

Sustainable Food Technology

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Sustainability Spotlight Statement

In the context of a growing global pursuit of sustainable food systems, the development of natural, efficient flavor precursors and antioxidants has become a critical direction for the food industry. The use of TTCA/ARP as natural flavor precursors and antioxidants can contribute to: (1) Reducing reliance on synthetic additives, supporting cleaner labels and potentially lowering the environmental burden associated with chemical synthesis; (2) Minimizing food waste through enhanced product stability and extended shelf-life; (3) Optimizing energy efficiency in food processing, as flavor formation can be regulated under varied, potentially milder thermal conditions. Utilizing natural and renewable precursors (cysteine and xylose) in a value-adding reaction.



Unlocking the Potential of Cysteine-Xylose Maillard Reaction Intermediates as Natural Flavor Precursor: A Comprehensive Study on Flavor Regulation, Storage Stability, and Antioxidant Activity

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Abstract

1 2-Threityl-thiazolidine-4-carboxylic acid (TTCA) and Amadori rearrangement
2 product (ARP), as intermediates of xylose-cysteine (Cys-Xyl) Maillard reaction system,
3 have been proven to form desirable flavor during thermal processing. The results showed
4 that both TTCA and ARP revealed excellent environmental stability at low temperature (\leq
5 25°C), neutral pH (7), and low water activity (0.113). Under storage conditions of 40°C ,
6 pH 9, and water activity of 0.843, the loss rates of TTCA and ARP reached 7.06%/12.17%,
7 11.19%/21.25% and 35.77%/60.61%, respectively. Further research found that TTCA and
8 ARP compounds have increased Fe^{2+} chelating ability, reducing power, and free radical
9 scavenging ability within the concentration range from 0.5 to 3.0 mg/mL. Additionally,
10 different heat treatment conditions as well as addition of different exogenous amino acids
11 could regulate the flavor formation profile and characteristics of TTCA, making it adapt to
12 different real food systems, broadening its processing adaptability. Especially when the pH
13 increased from 5.5 to 8, the meaty flavor of the system gradually lightened, while flavor
14 characteristics such as popcorn flavor, burnt flavor, and nutty flavor gradually strengthened.
15 This research explored the potential of natural flavor precursors MRIs as a supplement and
16 partial alternative to synthetic additives. The findings provide a theoretical basis for
17 developing more sustainable food systems and may help reduce the environmental
18 footprint of food processing.

19 **Key words:** sustainable food additives, Maillard reaction intermediate, flavor formation
20 rules, storage stability, antioxidant characteristics.

21 **Accession Codes:**



22 D-xylose (PubChem CID: 135191); L-Cysteine (PubChem CID: 5862); Acetonitrile
23 (PubChem CID: 6342); Methanol (PubChem CID: 887); Deuterioxide (PubChem CID:
24 24602); Ammonium formate (PubChem CID: 10904).

25 **1. Introduction**

26 In the context of a growing global pursuit of sustainable food systems, the
27 development of natural and efficient additives has become a critical direction for the food
28 industry. Maillard reaction intermediates (MRIs), Amadori/Heyns rearrangement
29 compounds (ARPs/HRPs), as naturally derived compounds, hold significant potential as
30 clean-label alternatives to synthetic flavor precursors and antioxidants.^{1,2} Compared to
31 complex and variable complete Maillard reaction products (MRPs), MRIs provide a stable
32 and controllable source for generating ideal flavor and color upon thermal processing.³
33 In particular, the Cys-Xyl system yields both the conventional ARP and a unique
34 thiazolidine derivative, TTCA.^{4,5} Current studies have investigated the fundamental
35 aspects of TTCA and ARP,^{6,7} including their aqueous-phase tracing preparation
36 mechanism, characteristic patterns of flavor formation and yield enhancement.^{8,9} These
37 investigations have preliminarily demonstrated their advantages in controlled flavor
38 formation, underscoring their potential as flavor precursors.

39 Existing studies confirmed that the flavor intensity in thermal processed MRIs
40 exhibits significant advantages, however the profile of characteristic volatile compounds
41 generated from MRIs remains less diverse compared to MRPs.^{8,9} While adjusting reaction
42 parameters is a common regulatory approach, its ability to fully compensate for the lack of
43 flavor complexity may be limited.^{10–13} The addition of exogenous amino acids presents a
44 promising alternative for modulating flavor, which could promote the Strecker degradation



45 between regenerated amino acids derived from MRIs degradation and dicarbonyl
46 compounds to form characteristic flavor. ^{14,15} Nevertheless, the systematic effect of
47 different exogenous amino acids on the flavor profiles derived from specific intermediates,
48 such as TTCA/ARP, is not yet clear. Moreover, comprehensive approaches for regulating
49 the flavor profile of TTCA/ARP are currently lacking. Besides, to enable flavor formation
50 exclusively during subsequent thermal processing, MRIs must maintain stability during
51 storage. ¹⁶ However, MRIs can further degrade through rearrangement and condensation
52 pathway. ^{17,18} Thus, understanding their behavior during storage and heating is essential
53 for commercial application. Concurrently, developing natural antioxidants is crucial for
54 food preservation. ¹⁹⁻²¹ MRPs exhibit confirmed antioxidant, antiallergic, and antibacterial
55 activities, ²²⁻²⁴ due primarily to the chromophoric rings of melanoidins. ^{22,25} Certain low-
56 molecular-weight compounds in MRPs, especially reductones and heterocycles, exhibit
57 greater antioxidant activity than pure melanoidins. ¹⁹ However, the specific contribution
58 and efficacy of defined intermediate compounds like TTCA and ARP have not been
59 comparatively evaluated against MRPs.

60 Therefore, to address the current research gap regarding TTCA and ARP, this study
61 adopts a unique perspective to investigate the regulatory mechanisms of their flavor
62 characteristics, storage stability, and antioxidant properties, and aims to comprehensively
63 evaluate the dual-function potential of TTCA and ARP as flavor precursors and
64 antioxidants through the following objectives: (1) Systematically assess the storage
65 stability of TTCA and ARP under different environmental stresses, providing essential data
66 for their commercial storage; (2) Elucidate the directional regulation of TTCA thermal
67 degradation flavor profiles by temperature, pH, and various exogenous amino acids,



68 thereby extending their processing adaptability; (3) Evaluate and compare the in vitro
69 antioxidant activities of TTCA/ARP with those of MRPs, clarifying their roles and
70 potential in different antioxidant pathways. This work provides essential data for the
71 commercial storage and application of TTCA/ARP, extends their processing adaptability,
72 and clarifies their dual-function potential as natural flavor precursors and potential
73 antioxidants, thereby supporting their development as sustainable food additives.

74 2. Materials and Methods

75 2.1 Chemicals.

76 Analytical grade L-cysteine, D-xylose, sodium hydroxide, 1,2,4-benzenetrio,
77 phenanthroline, potassium ferricyanide, trichloroacetic acid, FeCl₃, DPPH, ethanol, 1,10-
78 phenanthroline, FeSO₄, ascorbic acid, lecithin, thiobarbituric acid, H₂O₂, NaOH and HCl
79 were purchased from Sigma-Aldrich Chemical Co. (Shanghai, China). LC-MS grade water,
80 acetonitrile, deuterioxide ammonium formate and 1,2-dichlorobenzene were purchased
81 from Sinopharm Chemical Reagent Co. Ltd (Shanghai, China).

82 2.2. Synthesis, Purification and Characterization of Isomeric TTCA and ARP.

83 The preparation and characterization of TTCA and ARP purities were carried out
84 referring to the methods of our previous studies.^{4,26} An equimolar solution of Cys and Xyl
85 (0.0827 mol/L) was prepared using deionized water. The prepared solution was maintained
86 at pH 7.4 ± 0.01 using NaOH (6 mol/L), followed by the heat treatment for 40 min at 90 °C.
87 The reaction vessel was promptly transferred to an ice-bath to terminate the reaction
88 immediately. The obtained reactant was employed to get TTCA and ARP purities, which
89 were preliminarily separated using an ion exchange resin (H⁺ Dowex 50WX4, 200-400
90 mesh, Acros Organics, SERVA, J&K, Beijing, China) with ammonium hydroxide (0.2



91 mol/L) as the eluent, and the further purification was carried out through a semi-preparative
92 RP-HPLC equipped with an Xbridge Amide column (4.6 mm × 150 mm, 3.5 μm; Waters
93 Co., Milford, MA, USA). The identification and characterization of TTCA and ARP
94 purities were performed using UPLC-ESI-MS (Waters Synapt MALDI Q-TOF MS, USA)
95 and NMR (Bruker DRX 400 MHz spectrometer). The specific procedures of the methods
96 mentioned above were accomplished referring to our published study. The mass spectral
97 and NMR spectral information were listed in the Supporting Information (Figure S1, Table
98 S1).

99 *2.3. Assessment of the browning index of the Maillard reaction solution.*

100 To assess the browning degree of the melamine reaction solution under variable
101 temperatures, the absorbance at 420 nm (A_{420}) was typically measured using a UV-visible
102 spectrophotometer and served as the browning index. The absorbance was determined by
103 diluting the sample to a suitable number of times so that the A_{420} value fell as close as
104 possible to a reasonable range (0.05 to 1.0).

105 *2.4. Construction of Maillard thermal reaction model system using TTCA as the base* 106 *material.*

107 Maillard thermal reaction model systems were established with TTCA as the base
108 material. The pH of the TTCA solution (10 mmol/L) was adjusted to 5.5, 7, 8, respectively,
109 using NaOH (6 mol/L). The prepared samples were then treated at varying temperatures
110 (100°C, 120°C, and 140°C) for 120 minutes.

111 Additionally, an equimolar amount of different exogenous amino acids (Gly, Ala, Met,
112 Leu, Ser, Thr, Pro, Val, Phe, His, Lys, Glu, Asp) was added to the TTCA solution (10



113 mmol/L). The pH of the system was adjusted to 7.0 using NaOH (6 mol/L), and the thermal
114 reaction was conducted at 120°C for 120 minutes.

115 Using high-temperature/high-pressure resistant glass reaction vessels set in an oil bath,
116 the aforementioned thermal reactions were carried out. Following this, to halt the reactions,
117 all mixtures were immediately transferred to an ice bath. Afterwards, samples were
118 gathered and kept under 4 °C for further analysis.

119 2.5. Determination of storage stability of TTCA and ARP.

120 To investigate the effect of different storage temperatures on the stability of TTCA
121 and ARP intermediates in solid and solution forms, the solid pure lyophilized powder of
122 TTCA and ARP was placed in self-sealing bags or dissolved in deionized water and
123 configured as a solution with a concentration of 100 mmol/L, and placed in the ambient
124 temperatures of 4°C, 25°C and 40°C, and the changes in concentration, A_{420} , A_{294} , and color
125 were recorded at intervals of 10 days in the storage process. The concentration, A_{420} , A_{294}
126 and color changes during storage were recorded every 10 days, and the storage stability
127 was investigated over a 60-day period.

128 In addition, the storage stability of the TTCA and ARP system was observed by
129 varying the pH of the system while maintaining the same ambient temperature, and the
130 same metrics were also examined. For TTCA and ARP solid samples, the effect of ambient
131 humidity on their storage stability was investigated. A fixed mass of dried powder of the
132 intermediate was taken in an open glass petri dish and transferred to a sealed desiccator,
133 where the bottom of the desiccator was filled with different saturated salt solutions
134 configured to maintain the ambient moisture activity of the system, where the comparison



135 table of the moisture activity of different saturated salt solutions at 25°C is shown in Table
136 S2.

137 2.6. Determination of antioxidant properties of intermediates.

138 (1) Measurement of Fe²⁺ chelating capacity.

139 Determination of the Fe²⁺ chelating capacity was conducted based on a previously
140 established method.²⁷ 1 mL of sample was combined with 1.85 mL of deionized water,
141 followed by the addition of 0.05 mL of FeCl₂ phenanthroline solution (5 mmol/L). After
142 standing at 28 °C for 30 seconds, the mixture was allowed to react for an additional 10
143 minutes under the same temperature. It was then centrifuged at 3000 rpm for 5 minutes.
144 Absorbance of the supernatant was measured at 562 nm using a UV-Vis spectrophotometer.
145 A blank control was prepared by replacing the sample with deionized water and analyzed
146 identically. The chelating capacity was calculated using the following formula:

147
$$\text{Fe}^{2+} \text{ chelation Rate} = \frac{A_0 - A_1}{A_0} \times 100\%$$

148 Where A₀ and A₁ are the absorbance of the blank and sample, respectively.

149 (2) Measurement of reducing power.

150 The assay was initiated by mixing 1 mL of sample solution with 2.5 mL of phosphate
151 buffer (pH 6.6) and 2.5 mL of 1% potassium ferricyanide solution. The resulting mixture
152 was incubated in a 50 °C water bath for 20 minutes, followed by the addition of 2.5 mL of
153 10% trichloroacetic acid. After centrifugation at 3000 rpm for 10 minutes, 2.5 mL of the
154 supernatant was collected and combined with 2.5 mL of 0.1% FeCl₃ solution and 2.5 mL
155 of deionized water. Following a 10-minute reaction period, the absorbance at 700 nm (A₇₀₀)
156 was measured using a UV-Vis spectrophotometer to indicate the sample's reducing
157 power.²⁸



158 (3) Measurement of DPPH free radical scavenging capacity.

159 The assay was initiated by combining 3 mL of 100 $\mu\text{mol/L}$ DPPH ethanol solution
160 with 1 mL of the sample. The mixture was then incubated in the dark at room temperature
161 for 25 minutes. Its absorbance at 517 nm (A_i) was measured using a UV-visible
162 spectrophotometer, with ethanol serving as the blank. Similarly, the absorbance of a control
163 mixture containing 1 mL of ethanol and 3 mL of DPPH solution was recorded as A_0 .
164 Additionally, a sample background absorbance (A_j) was obtained by measuring a mixture
165 of 1 mL of sample and 3 mL of ethanol. The DPPH radical scavenging capacity was
166 calculated using the following formula:

$$167 \quad \text{DPPH}\cdot \text{ Clearance Rate} = \left(1 - \frac{A_i - A_j}{A_0}\right) \times 100\%$$

168 (4) Measurement of hydroxyl radical scavenging capacity.

169 For the assay, 1 mL of 2.5 mmol/L 1,10-phenanthroline was combined sequentially
170 with 2 mL of phosphate buffer (pH 7.40), 1 mL of 2.5 mmol/L FeSO_4 solution, 1 mL of
171 deionized water, and 1 mL of 20 mmol/L H_2O_2 . Following a 60-minute incubation at 37°C
172 in a water bath, the absorbance of the mixture at 536 nm was recorded immediately on a
173 UV-Vis spectrophotometer. The scavenging capacity against hydroxyl radicals was
174 calculated using the formula below:

$$175 \quad \cdot\text{OH Clearance Rate} = \frac{A_s - A_1}{A_0 - A_1} \times 100\%$$

176 where A_s represents the absorbance of the sample; A_1 denotes the absorbance of the
177 control reaction solution, which containing 1, 10-phenanthroline, FeSO_4 and H_2O_2 ; and A_0
178 corresponds to the absorbance of the blank solution, composed solely of FeSO_4 and 1, 10-
179 phenanthroline.⁷



180 (5) Measurement of superoxide anion radical scavenging capacity.

181 The method is grounded in the principle that antioxidants bind to superoxide anion
182 radicals, forming stable species and thereby terminating the radical chain reaction, which
183 releases chromogenic products. According to this principle, 9 mL of 50 mmol/L Tris-HCl
184 buffer (pH 8.2) was combined with 0.5 mL sample and incubated in a water bath at 25°C
185 for 20 minutes. Immediately thereafter, 0.04 mL of a pre-warmed (25°C) 1,2,4-
186 benzenetriol solution (prepared by dissolving 0.45 mmol of 1,2,4-benzenetriol in 10 mL of
187 10 mmol/L hydrochloric acid) was added. After a 3-minute standing period, one drop of
188 10 mmol/L ascorbic acid solution was introduced. Following an additional 5-minute
189 incubation, the mixture's absorbance at 327.2 nm (A_{327}) was measured using a UV-Vis
190 spectrophotometer. The superoxide anion radical scavenging capacity was calculated using
191 the following formula:

192
$$\text{O}_2^\bullet \text{ Clearance Rate} = \frac{A_0 - A_1}{A_0} \times 100\%$$

193 Where A_0 represents A_{327} of the blank and A_1 represents the A_{327} of sample.⁷

194 (6) Measurement of lipid peroxidation inhibition capacity.

195 A 10.0 mg/mL lecithin dispersion (denoted S_1) was prepared in 0.01 mol/L phosphate
196 buffer (pH 7.4) using a magnetic stirrer. Separately, solution S_2 was prepared by dissolving
197 15 g trichloroacetic acid, 0.37 g thiobarbituric acid, and 2 mL concentrated hydrochloric
198 acid in deionized water, followed by dilution to a final volume of 100 mL. For the assay,
199 1.0 mL of S_1 was mixed sequentially with 1.0 mL of 400 $\mu\text{mol/L}$ FeCl_3 solution, 1.0 mL
200 of 400 $\mu\text{mol/L}$ ascorbic acid solution, and 1.0 mL of sample solution. The mixture was
201 incubated at 37°C in a water bath for 60 minutes. Subsequently, 2.0 mL of S_2 was added,
202 and the mixture was heated at 95°C for 15 minutes. After cooling in an ice-water bath for



203 10 minutes and centrifugation, the absorbance of the supernatant at 532 nm (A_{532}) was
204 measured using a UV-Vis spectrophotometer. The lipid peroxidation inhibition rate was
205 calculated as follows:

$$206 \quad \text{Inhibition Rate} = \frac{A_0 - A_1}{A_0} \times 100\%$$

207 Where A_0 and A_1 denote the absorbance of the blank (deionized water as substitute)
208 and the sample, respectively.⁷

209 *2.7. Identification and Quantification of Characteristic Volatile Compounds.*

210 The analysis of volatile flavor compounds was conducted using headspace solid-phase
211 microextraction coupled with gas chromatography-mass spectrometry (HS-SPME-GC-
212 MS). The sample solution (3 g) was added with 3 μL internal standard (1,2-
213 dichlorobenzene, 0.018 $\mu\text{g}/\mu\text{L}$ in methanol), which was then placed in the 20 mL headspace
214 vial. The extraction and desorption procedures followed a previously published method,¹⁵
215 employing a divinylbenzene/carboxen/polydimethylsiloxane (CAR/PDMS/DVB, 75 μm)
216 SPME fiber. Compound separation and identification were achieved using an Agilent
217 7890B gas chromatograph interfaced with a 5977B mass selective detector. Following a
218 10-minute thermal desorption at 250 $^{\circ}\text{C}$ in the injection port, the volatiles were transferred
219 onto the analytical column. Chromatographic separation was performed on a 30 m DB-
220 Wax capillary column (0.25 mm i.d., 0.25 μm film thickness). Specific instrumental
221 conditions were adopted from our prior research.¹⁵

222 Compound identification was based on matching mass spectra against the NIST 17,
223 and WILEY 07 databases, comparing calculated Kovats retention indices (RI) with
224 literature values, and referencing literature data. To facilitate accurate RI matching, the
225 retention indices of target compounds were calculated by analyzing a $\text{C}_7\text{-C}_{30}$ *n*-alkanes



226 series under identical chromatographic conditions. For compounds with available reference
227 standards, their contents were determined by the standard curve method with internal
228 standard calibration (see Table S3); for those without reference standards, quantification
229 was performed just using the internal standard method.

230 2.8. Data analysis.

231 We performed all experiments in triplicate and report data as mean \pm standard
232 deviation. Data processing involved the following steps: 1) analysis using SPSS Statistics
233 22.0 and Microsoft Excel 2010; and 2) multiple comparison via Duncan's Multiple Range
234 Test, where a *p*-value of less than 0.05 indicated statistical significance.

235 3. Results and Discussion

236 3.1 Regulating the characteristic flavors formation of TTCA intermediate through
237 parameter regulation.

238 Given the established similarity in flavor formation pathways between TTCA and
239 ARP (differing primarily in rate), TTCA was selected to investigate the effects of
240 temperature and pH. As a key parameter, temperature is a key kinetic parameter in
241 controlling the reaction progress. The appropriate increase of temperature is beneficial for
242 accelerating effective collisions between substrates and the thermal degradation of
243 cysteine.²⁹ Comparatively studying of heated TTCA models at 100°C, 120°C, and 140°C
244 revealed that the total volatile flavor substances increased markedly, with sulfur-containing
245 compounds showing a leapfrog increase (Table S4, Figure 1a). The content of furfural (the
246 dominant furan) increased significantly, reaching 11.039 $\mu\text{g/L}$ at 140°C, which was 3.3
247 times that at 100°C. The concentrations of 2-methyl-3-furanthiol and 2-furfurylthiol also
248 increased accordingly. This is because high temperatures promote the enolization of ARPs,



249 leading to the rapid generation and accumulation of the direct precursors like 3-DX and 1-
250 DX for furfural and 4-hydroxy-5-methyl-3(2H)-furanone (Figure 1f).³⁰ They also increase
251 the generation of furan and 2-methylfuran, which are precursors to thiophene compounds
252 (Figure 1f). Furthermore, elevated temperatures intensified the thermal degradation of
253 TTCA and free Cys (Figure S2, S3), resulting in increased release of H₂S. The boosted
254 supply of these key precursors (α -dicarbonyls, furans, H₂S) strengthened their interactions,
255 thereby promoting the formation of corresponding thiols and thiophenes (Figure 1f).

256 Key steps such as enolization, retro-aldolization, and Strecker degradation are pH-
257 dependent, the activity of key precursors like H₂S and NH₃ also varies with pH values.³
258 Therefore, the influence of different reaction pH (5.5, 7, and 8) on the TTCA thermal
259 reaction system was studied. The characteristic volatile compounds were predominantly
260 sulfur-containing (Table S5). Figure 5b details the concentrations of key flavor classes.
261 Under acidic condition (pH 5.5), thiols were most abundant, followed by thiophenes; no
262 nitrogen-containing heterocycles were detected. As pH increased from 5.5 to 8, thiol and
263 furan contents decreased, while thiazoles increased sharply to 10.952 $\mu\text{g/L}$. Notably, 2-
264 acetylthiazole (nutty/popcorn odor) increased significantly with pH, comprising 27.01% of
265 total thiazoles at pH 8. Pyrazines were only detected at pH 8 (0.838 $\mu\text{g/L}$), a much lower
266 concentration compared to sulfur-containing flavors. Post-reaction, the system pH dropped
267 (to 5.1, 4.9, and 4.7, respectively), due to organic acid formation from accelerated TTCA
268 degradation at higher initial pH. Under acidic conditions, TTCA undergoes 1,2-enolization,
269 generating thiol and furan precursors (Figure 1f).¹⁵ The low thiazoles and pyrazines
270 contents under acidic conditions is due to the superior competitiveness of H₂S over
271 NH₃/amines in capturing reactive α -dicarbonyl intermediates including glyoxal,



272 methylglyoxal, 2,3-butandione, and so on.¹⁷ In contrast, alkaline conditions enhanced the
273 nucleophilicity of deprotonated NH₃/amines, enabling them to successfully compete for
274 intermediates and drive the formation of thiazoles and pyrazines, imparting roasted notes.
275 ¹⁷ Alkaline conditions also favor the regeneration of Cys during TTCA degradation,
276 promoting Strecker degradation with α -dicarbonyls and further amplifying
277 thiazoles/pyrazines formation (Figure 1f).³

278 In summary, increased temperature universally intensified flavor formation by
279 elevating precursor supply and reaction rates. In contrast, increased pH altered the
280 fundamental reaction pathway, shifting the flavor profile from sulfur-rich/meaty notes
281 toward nitrogen-containing heterocycles associated with roasted characteristics. These
282 findings confirmed that targeted parameter control (temperature, pH) can effectively steer
283 the flavor formation pathways of TTCA, demonstrating its adaptable potential for
284 generating diverse flavor profiles under different processing conditions.

285 3.2 Addition of exogenous amino acids regulating the formation of characteristic flavors
286 of TTCA reaction system.

287 While TTCA intermediates produce intense flavors surpassing those of typical MRPs,
288 their volatile compound diversity is limited.³¹ The formation of key flavor compounds such
289 as pyrazines, pyridines, pyrroles, thiazoles, and Strecker aldehydes may require sufficient
290 exogenous amino acids to participate in subsequent thermal reactions of the
291 intermediates.^{15,31} To further elucidate the effects of different amino acids, types with
292 varying isoelectric points and side chain groups were selected to combine with TTCA for
293 thermal reactions. Browning intensity was significantly higher ($p < 0.05$) in systems with



294 added amino acids than in the TTCA control (Figure 1c), likely because amino acids react
295 with carbonyl compounds in later stages to form colored polymers.³²

296 Volatile analysis revealed that all systems predominantly generated sulfur-containing
297 compounds (Table 1, Figure 1d, e), driven by the high reactivity of H₂S and NH₃ released
298 from TTCA degradation (Figure 1f).¹⁰ Results showed that side chains of the added amino
299 acids had a negligible influence on sulfur-containing flavor formation (Table 1). High
300 levels of 2-methyl-3-furanthiol and 2-furfuryl thiol were detected across all systems. This
301 consistent presence indicated that added amino acids did not inhibit the key pathway
302 leading to these furans—namely, the dehydration and cyclization of deoxyosones during
303 TTCA degradation (Figure 1f).³³ Furthermore, sulfur heterocycles requiring the
304 participation of α -dicarbonyls (e.g., 2-methyl-thiophene, thieno[3,2-b]thiophene, thiazole)
305 were detected and were common to all systems supplemented with Gly, Lys, or Glu (Figure
306 1f).³⁴ This reinforces that the core pathway for sulfur-containing flavor generation is
307 governed by α -dicarbonyl formation rather than by the specific side chains of the added
308 amino acids. The addition of the three exogenous amino acids can promote the retro-aldol
309 cleavage of deoxyosones, yielding more short-chain α -dicarbonyl compounds and
310 reductive ketones, which expanded flavor diversity (including sulfur compounds and
311 pyrazines), and contributed to enhanced browning.¹³ Gly supplementation produced the
312 greatest variety of flavor compounds, possibly due to its propensity to form short-chain
313 reactive aldehydes.³⁵ Lys significantly increased flavor variety ($p < 0.05$), likely because
314 its additional amino group boosted α -dicarbonyl formation via reactions with sugar
315 chains.³⁶



316 Compared to the TTCA control, systems with Gly, Lys, or Glu showed reduced
317 thiazole content/variety but increased nitrogenous heterocycles, confirming greater NH_3
318 participation in pyrazine formation (Figure 1f).³⁷ The TTCA-Lys thermal reaction system
319 could generate pyrazines. At the system pH (7.0), some Lys remained undissociated and
320 acts as a nucleophile, reacting with sugar residues to produce short-chain α -dicarbonyls
321 that subsequently formed pyrazines³⁸(Figure 1f). In contrast, TTCA-His produced less
322 meat or roasting aroma (Table 1), as its imino group is less reactive than amino group to
323 form carbonyl compounds as a nucleophile.²⁹ Moreover, the imidazole group with strong
324 basic properties has lower reactivity and is less likely to participate in the formation of
325 flavor compounds.³⁵ In addition to Lys, the TTCA thermal reaction system supplemented
326 with Glu also detected a considerable amount of pyrazine compounds, which is consistent
327 with previous research.³⁹ Beyond pyrazines, pyrroles were detected in all three systems,
328 linked to reactions between α -dicarbonyls (MGO/GO) and NH_3 (Figure 1f).⁴⁰ Among them,
329 2-methylpyrrole and pyrrole were synthesized via an aldol condensation between
330 MGO/GO and acetaldehyde, followed by nucleophilic addition with NH_3 and dehydration.
331 2-Pyrrolicarboxaldehyde was produced by nucleophilic addition of NH_3 to a 3-DX isomer,
332 followed by dehydration and cyclization (Figure 1f).⁴¹ The large amount of pyridine
333 compounds in the TTCA-Lys reaction system was generated by the Strecker degradation
334 reaction between Lys and α -dicarbonyl compounds, followed by subsequent dehydration
335 reactions (Figure 1f).⁴²

336 Furans (mainly 2-methylfuran and furfural) were formed in TTCA reaction systems
337 with Gly, Lys, or Glu but not in a TTCA-Cys system, indirectly confirming the critical role
338 of H_2S from Cys degradation in diverting precursors toward sulfur-containing flavors. In



339 summary, supplementing TTCA with Gly or Lys enriched the overall flavor profile (sulfur
340 compounds and pyrazines), while Glu specifically promoted pyrazine formation ($p < 0.05$).
341 Exogenous amino acids help shift the reaction equilibrium by ensuring sufficient α -
342 dicarbonyl precursors for key flavor pathways.⁴³ Simultaneously, incorporating various
343 exogenous amino acids can markedly enhance the browning development in the TTCA
344 thermal reaction system. These results suggest that combining Gly, Lys, or Glu with TTCA
345 could enhance TTCA's versatility for generating desired colors and flavors in diverse food
346 applications.

347 3.3 Extremely high environmental stability of TTCA and ARP intermediate to act as
348 natural flavoring additives.

349 The storage environment is a key factor affecting the structure, content and
350 physicochemical properties of all kinds of chemical products in the storage process.⁴⁴ In
351 this study, the storage stability of TTCA and ARP was investigated. The key finding is
352 that TTCA exhibits superior storage stability compared to ARP across all conditions tested,
353 attributable to its stable cyclic molecular structure.³ Both intermediates were highly stable
354 at 4°C and 25°C. Degradation became noticeable only under stress conditions: at 40°C,
355 TTCA concentration decreased by 7.06% after 60 days, less than ARP's 12.17% loss. The
356 increased degradation at elevated temperature is linked to water autoionization,⁴⁵ which
357 catalyzes the enolization and ring-opening of ARP.⁴⁴ Corresponding increases in A_{294} and
358 A_{420} values at 40°C (Figure 2c and 2d) indicate the formation of carbonyl intermediates
359 and browning products.²¹

360 The stability of TTCA and ARP solutions stored at pH 5.5, 7 and 9 was monitored for
361 60 days at room temperature (Figures 2e-h). The intermediates maintained stable



362 concentrations under acidic and neutral conditions (pH 5.5, 7). In contrast, at pH 9, the
363 concentrations of TTCA and ARP decreased by 11.19% and 21.25% after 60 days,
364 respectively. Meanwhile, the rise of A_{294} and A_{420} also indicate the alkaline-promoted
365 degradation into downstream products. The slight decrease in TTCA concentration at pH
366 5.5—accompanied by opposing trends in A_{420} (increase) and A_{294} (decrease)—suggested a
367 transformation distinct from degradation, potentially due to acid-catalyzed hydrolysis or
368 reversion to *N*-xylosylamine.⁴⁵ This phenomenon is attributed to the formation of
369 rearrangement reaction by deprotonation of imine positive ions, so the presence of H^+ under
370 acidic conditions could promote the reverse reaction.²¹

371 Moisture content and water activity values of food products are key indicators of their
372 shelf life and eating quality.⁴⁶ The stability of solid TTCA and ARP was assessed for 60
373 days under varying humidity levels. When stored under dry conditions (a_w 0.113) for 60
374 days, the TTCA content decreased by only 13.5% (Fig. 2i). This exceptional stability
375 implies lower storage losses, reduced packaging demands, and potentially extended shelf-
376 life in food applications. Elevated moisture activity significantly accelerated the
377 degradation and browning of both TTCA and ARP. Colorimetric analysis (L^* , a^* , b^* , ΔE)
378 showed that increased a_w led to darker samples (decreased L^*) and more pronounced
379 overall color change (ΔE) (Table 2). Notably, the ΔE values for ARP were consistently
380 higher than for TTCA under all humidity conditions, further corroborating TTCA's
381 superior stability. A semi-quantitative estimate indicates TTCA's degradation rate in high
382 humidity (a_w 0.843) is approximately 60% slower than that of ARP (Figure 2j).

383 In summary, TTCA possesses significantly greater environmental stability than ARP.
384 Optimal storage conditions are low temperature, acidic-to-neutral pH, and low humidity,



385 while high temperature, alkaline pH, and high moisture accelerate degradation. These
386 results confirm TTCA's extended effective shelf life and underscore the necessity of cool,
387 dry, and sealed storage for practical application as a stable flavor precursor.

388 3.4 Antioxidant properties of MRIs demonstrate promising potential as alternative to
389 synthetic antioxidants.

390 The antioxidant capacities of TTCA and ARP intermediates, complete Maillard
391 reaction products (MRPs) were evaluated in this study. The chelating ability of Fe^{2+} helps
392 to reduce the occurrence of lipid oxidation and is an important indicator for evaluating
393 antioxidants.⁷ A concentration-dependent Fe^{2+} chelating ability was observed for TTCA
394 and ARP (Figure 3a). Fe^{2+} can coordinate with the carbonyl oxygen on the xylose segment
395 of Amadori compounds, forming a five-membered chelating ring.⁷ Compared to the ARP
396 molecule, the TTCA compound can provide stronger overall molecular polarity and more
397 polar sites (including hydroxyl and carbonyl groups) in its possible tautomeric forms
398 (Table S1).³⁶ This enhanced polar characteristic allowed the TTCA molecule to interact
399 more effectively with solvent water molecules in aqueous solutions through electrostatic
400 interactions and dynamic hydrogen-bonding networks, thereby forming more stable
401 solvated complexes.⁴⁷ Therefore, TTCA compound molecules have more sites that can
402 form complexes with Fe^{2+} , and the complexes formed may be more stable than those
403 formed by ARP, which likely directly leads to the higher Fe^{2+} chelating ability of TTCA
404 (71.35%) compared to ARP (51.03%) (Figure 3a). Besides, MRPs always revealed a higher
405 Fe^{2+} chelating ability than TTCA and ARP compounds. Some melanoidins in MRPs can
406 be natural free radical scavengers in the food system.



407 The reducing power, reflecting a substance's electron transfer ability, was measured
408 to further evaluate antioxidant activity.⁷ A positive correlation was observed between the
409 concentration of MRIs (0.5-3 mg/mL) and their reducing power (Figure 3b). This can be
410 attributed to reductive ketones and hydroxyl-containing compounds in the intermediates,
411 which can donate electrons or hydrogen atoms to reduce the ferricyanide complex.²³
412 Notably, ARP exhibited stronger reducing properties than TTCA. This difference likely
413 stems from the role of free thiol groups in ARP, which possess strong electron-donating
414 ability due to their S–H bonds. In contrast, the thiol groups in TTCA are involved in
415 forming a stable five-membered ring, leaving no exposed free thiols available for
416 reduction.⁴⁸ Furthermore, MRPs exhibited stronger reducing power compared to
417 intermediates at all tested concentrations, with a rapid increase as concentration rose, which
418 is consistent with the existing research results.⁴⁷

419 The chelating action on transition metals and the reducing action can inhibit lipid
420 oxidation. Within the concentrations of 0.5 to 2.5 mg/mL, ARP/TTCA inhibited lipid
421 peroxidation more than the MRPs, indicating ARP's better efficacy (Figure 3c). As the
422 concentration increased to 3.0 mg/mL, the inhibition rate for lipid peroxidation of MRPs
423 increased from 2.18% to 36.39%, exceeding that of ARP (29.18%) and TTCA (31.30%) at
424 3.0 mg/mL (Figure 3c). The increased concentration of TTCA or ARP beyond a certain
425 point did not lead to a corresponding significant improvement in the inhibition of lipid
426 peroxidation. This observed plateau in efficacy could result from several factors, such as
427 potential solubility limitations of TTCA and ARP comparing to MRPs (rich in aldehydes
428 and ketones) in the emulsion system,^{27,30} or the attainment of a saturation point for their
429 antioxidant action under these specific conditions. The results confirmed that TTCA or



430 ARP could be one reason for the antioxidant activity of Maillard reaction during food
431 processing and storage. However, at higher concentrations, the MRPs demonstrated
432 superior performance compared to their precursors, suggesting that they possess better
433 emulsion system compatibility and contain a greater number of compounds with strong
434 antioxidant activity.

435 The DPPH free radical scavenging ability typically reflects the hydrogen-donating
436 ability of antioxidants.⁴⁹ The scavenging rates of TTCA and ARP increased with its
437 concentration during the range of 0.5 to 3.0 mg/mL (Figure 4a). Their effect was weaker
438 than that of MRPs, and considerably lower than ascorbic acid, which achieved nearly 100%
439 scavenging across the same range. Similar concentration-dependent trends were observed
440 for superoxide anion and hydroxyl radical scavenging (Figures 4a and 4b). At a
441 concentration of 3.0 mg/mL, the superoxide anion radical scavenging activities of TTCA
442 and ARP are functionally comparable to that of 0.5 mg/mL ascorbic acid, suggesting that
443 approximately 6 g of TTCA could theoretically replace 1 g of ascorbic acid in antioxidant
444 applications. MRPs consistently demonstrated stronger radical scavenging ability than the
445 intermediates, with activity rising sharply at higher concentrations (Figures 4a, 4b and 4c),
446 consistent with prior reports.⁴⁷ This enhanced activity is attributed to the complex mixture
447 of cross-linked polymers, aldehydes, ketones, and heterocyclic compounds formed in later
448 Maillard stages, which exert antioxidant effects through multiple pathways.²⁷ In summary,
449 while MRIs such as TTCA and ARP contribute to controlled flavor formation, the final
450 MRPs exhibit superior antioxidant properties. Therefore, a combination of MRIs and
451 MRPs holds potential as dual-function ingredients, serving as both natural flavor enhancers
452 and antioxidants in food systems.



453 TTCA and ARP compounds, derived from xylose and cysteine, exhibit antioxidant
454 properties distinct from a simple xylose-cysteine mixture, indicating that structural
455 differences at the molecular level are responsible.⁵⁰ While free sugars like pentoses,
456 hexoses, and ketoses predominantly exist in a closed-ring form in solution, with minimal
457 reactive open-chain carbonyls.⁵¹ For MRIs, the conjugated electron cloud structure
458 between the enol and aminium ion leads to an equilibrium of isomeric interconversion
459 between their open-ring and closed-ring forms, with the open-chain conformation
460 becoming a significant configuration in aqueous media.⁵² This provides abundant free
461 carbonyl groups capable of donating electrons to neutralize free radicals.⁷ Based on this, a
462 possible pathway for the intermediates to scavenge free radicals is proposed (Figure 4d).
463 TTCA and ARP can donate hydrogen atoms to free radicals, stabilizing them and
464 terminating chain reactions. The radical electron can localize on the carbon adjacent to the
465 carbonyl group on the xylose residue, where resonance and intramolecular hydrogen
466 bonding lead to a thermodynamically stable intermediate. Alternatively, the carbonyl group
467 itself may be attacked, with the radical electron transferring to the oxygen atom, followed
468 by stabilization via intramolecular hydrogen bonding.^{7,53,54} Research also found that the
469 free radical scavenging ability of ARP is consistent with its electron-donating or hydrogen-
470 donating properties, and specifically proposed that its metal chelating ability may also be
471 related to this principle.³⁹ Although this study indicates that TTCA and ARP exhibited
472 certain antioxidant activity, its long-term stability in complex food systems and the
473 production costs at scale still require further evaluation. Therefore, MRIs can serve as
474 natural flavor precursors to achieve controlled formation of processing flavors and provide
475 partial antioxidant function, suggesting potential practical value.



476 4. **Conclusion**

477 This study demonstrates that the Xyl-Cys MRIs including TTCA and ARP could act
478 as dual-functional, sustainable platform, which integrating efficient flavor precursor
479 properties with certain natural antioxidant activity. The flavor profile released from TTCA
480 can be regulated from "meaty" to "roasted nutty" notes through simply modulating thermal
481 processing parameters (pH, temperature). The addition of exogenous amino acids can
482 further enrich the diversity of roasted flavor compounds, such as pyrazines. This
483 significantly enhances the processing adaptability of the flavor precursors of MRIs.
484 Compared to MRPs, TTCA or ARP exhibits superior stability for commercial application,
485 which can retain over 90% of its content after 60 days of storage under ambient, neutral,
486 and dry conditions. This stability ensures consistent performance and reliability during the
487 storage, transportation, and use of TTCA or ARP as an ingredient, significantly reducing
488 the risks of raw material loss and flavor variability due to degradation. TTCA and ARP
489 compounds revealed increased Fe²⁺ chelating ability, reducing power, and free radical
490 scavenging ability with increasing concentration, demonstrating certain antioxidant
491 capabilities. By combining flavor enhancement with moderate oxidative stability
492 contributed by MRIs, this approach provides a new direction for "clean-label" food
493 development that may enable a partial reduction in reliance on synthetic antioxidants and
494 flavor additives. In summary, this work supports the potential of TTCA or ARP to act as a
495 sustainable platform for natural flavoring and supporting antioxidant properties. It offers a
496 feasible strategy for improving food quality and stability and aligns with the pursuit of
497 greener food processing practices.

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504 **AUTHOR CONTRIBUTIONS**

505 Di Kang and Songjin Zheng conceived the study, designed and supervised the research.
506 Yun Zhai and Yuying Fu outlined the study, performed the literature search, and prepared
507 the original draft and figures. Lin Jiang, Yuan Hu, Haifeng Wang and Teng Li helped to
508 prepare the figures and the draft. All authors have agreed to the content of the article and
509 approved the final version to be published

510 **CONFLICT OF INTEREST**

511 The authors declare no competing financial interest. This article does not contain any study
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690 TABLES

Table 1. Characteristic flavor compounds (µg/L) derived from reaction between different amino acids and TTCA

| Compounds | RI ^a | RI ^b | Control | Neutral | | | | Acidic | | Basic | |
|--|-----------------|-----------------|--------------|---------------|---------------|--------------|---------------|---------------|--------------|---------------|---------------|
| | | | | Cys | Gly | Ala | Met | Glu | Asp | His | Lys |
| Sulfur-containing compounds | | | | | | | | | | | |
| Thiols and sulfides | | | | | | | | | | | |
| Dimethyl disulfide | 1085 | 1037 | ND | ND | ND | ND | 0.956±0.013a | ND | ND | ND | ND |
| 3-Mercapto-2-butanone* | 1273 | 1283 | 1.798±0.025a | ND | ND | ND | ND | ND | ND | ND | ND |
| 2-Methyl-3-furanthiol* | 1302 | 1305 | 1.388±0.207f | 4.083±0.092d | 11.569±0.165b | 0.329±0.008g | - | 7.327±0.222c | 0.344±0.007g | 2.536±0.149e | 18.752±1.061a |
| 3-Mercapto-2-pentanone* | 1352 | 1343 | 1.298±0.021a | 0.438±0.01b | ND | ND | ND | ND | ND | ND | ND |
| Dimethyl trisulfide | 1392 | 1370 | ND | ND | ND | ND | 2.031±0.028a | ND | ND | ND | ND |
| 2-Furfurylthiol* | 1426 | 1402 | 4.107±0.152d | 11.591±0.26c | 15.254±0.217b | 2.133±0.003e | ND | 12.107±0.366c | 3.176±0.061d | 0.38±0.022f | 29.295±1.658a |
| 3-Thiophenethiol* | 1564 | 1530 | 0.178±0.011e | 12.892±0.289a | 0.74±0.011c | ND | ND | 0.405±0.012e | ND | ND | 2.31±0.131b |
| 2-Thiophenemethanethiol* | 1702 | 1713 | 0.196±0.012d | 0.483±0.011a | 0.267±0.004c | ND | ND | 0.207±0.006d | ND | ND | 0.41±0.023b |
| 2-methyl-3-[(2-methyl-3-thienyl)dithio]furan | 1732 | - | 0.098±0.006d | ND | 2.804±0.04b | ND | ND | 0.574±0.017c | ND | ND | 4.428±0.251a |
| 1,2,3-Trithiolane | 1794 | - | 0.378±0.026a | ND | ND | ND | ND | ND | ND | ND | ND |
| 3,3'-Dithiobis[2-methyl-furan] | 2139 | - | ND | ND | ND | ND | ND | 3.946±0.119b | ND | ND | 6.862±0.388a |
| Furfuryl sulfide | 2274 | 2223 | ND | ND | 0.159±0.002a | ND | ND | ND | 0.025±0b | ND | ND |
| Bis(2-furfuryl)sulfide | 2419 | - | 0.042±0.008b | 0.972±0.022a | - | ND | ND | ND | ND | ND | ND |
| Subtotal | | | 9.483±0.199d | 30.46±0.684b | 30.794±0.438b | 2.461±0.011f | 2.987±0.042ef | 24.566±0.743c | 3.545±0.068e | 2.916±0.171ef | 62.057±3.512a |
| Kinds | | | 9 | 6 | 6 | 2 | 2 | 6 | 4 | 2 | 5 |
| Thiophenes | | | | | | | | | | | |
| Thiophene* | 1016 | 1022 | ND | ND | ND | ND | ND | ND | ND | ND | ND |
| 3-Methyl-thiophene | 1119 | 1106 | ND | ND | 10.192±0.145a | ND | ND | ND | ND | ND | 0.083±0.005b |
| 2,3-Dihydro-5-methyl-thiophene | 1129 | 1156 | 0±0e | 0.058±0.001d | 0.167±0.002b | ND | ND | 0.126±0.004c | ND | ND | 0.204±0.012a |
| 2-Methyl-thiophene* | 1135 | 1095 | 1.128±0.159c | 0.237±0.005d | 0.147±0.002d | 0.098±0.002d | ND | 5.389±0.163b | 0.141±0.003d | 0.48±0.028d | 12.21±0.691a |
| 2-Ethyl-thiophene | 1179 | 1167 | ND | ND | ND | ND | ND | 0.062±0.002c | ND | ND | 0.083±0.005b |
| 2,5-Dimethyl-thiophene | 1216 | 1202 | ND | ND | 0.464±0.007b | 0.76±0.019a | ND | ND | ND | ND | 0.461±0.026b |



| | | | | | | | | | | | |
|--|------|------|---------------|---------------|---------------|--------------|--------------|---------------|--------------|--------------|---------------|
| 2,3-Dimethyl-thiophene | 1232 | 1212 | ND | ND | 0.038±0.001b | ND | ND | 0.207±0.006a | ND | ND | ND |
| Dihydro-2-methyl-3(2H)-thiophenone | 1511 | 1506 | ND | ND | 1.167±0.017b | ND | ND | ND | ND | ND | 1.77±0.1a |
| 2-Thiophenecarboxaldehyde* | 1674 | 1679 | 2.196±0.076b | 1.06±0.024c | 0.541±0.008e | ND | 3.109±0.043a | ND | 0.125±0.002f | 0.078±0.005f | 0.695±0.039d |
| 5-Methyl-2-thiophenecarboxaldehyde* | 1701 | 1785 | 3.649±0.075a | 2.535±0.057c | ND | ND | ND | ND | ND | ND | 2.808±0.159b |
| 2-Acetyl-3-methylthiophene | 1754 | 1760 | ND | ND | 0.488±0.007a | ND | ND | ND | ND | ND | 0.388±0.022b |
| Thieno[3,2-b]thiophene | 1868 | 1843 | 1.302±0.099b | 7.967±0.179a | 0.659±0.009d | 0.038±0.001f | 0.063±0.001f | 0.39±0.012e | 0.192±0.004f | 0.105±0.006f | 0.964±0.055c |
| 1-(2-Thienyl)-1-propanone | 1872 | - | ND | ND | 0.46±0.007b | ND | ND | ND | ND | ND | 0.584±0.033a |
| 2,5-Thiophenedicarboxaldehyde* | 1911 | - | 0.724±0.055d | 1.261±0.028a | 1.015±0.014c | ND | 1.079±0.015b | 0.292±0.009e | ND | ND | ND |
| 2-Methylthieno[2,3-b]thiophene | 1947 | - | 0.19±0.023c | 1.886±0.042a | ND | ND | 0.012±0.000d | ND | ND | ND | 0.24±0.014b |
| Subtotal | | | 9.189±0.392c | 15.003±0.337b | 15.338±0.218b | 0.895±0.022f | 4.263±0.059e | 6.466±0.196d | 0.457±0.009f | 0.663±0.039f | 20.49±1.16a |
| Kinds | | | 6 | 7 | 13 | 3 | 4 | 7 | 3 | 3 | 14 |
| Thiazoles | | | | | | | | | | | |
| Thiazole* | 1240 | 1262 | 1.796±0.128a | 0.461±0.01d | 0.556±0.008cd | 0.131±0.003f | - | 0.58±0.018c | 0.353±0.007e | 0.772±0.045b | 0.506±0.029cd |
| 2-Methyl-thiazole* | 1272 | 1278 | 0.194±0.017d | 0.579±0.013b | ND | ND | ND | 0±0e | ND | 0.612±0.036a | 0.277±0.016c |
| 2,5-Dimethyl-thiazole* | 1301 | 1326 | 2.203±0.143a | 0.127±0.003b | ND | ND | ND | 0.188±0.006b | ND | ND | ND |
| 2-Ethyl-thiazole* | 1319 | 1304 | 0.273±0.008a | 0.152±0.003b | ND | ND | ND | 0±0c | ND | ND | ND |
| 2,4,5-Trimethyl-thiazole* | 1373 | 1390 | 3.199±0.127a | 0.928±0.021b | 0.38±0.005c | ND | ND | 0.442±0.013c | ND | ND | 0.209±0.012d |
| 4,5-Dimethyl-thiazole* | 1378 | 1843 | ND | ND | ND | ND | ND | ND | ND | ND | ND |
| 5-Ethyl-2,4-dimethyl-thiazole* | 1437 | - | 0.587±0.014a | 0.174±0.004d | 0.339±0.005c | ND | ND | ND | ND | ND | 0.38±0.022b |
| 2-Ethyl-4-methylthiazole* | 1449 | 1410 | 1.278±0.117a | 0.284±0.006b | 0.219±0.003b | 0.014±0.000c | ND | ND | 0.018±0c | ND | ND |
| 2-Acetylthiazole | 1646 | 1643 | 8.795±0.213a | 0.538±0.012c | ND | ND | 2.837±0.04b | ND | ND | ND | ND |
| Subtotal | | | 18.325±0.744a | 3.255±0.073b | 1.494±0.021c | 0.145±0.004d | 2.837±0.04b | 1.211±0.037c | 0.371±0.007d | 1.385±0.081c | 1.371±0.078c |
| Kinds | | | 8 | 9 | 4 | 2 | 1 | 3 | 2 | 2 | 4 |
| Total contents of sulfur-containing compounds | | | 36.997±1.1c | 48.718±1.094b | 47.626±0.678b | 3.501±0.037g | 10.087±0.14e | 32.243±0.975d | 4.374±0.084f | 4.963±0.291f | 83.918±4.749a |
| Total kinds of sulfur-containing compounds | | | 23 | 22 | 23 | 6 | 6 | 16 | 9 | 7 | 23 |
| Nitrogen-containing heterocycles | | | | | | | | | | | |
| Pyridine | 1189 | 1202 | ND | ND | ND | ND | ND | ND | ND | ND | 0.033±0.002a |
| Methylpyrazine* | 1214 | 1263 | ND | 0.343±0.008d | 1.623±0.023b | ND | 0.423±0.006d | 1.512±0.046b | ND | 0.692±0.041c | 4.055±0.229a |
| 2-Pyridinecarboxaldehyde | 1186 | - | ND | ND | ND | ND | ND | ND | ND | ND | 1.239±0.07a |



| | | | | | | | | | | | |
|---------------------------------------|------|------|---------------|---------------|---------------|--------------|---------------|---------------|---------------|--------------|----------------|
| Pyrazine* | 1216 | 1209 | ND | 0.063±0.001cd | 0.208±0.003cd | 0.039±0.001d | 0.12±0.002cd | 0.119±0.004cd | 0.294±0.006bc | 0.498±0.029b | 5.699±0.323a |
| 3-Methyl-pyridine* | 1307 | 1346 | ND | 0.076±0.002a | ND | ND | ND | ND | ND | ND | ND |
| 2,5-Dimethyl-pyrazine* | 1320 | 1328 | ND | 0.679±0.015b | 0.423±0.006c | ND | 0.069±0.001e | 0.402±0.012c | ND | 0.129±0.008d | 0.927±0.052a |
| 2-(n-Propyl)-pyrazine | 1476 | 1428 | ND | ND | ND | ND | ND | ND | ND | ND | 0.378±0.021a |
| Pyrrrole | 1499 | 1518 | ND | ND | 0.228±0.003a | ND | ND | 0.12±0.004b | ND | ND | ND |
| 2-Methyl-1H-pyrrrole | 1534 | 1551 | ND | ND | 0.371±0.005b | 0.036±0.001d | ND | 0.139±0.004c | ND | ND | 0.411±0.023a |
| 1H-Pyrrrole-2-carboxaldehyde | 2002 | 2028 | ND | ND | ND | ND | 0.836±0.012a | ND | ND | ND | ND |
| Subtotal | | | ND | 1.161±0.026d | 2.853±0.041b | 0.076±0.002e | 1.448±0.02d | 2.292±0.069c | 0.294±0.006e | 1.319±0.077d | 12.741±0.721a |
| Kinds | | | 0 | 4 | 5 | 2 | 2 | 5 | 1 | 3 | 7 |
| Oxygen-containing heterocycles | | | | | | | | | | | |
| Furan* | 797 | 798 | 0.138±0.01a | ND | ND | ND | ND | 0±0b | 0±0b | ND | ND |
| 2-Methyl-furan* | 851 | 829 | 0.019±0.005f | ND | 6.884±0.098b | 0.505±0.012e | ND | 4.741±0.143c | 0.276±0.005ef | 1.589±0.093d | 8.225±0.465a |
| 2-Ethyl-furan | 913 | 949 | ND | ND | 0.041±0.001b | ND | ND | 0.042±0.001b | 0±0c | ND | 0.089±0.005a |
| Furfural* | 1457 | 1460 | 5.793±0.155a | ND | 1.05±0.015e | 0.242±0.006g | 1.968±0.027c | 1.019±0.031e | 3.589±0.069b | 1.369±0.08d | 0.664±0.038f |
| 2-Acetylfuran | 1491 | 1500 | ND | ND | ND | ND | ND | 1.189±0.036a | 0.378±0.007b | ND | ND |
| 1-(2-Furanyl)-ethanone* | 1497 | 1501 | 0.136±0.077a | ND | ND | ND | ND | ND | ND | ND | ND |
| 5-methylfurfural | 1542 | 1567 | ND | ND | 1.34±0.019a | 0.703±0.017b | ND | ND | ND | ND | ND |
| 2(5H)-Furanone* | 1748 | 1767 | 0.072±0.005a | ND | ND | ND | ND | ND | ND | ND | ND |
| 4-Hydroxy-5-methyl-3(2H)-furanone* | 2108 | 2124 | ND | ND | ND | ND | ND | 0.19±0.006a | ND | ND | ND |
| Subtotal | | | 6.158±0.231c | ND | 9.314±0.133a | 1.45±0.036g | 1.968±0.027f | 7.182±0.217b | 4.243±0.082d | 2.958±0.174e | 8.978±0.508a |
| Kinds | | | 5 | 0 | 4 | 3 | 1.000 | 5 | 3 | 2 | 3 |
| Total | | | 43.155±1.223d | 49.878±1.12c | 59.793±0.851b | 5.027±0.075f | 13.503±0.188e | 41.717±1.262d | 8.911±0.172e | 9.24±0.542e | 105.637±5.978a |
| Total kinds | | | 28 | 26 | 32 | 11 | 9 | 26 | 13 | 12 | 33 |

Notes: Results were presented as means ± standard deviation, data within a row with different letters are significantly different ($p < 0.05$) using Duncan's multiple comparison test ($n = 3$).

a: Linear retention indices calculated using a series of n-alkanes on the DB-W AX column ($30 \text{ m} \times 0.25 \text{ mm} \times 0.25 \text{ } \mu\text{m}$).

b: Linear retention indices searched from <http://www.flavornet.org> and <http://webbook.nist.gov/chemistry/>.

c: ID: Identification methods. d: "ND", not detected.

e: Compounds marked with an asterisk (*) were quantified using the standard curve method with internal standard calibration.

Table 2.

Variations of TTCA (a) and ARP (b) contents in their solid samples over time after storage for 60 days at different water activities

| MRI | Indicator | a_w | | | |
|------|------------|-------------|-------------|-------------|-------------|
| | | Control | 0.113 | 0.432 | 0.843 |
| TTCA | L^* | 88.39±0.03a | 81.36±0.02b | 77.39±0.04c | 72.34±0.02d |
| | a^* | 5.19±0.01d | 7.71±0.03c | 9.19±0.02b | 11.63±0.04a |
| | b^* | 11.13±0.02d | 16.78±0.04c | 22.36±0.03b | 27.63±0.01a |
| | ΔE | - | 9.36c | 16.22b | 23.90a |
| ARP | L^* | 87.68±0.03a | 79.36±0.03b | 62.18±0.03c | 51.03±0.03d |
| | a^* | 6.21±0.03d | 11.39±0.03c | 17.68±0.03b | 21.39±0.03a |
| | b^* | 13.19±0.03d | 20.38±0.03c | 29.63±0.03b | 37.68±0.03a |
| | ΔE | - | 12.16c | 32.44b | 46.62a |



691 **FIGURE CAPTIONS**

692 **Fig. 1** Flavor formation from the TTCA model system under different conditions: (a) Effect
693 of temperature (100, 120, 140°C) at pH 7.0 on volatile compounds; (b) Effect of pH (5.5,
694 7, 8) at 120°C on volatile compounds; (c) Browning intensity of TTCA systems with
695 different added amino acids; (d) Contents and (e) types of characteristic flavor compounds
696 from TTCA with Gly, Lys, or Glu; (f) Formation pathways for characteristic flavor
697 formation from TTCA with amino acids.

698 **Fig. 2** Residual content of TTCA (a) and ARP (b) in solution at different temperatures;
699 Absorbance at 420 nm and 294 nm for TTCA (c) and ARP (d) solutions at different
700 temperatures; Residual content of TTCA (e) and ARP (f) in solution at different pH levels;
701 Absorbance at 420 nm and 294 nm for TTCA (g) and ARP (h) solutions at different pH
702 levels; Residual content of solid TTCA (i) and ARP (j) after 60 days at different water
703 activities.

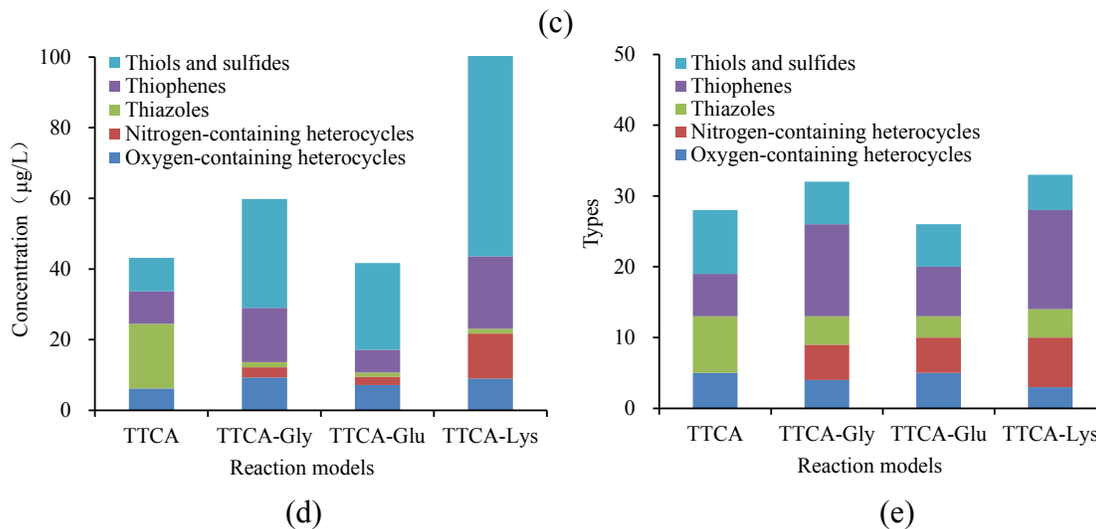
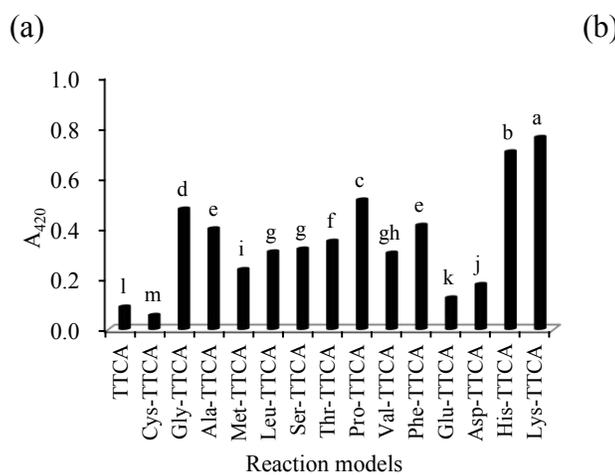
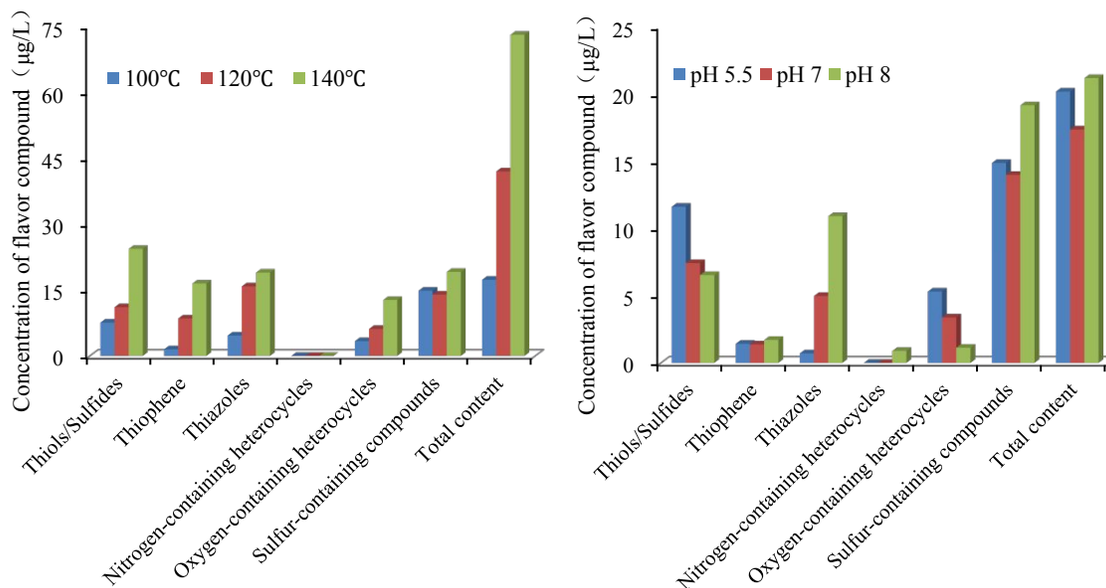
704 **Fig. 3** Fe²⁺ Chelating activity (a) and reducing power (b) of TTCA, ARP and MRPs;
705 Inhibition of TTCA, ARP, MRPs and ascorbic acid on lipid peroxidation induced by iron
706 (c).

707 **Fig. 4** DPPH radical scavenging activity (a), hydroxyl radical scavenging activity (b) and
708 superoxide anion radical scavenging activity (c) of TTCA, ARP, MRPs and ascorbic acid;
709 Possible pathway of the reaction between free radicals and ARP (R • is the free radical)
710 (d).

711



Fig. 1



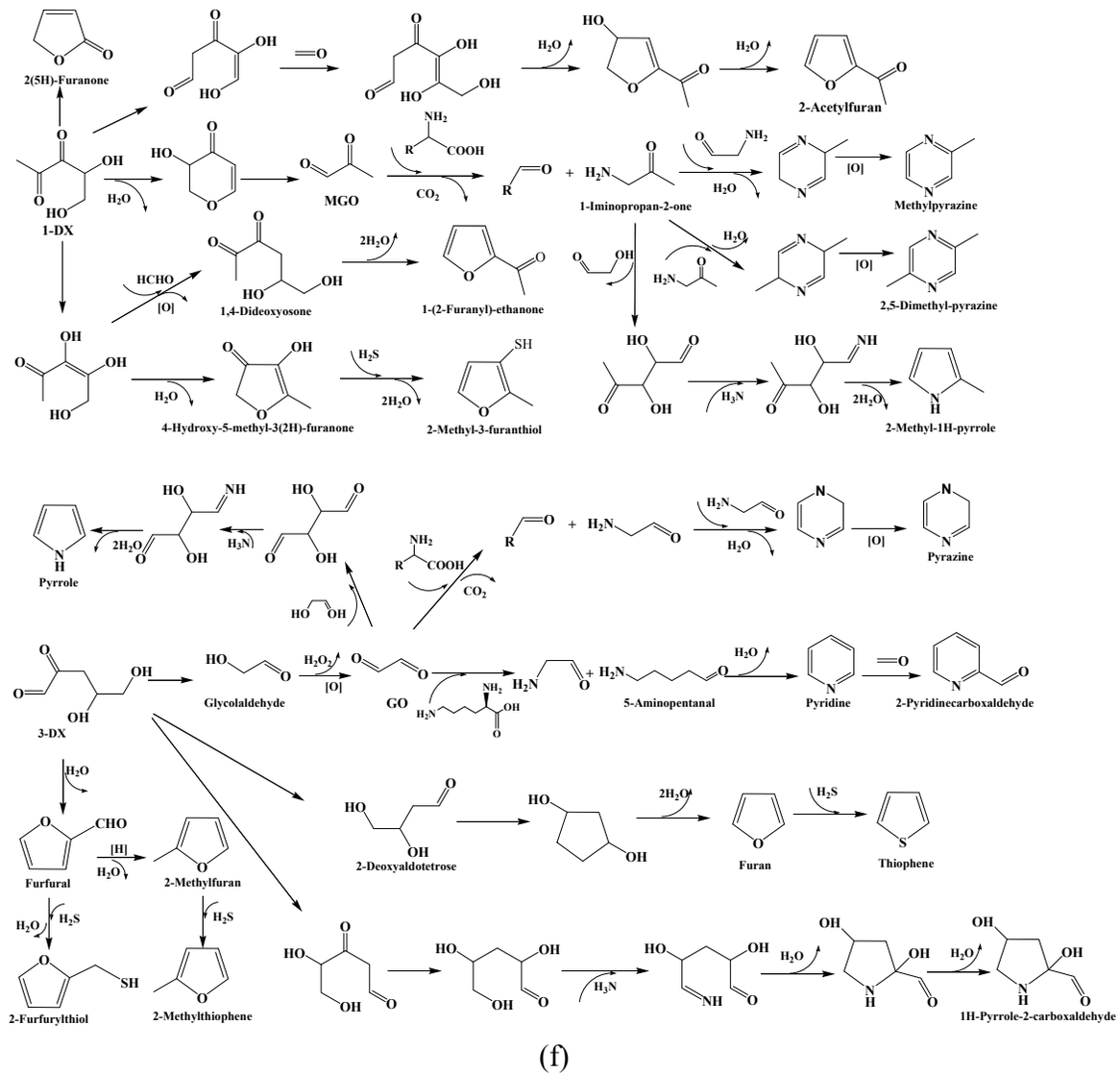
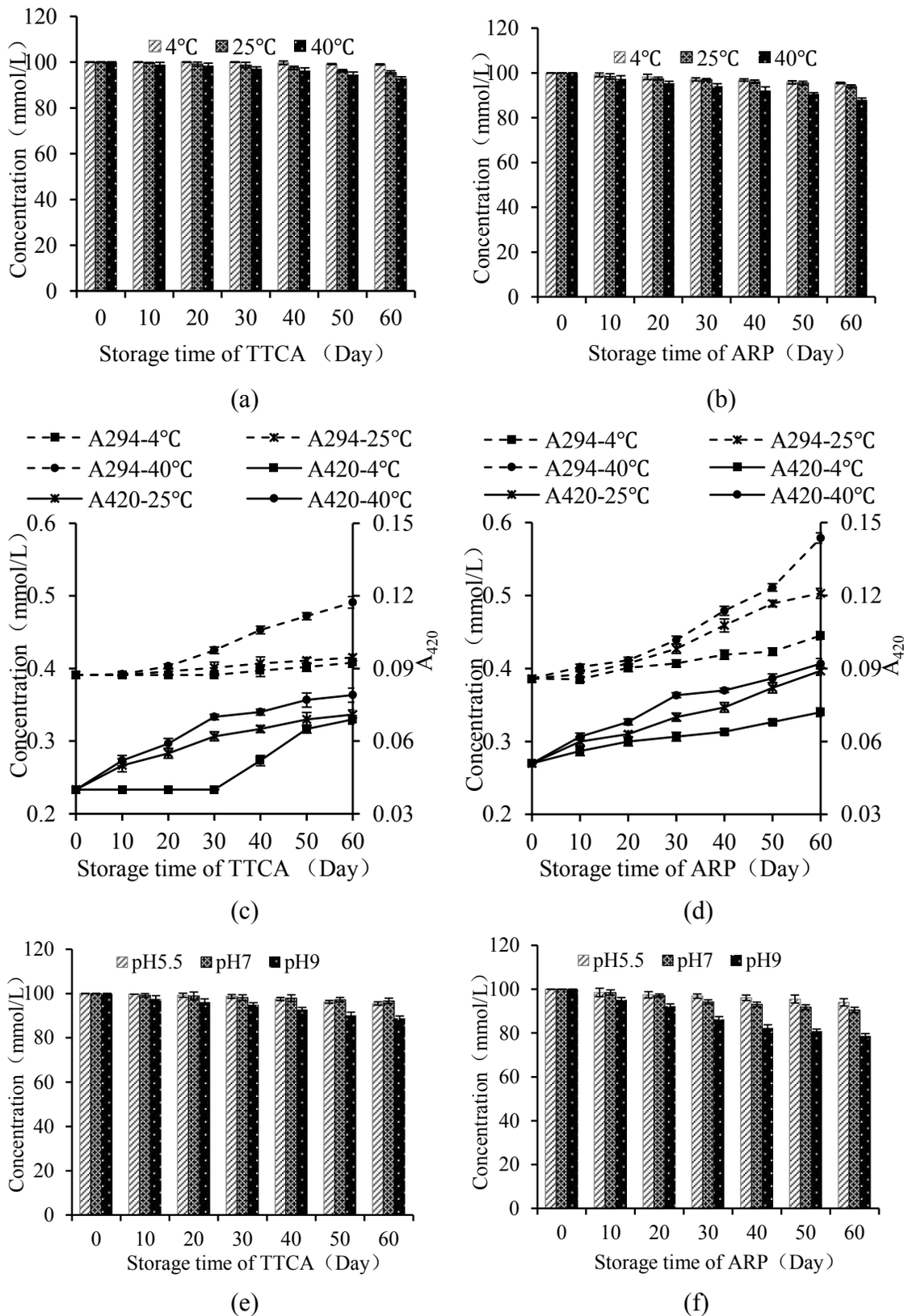
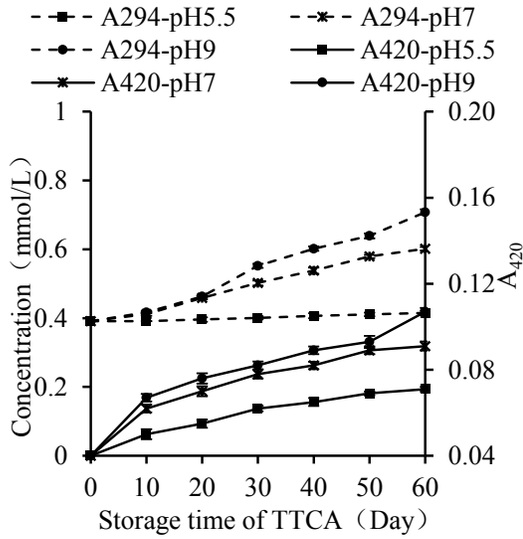
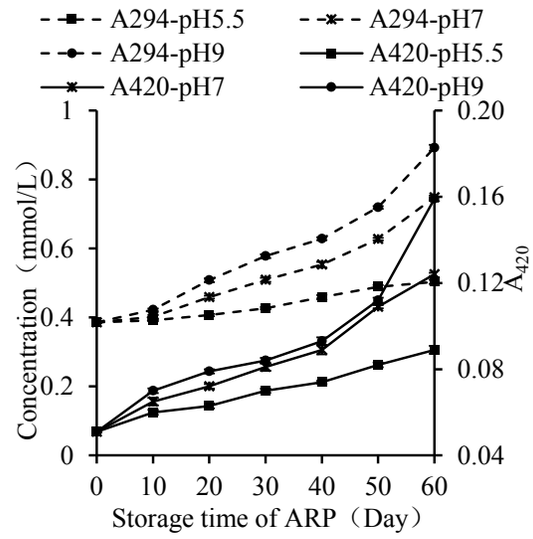


Fig. 2

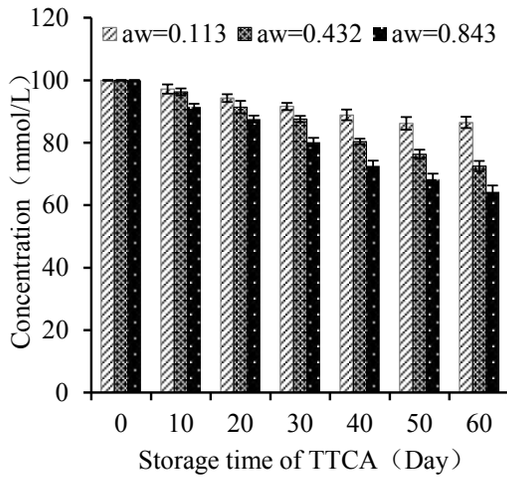




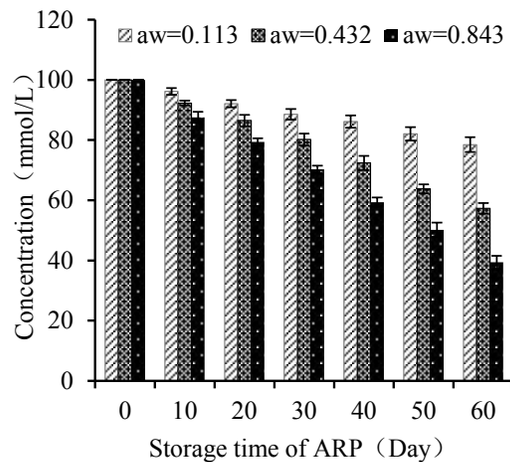
(g)



(h)



(i)



(j)



Fig. 3

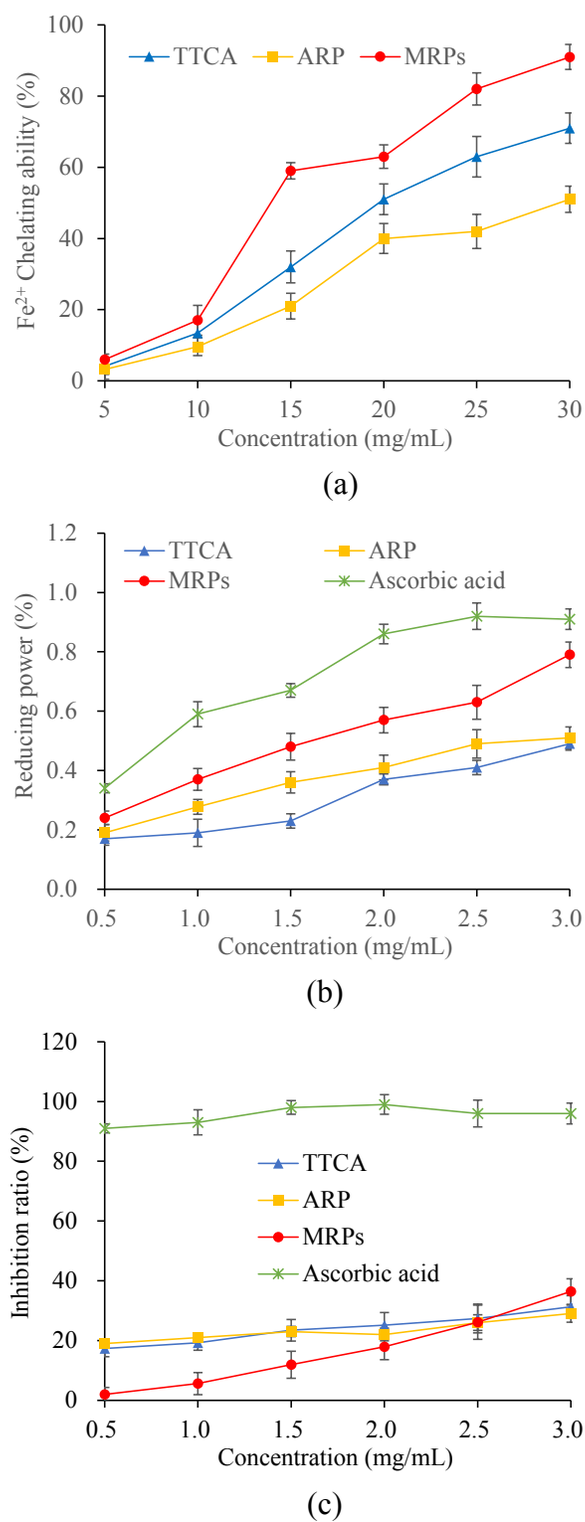
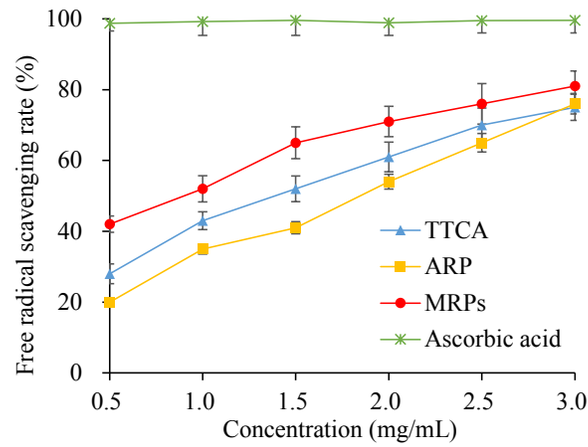
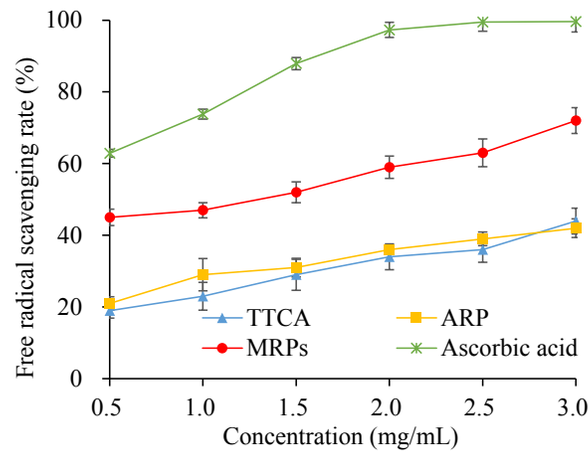


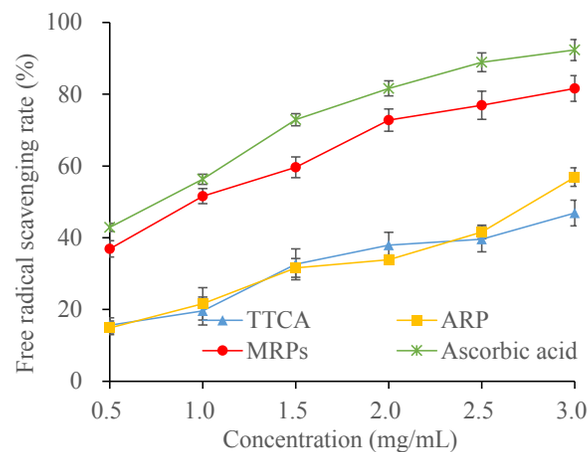
Fig. 4



(a)

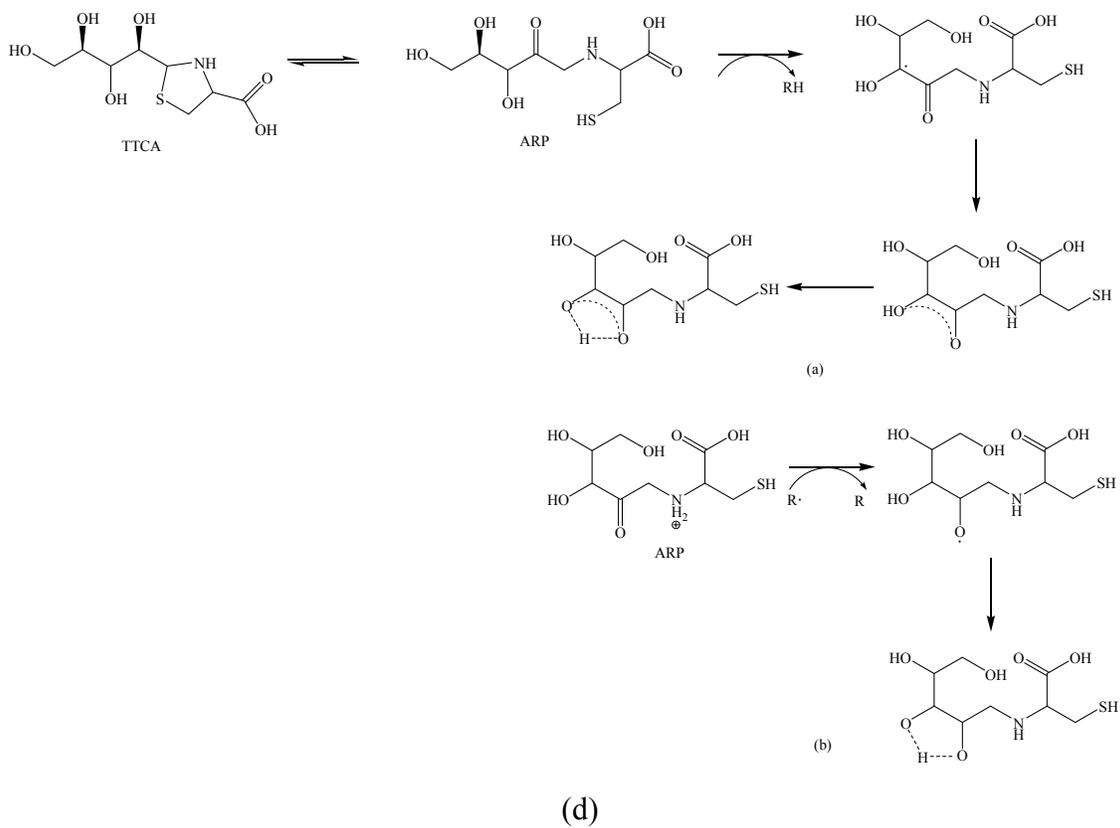


(b)



(c)





Data availability

All the data generated or analyzed during this study are included in the manuscript.

