances chromatogram, as shown in Figure 2, UV absorption spectrum and retention time for each compound in the sample were consistent with those in control substances.

**3.2 Taste judgments of 15 main components in XXZOL by electronic tongue**

**Analysis of taste resolution ability of electronic tongue**

Prepare 50% ethanol solutions with high (8.0 mg/mL), medium (5.0 mg/mL), and low (1.0 mg/mL) concentrations of

**Figure 1.** The negative (A) and positive (B) total ion chromatogram of XXZOL. 1-mannitol, 2-trigonelline, 3-sucrose, 4-malic acid, 5-synephrine, 6-citric acid, 7-L-phenylalanine, 8-forsythoside E, 9-caffeic acid, 10-sinapine, 11-forsythoside I, 12-forsythoside A, 13-pinoresinol-4-O-β-D-glucopyranoside, 14-naringin, 15-neohesperidin.
citric acid monohydrate (sour), quinine (bitter), sucrose (sweet), zingerone (pungent) and sodium chloride (salty), and collect the data of the above 15 solutions by the e-tongue, 3 replicates for each solution. PCA and DFA were applied to process the e-tongue data of 15 samples, as shown in Figure 3, 3 collected data of the same solution well-replicated, which indicated the stability of the e-tongue platform and the robustness of the e-tongue data.

In the two-dimensional PCA scatter plot of 5 reference components with different tastes, the accumulative contribution rate for the two PCs was 91.264%, in which PC1 accounted for 69.075%, and PC2 contributed 22.189%, and five reference components with different tastes were differentiated (Figure 3A). Three different concentration samples from the same reference component had small changes in e-tongue data and were all clustered into the same category in the two-dimensional PCA plot. The results showed that the e-tongue combined with PCA could distinguish the reference components with different tastes, and concentration of the sample solution had little effect on the classification of the PCA model.

In the two-dimensional DFA scatter plot of 5 reference components with different tastes, the accumulative contribution rate with the two DFs was 97.395%, in which DF1 accounted for 91.131%, and DF2 contributed 6.264%. Five reference components with different tastes were differentiated, and reference components with the same taste were clustered into one category (Figure 4). The results showed that the e-tongue combined with DFA could effectively distinguish reference components with different tastes.

Figure 2. The sample chromatogram (A) and the mixed control substances chromatogram (B) at the detection wavelength of 254 nm. 2-trigonelline, 5-synephrine, 6-citric acid, 7-L-phenylalanine, 8-forsythoside E, 9-caffeic acid, 10-sinapine, 11-forsythoside I, 12-forsythoside A, 13-pinoresinol-4-O-β-D-glucopyranoside, 14-naringin, 15-neohesperidin.

The classification model of the e-tongue data had better classification results for five reference components with different tastes than the PCA model.

The classification model of the e-tongue data could effectively distinguish the reference components with different tastes. The different concentration samples of the same reference component overlapped together in the two-dimensional PCA or DFA scatter plot, indicating that the concentration of the sample solution had little effect on the classification ability of the classification model. So 1.0 mg/mL reference component solutions were prepared in the following research, and DFA was applied to the sensor data of these solutions collected by the e-tongue platform.

DFA classification model for distinguishing components with different tastes

Prepare 1.0 mg/mL 50% ethanol solutions of reference components in Table 1, collect data according to the parameters of the e-tongue in section 2.3, and DFA was used to recognize the taste characteristics of reference component solutions with five different tastes.

In the two-dimensional DFA scatter plot of these reference component samples with 5 kinds of tastes, the accumulative contribution rate with the two DFs was 75.023%, in which DF1 was 44.389%, and DF2 was 30.634%. Reference components with different tastes were differentiated, and reference components with the same taste were clustered into one category (Figure 4). The results showed that the e-tongue combined with DFA could effectively distinguish reference components with different tastes.
Taste judgments of 15 main components in XXZOL

Prepare 1.0 mg/mL 50% ethanol solutions of 15 main components in XXZOL. The established DFA model was used to discriminate the tastes of these solutions. The taste information of 15 main components was shown in Table 3. Authentic taste of each component was evaluated by three assessors, predicted taste of each component was judged by the DFA model of the reference components in Table 1, the originated Chinese medicinal piece was obtained by the reported works of literature, and the piece taste was obtained by the Chinese Pharmacopoeia (2020 version a).

As shown in Table 3, the tastes of 12 main components were effectively recognized by the established DFA model. Except for forsythoside I, the predicted tastes of mannitol, trigonelline, sucrose, malic acid, citric acid, L-phenylalanine, forsythoside E, caffeic acid, forsythoside A, naringin, and neohesperidin were
The authentic tastes of mannitol, trigonelline, malic acid, citric acid, L-phenylalanine, caffeic acid, sinapine, forsythoside I, pinoresinol-4-O-β-D-glucopyranoside, naringin, and neohesperidin were similar to the main tastes of their respective originated Chinese medicinal pieces. The concordance rate of authentic tastes of 14 compounds with the main tastes of their respective originated Chinese medicinal pieces was 78.57%, the false rate was 21.43%.

The originated Chinese medicinal pieces were the main herbal decoctions in the prescription of XXZOL, including Stir-baked *Crataegi Fructus*, *Trichosanthis Fructus*, *Aurantii Fructus Immaturus*, *Platycodonis Radix*, *Forsythiae Fructus*, consistent with their authentic tastes. The accurate recognition rate for the above 15 main components was 73.33%, the false rate was 6.67%, and the unrecognized rate was 20%. When the number and representativeness of the reference components for the established DFA model increase, the unrecognized rate will further decrease. So the e-tongue combined with classification model can be used for predicting the tastes of different components in Chinese medicinal pieces.

For XXZOL, exclude sucrose as the additive of the preparation, the remaining 14 ingredients with high responses in the mass spectrometer, were all from Chinese medicinal pieces in XXZOL.

### Table 3. The taste information of 15 main components in XXZOL.

| Compound                   | Authentic taste | Predicted taste | Originated Chinese medicinal piece                          | Main taste of Chinese medicinal piece |
|----------------------------|-----------------|-----------------|-------------------------------------------------------------|---------------------------------------|
| Mannitol                   | Sweet           | Sweet           | Stir-baked *Crataegi Fructus*                              | Sour, sweet                           |
| Trigonelline               | Sour            | Sour            | *Trichosanthis Fructus*                                    | Sour, sweet                           |
| Sucrose                    | Sweet           | Sweet           | Additive                                                   | Sweet                                 |
| Malic acid                 | Sour            | Sour            | Stir-baked *Crataegi Fructus*                              | Sour, sweet                           |
| Synephrine                 | Pungent         | —               | *Aurantii Fructus Immaturus*                                | Bitter, sour                          |
| Citric acid                | Sour            | Sour            | Stir-baked *Crataegi Fructus*                              | Sour, sweet                           |
| L-phenylalanine            | Bitter          | Bitter          | *Platycodonis Radix*                                       | Sweet, bitter                         |
| Forsythoside E             | Sweet           | Sweet           | *Forsythiae Fructus*                                       | Bitter                                |
| Caffeic acid               | Sour            | Sour            | Stir-baked *Crataegi Fructus*                              | Sour, sweet                           |
| Sinapine                   | Bitter          | —               | Stir-baked *Descurainiae Semen Lepidii Semen*, Stir-baked *Raphani Semen* | Bitter, pungent                       |
| Forsythoside I             | Bitter          | Sweet           | *Forsythiae Fructus*                                       | Bitter                                |
| Forsythoside A             | Sweet           | Sweet           | *Forsythiae Fructus*                                       | Bitter                                |
| Pinoresinol-4-O-β-D-glucopyranoside | Bitter          | —               | *Forsythiae Fructus*                                       | Bitter                                |
| Naringin                   | Bitter          | Bitter          | *Aurantii Fructus Immaturus*                                | Bitter, sour                          |
| Neohesperidin              | Bitter          | Bitter          | *Aurantii Fructus Immaturus*                                | Bitter, sour                          |

* indicated that the taste of the compound was not effectively recognized by the established DFA model because of the far distances from the 5 taste regions in the DFA scatter plot.
Stir-baked *Descurainia Semen Lepidii Semen* and Stir-baked *Raphani Semen*. The high content compounds of the above Chinese medicinal pieces included 14 compounds in Table 3, from which 11 compounds were chosen because of consistent tastes with the main tastes of their respective originated Chinese medicinal pieces, and these compounds were more likely to be the quality markers of XXZOL. Trigonelline, malic acid, citric acid, and caffeic acid were the potential sour material bases of XXZOL. Mannitol was the potential sweet material basis of XXZOL. L-phenylalanine, sinapine, forsythoside I, pinoresinol-4-O-β-D-glucopyranoside, naringin, and neohesperidin were the potential bitter material bases of XXZOL.

4 Conclusions

In this work, 15 high content compounds in Xiao’er Xiaoji Zhike oral liquid were identified according to the high-resolution mass data and the MS/MS fragments. The e-tongue collected the sour, bitter, sweet, pungent, and salty samples, which were 1.0 mg/mL solutions of the reference compounds with above tastes, and DFA model to recognize the taste characteristics of 23 liquids samples was established to predict the tastes of 15 main components in XXZOL, and the accurate recognition rate was 73.33%, the false rate was 6.67%, and the unrecognized rate was 20%. The concordance rate of their authentic tastes with the tastes of their respective originated Chinese medicinal pieces was 78.57%, and 11 compounds were chosen because of consistent tastes with the main tastes of their respective originated Chinese medicinal pieces. Trigonelline, malic acid, citric acid, and caffeic acid were the potential sour material bases of XXZOL. Mannitol was the potential sweet material basis of XXZOL. L-phenylalanine, sinapine, forsythoside I, pinoresinol-4-O-β-D-glucopyranoside, naringin, and neohesperidin were the potential bitter material bases of XXZOL.

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