

1 **The Use of Chromatographic-Based Techniques and**
2 **chemometrics for Halal Authentication of Food Products: A**
3 **Review**

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19
20 **Abstract**

21 Halal food and halal pharmaceutical products are requisite to be consumed by Muslim
22 communities in the world. The standard methods capable of quantifying non-halal components
23 are very urgent. This review highlights chromatography and chemometric based techniques
24 that offer reliable techniques to provide separation capacity in halal authentication analysis.

25 Methods: This review article was written from reputable worldwide databases including Web
26 of Science, Scopus, and PubMed, between January and February 2022. The keywords were
27 “halal research”, “food analysis”, “pharmaceutical analysis”, “chromatography”,
28 “chemometrics”, and “authentication”. Chromatographic-based techniques combination with
29 chemometrics of multivariate analysis, a powerful statistical analysis to manage huge data
30 generated from analytical measurement, could be used to identify potential markers to
31 differentiate halal and non-halal samples. Chromatogram and peak separation profiles resulted
32 as the instrument responses can be further evaluated for determination as well as quantification
33 for halal and non-halal components in food and pharmaceutical products.

34 Combination of chromatographic-based method and chemometrics techniques with some
35 scenarios can be applied for halal research on food and pharmaceutical products.

36

37 **Keywords:** halal authentication, chemometrics, chromatography, pig derivatives,
38 pharmaceutical.

39

40 **1. Introduction**

41 Food, cosmetics, drugs and other pharmaceutical products are important needs for
42 human beings. In line with the development of science and technology, industrialization and
43 globalization, the halal products may be added or substituted and contaminated with non-halal
44 components such as pig derivatives and alcohols as ingredients or additives to reduce the
45 production cost make the products non-halal [1]. In addition, the products available in markets
46 may contain incorrect labelling in terms of ingredient sources making the consumers lost on
47 composition information, therefore the use of analytical tools to check the presence of non-
48 halal components in the products is a must for assisting the certification processes [2]. In
49 Indonesia, the halal certification is mandatory which means that all halal declared products sold
50 and distributed in Indonesia must be halal certified. In addition, the analysis of non-halal
51 components in post-marketed products is also needed to confirm that the marketed products
52 are not adulterated with non-halal components [3].

53 According to Indonesian Act No. 33 (2014), the certification process is carried out by
54 Halal Product Assurance Organizing Agency (BPJPH) and the auditing process was carried
55 out by Halal Examination Agency (LPH). During audit, if the products are supposed to contain
56 non-halal components (pork derivatives and alcohols), the laboratory testing using standard
57 analytical methods is needed to confirm that the audited products are free from any non-halal
58 components [4,5]. Today, the Muslim community constitute for approximately of 25% world's
59 population and is expected to increase further. With the increased awareness among Muslim
60 community to consume the only halal products, the global market of halal products could reach
61 exponentially [6]. Halal is Arabic terms used to any products permissible to be consumed by
62 Muslim community. Today, the term of halal has widely used not only Muslim but also non-
63 Muslim because Non-Muslim community intended to export the products into Muslim
64 community, especially in halal certification issues [7]. Therefore, it is not surprising that halal-
65 related studies are performed not only in majority Muslim countries like Indonesia and
66 Malaysia but also in countries whose Muslims are minority such as the Netherlands, the United
67 States, France and the European Union [8].

68 Halal food and Halal pharmaceuticals must be free from non-halal components which
69 are pig and all pig derivatives such as pork, lard and porcine gelatines, carrion or dead animals,
70 blood (flowing or congealed), animals slaughtered not according to Islamic law, animals that
71 were killed accidentally or on purpose through means such as strangling or beating, intoxicants
72 including alcohol and drugs [9], carnivorous animals, predator birds, and certain land animals
73 [10]. Among these, pig derivatives and alcohols are typically found in halal and pharmaceutical
74 products, therefore some scientists are continuously researches on halal including developing
75 instrumental analytical methods for detecting of non-halal components intended for halal
76 certification [11]. Some countries have obligated the products to be halal certified which can
77 be understood that the products are free from prohibited components. Besides, the products are
78 manufactured using equipment dedicated for halal food and halal pharmaceuticals [12]. Pork
79 is typically met in meat-based food products such as meatball, sausages, etc.; while lard can be
80 good vehicle in some cosmetics products such as cream, lipstick and lotion. Porcine gelatines
81 are common materials used in food (in candies) and pharmaceutical products (capsule shells)
82 [13]. The objective of this review was to provide integrative information on identification and
83 quantification of non-halal components in food and pharmaceutical products by
84 chromatographic methods. In addition, chemometrics techniques were reported to be applied
85 to employ the big data evaluation as resulted from the chromatographic detection.

86

87 **2. Methods**

88 This review article was written by identifying, investigating, and assembling several
89 review articles, original articles, books, and relevant sources on metabolite fingerprintings from
90 reputable worldwide databases including Web of Science, Scopus, and PubMed. Literature
91 searching was carried out between January and February 2022. The keywords explored during
92 literature investigation were “halal research”, “food analysis”, “pharmaceutical analysis”,
93 “chromatography”, “chemometrics”, and “authentication”.

94

95 **3. Chromatographic-based techniques and chemometrics for analysis of non-halal** 96 **components**

97 For many years, chromatography has been known as the method of choice to assess the
98 purity and levels of analytes in the laboratories of research, industry, and quality control [14].
99 Gas chromatography (GC) and liquid chromatography (LC) techniques are often used for the
100 analysis of non-halal components in food and pharmaceutical products. In terms of compound
101 types, GC is more suitable for the analysis of smaller, volatile and stable compounds to heat,

102 while LC is more robust and suitable for larger and less/non-volatile compounds [15]. Some
103 derivatization techniques are needed in LC in order to convert analytes into detectable derivatives
104 with certain detectors, while derivatization in GC for fewer volatile compounds is intended to
105 provide more volatile and stable derivative products, although this derivatization process
106 increases the method complexity and lengthens the sample preparation. In addition, the
107 availability of derivative agents and its steric hindrance in the analyte, and the stability of the
108 derivatized compounds must also be considered [16].

109 One-dimensional gas or liquid chromatography using one column is considered as
110 simple and powerful separation techniques for simple and un-complex samples. When the
111 analyzed samples are complex enough, the application of just one-dimension chromatography
112 leads to peak co-elution as well as overlapping and non-resolved peaks, therefore one
113 dimension chromatography technique is not suitable for separation of large analytes because
114 the peak capacity of one-dimensional analysis is not large enough to achieve the complete
115 separation with acceptable resolution [17]. In last decades, two-dimensional gas
116 chromatography (GC x GC) and liquid chromatography (LC x LC) has been applied in analysis
117 of complex mixture in order to increase the separation speed [18].

118 In two-dimensional chromatography, the separation is carried out in two columns with
119 different polarity connected in series by a modulator, as a consequence, the separation capacity
120 of regular one-column in one dimensional chromatography can be considerably increased. The
121 effluent from the first column is transferred to the second column using modulator so that the
122 analytical information obtained (such as retention times, t_R) from the first column can be
123 combined with that from second column, leading to a plot of two retention times [19]. Because
124 of the excellent separation capacity of GC x GC and LC x LC combined with mass
125 spectrometry (MS), both techniques are applied for separation all components in the complex
126 mixtures, especially for metabolomics studies [18]. GC x GC has been widely applied for
127 analysis of metabolites (all fatty acid types) of lard in food samples [20], while LC x LC is
128 typically used for analysis of peptides [21], which can be used for identification of pork and
129 porcine gelatines.

130 Chromatographic-based techniques offered reliable technique in halal authentication
131 analysis. However, due to high number of data covered, the application of chemometrics to
132 treat big data is unavoidable. Chemometrics can be defined as the employment of statistical
133 and mathematical methods to obtain the objective data evaluation by extracting the relevant
134 and meaningful information from related and unrelated responses from chemical
135 measurements. Chemometrics or multivariate data analysis (MDA) is typically applied in

136 numerous aspects including the quality control of halal products, qualitative and quantitative
137 determination of chemical parameters for assessing the products authenticity [22].

138 Chemometrics can provide the powerful tools in giving important information extracted
139 from big data obtained from instrumental analyses such as methods based on spectroscopic and
140 chromatographic. The common chemometrics techniques applied in products authentication
141 could be grouped into exploratory data analysis, data pre-processing, description and
142 visualization, discrimination and pattern recognition (classification), regression and prediction
143 and experimental design [23]. Some chromatographic problems encountered during halal
144 authentication analysis included the assessment of separation quality, the evaluation of peak
145 alignment using pre-processing, the optimization of chromatographic systems providing the
146 good separation of all peaks using experimental design, the accuracy of discrimination and
147 classification using pattern recognition, and quantitative analysis applying multivariate
148 calibration. Figure 1 showed the correlation between chromatographic responses and
149 chemometrics for certain analytical purposes. In scenario (a), peaks with good separation (good
150 selectivity) in chromatogram was used as variable for the evaluation of compositional analysis
151 (concentration) of analytes assisted by multivariate calibrations. In (b), certain peaks with lack
152 selectivity was used as variable during chromatographic profiling of objects (samples) using
153 discrete datasets (peak area or peak height), while in scenario (c), whole datasets in
154 chromatograms were used as variables during chromatographic fingerprinting of objects.
155 Indeed, the chemometrics of pre-processing was widely applied to obtain the desired analytical
156 modelling.

157 The classification chemometrics was typically carried using (1) exploratory data
158 analysis including principal component analysis (PCA) and cluster analysis (hierarchical
159 cluster analysis and non-hierarchical such as k-means and k-medians), and this technique is
160 typically called as unsupervised pattern recognition and (2) classification and discrimination
161 methods known supervised pattern recognition. There are two types of classification
162 chemometrics methods regardless of the statistical background. The first type is typically
163 employed to assess to which of various pre-defined classes of samples (objects). The example
164 of this technique is linear discriminant analysis (LDA), orthogonal projection to latent
165 structures – discriminant analysis (OPLS-DA), k-nearest neighbors (KNN) and many others.
166 The second type of classification chemometrics is called as class modelling or one class
167 classifier (OCC), and the example for this group data driven soft independent modeling of class
168 analogy (DD-SIMCA) and Unequal Class-Modeling (UNEQ) [25]. Using these chemometrics,

169 someone can answer the question: is the meat belong to pork (non-halal) or beef (halal)? or the
170 question: is the meatball authentic or adulterated? [26,27].

171

172 **4. Analysis of non halal components using liquid chromatography**

173 High performance liquid chromatography (HPLC) using certain detectors have been
174 widely applied for analysis of specific components in non-halal components. HPLC using
175 fluorescence detector has been successfully used for analysis of Hydroxyproline and other
176 amino acids in gelatin and collagen samples as initial screening for identification of gelatin
177 types. Hydroxyproline has been known as signature amino acid for gelatin and collagen. The
178 level of hydroxyproline is typically higher in the gelatin samples than that in the collagen
179 samples, except for the samples of fish skin gelatin, and this result could be used as screening
180 tools for identification of non-halal gelatin and collagen in the analyzed samples [28]. Table 1
181 listed the application of HPLC and LC-MS/MS for analysis of halal components in the
182 products. Liquid chromatography using fluorescence detector was also successfully applied for
183 analysis of amino acid (AA) composition non-halal (porcine) and halal (bovine and fish)
184 gelatins. The classification between halal and non-halal gelatins was carried using PCA
185 applying amino acid compositions as variable. AAs with strong fluorescence (Hyp, His, Ser,
186 Arg, Gly, Thr, Pro, Tyr, Met, Val, Leu and Phe) contribute to the classification and become the
187 biomarkers to identify the gelatine sources [29]. Gelatin from three mammalian species
188 including bovine gelatin, porcine gelatin, and donkey gelatin has been successfully identified
189 using liquid chromatography-linear ion-trap high resolution mass spectrometry. Hemoglobin
190 was just found in donkey gelatin. The unique peptide obtained from donkey, bovine, and
191 porcine gelatin was GEAGPAGPAGPIGPVGAR, GETGPAGPAGPIGPVGAR, and
192 GETGPAGPAGPVGPGVGAR, respectively. The unique peptides could be detected either in
193 individual gelatin or in the mixtures of three mammalian gelatins [30].

194 Liquid chromatography especially combined with mass spectrometer (LC/MS) is
195 widely applied for identification of non-halal component in food and pharmaceutical products
196 including porcine gelatin and pork. Gel-enhanced liquid chromatography-mass spectrometry
197 (GeLCMS) in combination with chemometrics of PCA has been developed for identification
198 of potential protein markers in pork and other meats along with its classification. The
199 myofibrillar protein with weight of 40-kDa such as troponin T, Tropomyosin alpha-1 chain,
200 and actin cytoplasmic 1 as well as the thin filament proteins such as actin, troponin, and
201 Tropomyosin had molecular weights ranging from 40 to 45 kDa could be used as markers for
202 differentiation of pork from chicken and beef. PCA using PC1 and PC2 accounting of 62% and

203 35% variances could classify meat types. From MS studies, the potential protein markers for
204 pork meat samples are Troponin T with peptide sequences of [(R)KPLNIDHLSSEDK(L)],
205 Tropomyosin alpha-1 chain [(K)EAETRAEFAER(S)], Actin cytoplasmic 1
206 [(R)HQGVMVGMGQK(D)], COP9 signalosome complex subunit 4 [(R)VLDYRR(K)], and
207 Ribonuclease inhibitor [(R)VLGQGLADSACQLETLR(L)][45].

208 The identification of potential biomarkers of gelatin from several sources could be
209 performed using UPLC-MS/MS. Samples used were gelatin from pig, cow, chicken, and fish.
210 After the extraction process of proteins from gelatin, proteins were then digested using
211 proteomic grade trypsin for 12 h to obtain peptides. Chemometrics of PCA was used to
212 differentiate partial hydrolysis of gelatin from cow and pig. Result from PCA score plot showed
213 that the sample of cow and pig obtained from digestion process could be well separated. For
214 identification of potential biomarkers from pig, cow, fish, and chicken gelatin, PCA employing
215 MPP (Mass Profiler Professional) was applied. Results showed that three unique peptides
216 found only in pig gelatin, seven unique peptides found in bovine/cow gelatin, 22 peptides found
217 only in chicken gelatin, and only 1 unique peptide found in fish gelatin. The developed method
218 was also successfully applied to identify species origin of commercial gelatin samples. It
219 indicated that UPLC-MS/MS offers a powerful analytical technique to identify gelatin from
220 different species in food and pharmaceutical products [46].

221 Targeted tandem liquid chromatography-mass spectrometry (LC-MS) using decoy,
222 randomized and concatenated database search program comprising MS-Fit and MS-Tag in
223 combination with chemometrics of principal component analysis and orthogonal partial least
224 square-discriminant analysis (OPLS-DA) was applied for identification of potential peptide
225 markers in non-Halal meat (pork) and halal meats (chicken and beef). The peptide markers
226 which are specific to certain species were identified through shot-gun proteomics. Potential
227 peptide marker identified for raw pork is myosin-2 having sequence of peptide marker of
228 (F)DFNSLE(Q). OPLS-DA using variable of identified peptides could separate halal and non-
229 halal meats [47].

230 Targeted proteomic analysis using LC-MS has been developed to investigate the heat
231 stable protein in pork meat. Five heat treatments were applied such as (1) water bath heating at
232 78°C for 30 min; (2) boiling at 100°C for 30 min; (3) sterilizing at 121°C for 30 min; (4) frying
233 using oil until golden brown colour; and (5) baking at 200°C for 30 min. Protein extraction
234 from samples was performed using buffer solution containing 2 M thiourea, 7 M urea, and 50
235 mM Tris-HCl (pH 8.0). Proteins were digested using proteomic grade trypsin added with DTT
236 to reduce disulphide bonds and IAA for alkylation. Incubation was carried out for at least 12 h

237 at 37°C. Result showed that seven heat-stable specific peptides of pork were found such as
238 DQLIHNLLK from l-lactate dehydrogenase A chain, HDPSLLPWTASYDPGSAK from
239 carbonic anhydrase 3, EPITVSSDQMAK from carbonic anhydrase 3, VNVDEVGGEALGR
240 from haemoglobin subunit beta, HPGDFGADAQGAMSK from myoglobin,
241 SLYSSAENEPPVPLVR from carbonic anhydrase 3, and YLEFISEAIIQVLQSK from
242 myoglobin. Commercial samples such as Iberian dried ham, Pasteur dry sausage, import dried
243 ham, lunch meat canned, sandwich sausage, and Thuringia flavour sausage were used to
244 identify the presence one or more pig heat-stable peptides. Results showed that the heat-stable
245 peptides of pig could be found in various types of food products with different cooking process
246 methods. It suggested that targeted proteomics analysis using seven heat stable peptides of pig
247 could be used for halal authentication of food products especially meat-based food products
248 containing pork [48].

249 Analysis using LC-MS employing MRM (multiple reaction monitoring) technique was
250 successfully used to detect heat-stable peptides in cooked meats including pork meat. Thermal
251 treatment applied was boiling at 100°C, grilling at 150°C, and grilling at 180°C. After the
252 protein was extracted, digestion process was performed using proteomic grade trypsin.
253 Identification of homologues protein and potential biomarkers of pork peptide was carried out
254 using UPLC Triple TOF-MS equipped with a C-18 column (2.1 × 100 mm, 1.7 µm; Waters
255 Corporation, Taunton, MA, USA and Wexford, Ireland). The mobile phase used was water
256 containing 0.1% formic acid (A) and acetonitrile containing 0.1% formic acid (B) with flow
257 rate of 0.3 mL/min. On the other hand, MRM analysis was performed using a SCIEX ExionLC
258 AD system (AB SCIEX, Framingham, MA, USA) and an AB SCIEX QTRAP 4500 mass
259 spectrometry system (AB SCIEX PTE. LTD., Marsiling, Singapore) equipped with a column
260 of Waters ACQUITY UPLC BEH C18 (2.1 × 50 mm, 1.7 µm). Results showed that the
261 potential peptide biomarkers in raw pork meat found were GHHEAELTPLAQSHATK from
262 myoglobin, FAGGNLDVVK; ADMVIEAVFEELSLK; TVLGAPEVLLGILPGAGGTQR
263 from trifunctional enzyme subunit alpha, mitochondrial, and
264 WGDAGATYVVESTGVFTTMEK from glyceraldehyde-3-phosphate dehydrogenase.
265 Meanwhile, the heat-stable peptide biomarkers of pork were FAGGNLDVVK and
266 TVLGAPEVLLGILPGAGGTQR from trifunctional enzyme subunit alpha, mitochondrial as
267 well as WGDAGATYVVESTGVFTTMEK from glyceraldehyde-3-phosphate dehydrogenase.
268 The MRM analysis confirmed the heat-stable peptide of pork in meat product samples. It
269 suggested that LC-MS employing MRM method could be used as promising analytical
270 technique for halal authentication of meat products [49].

271

272 **5. Application of gas chromatography for analysis of non-halal components**

273 The use of Herbal medicines (HMs) as complementary and alternative medicine is
274 becoming popular in the general population worldwide. Parallel to the increased trends on
275 application of HMs as alternative therapies either for preventive or promotive, some research
276 activities dealing with the quality control, standardization, and authentication of HMs also
277 increased. The efficacy of HMs depends on their quality and its authenticity. Fingerprint
278 profiling based on spectroscopic especially ¹H-NMR and chromatographic techniques
279 hyphenated with mass spectrometers (LC-MS/MS) in combination with classification
280 chemometrics has emerged as powerful tools for standardization and authentication of HMs.
281 Table 2 listed the application of gas chromatography for analysis of halal components in the
282 food and pharmaceutical products.

283 GC-MS combined with chemometrics has been proposed as tools for detection of lard
284 as adulterant in olive oil using metabolomic approach. GC separation of fatty acid methyl esters
285 (FAME) was achieved using HP-5MS nonpolar capillary column. The identification of
286 metabolites of FAMEs was carried out using standard FAMEs and mass spectrometer detector
287 using the WILEY 2007 library. Some FAMEs are specific, i.e., methyl behenate was only
288 present in olive oil and methyl myristate was only detected in lard. PCA using identified
289 FAMEs was successful for separating lard, olive oil and olive oil adulterated with lard for halal
290 authentication study [50].

291 Two dimensional GC combined with time-of-flight mass spectrometer (GC x GC-
292 TOF/MS) is successfully used for analysis of lard as adulterant in virgin coconut oil (VCO)
293 through analysis of sterols. GC x GC system could perform the complete baseline separation
294 of sterol trimethylsilyl ethers derived from cholesterol and cholestanol, which facilitate the
295 detection of lard in VCO. Using GC x GC-TOF/MS Cholestanol trimethylsilyl ether (Cha-
296 TME) and cholesterol trimethylsilyl ether (Che-TME) were separated into some peaks,
297 identified as CHE₁, CHE_{bI}, CHE_{bII}, CHE₂ (Che-TME), and Cha₁, CHA_{bI}, CHA_{bII}, and CHA₂ for
298 Cha-TME. Quantification of these compounds could be used as tools for quantification of
299 adulteration levels of lard in VCO [20].

300 GC-MS coupled with headspace solid-phase micro-extraction (HS-SPME) is successful
301 for the analysis of volatile compounds in pork. The profiles of volatile compounds from
302 different meats are different, therefore, the volatile compounds analysed by GC-HS-SPME/MS
303 could be used as fingerprinting tools for specific meats [51]. In addition, VOCs also contribute
304 to the aroma which can be used for the discrimination tools among animal meats [52]. Analysis

305 of VOCs is very challenging because of different factors, including the high number of volatile
306 compounds, differences in volatility degree and the great amount of functional groups [53].
307 Chen et al. [54] have identified the key volatile compounds for differentiation of pork from
308 different pig breeding. The volatile compounds contributing to the pork flavour identified
309 during this study were 3-methyl-1-butanol, 1-nonanal, octanal, hexanal, 2-pentyl- furan, 1-
310 penten-3-one, N-morpholinomethyl-isopropyl-sulphide, methyl butyrate, and (E,E)-2, 4-
311 decadienal. Kosowska et al. [55] reported that some volatile compounds namely octanal,
312 nonanal, (E,E)-2,4-decadienal, methanethiol, methional, 2-furfurylthiol, 2-metyl-3-furanthiol,
313 3-mercapto-2-pentanone, and 4-hydroxy-2,5-dimethyl-3-(2H)- furanone are key features in
314 cooked pork. Thus, the identification of marker volatile compounds in pork can be meaningful
315 for pork identification during halal authentication analysis of products. GC-HS-SPME/MS and
316 GC-MS using simultaneous distillation and extraction (SDE) are also successful for
317 identification of volatile compounds used for the identification of cooking braised pork. There
318 are 109 aroma compounds identified, in which aldehydes were the most predominant in
319 number, followed by alcohols, oxygen-containing heterocyclic compounds, acids, and ketones.
320 Methanethiol was the most abundant aroma substance in SPME, while anethole was the most
321 abundant in SDE [56].

322 GC-HS-SPME/MS has been developed and validated as reliable analytical method for
323 analysis of volatile organic compounds (VOCs) of minced pork meat during storage. The origin
324 of aromatic hydrocarbons in pork was verified using migration test. Two chemometrics
325 techniques namely PCA and OPLS-DA were employed for characterizing and profiling VOCs
326 in pork meat and for identifying the marker VOCs associated with the spoilage of pork. There
327 are 41 VOCs (consisting of 10 alcohols, 7 aldehydes, 7 ketones, 6 aromatic hydrocarbons, 6
328 linear hydrocarbons, 2 terpenes, 1 acid, 1 ester, 1 furan) were identified during this study. The
329 major VOCs of minced pork are aromatic hydrocarbons, alcohols, aldehydes, linear
330 hydrocarbons, and ketones). From loading plot study, three VOCs namely ethanol, 2,3-
331 butanediol and 2-ethyl-1-hexanol were selected and considered as important variables in the
332 projection values, because these VOCs contribute to the discrimination of pork with different
333 storage times [72].

334 Analysis of volatile organic compounds (VOCs) as fingerprinting profiles for
335 identification of dried pork slices from different processing stages have been done using GC
336 coupled with ion mobility spectrometry (GC-IMS). Using LAV software, 54 peaks were
337 selected. During this study, thirty seven VOCs were detected in the evaluated samples, in which
338 aldehydes and alcohols accounted for the largest proportion. 1-octene-3-ol has the flavour of

339 cooked mushroom, is important compound contributing to the VOCs of pork. This compound
340 is considered as the autoxidation product of linoleic acid [73]. GC-MS has been employed for
341 identification of key aroma in pork broth. The multivariate calibration of PLS is used for
342 screening the relatively better flavour of pork broth among different stewing time and applied
343 for assisting the quantitative analysis of VOCs using standard internal of 1,2-dichlorobenzene.
344 From this study, the key odorants of the aroma profile of pork broth were identified namely 4-
345 hydroxy-2,5-dimethyl-3(2H)-furanone, hexanal, 1-octen-3-ol, (E)-2-octenal, (E)-2-decenal,
346 (E)-2-undecenal, (E, E)-2,4-decadienal, nonanoic acid, decanoic acid, 2-heptanone, 3-hydroxy-
347 2-butanone, δ -decanolactone, and 2-acetylpyrrole [74].

348 GC-MS coupled with olfactometry (GC-MS/O) and in combination with chemometrics
349 of PCA and PLS-DA was reported to differentiate Chinese marinated pork hocks from four
350 different local brands. The results of PCA and PLS-DA indicated that both chemometrics using
351 variable of VOCs could clearly separated marinated pork hocks according to its groups. There
352 are nine odour-active compounds having the high loading capability for discrimination namely
353 heptanal, nonanal, 3-carene, D-limonene, β -phellandrene, p-cymene, eugenol, 2-ethylfuran and
354 2-pentylfuran. This study concluded that the validated GC-MS/O offered an alternative tools
355 for the differentiation of VOCs profile in different brands of marinated pork hocks [75].

356

357 **6. Analysis alcoholic compounds in products using chromatographic techniques**

358 GC-MS is an excellent method for analysis of alcoholic compounds in foods. Park et
359 al. have validated and reported GC-MS for the simultaneous analysis of five alcohols
360 (methanol, ethanol, propanol, butanol and pentanol) in fermented Korean foods. The separation
361 of alcohols was carried out using silica-based INNOWAX column (film thickness 0.25 μ m,
362 i.d. 250 μ m, length 30 m) coated with poly-ethylene glycol and applying mass selective
363 detector set to determine the specific selected ions for each alcohol. The LoD and LoQ values
364 ranged from 0.25 to 1.16 mg/kg. The precision and accuracy of GC-MS are acceptable as
365 indicated by Intra-day and inter-day RSDs for individual alcohols of below 7%, with recovery
366 values of 90.79 -101.50%. The method is valid, therefore, the developed method is suitable for
367 analysis of alcohols in food samples intended in Halal food authentication supporting the
368 certification processes [76].

369 Mahama et al. has applied GC with flame ionization detector (GC-FID) for analysis of
370 alcohol (ethanol) in marketed post samples (Fruit and vegetable juices from concentrate,
371 syrups, sauce samples etc.) in Thailand for identification of non-halal components suspected
372 to be present in the products. The internal standard used is n-propanol. Ethanol, internal

373 standard and others were separated using capillary columns DB-WAXTER (Agilent
374 Technologies, 30 m by 0.32 mm by 1.00 μm) with temperature of FID was set at 250°C. Some
375 certification bodies have different regulation related to the maximum limits of ethanol, and the
376 majority allowed the maximum limit is 1%. The surveillance results indicated that 1 of 24 sauce
377 samples showed an ethanol concentration of 1.0%. Furthermore, an about of 4% of all the
378 concentrated syrup samples exhibited a higher percentage of ethanol than that permitted for
379 Halal products. GC-FID method using a column HP-5 (5% Phenyl 95% Methyl Siloxane) is
380 also valid for analysis of vinegar samples from Indonesia and Saudi Arabia offering reliable
381 technique for alcohol determination [57].

382 Šorgić et al. developed gas chromatography coupled with the flame ionization detector
383 and headspace autosampler (HSS-GC/FID) method for analyzing volatile compounds in the
384 wine samples. The HSS-GC/FID method was developed, validated, and verified for
385 determining content of methanol, higher alcohols, and esters. The developed method was met
386 the validation requirement for linearity, range, sensitivity, accuracy, and precision parameters.
387 Two grape varieties namely Merlot and Cabernet Sauvignon were analyzed. It was found that
388 contents of the methanol were 198.0 mg/L and 150.5 mg/L, higher alcohols were 398.5 mg/L
389 and 335.8 mg/L, ethyl acetate were 42.0 mg/L and 55.6 mg/L, and acetaldehyde were 23.3
390 mg/L and 16.1 mg/L for Merlot and Cabernet Sauvignon varieties, respectively. This study
391 revealed that the higher content of methanol was influenced by type of grape used for
392 preparation as well as maceration duration. Further evaluation were carried out using PCA to
393 assess the effect of genotypes variation and extraction methods on wine samples [77].

394 Gas chromatography combined with PCA and cluster analysis (CA) were successfully
395 applied in determining content of alcoholic compound in Chinese beverages. According to the
396 study, twenty one aroma components were found to be important in the aroma profiles of
397 Chinese liquor. Among all the compounds, seven alcoholic compound including methanol, 2-
398 butanol, 1-propanol, isobutanol, *n*-butanol, isoamylol and phenylethanol were detected by
399 validated GC analysis method. Isoamylol, isobutanol, and 1-propanol were found as the
400 dominant alcoholic compound with the content of 800.53, 637.67, and 338.84 mg/L,
401 respectively. The dimensionality reduction of PCA were employed in this study to statistically
402 separated young liquor (fresh) and aged liquors. Individual plot was generated as two
403 dimensional visualization constructed by PC1 and PC2 with total variance of 98.27%. Further
404 separation using CA was built using the Euclidean distance. All liquor samples were clustered
405 into two big groups of young liquor and aged liquors. This results proved the ability of PCA
406 and CA to successfully separate and classify the different ages Chinese liquor samples [78].

407 In Indonesia, a majority Muslim country, it was stated by the government that the
408 alcohol content (in percentage) of alcohol-containing drugs, traditional medicines, and
409 supplements have to be declared on the label. Halal evaluation of alcohol content in noni
410 (*Morinda citrifolia* L.) can be performed using gas chromatography method. The GC
411 instrumentation was set as the inlet injection mode split of 2.5:1, injection temperature of
412 140°C, oven initial temperature FID detector of 40 °C, and hold for 5 minutes. The sample of
413 noni herbal medicines were collected from herbal drugstores or online shops in Jakarta,
414 Indonesia. Twenty samples were evaluated and categorized as beverages (18 samples) and
415 herbal medicines (2 samples). It was found that thirteen out of twenty samples contained
416 alcohol in the range of 0.04 - 1.07%. Unfortunately, none of them were labelled properly
417 according to the regulation [79].

418 GC-FID has been used for analysis of ethanol in foods and beverages such as tea-based,
419 fruit-based, cheese-based, milk-based, seaweed-based, instant dried noodle, etc. Ethanol stock
420 solution was prepared (1mg/mL) and internal standard of 0.1% v/v 1-propanol was used for
421 sample preparation. Sample preparation was carried out using magnetic stirring aqueous
422 extraction. Analysis was performed out using an HP-Innowax (Agilent technologies) column
423 (30 m x 0.25 mm x 0.25 µm). The sample injection volume was 1 µL using split ratio of 13:1.
424 The developed method was validated according to the requirements of ISO/IEC 17025:2017.
425 Validation result showed that the method had good linearity ($R^2 > 0.999$), good accuracy
426 (recoveries of 96-105%), and good precision (RSD < 5%). The detection limit was low (0.006
427 mg/g). The determination of ethanol concentration was successfully applied in 108 samples of
428 processed foods and beverages. Therefore, this method could be used as valid method for halal
429 authentication of processed foods and beverages [58].

430 GC-MS using static headspace has been applied for determination of ethanol in solid
431 and semi-solid consumer goods such as cakes, ice creams, sauces, and powders. Sample
432 preparation was carried out using mechanical homogenization and aqueous dilution of the
433 products. Subsequently, the sample was analysed using headspace GC-MS. Separation of
434 analytes was performed using a capillary column DB-624 (30 m x 0.25 mm x 1.4 µm) and
435 sample was injected in split mode employing ratio of 1:200. Identification and quantification
436 of ethanol and ethanol-d6 was performed at scan range of 29-250 m/z with a rate of 6.1 scans/s.
437 Result showed that the developed method was specific to detect ethanol and ethanol-d6 at the
438 retention time of 2.65 and 2.61, respectively. The method demonstrated good linearity at the
439 concentration range of 0.1-2.0% v/v showed by its high R^2 value (>0.998). Additionally, good

440 accuracy as well as good precision was obtained. The accuracy was represented by recoveries
441 value (average recoveries of 99.7%). The precision was demonstrated by its lower RSD value
442 (<1.5%). From the above results, it suggested that headspace GC-MS could be used for
443 identification and quantification of ethanol in a various solid and semi solid food products for
444 halal authentication [80].

445 Identification of ethanol using headspace GC-MS has also been applied in Kombucha
446 products. Kombucha is one of fermented beverages consist of sugar, tea, a symbiotic of bacteria
447 and yeast which is commonly known as non-alcoholic beverage. The United States and Canada
448 state that the content of alcoholic compounds in product must be <0.5% and <1.1% alcohol by
449 volume, respectively to be categorized as non-alcoholic drink. Propan-1-ol was used as internal
450 standard for ethanol quantification. The condition of headspace was incubation temperature at
451 70°C, syringe temperature at 70°C, incubation time of 300s, agitator speed at 500 rpm, injection
452 volume of 500 µL, and split ratio of 10:1. Analysis was performed using an Agilent J&W DB-
453 624 UI (30 m x 0.25 mm x 1.4 µm) applying flow rate of 1.4 mL/min (constant flow). The
454 developed method was linear ($R^2>0.995$) obtained at concentration range of 0.025%-2.47%.
455 The accuracy result was good demonstrated by its recovery value (102%) and good precision
456 was also obtained (RSD<4%). The LOD and LOQ values were 0.0002% and 0.002%,
457 respectively. It can be concluded that the method is suitable for identification and quantification
458 of ethanol in Kombucha product. It indicated a rapid and easy integration of analytical method
459 for halal authentication of Kombucha [81].

460 The development of GC-MS coupled with headspace and multidimensional (heart-cut)
461 chromatography has been successfully applied to determine ethanol content in medicinal
462 syrups. The aim was to ensure and guarantee the safety of the syrups. Samples used for analysis
463 consist of adult and paediatric syrups. Monitoring and quality control of ethanol content in
464 pharmaceutical products were important due to the efforts of industry to reduce the ethanol
465 content in the pharmaceutical and medicinal products. Sample preparation was directly
466 performed using headspace with condition as follows: heating syringe temperature of 90°C,
467 incubator temperature of 100°C, incubation time 15 min at 500 rpm, sample volume of 500
468 µL with split mode using ratio of 1:20. Two dimensional GC analysis was carried out using
469 GC-MS equipped with analytical column of RTX-5 capillary column (Crossbond® 5%
470 diphenyl/95% dimethyl polysiloxane, 30 m × 0.25 mm × 0.25 µm) for the first dimension then
471 for the second dimension used an NST 100 MS column (Carbowax polyethylene glycol, 30 m
472 × 0.25 mm × 2.00 µm). The method was validated according to National Agency of Sanitary
473 Surveillance (ANVISA) with validation parameters of selectivity, linearity, precision,

474 accuracy, LOD, LOQ, and robustness. Selectivity test found that isopropyl alcohol was an
475 interfering compound of ethanol determination in syrups. Linearity assay demonstrated linear
476 model at concentration range of 0.25% to 10.00% v/v ($R^2 > 0.999$). The developed method was
477 sensitive enough as shown by its LOD value (0.03% v/v) and LOQ value (0.06% v/v). The
478 precision was measured for repeatability (CV=3.04%) and intermediate precision
479 (CV=3.03%). The recoveries value obtained ranged from 97.28%-101.38% indicating good
480 accuracy. The robustness test showed that the method remains unchanged with the small
481 changes of several parameters. This developed method could be used as rapid and easy
482 analytical technique for halal authentication of syrups by determining of the ethanol content
483 [82].

484

485 **7. Conclusion**

486 Chromatography-based method consist of liquid chromatography and gas
487 chromatography using various detectors has been widely applied for food and pharmaceutical
488 products authentication including halal analysis due to its advantages. Combination with
489 chemometrics of multivariate analysis, a powerful statistical analysis to manage huge data
490 generated from analytical measurement, could be used to identify potential markers to
491 differentiate halal and non-halal samples. It will be very useful for the institutions which have
492 responsibility for halal quality assurance. Chromatogram and peak separation profiles resulted
493 as the instrument responses can be further evaluated for determination as well as quantification
494 for halal and non-halal components in food and pharmaceutical products. Chromatographic-
495 based method methods were successfully carried out to analyze products containing non-halal
496 material such as pork and alcoholic compound. Combination of chromatographic-based
497 method and chemometrics techniques with some scenarios can be applied for halal research on
498 food and pharmaceutical products.

499

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518

519 **Conflict of interest**

520 The authors declare no conflict of interest.

521

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788 **Figure and Scheme captions**

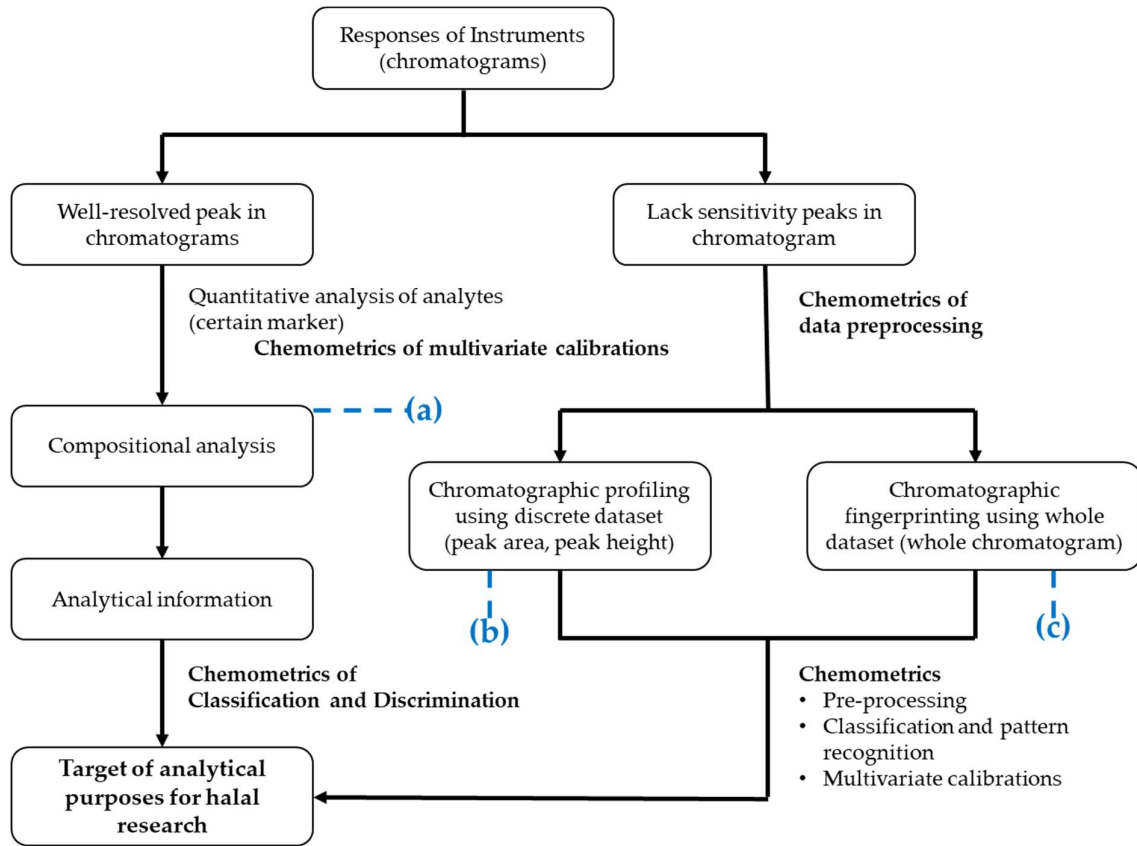
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790 **Figure 1:** Three different scenarios (a, b, and c) of chemometrics applications employing the
791 chromatograms as variable for obtaining the analytical purposes (classification of halal and
792 non-halal products as well as prediction the levels of non-halal components in the products).

793 Adapted from [24].

794

796 **Figure 1**



799 **Tables and Table captions**

800

801 **Table 1.** The application of liquid chromatography (HPLC and LC-MS/MS) for analysis of
 802 halal components in the food and pharmaceutical products

Methods	Issues	Results	References
HPLC-UV detection	Identification of pork in meatball products	HPLC-UV in combination with PCA could classify meatballs containing pork and beef in the products using variable of hydrolysis of Triacylglycerols (TAG). However, the authors did not mention which TAG markers contributing to this classification.	[31]
HPLC-Fluorescence detector	Identification of pork through amino acid composition	HPLC using fluorescence detector has been successfully applied for differentiation of pork and other animal meats based on analysis of derivatized amino acids with orto-phtalaldehyde. The amino acid VAL can be identified as marker for differentiating pork from the other meats studied (beef, chicken mutton, and chevon).	[32]
HPLC-Fluorescence detector	Detection of pig collagen using D,L-amino acids	Pre column derivatization using R(-)-4-(3-isothiocyanatopyrrolidin-1-yl)-7-(<i>N,N</i> -dimethylaminosulfonyl)-2,1,3-benzoxadiazole [R(-)-DBD-PyNCS] could be used to determine D,L-amino acids in pork collagen. Three amino acids of D-Asp, D-Pro, and D-Hyp were first detected in pork collagen samples.	[33]
LC-MS/MS with multiple reaction monitoring (MRM)	Detection of Horse and Pork in Halal Beef	Biomarker peptides were successfully identified by a shotgun proteomic approach using tryptic digests of protein extracts. Pork was identified by peptide markers: TLAFLFAER (from myosin-4), SALAHAVQSSR (from myosin-1 and myosin-4). The detection limit is 0.55% horse or pork in a beef matrix.	[34]
HPLC-MS/MS with MRM	Detection of Pork in Highly Processed Food by analysis of specific tryptic marker peptides	HPLC-MS/MS using MRM has been successfully applied for analysis of pork in some processed food products (cooking, frying and baking) based on peptide markers which are specific for pork. The peptide markers of pork identified based on MRM experiment were: marker 1 (YDIINLR) markers 2 (TLAFLFAER) and 3 (SALAHAVQSSR).	[35]
LC-MS/MS	Differentiation of porcine gelatine and bovine gelatine	LC-MS/MS in combination with exploratory data analysis of PCA could discriminate porcine and bovine gelatines. Based on loading plot PCA, peptides appearing in retention time (t_R) 32 min could be identified as peptide markers	[36]
Nano UPLC-Q-TOF-MS	Differentiation of porcine and bovine gelatin in food products	Marker peptide of bovine and porcine gelatin could be detected using nano UPLC-Q-TOF-MS based data dependent technique in yoghurt, cheese, and ice cream. The method could be used to detect bovine and porcine gelatin in the mixtures.	[37]

Nano UPLC-Q-TOF-MS	Differentiation of porcine and bovine gelatin in food products	Marker peptide of bovine and porcine gelatin could be detected using nano UPLC-Q-TOF-MS based data dependent technique in yoghurt, cheese, and ice cream. The method could be used to detect bovine and porcine gelatin in the mixtures.	[37]
LC-MS QTRAP	Gelatin speciation (bovine, porcine, and fish)	LC-MS in combination with PCA could differentiate bovine, porcine, and fish gelatin. PLS-DA could be used for classification of pure gelatin and adulterated gelatin (fish and bovine) with porcine gelatin using several concentration levels of porcine gelatin.	[38]
LC-MS/MS	Discrimination of raw beef, pork, poultry and their mixtures	Protein of troponin I (TnI), enolase 3, L-lactate dehydrogenase (LDH), triose-phosphate isomerase (TPI), Tropomyosin 1 and carbonic anhydrase 3 could be used as potential markers to distinguish mammals and poultry.	[39]
LC-Q-TOF-MS	Differentiation between dead-on arrival and normally slaughtered of poultry meat	LC-Q-TOF-MS could be used to differentiate between normally slaughtered and dead-on arrival poultry meat based on metabolic profiles analysed using multivariate analysis. Using METLIN and analysis of chemical standards, metabolite of sphingosine was found to be potential marker for dead-on arrival poultry meat.	[40]
UPLC-TOF-MS	Metabolite's differentiation of broiler chicken slaughtered using different techniques	UPLC-TOF-MS could be used to distinguish between halal slaughtering method and non-halal slaughtering method of broiler chicken based on their metabolite profiles. Non-halal slaughtered method demonstrated high amino acid and high glucose breakdown.	[41]
LC-HRMS	Analysis of pork meat in meat mixtures using PRM	Five peptides of myosin were screened and used for PRM analysis using LC-Orbitrap HRMS. Peptide of KLETDISIQGEMEDIVQEAR was found to be the most sensitive peptide with LOD value of 0.5% in meat mixtures.	[42]
UPLC-MS	Detection of pork adulteration in beef using metabolomics approach	PLS-DA using metabolomics data obtained from untargeted measurement could classify pure and adulterated beef samples with pork. There was a significant difference in the metabolism of inositol, glutathione, and sphingolipid between beef and pork.	[43]
LC-MS/MS	Detection of pork adulteration in meat samples using carbonic anhydrase 3 as a marker	Three peptides from carbonic anhydrase 3 were found as marker of pork (EPITVSSDQMAK, GGPLTAAAYR, HDPSLLPWTASYDPGSAK). Quantification analysis could be performed using those three peptides with perfect quantitative ability and provided good correlation and recovery results.	[44]

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809 **Table 2.** The application of gas chromatography (GC-FID and GC-MS) for analysis of halal
 810 components in the food and pharmaceutical products

Methods	Issues	Results	References
GC-FID for analysis of alcohol	Determination of ethanol contents in vinegar	The maximum contents of ethanol in vinegar is 1.0%. GC-FID could determine the levels of ethanol (alcohol) in the marketed vinegar samples. The detection level of ethanol was about 0.4 mg%.	[57]
GC-FID for analysis of ethanol in foods	Determination of ethanol in different processed foods and beverages	Extraction technique using aqueous extraction assisted magnetic-stirring could be used to extract ethanol from different foods and beverages. GC-FID successfully used to determine ethanol with good validity. The validated method was successfully used to determine ethanol in 108 food and beverage products.	[58]
GC-MS for analysis of alcohol	Determination of alcohol in fermented black tape ketan using GC-MS	GC-MS could be used for quantitative analysis of alcohol content in fermented black tape ketan with good recovery (89%). The alcohol concentrations determined at 3, 10, 17, 24, and 31 days were 4.295, 4.23, 5.005, 4.747, and 5.344 % v/v, respectively.	[59]
GC-FID for analysis of lard	Differentiation of lard from other edible fats using GC-FID and chemometrics	Lard contains high amount of C18: 2cis and low amount of C16:0. Chemometrics of PCA and K-mean cluster analysis could differentiate lard adulteration on chicken fat and beef tallow at low concentrations (0.5%-10%).	[60]
GC-MS for analysis of pork	Analysis of fatty acids and fatty acid methyl esters of pork (non-halal meats) in sausages compared with beef sausages (halal meat)	The dominant fatty acids in pork sausage are palmitic, myristic, oleic acid, and lauric acids. While fatty acids dominating in beef sausage are palmitic, oleic, stearic and myristic acids. The chemometrics of PCA could classify sausages according to meat sources (beef and pork)	[61]
GC-MS for analysis of rat meat	Analysis of rat meat (non-halal meat) and its classification with other meats using chemometrics of PCA	Six fatty acids, i.e. myristic, palmitoleic, palmitic, linoleic, oleic and stearic acids combined with PCA could classify rat meat and other meats.	[62]
Headspace GC-MS for analysis of pork	Differentiation of pork (non-halal meat) and pork sausages from beef, mutton and chicken meats	The samples were introduced into GC instrument using headspace, and volatile compounds present in the evaluated samples were separated using GC and detected by MS. The chemometrics of PCA provided good separation between pork-based sausages and halal meat-based sausages.	[63]
GC-MS for analysis of lard	Analysis of lard (non-halal fat derived from adipose lard)	The fatty acid of 11,14-eicosadienoic acid is used as fatty acid marker for identification of lard.	[64]

	tissue of pig) in chocolate products		
GC-MS-SPME for analysis of wild boar	Volatilomics analysis of non-halal (wild boar) meat ball using GC-MS-SPME and chemometrics	PLS-DA could be used to differentiate volatile compounds of halal meatball and non-halal meatball. Compounds of β -cymene, 3-methylbutanal, and 2-pentanol were found to be potential markers for chicken meatball. Compounds of 5-ethyl-m-xylene, benzaldehyde, and 3-ethyl-2-methyl-1,3-hexadiene were associated to the potential markers of beef meatball. Compounds of pentanal, 2,6-dimethylcyclohexanone, 1-undecanol, cyclobutanol, 2,4,5-trimethylthiazole, and 5-ethyl-3-(3-methyl-5-phenylpyrazol-1-yl)-1,2,4-triazol-4-amine could be used as potential markers as wild boar meatball.	[65]
HS-SPME-GC-MS for analysis of minced beef and pork meat	Volatilomics analysis using HS-SPME-GC-MS combined with multivariate analysis to differentiate minced beef and pork meat	GC-MS based on volatilomics analysis and chemometrics of PCA and PLS-DA could be used to differentiate minced beef and pork meat. Heptanal, octanal, butanol, pentanol, hexanol, 1-penten-3-ol, 2-octen-1-ol, 3-hydroxy-2-butanone were associated to the potential markers of beef whereas pentanal, hexanal, decanal, nonanal, benzaldehyde, trans-2-hexenal, trans-2-heptenal could be used as potential volatile compound markers of pork meat.	[66]
GC-MS for analysis of pork	Detection of pork in beef meatball using GC-MS and chemometrics	PCA using fatty acid compositions of pure beef meatball and adulterated beef meatball using pork as the variables successfully differentiate pure and adulterated beef meatball. The ratio of SFA:MUFA of pork meatball was 1.0.	[67]
GC-MS for analysis of house rat	Detection of rat house in beef meatball by analysis of fat using G-CMS	The fatty acids composition of house rats were myristate (0.19 \pm 0.03)%, palmitoleat (2.40 \pm 0.29)%, methyl palmitate (27.65 \pm 0.32)%, oleate (45.81 \pm 3.25)%, and stearate (4.65 \pm 0.28)%. Analysis using PCA could differentiate beef meatball and beef meatball containing rat house meat. Further analysis using PCA demonstrated that fatty acids of house rats have high similarity to chicken fatty acids.	[68]
GC-MS for analysis of lard	Detection of lard in wheat biscuits using GC-MS and chemometrics	PCA using fatty acids composition could differentiate lard, wheat biscuits, and adulterated wheat biscuits with lard. PLS-DA could be used to find potential marker for differentiation. Fatty acid of C18:3n6 is suggested as potential marker to distinguish	[69]

			pure wheat biscuits and adulterated wheat biscuits with lard.	
GC-MS for analysis of dog fat	for Detection of dog fat from other animal fats using GC-MS and chemometrics		Nine types of fatty acids in dog fat were found such as lauric, myristate, pentadecanoate, palmitoleate, palmitate, margarate, oleat, stearic, and arachidonic. Analysis PCA showed that dog fat is close to lard.	[70]
GC-MS for analysis of rat fat	for Detection of Sprague Dawley rat fat in meatball using GC-MS and chemometrics		PCA could differentiate meatball and adulterated meatball with Sprague Dawley rat meats. Further analysis revealed that the Sprague Dawley rat fat is close to beef fat.	[71]

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The screenshot shows a Gmail interface with an email from L.JFP-peerreview@journals.tandf.co.uk. The email subject is "Submission received for International Journal of Food Properties (Submission ID: 221850525)". The email content includes the Taylor & Francis logo and a message to Abdul Rohman. The message text is as follows:

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Thank you for your submission.

Submission ID: 221850525
Manuscript Title: **The Use of Chromatographic Based Techniques and chemometrics for Halal Authentication of Food Products: A review**
Journal: International Journal of Food Properties
Article Publishing Charge: USD \$1160.00 (plus VAT or other local taxes where applicable in your country)

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The screenshot also shows the Windows taskbar at the bottom with the date 16/06/2022 and time 13:45.

Masukan reviewer

The screenshot shows a Gmail interface with an email from the International Journal of Food Properties. The email subject is "221850525 (International Journal of Food Properties) A revise decision has been made on your submission". The email content includes the following text:

Dear Dr Rohman:

Your manuscript entitled "The Use of Chromatographic-Based Techniques and chemometrics for Halal Authentication of Food Products: A Review", which you submitted to International Journal of Food Properties, has been reviewed. The reviewer comments are included at the bottom of this letter.

The reviews are in general favourable and suggest that, subject to minor revisions, your paper could be suitable for publication. Please consider these suggestions, and I look forward to receiving your revision.

When you revise your manuscript please highlight the changes you make in the manuscript by using the track changes mode in MS Word or by using bold or colored text.

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Editor, International Journal of Food Properties
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Reviewer: 1

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Comments from the Editors and Reviewers

Reviewer: 1

Comments to the Author

This is a good review to see the outcome of analysis in various settings using **chromatographic** and chromatomatic analysis. Some English errors are found but minimum.

The literature search took over 2 months period (but the statement in abstract might be understood as written within 2 months?). It would be nice to know how many articles were retrieved and how was the selection made. I noticed some long paragraph discussing in detail the outcome from a single study in the respective paragraph citing only one single reference each while there are >80 articles being listed in the reference list, some paragraph cited a single reference at the very end making the starting point of the paragraph ambiguous on whose study it was referring to.

Reviewer: 2

Comments to the Author

The manuscript "The Use of **Chromatographic Based Techniques and chemometrics for Halal Authentication of Food Products**: A Review" that has been sent for publication in the journal "International Journal of Food Properties" provides an interesting review on **Halal Authentication of Food Products**.

The article needs some minor corrections, which are indicated below:

- 1- The keywords on the home page and those on lines 37 and 38 do not match. I think the ones on lines 37 and 38 are more correct, but the term pharmaceutical should be deleted.
- 2- Lines 148-156: Include the following reference "Chromatographic Fingerprinting: An innovative approach for **food** identification and **food authentication**". A *Journal Analytica Chimica Acta* 909 (2016) 9-23 which clearly explains what a marker, a profile and a **chromatographic** fingerprint are. Include the term marker in the paragraph.
- 3- Lines 273-280: This paragraph should be deleted, medicinal herbs are not the subject of the study.
- 4- In tables 1 and 2, there is no specific application dedicated to **halal** pharmaceutical. It only appears in the main discussion. I think it should be removed from the main discussion or include specific applications in **halal** pharmaceutical. In fact in the title the term **halal** pharmaceutical does not appear.

General comment: The authors have two options: To include in tables 1 and 2 specific references to **halal** pharmaceuticals or to remove all comments on **halal** pharmaceuticals.

Editor's Comments

Comments are marked in the attached file. Please work in the attached file so that format remains the same.

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Sincerely,
Dr Rohman
Editor, International Journal of Food Properties
shafiq@tqf.edu.au

Editor's Comments to Author

A copy of the final version is enclosed.

22-may-2022-4-R1

1 **The Use of Chromatographic-Based Techniques and**
2 **chemometrics for Halal Authentication of Food Products: A**
3 **Review**

4
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19
20 **ABSTRACT**

21
22 Halal food ~~and halal pharmaceutical~~ products are requisite to be consumed by Muslim
23 communities in the world. The standard methods capable of quantifying non-halal components
24 are very urgent. This review highlights ~~the chromatography~~ [chromatographic methods](#) and
25 chemometric ~~based techniques~~ [or multivariate data analysis](#) that offer reliable techniques to
26 provide [the](#) separation capacity in halal authentication analysis.

27 Methods: This review article was written from reputable worldwide databases including Web
28 of Science, Scopus, and PubMed, between January and February 2022. The keywords were
29 “halal research”, “food analysis”, ~~“pharmaceutical analysis”~~, “chromatography”,
30 “chemometrics”, and “authentication”. Chromatographic-based techniques [in](#) combination
31 with chemometrics of multivariate analysis, a powerful statistical analysis to manage huge data
32 generated from analytical measurement, could be used to identify potential markers to
33 differentiate halal and non-halal samples. Chromatogram and peak separation profiles resulted

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34 as the instrument responses can be further evaluated for determination as well as quantification
35 ~~for of~~ halal and non-halal components in food ~~and pharmaceutical~~ products.

36 Combination of chromatographic-based method and chemometrics techniques with some
37 scenarios can be applied for halal research on food ~~and pharmaceutical~~ products.

38
39 **Keywords:** halal authentication, chemometrics, chromatography, pig derivatives, ~~food~~
40 ~~products~~ ~~pharmaceutical~~.

42 ~~1.~~ INTRODUCTION

43
44 Food, ~~cosmetics, drugs and other pharmaceutical products~~ ~~and pharmaceutical products~~ are
45 important needs for human beings. In line with the development of science and technology,
46 industrialization and globalization, the halal products may be added or substituted and
47 contaminated with non-halal components such as pig derivatives and alcohols as ingredients or
48 additives to reduce the production cost ~~make the products non-halal~~ [1]. In addition, the
49 products available in markets may contain incorrect labelling in terms of ingredient sources
50 making the consumers lost on composition information, therefore the use of analytical tools to
51 check the presence of non-halal components in the products is a must for assisting the
52 certification processes [2]. In Indonesia, the halal certification is mandatory which means that
53 all halal declared products sold and distributed in Indonesia must be halal certified. In addition,
54 the analysis of non-halal components in post-marketed products is also needed to confirm that
55 the marketed products are not adulterated with non-halal components [3].

56
57 According to Indonesian Act No. 33 (2014), the certification process is carried out by Halal
58 Product Assurance Organizing Agency (BPJPH) and the auditing process was carried out by
59 Halal Examination Agency (LPH). During audit, if the products are supposed to contain non-
60 halal components (pork derivatives and alcohols), the laboratory testing using standard
61 analytical methods is needed to confirm that the audited products are free from any non-halal
62 components [4,5]. Today, the Muslim community constitute for approximately of 25% world's
63 population and is expected to increase further. With the increased awareness among Muslim
64 community to consume the only halal products, the global market of halal products could reach
65 exponentially [6]. Halal is Arabic terms used to any products permissible to be consumed by
66 Muslim community. Today, the term of halal has widely used not only Muslim but also non-
67 Muslim because Non-Muslim community intended to export the products into Muslim

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68 community, especially in halal certification issues [7]. Therefore, it is not surprising that halal-
69 related studies are performed not only in majority Muslim countries like Indonesia and
70 Malaysia but also in countries whose Muslims are minority such as the Netherlands, the United
71 States, France and the European Union [8].

72
73 Halal food ~~and Halal pharmaceuticals~~ must be free from non-halal components which are pig
74 and all pig derivatives such as pork, lard and porcine gelatines, carrion or dead animals, blood
75 (flowing or congealed), animals slaughtered not according to Islamic law, animals that were
76 killed accidentally or on purpose through means such as strangling or beating, intoxicants
77 including alcohol and drugs [9], carnivorous animals, predator birds, and certain land animals
78 [10]. Among these, pig derivatives and alcohols are typically found in halal ~~and~~
79 ~~pharmaceutical~~food products, therefore some scientists are continuously researches on halal-
80 related issues including developing instrumental analytical methods for detecting of non-halal
81 components intended for halal certification [11]. Some countries have obligated the products
82 to be halal certified which can be understood that the products are free from prohibited
83 components. Besides, the products are manufactured using equipment dedicated for halal food
84 ~~and halal pharmaceuticals~~ [12]. Pork is typically met in meat-based food products such as
85 meatball, sausages, etc.; while lard can be good vehicle in some cosmetics products such as
86 cream, lipstick and lotion. Porcine gelatines are common materials used in food (in candies)
87 and pharmaceutical products (capsule shells) [13]. The objective of this review was to provide
88 the integrative information on identification and quantification of non-halal components in
89 food ~~and pharmaceutical~~ products by chromatographic methods. In addition, chemometrics
90 techniques were reported to be applied to employ the big data evaluation as resulted from the
91 chromatographic detection.

92 93 ~~2.~~ **METHODS**

94
95 This review article was written by identifying, investigating, and assembling several review
96 articles, original articles, books, and relevant sources on metabolite
97 ~~fingerprintings~~fingerprinting from reputable worldwide databases including Web of Science,
98 Scopus, and PubMed. Literature searching was carried out between January and February 2022.
99 The keywords explored during literature investigation were “halal research”, “food analysis”,
100 ~~“pharmaceutical analysis”~~, “chromatography”, “chemometrics”, and “authentication”. First, to
101 select the suitable papers, 250 articles were reviewed through the title and abstract. The

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102 inclusion criteria to select the papers were (1) studies regarding halal authentication of food
103 products using chromatographic technique between 2005-2022; (2) studies on analysis of non-
104 halal components in food products using liquid chromatography and gas chromatography
105 conducted between 2005-2022; (3) studies on the employment of chemometrics in combination
106 with chromatographic technique for halal authentication of food products; (4) all papers written
107 in English. The exclusion criteria of the papers were (1) studies on halal authentication of food
108 products using chromatographic techniques published before 2005; (2) all articles written using
109 language other than English.

110 During The criteria

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113 **3. Chromatographic-based techniques and chemometrics for analysis of non-halal** 114 **components**

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115
116 For many years, chromatography has been known as the method of choice to assess the purity
117 and levels of analytes in the laboratories of research, industry, and quality control [14]. Gas
118 chromatography (GC) and liquid chromatography (LC) techniques are often used for the
119 analysis of non-halal components in food ~~and pharmaceutical~~ products. In terms of compound
120 types, GC is more suitable for the analysis of smaller, volatile and stable compounds to heat,
121 while LC is more robust and suitable for larger and less/non-volatile compounds [15]. Some
122 derivatization techniques are needed in LC in order to convert analytes into detectable derivates
123 with certain detectors, while derivatization in GC for fewer volatile compounds is intended to
124 provide more volatile and stable derivate products, although this derivatization process
125 increases the method complexity and lengthens the sample preparation. In addition, the
126 availability of derivative agents and its steric hindrance in the analyte, and the stability of the
127 derivatized compounds must also be considered [16].

128
129 One-dimensional gas or liquid chromatography using one column is considered as simple and
130 powerful separation techniques for simple and un-complex samples. When the analyzed
131 samples are complex enough, the application of just one-dimension chromatography leads to
132 peak co-elution as well as overlapping and non-resolved peaks, therefore one dimension
133 chromatography technique is not suitable for separation of large analytes because the peak
134 capacity of one-dimensional analysis is not large enough to achieve the complete separation
135 with acceptable resolution [17]. In last decades, two-dimensional gas chromatography (GC x

136 GC) and liquid chromatography (LC x LC) has been applied in analysis of complex mixture in
137 order to increase the separation speed [18].

138
139 In two-dimensional chromatography, the separation is carried out in two columns with different
140 polarity connected in series by a modulator, as a consequence, the separation capacity of
141 regular one-column in one dimensional chromatography can be considerably increased. The
142 effluent from the first column is transferred to the second column using modulator so that the
143 analytical information obtained (such as retention times, t_R) from the first column can be
144 combined with that from second column, leading to a plot of two retention times [19]. Because
145 of the excellent separation capacity of GC x GC and LC x LC combined with mass
146 spectrometry (MS), both techniques are applied for separation all components in the complex
147 mixtures, especially for metabolomics studies [18]. GC x GC has been widely applied for
148 analysis of metabolites (all fatty acid types) of lard in food samples [20], while LC x LC is
149 typically used for analysis of peptides [21], which can be used for identification of pork and
150 porcine gelatines.

151
152 Chromatographic-based techniques offered reliable technique in halal authentication analysis.
153 However, due to high number of data covered, the application of chemometrics to treat big data
154 is unavoidable. Chemometrics can be defined as the employment of statistical and
155 mathematical methods to obtain the objective data evaluation by extracting the relevant and
156 meaningful information from related and unrelated responses from chemical measurements.
157 Chemometrics or multivariate data analysis (MDA) is typically applied in numerous aspects
158 including the quality control of halal products, qualitative and quantitative determination of
159 chemical parameters for assessing the products authenticity [22].

160
161 Chemometrics can provide the powerful tools in giving important information extracted from
162 big data obtained from instrumental analyses such as methods based on spectroscopic and
163 chromatographic. The common chemometrics techniques applied in products authentication
164 could be grouped into exploratory data analysis, data pre-processing, description and
165 visualization, discrimination and pattern recognition (classification), regression and prediction
166 and experimental design [23]. Some chromatographic problems encountered during halal
167 authentication analysis included the assessment of separation quality, the evaluation of peak
168 alignment using pre-processing, the optimization of chromatographic systems providing the
169 good separation of all peaks using experimental design, the accuracy of discrimination and

170 classification using pattern recognition, and quantitative analysis applying multivariate
171 calibration. Figure 1 showed the correlation between chromatographic responses and
172 chemometrics for certain analytical purposes. In scenario (a), peaks with good separation (good
173 selectivity) in chromatogram was used as variable for the evaluation of compositional analysis
174 (concentration) of analytes assisted by multivariate calibrations. In (b), certain peaks with lack
175 selectivity was used as variable during chromatographic profiling of objects (samples) using
176 discrete datasets (peak area or peak height), while in scenario (c), whole datasets in
177 chromatograms were used as variables during chromatographic fingerprinting of objects.
178 Indeed, the chemometrics of pre-processing was widely applied to obtain the desired analytical
179 modelling.

180
181 The classification chemometrics was typically carried using (1) exploratory data analysis
182 including principal component analysis (PCA) and cluster analysis (hierarchical cluster
183 analysis and non-hierarchical such as k-means and k-medians), and this technique is typically
184 called as unsupervised pattern recognition and (2) classification and discrimination methods
185 known supervised pattern recognition. There are two types of classification chemometrics
186 methods regardless of the statistical background. The first type is typically employed to assess
187 to which of various pre-defined classes of samples (objects). The example of this technique is
188 linear discriminant analysis (LDA), orthogonal projection to latent structures – discriminant
189 analysis (OPLS-DA), k-nearest neighbors (KNN) and many others. The second type of
190 classification chemometrics is called as class modelling or one class classifier (OCC), and the
191 example for this group data driven soft independent modeling of class analogy (DD-SIMCA)
192 and Unequal Class-Modeling (UNEQ) [25]. Using these chemometrics, someone can answer
193 the question: is the meat belong to pork (non-halal) or beef (halal)? or the question: is the
194 meatball authentic or adulterated? [26,27].

195

196 **4-Analysis of non halal components using liquid chromatography**

197

198 High performance liquid chromatography (HPLC) using certain detectors have been widely
199 applied for analysis of specific components in non-halal components. HPLC using fluorescence
200 detector has been successfully used for analysis of Hydroxyproline and other amino acids in
201 gelatin and collagen samples as initial screening for identification of gelatin types.
202 Hydroxyproline has been known as signature amino acid for gelatin and collagen. The level of
203 hydroxyproline is typically higher in the gelatin samples than that in the collagen samples,

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204 except for the samples of fish skin gelatin, and this result could be used as screening tools for
205 identification of non-halal gelatin and collagen in the analyzed samples [28].

206 There are three approaches to detect and to identify the presence of non-halal components in
207 food products using chromatographic based methods. The first approach is based on searching
208 the specific markers through analysis of the separated specific components. Indeed, the
209 availability of reference standards is a must. The second approach is used fingerprinting
210 profiles in which the chromatogram profiles of samples with and without non-halal components
211 are compared and evaluated. The third approach involved metabolomics studies either targeting
212 and untargeted techniques by analysis of all metabolites in the analyzed samples. The second
213 and third approaches involved the large datasets, therefore, the chemometrics is employed to
214 perform the analytical tasks (discrimination, classification, etc.) [29].

215
216 Table 1 listed the application of HPLC and LC-MS/MS for analysis of halal components in the
217 products. Liquid chromatography using fluorescence detector was also successfully applied for
218 analysis of amino acid (AA) composition non-halal (porcine) and halal (bovine and fish)
219 gelatins. The classification between halal and non-halal gelatins was carried using PCA
220 applying amino acid compositions as variable. AAs with strong fluorescence (Hyp, His, Ser,
221 Arg, Gly, Thr, Pro, Tyr, Met, Val, Leu and Phe) contribute to the classification and become the
222 biomarkers to identify the gelatine sources [2930]. Gelatin from three mammalian species
223 including bovine gelatin, porcine gelatin, and donkey gelatin has been successfully identified
224 using liquid chromatography-linear ion-trap high resolution mass spectrometry. Hemoglobin
225 was just found in donkey gelatin. The unique peptide obtained from donkey, bovine, and
226 porcine gelatin was GEAGPAGPAGPIGPVGAR, GETGPAGPAGPIGPVGAR, and
227 GETGPAGPAGPVGPVGAR, respectively. The unique peptides could be detected either in
228 individual gelatin or in the mixtures of three mammalian gelatins [3031].

229
230 Liquid chromatography especially combined with mass spectrometer tandem mass
231 spectrometer (LC/MS-MS) is widely applied for identification of non-halal component in food
232 and pharmaceutical products including porcine gelatin and pork. Gel-enhanced liquid
233 chromatography-mass spectrometry (GeLCMS) in combination with chemometrics of PCA
234 has been developed for identification of potential protein markers in pork and other meats along
235 with its classification. The myofibrillar protein with weight of 40-kDa such as troponin T,
236 Tropomyosin alpha-1 chain, and actin cytoplasmic 1 as well as the thin filament proteins such
237 as actin, troponin, and Tropomyosin had molecular weights ranging from 40 to 45 kDa could

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238 be used as markers for differentiation of pork from chicken and beef. PCA using PC1 and PC2
239 accounting of 62% and 35% variances could classify meat types. From MS studies, the
240 potential protein markers for pork meat samples are Troponin T with peptide sequences of
241 [(R)KPLNIDHLSSEDK(L)], Tropomyosin alpha-1 chain [(K)EAETRAEFAER(S)], Actin
242 cytoplasmic 1 [(R)HQGVMVGMGQK(D)], COP9 signalosome complex subunit 4
243 [(R)VLDYRR(K)], and Ribonuclease inhibitor [(R)VLGQGLADSACQLETLR(L)][4546].
244

245 The identification of potential biomarkers of gelatin from several sources could be performed
246 using UPLC-MS/MS. Samples used were gelatin from pig, cow, chicken, and fish. After the
247 extraction process of proteins from gelatin, proteins were then digested using proteomic grade
248 trypsin for 12 h to obtain peptides. Chemometrics of PCA was used to differentiate partial
249 hydrolysis of gelatin from cow and pig. Result from PCA score plot showed that the sample of
250 cow and pig obtained from digestion process could be well separated. For identification of
251 potential biomarkers from pig, cow, fish, and chicken gelatin, PCA employing MPP (Mass
252 Profiler Professional) was applied. Results showed that three unique peptides found only in pig
253 gelatin, seven unique peptides found in bovine/cow gelatin, 22 peptides found only in chicken
254 gelatin, and only 1 unique peptide found in fish gelatin. The developed method was also
255 successfully applied to identify species origin of commercial gelatin samples. It indicated that
256 UPLC-MS/MS offers a powerful analytical technique to identify gelatin from different species
257 in food and pharmaceutical products [4647].
258

259 Targeted tandem liquid chromatography-mass spectrometry (LC-MS) using decoy,
260 randomized and concatenated database search program comprising MS-Fit and MS-Tag in
261 combination with chemometrics of principal component analysis and orthogonal partial least
262 square-discriminant analysis (OPLS-DA) was applied for identification of potential peptide
263 markers in non-Halal meat (pork) and halal meats (chicken and beef). The peptide markers
264 which are specific to certain species were identified through shot- gun proteomics. Potential
265 peptide marker identified for raw pork is myosin-2 having sequence of peptide marker of
266 (F)DFNSLE(Q). OPLS-DA using variable of identified peptides could separate halal and non-
267 halal meats [4748].
268

269 Targeted proteomic analysis using LC-MS has been developed to investigate the heat stable
270 protein in pork meat. Five heat treatments were applied such as (1) water bath heating at 78°C
271 for 30 min; (2) boiling at 100°C for 30 min; (3) sterilizing at 121°C for 30 min; (4) frying using

272 oil until golden brown colour; and (5) baking at 200°C for 30 min. Protein extraction from
273 samples was performed using buffer solution containing 2 M thiourea, 7 M urea, and 50 mM
274 Tris-HCl (pH 8.0). Proteins were digested using proteomic grade trypsin added with DTT to
275 reduce disulphide bonds and IAA for alkylation. Incubation was carried out for at least 12 h at
276 37°C. Result showed that seven heat-stable specific peptides of pork were found such as
277 DQLIHNLLK from l-lactate dehydrogenase A chain, HDPSLLPWTASYDPGSAK from
278 carbonic anhydrase 3, EPITVSSDQMAK from carbonic anhydrase 3, VNVDEVGGEALGR
279 from haemoglobin subunit beta, HPGDFGADAQGAMSK from myoglobin,
280 SLYSSAENEPPVPLVR from carbonic anhydrase 3, and YLEFISEAIIQVLQSK from
281 myoglobin. Commercial samples such as Iberian dried ham, Pasteur dry sausage, import dried
282 ham, lunch meat canned, sandwich sausage, and Thuringia flavour sausage were used to
283 identify the presence one or more pig heat-stable peptides. Results showed that the heat-stable
284 peptides of pig could be found in various types of food products with different cooking process
285 methods. It suggested that targeted proteomics analysis using seven heat stable peptides of pig
286 could be used for halal authentication of food products especially meat-based food products
287 containing pork [4849].

288 ———
289 Analysis using LC-MS employing MRM (multiple reaction monitoring) technique was
290 successfully used to detect heat-stable peptides in cooked meats including pork meat. Thermal
291 treatment applied was boiling at 100°C, grilling at 150°C, and grilling at 180°C. After the
292 protein was extracted, digestion process was performed using proteomic grade trypsin.
293 Identification of homologues protein and potential biomarkers of pork peptide was carried out
294 using UPLC Triple TOF-MS equipped with a C-18 column (2.1 × 100 mm, 1.7 µm; Waters
295 Corporation, Taunton, MA, USA and Wexford, Ireland). The mobile phase used was water
296 containing 0.1% formic acid (A) and acetonitrile containing 0.1% formic acid (B) with flow
297 rate of 0.3 mL/min. On the other hand, MRM analysis was performed using a SCIEX ExionLC
298 AD system (AB SCIEX, Framingham, MA, USA) and an AB SCIEX QTRAP 4500 mass
299 spectrometry system (AB SCIEX PTE. LTD., Marsiling, Singapore) equipped with a column
300 of Waters ACQUITY UPLC BEH C18 (2.1 × 50 mm, 1.7 µm). Results showed that the
301 potential peptide biomarkers in raw pork meat found were GHHEAELTPLAQSHATK from
302 myoglobin, FAGGNLDVLK; ADMVIEAVFEELSLK; TVLGAPEVLLGILPGAGGTQR
303 from trifunctional enzyme subunit alpha, mitochondrial, and
304 WGDAGATYVVESTGVFTTMEK from glyceraldehyde-3-phosphate dehydrogenase.
305 Meanwhile, the heat-stable peptide biomarkers of pork were FAGGNLDVLK and

306 TVLGAPEVLLGILPGAGGTQR from trifunctional enzyme subunit alpha, mitochondrial as
307 well as WGDAGATYVVESTGVFTTMEK from glyceraldehyde-3-phosphate dehydrogenase.
308 The MRM analysis confirmed the heat-stable peptide of pork in meat product samples. It
309 suggested that LC-MS employing MRM method could be used as promising analytical
310 technique for halal authentication of meat products [4950].

311

312 **5. Application of gas chromatography for analysis of non-halal components**

313

314 ~~The use of Herbal medicines (HMs) as complementary and alternative medicine is becoming~~
315 ~~popular in the general population worldwide. Parallel to the increased trends on application of~~
316 ~~HMs as alternative therapies either for preventive or promotive, some research activities~~
317 ~~dealing with the quality control, standardization, and authentication of HMs also increased.~~
318 ~~The efficacy of HMs depends on their quality and its authenticity. Fingerprint profiling based~~
319 ~~on spectroscopic especially ¹H-NMR and chromatographic techniques hyphenated with mass~~
320 ~~spectrometers (LC-MS/MS) in combination with classification chemometrics has emerged as~~
321 ~~powerful tools for standardization and authentication of HMs.~~ Table 2 listed the application of
322 gas chromatography for analysis of halal components in the food and pharmaceutical products.

323 GC-MS combined with chemometrics has been proposed as tools for detection of lard as
324 adulterant in olive oil using metabolomic approach. GC separation of fatty acid methyl esters
325 (FAME) was achieved using HP-5MS nonpolar capillary column. The identification of
326 metabolites of FAMEs was carried out using standard FAMEs and mass spectrometer detector
327 using the WILEY 2007 library. Some FAMEs are specific, i.e., methyl behenate was only
328 present in olive oil and methyl myristate was only detected in lard. PCA using identified
329 FAMEs was successful for separating lard, olive oil and olive oil adulterated with lard for halal
330 authentication study [5051].

331

332 Two dimensional GC combined with time-of-flight mass spectrometer (GC x GC-TOF/MS) is
333 successfully used for analysis of lard as adulterant in virgin coconut oil (VCO) through analysis
334 of sterols. GC x GC system could perform the complete baseline separation of sterol
335 trimethylsilyl ethers derived from cholesterol and cholestanol, which facilitate the detection of
336 lard in VCO. Using GC x GC-TOF/MS Cholestanol trimethylsilyl ether (Cha-TME) and
337 cholesterol trimethylsilyl ether (Che-TME) were separated into some peaks, identified as Che₁,
338 Che_{bI}, Che_{bII}, Che₂ (Che-TME), and Cha₁, Cha_{bI}, Cha_{bII}, and Cha₂ for Cha-TME.

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339 Quantification of these compounds could be used as tools for quantification of adulteration
340 levels of lard in VCO [20].

341
342 GC-MS coupled with headspace solid-phase micro-extraction (HS-SPME) is successful for the
343 analysis of volatile compounds in pork. The profiles of volatile compounds from different
344 meats are different, therefore, the volatile compounds ~~analysed~~analyzed by GC-HS-SPME/MS
345 could be used as fingerprinting tools for specific meats [5152]. In addition, VOCs also
346 contribute to the aroma which can be used for the discrimination tools among animal meats
347 [5253]. Analysis of VOCs is very challenging because of different factors, including the high
348 number of volatile compounds, differences in volatility degree and the great amount of
349 functional groups [5354]. Chen et al. [5455] have identified the key volatile compounds for
350 differentiation of pork from different pig breeding. The volatile compounds contributing to the
351 pork flavour identified during this study were 3-methyl-1-butanol, 1-nonanal, octanal, hexanal,
352 2-pentyl- furan, 1-penten-3-one, N-morpholinomethyl-isopropyl-sulphide, methyl butyrate,
353 and (E,E)-2, 4-decadienal. Kosowska et al. [5556] reported that some volatile compounds
354 namely octanal, nonanal, (E,E)-2,4-decadienal, methanethiol, methional, 2-furfurylthiol, 2-
355 metyl-3-furanthiol, 3-mercapto-2-pentanone, and 4-hydroxy-2,5-dimethyl-3-(2H)- furanone
356 are key features in cooked pork. Thus, the identification of marker volatile compounds in pork
357 can be meaningful for pork identification during halal authentication analysis of products. GC-
358 HS-SPME/MS and GC-MS using simultaneous distillation and extraction (SDE) are also
359 successful for identification of volatile compounds used for the identification of cooking
360 braised pork. There are 109 aroma compounds identified, in which aldehydes were the most
361 predominant in number, followed by alcohols, oxygen-containing heterocyclic compounds,
362 acids, and ketones. Methanethiol was the most abundant aroma substance in SPME, while
363 anethole was the most abundant in SDE [5657].

364
365 GC-HS-SPME/MS has been developed and validated as reliable analytical method for analysis
366 of volatile organic compounds (VOCs) of minced pork meat during storage. The origin of
367 aromatic hydrocarbons in pork was verified using migration test. Two chemometrics
368 techniques namely PCA and OPLS-DA were employed for characterizing and profiling VOCs
369 in pork meat and for identifying the marker VOCs associated with the spoilage of pork. There
370 are 41 VOCs (consisting of 10 alcohols, 7 aldehydes, 7 ketones, 6 aromatic hydrocarbons, 6
371 linear hydrocarbons, 2 terpenes, 1 acid, 1 ester, 1 furan) were identified during this study. The
372 major VOCs of minced pork are aromatic hydrocarbons, alcohols, aldehydes, linear

373 hydrocarbons, and ketones). From loading plot study, three VOCs namely ethanol, 2,3-
374 butanediol and 2-ethyl-1-hexanol were selected and considered as important variables in the
375 projection values, because these VOCs contribute to the discrimination of pork with different
376 storage times [7273].

377
378 Analysis of volatile organic compounds (VOCs) as fingerprinting profiles for identification of
379 dried pork slices from different processing stages have been done using GC coupled with ion
380 mobility spectrometry (GC-IMS). Using LAV software, 54 peaks were selected. During this
381 study, thirty seven VOCs were detected in the evaluated samples, in which aldehydes and
382 alcohols accounted for the largest proportion. 1-octene-3-ol has the flavour of cooked
383 mushroom, is important compound contributing to the VOCs of pork. This compound is
384 considered as the autoxidation product of linoleic acid [7374]. GC-MS has been employed for
385 identification of key aroma in pork broth. The multivariate calibration of PLS is used for
386 screening the relatively better flavour of pork broth among different stewing time and applied
387 for assisting the quantitative analysis of VOCs using standard internal of 1,2-dichlorobenzene.
388 From this study, the key odorants of the aroma profile of pork broth were identified namely 4-
389 hydroxy-2,5-dimethyl-3(2H)- furanone, hexanal, 1-octen-3-ol, (E)-2-octenal, (E)-2-decenal,
390 (E)-2-undecenal, (E, E)-2,4-decadienal, nonanoic acid, decanoic acid, 2-heptanone, 3-hydroxy-
391 2- butanone, δ -decanolactone, and 2-acetylpyrrole [7475].

392
393 GC-MS coupled with olfactometry (GC-MS/O) and in combination with chemometrics of PCA
394 and PLS-DA was reported to differentiate Chinese marinated pork hocks from four different
395 local brands. The results of PCA and PLS-DA indicated that both chemometrics using variable
396 of VOCs could clearly separate marinated pork hocks according to its groups. There are nine
397 odour-active compounds having the high loading capability for discrimination namely
398 heptanal, nonanal, 3-carene, D-limonene, β -phellandrene, p-cymene, eugenol, 2-ethylfuran and
399 2-pentylfuran. This study concluded that the validated GC-MS/O offered an alternative tools
400 for the differentiation of VOCs profile in different brands of marinated pork hocks [7576].

402 **6- Analysis alcoholic compounds in products using chromatographic techniques**

403
404 GC-MS is an excellent method for analysis of alcoholic compounds in foods. Park et al. have
405 validated and reported GC-MS for the simultaneous analysis of five alcohols (methanol,
406 ethanol, propanol, butanol and pentanol) in fermented Korean foods. The separation of alcohols

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407 was carried out using silica-based INNOWAX column (film thickness 0.25 mm, i.d. 250 mm,
408 length 30 m) coated with poly- ethylene glycol and applying mass selective detector set to
409 determine the specific selected ions for each alcohol. The LoD and LoQ values ranged from
410 0.25 to 1.16 mg/kg. The precision and accuracy of GC-MS are acceptable as indicated by Intra-
411 day and inter-day RSDs for individual alcohols of below 7%, with recovery values of 90.79 -
412 01.50%. The method is valid, therefore, the developed method is suitable for analysis of
413 alcohols in food samples intended in Halal food authentication supporting the certification
414 processes [7677].

415
416 Mahama et al. has applied GC with flame ionization detector (GC-FID) for analysis of alcohol
417 (ethanol) in marketed post samples (Fruit and vegetable juices from concentrate, syrups, sauce
418 samples etc.) in Thailand for identification of non-halal components suspected to be present in
419 the products. The internal standard used is n-propanol. Ethanol, internal standard and others
420 were separated using capillary columns DB-WAXTER (Agilent Technologies, 30 m by 0.32
421 mm by 1.00 µm) with temperature of FID was set at 250°C. Some certification bodies have
422 different regulation related to the maximum limits of ethanol, and the majority allowed the
423 maximum limit is 1%. The surveillance results indicated that 1 of 24 sauce samples showed an
424 ethanol concentration of 1.0%. Furthermore, an about of 4% of all the concentrated syrup
425 samples exhibited a higher percentage of ethanol than that permitted for Halal products. GC-
426 FID method using a column HP-5 (5% Phenyl 95% Methyl Siloxane) is also valid for analysis
427 of vinegar samples from Indonesia and Saudi Arabia offering reliable technique for alcohol
428 determination [5758].

429
430 Šorgić et al. developed gas chromatography coupled with the flame ionization detector and
431 headspace autosampler (HSS-GC/FID) method for analyzing volatile compounds in the wine
432 samples. The HSS-GC/FID method was developed, validated, and verified for determining
433 content of methanol, higher alcohols, and esters. The developed method was met the validation
434 requirement for linearity, range, sensitivity, accuracy, and precision parameters. Two grape
435 varieties namely Merlot and Cabernet Sauvignon were analyzed. It was found that contents of
436 the methanol were 198.0 mg/L and 150.5 mg/L, higher alcohols were 398.5 mg/L and 335.8
437 mg/L, ethyl acetate were 42.0 mg/L and 55.6 mg/L, and acetaldehyde were 23.3 mg/L and 16.1
438 mg/L for Merlot and Cabernet Sauvignon varieties, respectively. This study revealed that the
439 higher content of methanol was influenced by type of grape used for preparation as well as

440 maceration duration. Further evaluation were carried out using PCA to assess the effect of
441 genotypes variation and extraction methods on wine samples [7778].

442
443 Gas chromatography combined with PCA and cluster analysis (CA) were successfully applied
444 in determining content of alcoholic compound in Chinese beverages. According to the study,
445 twenty one aroma components were found to be important in the aroma profiles of Chinese
446 liquor. Among all the compounds, seven alcoholic compound including methanol, 2-butanol,
447 1-propanol, isobutanol, *n*-butanol, isoamylol and phenylethanol were detected by validated GC
448 analysis method. Isoamylol, isobutanol, and 1-propanol were found as the dominant alcoholic
449 compound with the content of 800.53, 637.67, and 338.84 mg/L, respectively. The
450 dimensionality reduction of PCA were employed in this study to statistically separated young
451 liquor (fresh) and aged liquors. Individual plot was generated as two dimensional visualization
452 constructed by PC1 and PC2 with total variance of 98.27%. Further separation using CA was
453 built using the Euclidean distance. All liquor samples were clustered into two big groups of
454 young liquor and aged liquors. This results proved the ability of PCA and CA to successfully
455 separate and classify the different ages Chinese liquor samples [7879].

456
457 In Indonesia, a majority Muslim country, it was stated by the government that the alcohol
458 content (in percentage) of alcohol-containing drugs, traditional medicines, and supplements
459 have to be declared on the label. Halal evaluation of alcohol content in noni (*Morinda citrifolia*
460 L.) can be performed using gas chromatography method. The GC instrumentation was set as
461 the inlet injection mode split of 2.5:1, injection temperature of 140°C, oven initial temperature
462 FID detector of 40 °C, and hold for 5 minutes. The sample of noni herbal medicines were
463 collected from herbal drugstores or online shops in Jakarta, Indonesia. Twenty samples were
464 evaluated and categorized as beverages (18 samples) and herbal medicines (2 samples). It was
465 found that thirteen out of twenty samples contained alcohol in the range of 0.04 - 1.07%.
466 Unfortunately, none of them were labelled properly according to the regulation [7980].

467
468 GC-FID has been used for analysis of ethanol in foods and beverages such as tea-based, fruit-
469 based, cheese-based, milk-based, seaweed-based, instant dried noodle, etc. Ethanol stock
470 solution was prepared (1mg/mL) and internal standard of 0.1% v/v 1-propanol was used for
471 sample preparation. Sample preparation was carried out using magnetic stirring aqueous
472 extraction. Analysis was performed out using an HP-Innowax (Agilent technologies) column

473 (30 m x 0.25 mm x 0.25 μ m). The sample injection volume was 1 μ L using split ratio of 13:1.
474 The developed method was validated according to the requirements of ISO/IEC 17025:2017.
475 Validation result showed that the method had good linearity ($R^2 > 0.999$), good accuracy
476 (recoveries of 96-105%), and good precision (RSD < 5%). The detection limit was low (0.006
477 mg/g). The determination of ethanol concentration was successfully applied in 108 samples of
478 processed foods and beverages. Therefore, this method could be used as valid method for halal
479 authentication of processed foods and beverages [5859].

480
481 GC-MS using static headspace has been applied for determination of ethanol in solid and semi-
482 solid consumer goods such as cakes, ice creams, sauces, and powders. Sample preparation was
483 carried out using mechanical homogenization and aqueous dilution of the products.
484 Subsequently, the sample was analysed using headspace GC-MS. Separation of analytes was
485 performed using a capillary column DB-624 (30 m x 0.25 mm x 1.4 μ m) and sample was
486 injected in split mode employing ratio of 1:200. Identification and quantification of ethanol
487 and ethanol-d6 was performed at scan range of 29-250 m/z with a rate of 6.1 scans/s. Result
488 showed that the developed method was specific to detect ethanol and ethanol-d6 at the retention
489 time of 2.65 and 2.61, respectively. The method demonstrated good linearity at the
490 concentration range of 0.1-2.0% v/v showed by its high R^2 value (>0.998). Additionally, good
491 accuracy as well as good precision was obtained. The accuracy was represented by recoveries
492 value (average recoveries of 99.7%). The precision was demonstrated by its lower RSD value
493 (<1.5%). From the above results, it suggested that headspace GC-MS could be used for
494 identification and quantification of ethanol in a various solid and semi solid food products for
495 halal authentication [8081].

496
497 Identification of ethanol using headspace GC-MS has also been applied in Kombucha products.
498 Kombucha is one of fermented beverages consist of sugar, tea, a symbiotic of bacteria and
499 yeast which is commonly known as non-alcoholic beverage. The United States and Canada
500 state that the content of alcoholic compounds in product must be <0.5% and <1.1% alcohol by
501 volume, respectively to be categorized as non-alcoholic drink. Propan-1-ol was used as internal
502 standard for ethanol quantification. The condition of headspace was incubation temperature at
503 70°C, syringe temperature at 70°C, incubation time of 300s, agitator speed at 500 rpm, injection
504 volume of 500 μ L, and split ratio of 10:1. Analysis was performed using an Agilent J&W DB-
505 624 UI (30 m x 0.25 mm x 1.4 μ m) applying flow rate of 1.4 mL/min (constant flow). The
506 developed method was linear ($R^2 > 0.995$) obtained at concentration range of 0.025%-2.47%.

507 The accuracy result was good demonstrated by its recovery value (102%) and good precision
508 was also obtained (RSD<4%). The LOD and LOQ values were 0.0002% and 0.002%,
509 respectively. It can be concluded that the method is suitable for identification and quantification
510 of ethanol in Kombucha product. It indicated a rapid and easy integration of analytical method
511 for halal authentication of Kombucha [8182].

512 _____
513 The development of GC-MS coupled with headspace and multidimensional (heart-cut)
514 chromatography has been successfully applied to determine ethanol content in medicinal
515 syrups. The aim was to ensure and guarantee the safety of the syrups. Samples used for analysis
516 consist of adult and paediatric syrups. Monitoring and quality control of ethanol
517 content in pharmaceutical products were important due to the efforts of industry to reduce
518 the ethanol content in the pharmaceutical and medicinal products. Sample preparation was
519 directly performed using headspace with condition as follows: heating syringe temperature of
520 90°C, incubator temperature of 100°C, incubation time 15 min -at 500 rpm, sample volume of
521 500 µL with split mode using ratio of 1:20. Two dimensional GC analysis was carried out using
522 GC-MS equipped with analytical column of RTX-5 capillary column (Crossbond® 5%
523 diphenyl/95% dimethyl polysiloxane, 30 m × 0.25 mm × 0.25 µm) for the first dimension then
524 for the second dimension used an NST 100 MS column (Carbowax polyethylene glycol, 30 m
525 × 0.25 mm × 2.00 µm). The method was validated according to National Agency of Sanitary
526 Surveillance (ANVISA) with validation parameters of selectivity, linearity, precision,
527 accuracy, LOD, LOQ, and robustness. Selectivity test found that isopropyl alcohol was an
528 interfering compound of ethanol determination in syrups. Linearity assay demonstrated linear
529 model at concentration range of 0.25% to 10.00% v/v (R²>0.999). The developed method was
530 sensitive enough as shown by its LOD value (0.03% v/v) and LOQ value (0.06% v/v). The
531 precision was measured for repeatability (CV=3.04%) and intermediate precision
532 (CV=3.03%). The recoveries value obtained ranged from 97.28%-101.38% indicating good
533 accuracy. The robustness test showed that the method remains unchanged with the small
534 changes of several parameters. This developed method could be used as rapid and easy
535 analytical technique for halal authentication of syrups by determining of the ethanol content
536 [8283].

538 **7. CONCLUSION**

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540 Chromatography-based method consist of liquid chromatography and gas chromatography
541 using various detectors has been widely applied for food ~~and pharmaceutical~~ products
542 authentication including halal analysis due to its advantages. The Ccombination of
543 chromatographic methods with chemometrics of multivariate analysis, a powerful statistical
544 analysis to manage huge data generated from analytical measurement, could be used to identify
545 potential markers to differentiate halal and non-halal samples. It will be very useful for the
546 institutions which have responsibility for halal quality assurance. Chromatogram and peak
547 separation profiles resulted as the instrument responses can be further evaluated for
548 determination as well as quantification for halal and non-halal components in food ~~and~~
549 ~~pharmaceutical~~ products. Chromatographic-based ~~method~~ methods were successfully carried
550 out to analyze products containing non-halal material such as pork and alcoholic compound.
551 Combination of chromatographic-based method and chemometrics techniques with some
552 scenarios can be applied for halal research on food ~~and pharmaceutical~~ products.

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572 **Author contribution**

574 **Laela Hayu Nurani:** Writing—original draft preparation, writing—review and editing,
575 funding acquisition; **Florentinus Dika Octa Riswanto:** Writing—original draft preparation,
576 writing—review and editing; **Anjar Windarsih:** Writing—original draft preparation,
577 writing—review and editing; **Citra Ariani Edityaningrum:** Writing—original draft
578 preparation, writing—review and editing; **Any Guntarti:** Writing—original draft preparation,
579 writing—review and editing; **Abdul Rahman:** Conceptualization, methodology, writing—
580 original draft preparation, funding acquisition.

581 All authors have read and agreed to the published version of the manuscript.

582

583 **Conflict of interest**

584 The authors declare no conflict of interest.

585

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859 **Figure and Scheme captions**

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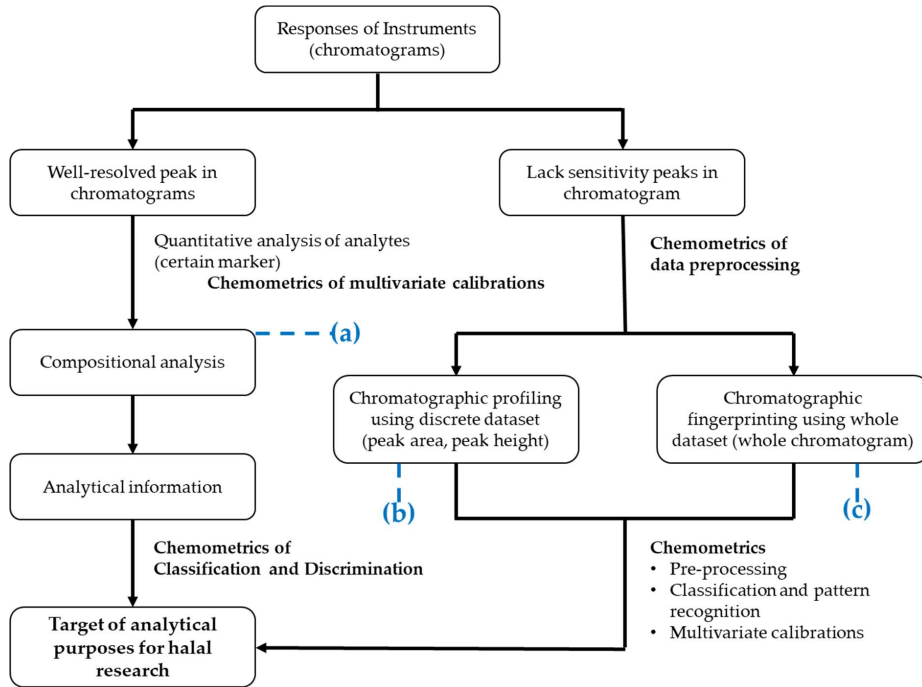
861 **Figure 1:** Three different scenarios (a, b, and c) of chemometrics applications employing the
862 chromatograms as variable for obtaining the analytical purposes (classification of halal and
863 non-halal products as well as prediction the levels of non-halal components in the products).

864 Adapted from [24].

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866 **Figures and Schemes**

867 **Figure 1**



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870 **Tables and Table captions**

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872 **Table 1.** The application of liquid chromatography (HPLC and LC-MS/MS) for analysis of

873 halal components in the food and pharmaceutical products

Methods	Issues	Results	References
HPLC-UV detection	Identification of pork in meatball products	HPLC-UV in combination with PCA could classify meatballs containing pork and beef in the products using variable of hydrolysis of Triacylglycerols (TAG). However, the authors did not mention which TAG markers contributing to this classification.	[3432]
HPLC-Fluorescence detector	Identification of pork through amino acid composition	HPLC using fluorescence detector has been successfully applied for differentiation of pork and other animal meats based on analysis of derivatized amino acids with orto-phtalaldehyde. The amino acid VAL can be identified as marker for differentiating pork from the other meats studied (beef, chicken mutton, and chevon).	[3233]
HPLC-Fluorescence detector	Detection of pig collagen using D,L-amino acids	Pre column derivatization using R(-)-4-(3-isothiocyanatopyrrolidin-1-yl)-7-(N,N-dimethylaminosulfonyl)-2,1,3-benzoxadiazole [R(-)-DBD-PyNCS] could be used to determine D,L-amino acids in pork collagen. Three amino acids of D-Asp, D-Pro, and D-Hyp were first detected in pork collagen samples.	[3334]
LC-MS/MS with multiple reaction monitoring (MRM)	Detection of Horse and Pork in Halal Beef	Biomarker peptides were successfully identified by a shotgun proteomic approach using tryptic digests of protein extracts. Pork was identified by peptide markers: TLAFLFAER (from myosin-4), SALAHAVQSSR (from myosin-1 and myosin-4). The detection limit is 0.55% horse or pork in a beef matrix.	[3435]
HPLC-MS/MS with MRM	Detection of Pork in Highly Processed Food by analysis of specific tryptic marker peptides	HPLC-MS/MS using MRM has been successfully applied for analysis of pork in some processed food products (cooking, frying and baking) based on peptide markers which are specific for pork. The peptide markers of pork identified based on MRM experiment were: marker 1 (YDIINLR) markers 2 (TLAFLFAER) and 3 (SALAHAVQSSR).	[3536]
LC-MS/MS	Differentiation of porcine gelatine and bovine gelatine	LC-MS/MS in combination with exploratory data analysis of PCA could discriminate porcine and bovine gelatines. Based on loading plot PCA, peptides appearing in retention time (t_R) 32 min could be identified as peptide markers	[3637]
Nano UPLC-Q-TOF-MS	Differentiation of porcine and bovine gelatin in food products	Marker peptide of bovine and porcine gelatin could be detected using nano UPLC-Q-TOF-MS based data dependent technique in yoghurt, cheese, and ice cream. The method could be used to detect bovine and porcine gelatin in the mixtures.	[3738]

Nano UPLC-Q-TOF-MS	Differentiation of porcine and bovine gelatin in food products	Marker peptide of bovine and porcine gelatin could be detected using nano UPLC-Q-TOF-MS based data dependent technique in yoghurt, cheese, and ice cream. The method could be used to detect bovine and porcine gelatin in the mixtures.	[3738]
LC-MS QTRAP	Gelatin speciation (bovine, porcine, and fish)	LC-MS in combination with PCA could differentiate bovine, porcine, and fish gelatin. PLS-DA could be used for classification of pure gelatin and adulterated gelatin (fish and bovine) with porcine gelatin using several concentration levels of porcine gelatin.	[3839]
LC-MS/MS	Discrimination of raw beef, pork, poultry and their mixtures	Protein of troponin I (TnI), enolase 3, L-lactate dehydrogenase (LDH), triose-phosphate isomerase (TPI), Tropomyosin 1 and carbonic anhydrase 3 could be used as potential markers to distinguish mammals and poultry.	[3940]
LC-Q-TOF-MS	Differentiation between dead-on arrival and normally slaughtered of poultry meat	LC-Q-TOF-MS could be used to differentiate between normally slaughtered and dead-on arrival poultry meat based on metabolic profiles analysed using multivariate analysis. Using METLIN and analysis of chemical standards, metabolite of sphingosine was found to be potential marker for dead-on arrival poultry meat.	[4041]
UPLC-TOF-MS	Metabolite's differentiation of broiler chicken slaughtered using different techniques	UPLC-TOF-MS could be used to distinguish between halal slaughtering method and non-halal slaughtering method of broiler chicken based on their metabolite profiles. Non-halal slaughtered method demonstrated high amino acid and high glucose breakdown.	[4142]
LC-HRMS	Analysis of pork meat in meat mixtures using PRM	Five peptides of myosin were screened and used for PRM analysis using LC-Orbitrap HRMS. Peptide of KLETDISIQGEMEDIVQEAR was found to be the most sensitive peptide with LOD value of 0.5% in meat mixtures.	[4243]
UPLC-MS	Detection of pork adulteration in beef using metabolomics approach	PLS-DA using metabolomics data obtained from untargeted measurement could classify pure and adulterated beef samples with pork. There was a significant difference in the metabolism of inositol, glutathione, and sphingolipid between beef and pork.	[4344]
LC-MS/MS	Detection of pork adulteration in meat samples using carbonic anhydrase 3 as a marker	Three peptides from carbonic anhydrase 3 were found as marker of pork (EPITVSSDQMAK, GGPLTAAYR, HDPSLLPWTASYDPGSAK). Quantification analysis could be performed using those three peptides with perfect quantitative ability and provided good correlation and recovery results.	[4445]

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880 **Table 2.** The application of gas chromatography (GC-FID and GC-MS) for analysis of halal
 881 components in the food and pharmaceutical products

Methods	Issues	Results	References
GC-FID for analysis of alcohol	Determination of ethanol contents in vinegar	The maximum contents of ethanol in vinegar is 1.0%. GC-FID could determine the levels of ethanol (alcohol) in the marketed vinegar samples. The detection level of ethanol was about 0.4 mg%.	[5758]
GC-FID for analysis of ethanol in foods	Determination of ethanol in different processed foods and beverages	Extraction technique using aqueous extraction assisted magnetic-stirring could be used to extract ethanol from different foods and beverages. GC-FID successfully used to determine ethanol with good validity. The validated method was successfully used to determine ethanol in 108 food and beverage products.	[5859]
GC-MS for analysis of alcohol	Determination of alcohol in fermented black tape ketan using GC-MS	GC-MS could be used for quantitative analysis of alcohol content in fermented black tape ketan with good recovery (89%). The alcohol concentrations determined at 3, 10, 17, 24, and 31 days were 4.295, 4.23, 5.005, 4.747, and 5.344 % v/v, respectively.	[5960]
GC-FID for analysis of lard	Differentiation of lard from other edible fats using GC-FID and chemometrics	Lard contains high amount of C18: 2 <i>cis</i> and low amount of C16:0. Chemometrics of PCA and K-mean cluster analysis could differentiate lard adulteration on chicken fat and beef tallow at low concentrations (0.5%-10%).	[6061]
GC-MS for analysis of pork	Analysis of fatty acids and methyl esters of pork (non-halal meats) in sausages compared with beef sausages (halal meat)	The dominant fatty acids in pork sausage are palmitic, myristic, oleic acid, and lauric acids. While fatty acids dominating in beef sausage are palmitic, oleic, stearic and myristic acids. The chemometrics of PCA could classify sausages according to meat sources (beef and pork)	[6162]
GC-MS for analysis of rat meat	Analysis of rat meat (non-halal meat) and its classification with other meats using chemometrics of PCA	Six fatty acids, i.e. myristic, palmitoleic, palmitic, linoleic, oleic and stearic acids combined with PCA could classify rat meat and other meats.	[6263]
Headspace GC-MS for analysis of pork	Differentiation of pork (non-halal meat) and pork sausages from beef, mutton and chicken meats	The samples were introduced into GC instrument using headspace, and volatile compounds present in the evaluated samples were separated using GC and detected by MS. The chemometrics of PCA provided good separation between pork-based sausages and halal meat-based sausages.	[6364]
GC-MS for analysis of lard	Analysis of lard (non-halal fat derived from adipose lard)	The fatty acid of 11,14-eicosadienoic acid is used as fatty acid marker for identification of lard.	[6465]

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	tissue of pig) in chocolate products		
GC-MS-SPME for analysis of wild boar	Volatilomics analysis of non-halal (wild boar) meat ball using GC-MS-SPME and chemometrics	PLS-DA could be used to differentiate volatile compounds of halal meatball and non-halal meatball. Compounds of β -cymene, 3-methylbutanal, and 2-pentanol were found to be potential markers for chicken meatball. Compounds of 5-ethyl-m-xylene, benzaldehyde, and 3-ethyl-2-methyl-1,3-hexadiene were associated to the potential markers of beef meatball. Compounds of pentanal, 2,6-dimethylcyclohexanone, 1-undecanol, cyclobutanol, 2,4,5-trimethylthiazole, and 5-ethyl-3-(3-methyl-5-phenylpyrazol-1-yl)-1,2,4-triazol-4-amine could be used as potential markers as wild boar meatball.	[6566]
HS-SPME-GC-MS for analysis of minced beef and pork meat	Volatilomics analysis using HS-SPME-GC-MS combined with multivariate analysis to differentiate minced beef and pork meat	GC-MS based on volatilomics analysis and chemometrics of PCA and PLS-DA could be used to differentiate minced beef and pork meat. Heptanal, octanal, butanol, pentanol, hexanol, 1-penten-3-ol, 2-octen-1-ol, 3-hydroxy-2-butanone were associated to the potential markers of beef whereas pentanal, hexanal, decanal, nonanal, benzaldehyde, trans-2-hexenal, trans-2-heptenal could be used as potential volatile compound markers of pork meat.	[6667]
GC-MS for analysis of pork	Detection of pork in beef meatball using GC-MS and chemometrics	PCA using fatty acid compositions of pure beef meatball and adulterated beef meatball using pork as the variables successfully differentiate pure and adulterated beef meatball. The ratio of SFA:MUFA of pork meatball was 1.0.	[6768]
GC-MS for analysis of house rat	Detection of rat house in beef meatball by analysis of fat using G-CMS	The fatty acids composition of house rats were myristate (0.19 \pm 0.03)%, palmitoleat (2.40 \pm 0.29)%, methyl palmitate (27.65 \pm 0.32)%, oleate (45.81 \pm 3.25)%, and stearate (4.65 \pm 0.28)%. Analysis using PCA could differentiate beef meatball and beef meatball containing rat house meat. Further analysis using PCA demonstrated that fatty acids of house rats have high similarity to chicken fatty acids.	[6869]
GC-MS for analysis of lard	Detection of lard in wheat biscuits using GC-MS and chemometrics	PCA using fatty acids composition could differentiate lard, wheat biscuits, and adulterated wheat biscuits with lard. PLS-DA could be used to find potential marker for differentiation. Fatty acid of C18:3n6 is suggested as potential marker to distinguish	[6970]

GC-MS for analysis of dog fat	Detection of dog fat from other animal fats using GC-MS and chemometrics	pure wheat biscuits and adulterated wheat biscuits with lard. Nine types of fatty acids in dog fat were found such as lauric, myristate, pentadecanoate, palmitoleate, palmitate, margarate, oleat, stearic, and arachidonic. Analysis PCA showed that dog fat is close to lard.	[7071]
GC-MS for analysis of rat fat	Detection of Sprague Dawley rat fat in meatball using GC-MS and chemometrics	PCA could differentiate meatball and adulterated meatball with Sprague Dawley rat meats. Further analysis revealed that the Sprague Dawley rat fat is close to beef fat.	[7472]

RESPONSE TO REVIEWER COMMENTS

Reviewer comments	Response to reviewer comments
<p>Reviewer: 1</p> <p>Comments to the Author This is a good review to see the outcome of analysis in various settings using chromatographic and chemometric analysis. Some English errors are found but minimum.</p>	<p>Thanks for this comment and appreciation. We have corrected English accordingly</p>
<p>The literature search took over 2 months period (but the statement in abstract might be understood as written within 2 months?); it would be nice to know how many articles were retrieved and how was the selection made. I noticed some long paragraph discussing in detail the outcome from a single study in the respective paragraph citing only one single reference each while there are >80 articles being listed in the reference list; some paragraph cited a single reference at the very end making the starting point of the paragraph ambiguous on whose study it was referring to.</p>	<p>We have added this information related to selection of articles used during performing this review (inclusion criteria, exclusion criteria). This information can be seen in section Methods.</p> <p>For some long paragraph citing one single reference, because we are trying to provide detail/in-depth explanation of a study that related to the criteria of our review paper. Therefore, we put the reference at the end of each paragraph.</p>
<p>Reviewer: 2</p> <p>The article needs some minor corrections, which are indicated below: 1.- The keywords on the home page and those on lines 37 and 38 do not match. I think the ones on lines 37 and 38 are more correct, but the term pharmaceutical should be deleted.</p>	<p>We have corrected this matter by matching keywords in home page in the revised manuscript.</p>
<p>2.- Lines 148-156: Include the following reference “Chromatographic fingerprinting: An innovative approach for food identification and food authentication - A tutorial. Analytica Chimica Acta 909 (2016) 9-23' which clearly explains what a marker, a profile and a chromatographic fingerprint are. Include the term marker in the paragraph.</p>	<p>Thanks for this comment, we have revised accordingly using the suggested reference as “There are three approaches to detect and to identify the presence of non-halal components in food and pharmaceutical products using chromatographic based methods. The first approach is based on searching the specific markers through analysis of the separated specific components. Indeed, the availability of reference standards is a must. The second approach is used fingerprinting profiles in which the chromatogram profiles of samples with and without non-halal components are compared and evaluated. The third approach involved metabolomics studies either targeting and untargeted techniques by analysis of all</p>

	<p>metabolites in the analyzed samples. The second and third approaches involved the large datasets, therefore, the chemometrics is employed to perform the analytical tasks (discrimination, classification, etc.)” along with reference as [29] Cuadros-Rodríguez L, Ruiz-Samblás C, Valverde-Som L, Pérez-Castaño E, González-Casado A. Chromatographic fingerprinting: An innovative approach for food “identification” and food authentication - A tutorial. Anal Chim Acta. 2016; 909; 9–23. https://doi.org/10.1016/j.aca.2015.12.042</p>
<p>3.- Lines 273-280: This paragraph should be deleted, medicinal herbs are not the subject of the study.</p>	<p>Thanks for this comment. We have removed this.</p>
<p>4.- In tables 1 and 2, there is no specific application dedicated to halal pharmaceutical. It only appears in the main discussion. I think it should be removed from the main discussion or include specific applications in halal pharmaceutical. In fact in the title the term "halal pharmaceutical" does not appear.</p>	<p>Thanks for this. We have removed the terms of pharmaceutical fields.</p>
<p>General comment: The authors have two options: To include in tables 1 and 2 specific references to halal pharmaceuticals or to remove all comments on halal pharmaceuticals.</p>	<p>We have chosen to remove the pharmaceutical applications. Thanks for this comment.</p>
<p>Editor’s Comments: Comments are marked in the attached file. Please work in the attached file so that format remains the same. Please make all changes with editing mode or different font color. In addition, you need to include a list of point by point responses against each comments from referees and editor, first include one comment and then your response. Please keep in mind that I will not correct your mistakes, but I will take decision on your efforts for a careful revision.</p>	<p>Thanks, with these comments. We have followed these instructions accordingly.</p>

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Use of chromatographic-based techniques and chemometrics for halal authentication of food products: A review

Laela Hayu Nurani, Florentinus Dika Octa Riswanto, Anjar Windarsih,
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Use of chromatographic-based techniques and chemometrics for halal authentication of food products: A review

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ABSTRACT

Halal food products are requisite to be consumed by Muslim communities in the world. The standard methods capable of quantifying non-halal components are very urgent. This review highlights the chromatographic methods and chemometric or multivariate data analysis that offer reliable techniques to provide the separation capacity in halal authentication analysis. This review article was written from reputable worldwide databases including Web of Science, Scopus, and PubMed, between January and February 2022. The keywords were “halal research,” “food analysis,” “chromatography,” “chemometrics” and “authentication.” Chromatographic-based techniques in combination with chemometrics of multivariate analysis, a powerful statistical analysis to manage huge data generated from analytical measurement, could be used to identify potential markers to differentiate halal and non-halal samples. Chromatogram and peak separation profiles resulted as the instrument responses can be further evaluated for determination as well as quantification of halal and non-halal components in food products. Combination of chromatographic-based method and chemometrics techniques with some scenarios can be applied for halal research on food products.

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INTRODUCTION

Food and pharmaceutical products are important needs for human beings. In line with the development of science and technology, industrialization and globalization, the halal products may be added or substituted and contaminated with non-halal components such as pig derivatives and alcohols as ingredients or additives to reduce the production cost.^[1] In addition, the products available in markets may contain incorrect labeling in terms of ingredient sources making the consumers lost on composition information; therefore, the use of analytical tools to check the presence of non-halal components in the products is a must for assisting the certification processes.^[2] In Indonesia, the halal certification is mandatory which means that all halal declared products sold and distributed in Indonesia must be halal certified. In addition, the analysis of non-halal components in post-marketed products is also needed to confirm that the marketed products are not adulterated with non-halal components.^[3]

According to Indonesian Act No. 33 (2014), the certification process is carried out by Halal Product Assurance Organizing Agency (BPJPH) and the auditing process is carried out by Halal Examination Agency (LPH). During audit, if the products are supposed to contain non-halal components (pork

derivatives and alcohols), the laboratory testing using standard analytical methods is needed to confirm that the audited products are free from any non-halal components.^[4,5] Today, the Muslim community constitutes for approximately of 25% world's population and is expected to increase further. With the increased awareness among Muslim community to consume the only halal products, the global market of halal products could reach exponentially.^[6] Halal is Arabic terms used to any products permissible to be consumed by Muslim community. Today, the term of halal has widely used not only Muslim but also non-Muslim because Non-Muslim community intended to export the products into Muslim community, especially in halal certification issues.^[7] Therefore, it is not surprising that halal-related studies are performed not only in majority Muslim countries like Indonesia and Malaysia but also in countries whose Muslims are minority such as the Netherlands, the United States, France and the European Union.^[8]

Halal food must be free from non-halal components which are pig and all pig derivatives such as pork, lard and porcine gelatines, carrion or dead animals, blood (flowing or congealed), animals slaughtered not according to Islamic law, animals that were killed accidentally or on purpose through means such as strangling or beating, intoxicants including alcohol and drugs,^[9] carnivorous animals, predator birds and certain land animals.^[10] Among these, pig derivatives and alcohols are typically found in halal food products; therefore, some scientists are continuously research works on halal-related issues including developing instrumental analytical methods for detecting of non-halal components intended for halal certification.^[11] Some countries have obligated the products to be halal certified which can be understood that the products are free from prohibited components. Besides, the products are manufactured using equipment dedicated for halal food.^[12] Pork is typically met in meat-based food products such as meatball, sausages, etc.; while lard can be good vehicle in some cosmetics products such as cream, lipstick and lotion. Porcine gelatines are common materials used in food (in candies) and pharmaceutical products (capsule shells).^[13] The objective of this review was to provide the integrative information on identification and quantification of non-halal components in food products by chromatographic methods. In addition, chemometrics techniques were reported to be applied to employ the big data evaluation as resulted from the chromatographic detection.

METHODS

This review article was written by identifying, investigating and assembling several review articles, original articles, books and relevant sources on metabolite fingerprinting from reputable worldwide databases including Web of Science, Scopus and PubMed. Literature searching was carried out between January and February 2022. The keywords explored during literature investigation were "halal research," "food analysis," "chromatography," "chemometrics" and "authentication." First, to select the suitable papers, 250 articles were reviewed through the title and abstract. The inclusion criteria to select the papers were (1) studies regarding halal authentication of food products using chromatographic technique between 2005–2022; (2) studies on analysis of non-halal components in food products using liquid chromatography and gas chromatography conducted between 2005–2022; (3) studies on the employment of chemometrics in combination with chromatographic technique for halal authentication of food products; and (4) all papers written in English. The exclusion criteria of the papers were (1) studies on halal authentication of food products using chromatographic techniques published before 2005 and (2) all articles written using language other than English.

Chromatographic-based techniques and chemometrics for analysis of non-halal components

For many years, chromatography has been known as the method of choice to assess the purity and levels of analytes in the laboratories of research, industry and quality control.^[14] Gas chromatography (GC) and liquid chromatography (LC) techniques are often used for the analysis of non-halal components in food products. In terms of compound types, GC is more suitable for the analysis of smaller, volatile and stable compounds to heat, while LC is more robust and suitable for larger and

less/nonvolatile compounds.^[15] Some derivatization techniques are needed in LC in order to convert analytes into detectable derivatives with certain detectors, while derivatization in GC for fewer volatile compounds is intended to provide more volatile and stable derivative products, although this derivatization process increases the method complexity and lengthens the sample preparation. In addition, the availability of derivative agents and its steric hindrance in the analyte, and the stability of the derivatized compounds must also be considered.^[16]

One-dimensional gas or liquid chromatography using one column is considered as simple and powerful separation techniques for simple and un-complex samples. When the analyzed samples are complex enough, the application of just one-dimension chromatography leads to peak co-elution as well as overlapping and non-resolved peaks; therefore, one dimension chromatography technique is not suitable for separation of large analytes because the peak capacity of one-dimensional analysis is not large enough to achieve the complete separation with acceptable resolution.^[17] In last decades, two-dimensional gas chromatography (GC x GC) and liquid chromatography (LC x LC) has been applied in analysis of complex mixture in order to increase the separation speed.^[18]

In two-dimensional chromatography, the separation is carried out in two columns with different polarity connected in series by a modulator; as a consequence, the separation capacity of regular one-column in one dimensional chromatography can be considerably increased. The effluent from the first column is transferred to the second column using modulator so that the analytical information obtained (such as retention times, t_R) from the first column can be combined with that from second column, leading to a plot of two retention times.^[19] Because of the excellent separation capacity of GC x GC and LC x LC combined with mass spectrometry (MS), both techniques are applied for separation all components in the complex mixtures, especially for metabolomics studies.^[18] GC x GC has been widely applied for analysis of metabolites (all fatty acid types) of lard in food samples,^[20] while LC x LC is typically used for analysis of peptides,^[21] which can be used for identification of pork and porcine gelatines.

Chromatographic-based techniques offered reliable technique in halal authentication analysis. However, due to high number of data covered, the application of chemometrics to treat big data is unavoidable. Chemometrics can be defined as the employment of statistical and mathematical methods to obtain the objective data evaluation by extracting the relevant and meaningful information from related and unrelated responses from chemical measurements. Chemometrics or multivariate data analysis (MDA) is typically applied in numerous aspects including the quality control of halal products, qualitative and quantitative determination of chemical parameters for assessing the products authenticity.^[22]

Chemometrics can provide the powerful tools in giving important information extracted from big data obtained from instrumental analyses such as methods based on spectroscopic and chromatographic. The common chemometrics techniques applied in product authentication could be grouped into exploratory data analysis, data pre-processing, description and visualization, discrimination and pattern recognition (classification), regression and prediction and experimental design.^[23] Some chromatographic problems encountered during halal authentication analysis included the assessment of separation quality, the evaluation of peak alignment using pre-processing, the optimization of chromatographic systems providing the good separation of all peaks using experimental design, the accuracy of discrimination and classification using pattern recognition and quantitative analysis applying multivariate calibration. **Figure 1** showed the correlation between chromatographic responses and chemometrics for certain analytical purposes. In scenario (a), peaks with good separation (good selectivity) in chromatogram was used as variable for the evaluation of compositional analysis (concentration) of analytes assisted by multivariate calibrations. In (b), certain peaks with lack selectivity was used as variable during chromatographic profiling of objects (samples) using discrete datasets (peak area or peak height), while in scenario (c), whole datasets in chromatograms were used as variables during chromatographic fingerprinting of objects. Indeed, the chemometrics of pre-processing was widely applied to obtain the desired analytical modeling.

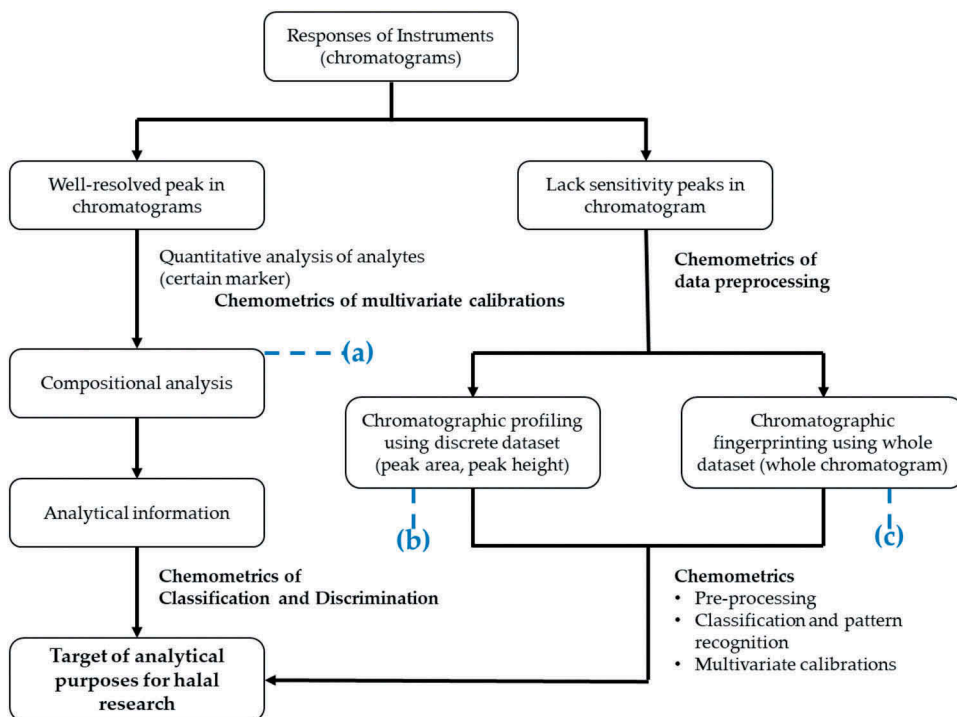


Figure 1. Three different scenarios (a, b, and c) of chemometrics applications employing the chromatograms as variable for obtaining the analytical purposes (classification of halal and non-halal products as well as prediction the levels of non-halal components in the products). Adapted from Ref.^[24].

The classification chemometrics was typically carried using (1) exploratory data analysis including principal component analysis (PCA) and cluster analysis (hierarchical cluster analysis and nonhierarchical such as k-means and k-medians), and this technique is typically called as unsupervised pattern recognition and (2) classification and discrimination methods known supervised pattern recognition. There are two types of classification chemometrics methods regardless of the statistical background. The first type is typically employed to assess to which of various predefined classes of samples (objects). The example of this technique is linear discriminant analysis (LDA), orthogonal projection to latent structures – discriminant analysis (OPLS-DA), k-nearest neighbors (KNN) and many others. The second type of classification chemometrics is called as class modeling or one class classifier (OCC) and the example for this group data driven soft independent modeling of class analogy (DD-SIMCA) and Unequal Class-Modeling (UNEQ).^[25] Using these chemometrics, someone can answer the question: is the meat belong to pork (non-halal) or beef (halal)? or the question: is the meatball authentic or adulterated?^[26,27]

Analysis of non halal components using liquid chromatography

High performance liquid chromatography (HPLC) using certain detectors have been widely applied for analysis of specific components in non-halal components. HPLC using fluorescence detector has been successfully used for analysis of hydroxyproline and other amino acids in gelatin and collagen samples as initial screening for identification of gelatin types. Hydroxyproline has been known as signature amino acid for gelatin and collagen. The level of hydroxyproline is typically higher in the gelatin samples than that in the collagen samples, except for the samples of fish skin gelatin, and this result could be used as screening tools for identification of non-halal gelatin and collagen in the analyzed samples.^[28]

There are three approaches to detect and to identify the presence of non-halal components in food products using chromatographic based methods. The first approach is based on searching the specific markers through analysis of the separated specific components. Indeed, the availability of reference standards is a must. The second approach is used fingerprinting profiles in which the chromatogram profiles of samples with and without non-halal components are compared and evaluated. The third approach involved metabolomics studies either targeting and untargeted techniques by analysis of all metabolites in the analyzed samples. The second and third approaches involved the large datasets; therefore, the chemometrics is employed to perform the analytical tasks (discrimination, classification, etc.) [29]

Table 1 listed the application of HPLC and LC-MS/MS for analysis of halal components in the products. Liquid chromatography using fluorescence detector was also successfully applied for analysis of amino acid (AA) composition non-halal (porcine) and halal (bovine and fish) gelatins. The classification between halal and non-halal gelatins was carried using PCA applying amino acid compositions as variable. AAs with strong fluorescence (Hyp, His, Ser, Arg, Gly, Thr, Pro, Tyr, Met, Val, Leu and Phe) contribute to the classification and become the biomarkers to identify the gelatine sources.^[44] Gelatin from three mammalian species including bovine gelatin, porcine gelatin, and donkey gelatin has been successfully identified using liquid chromatography-linear ion-trap high resolution mass spectrometry. Hemoglobin was just found in donkey gelatin. The unique peptide obtained from donkey, bovine and porcine gelatin was GEAGPAGPAGPIGPVGAR, GETGPAGPAGPIGPVGAR and GETGPAGPAGPVGAR, respectively. The unique peptides could be detected either in individual gelatin or in the mixtures of three mammalian gelatins.^[45]

Liquid chromatography especially combined with mass spectrometer tandem mass spectrometer (LC/MS-MS) is widely applied for identification of non-halal component in food products including porcine gelatin and pork. Gel-enhanced liquid chromatography-mass spectrometry (GeLCMS) in combination with chemometrics of PCA has been developed for identification of potential protein markers in pork and other meats along with its classification. The myofibrillar protein with weight of 40-kDa such as troponin T, Tropomyosin alpha-1 chain and actin cytoplasmic 1 as well as the thin filament proteins such as actin, troponin and Tropomyosin had molecular weights ranging from 40 to 45 kDa could be used as markers for differentiation of pork from chicken and beef. PCA using PC1 and PC2 accounting of 62% and 35% variances could classify meat types. From MS studies, the potential protein markers for pork meat samples are Troponin T with peptide sequences of [(R)KPLNIDHLSSEDK(L)], Tropomyosin alpha-1 chain [(K)EAETRAEFAER(S)], Actin cytoplasmic 1 [(R)HQGVVMGMGQK(D)], COP9 signalosome complex subunit 4 [(R)VLDYRR(K)] and Ribonuclease inhibitor [(R)VLGQGLADSACQLETLR(L)].^[46]

The identification of potential biomarkers of gelatin from several sources could be performed using UPLC-MS/MS. Samples used were gelatin from pig, cow, chicken and fish. After the extraction process of proteins from gelatin, proteins were then digested using proteomic grade trypsin for 12 h to obtain peptides. Chemometrics of PCA was used to differentiate partial hydrolysis of gelatin from cow and pig. Result from PCA score plot showed that the sample of cow and pig obtained from digestion process could be well separated. For identification of potential biomarkers from pig, cow, fish and chicken gelatin, PCA employing MPP (Mass Profiler Professional) was applied. Results showed that three unique peptides found only in pig gelatin, seven unique peptides found in bovine/cow gelatin, 22 peptides found only in chicken gelatin and only 1 unique peptide found in fish gelatin. The developed method was also successfully applied to identify species origin of commercial gelatin samples. It indicated that UPLC-MS/MS offers a powerful analytical technique to identify gelatin from different species in food products.^[47]

Targeted tandem liquid chromatography-mass spectrometry (LC-MS) using decoy, randomized and concatenated database search program comprising MS-Fit and MS-Tag in combination with chemometrics of principal component analysis and orthogonal partial least square-discriminant analysis (OPLS-DA) was applied for identification of potential peptide markers in non-halal meat (pork) and halal meats (chicken and beef). The peptide markers which are specific to certain species

Table 1. The application of liquid chromatography (HPLC and LC-MS/MS) for analysis of halal components in the food and pharmaceutical products.

Methods	Issues	Results	References
HPLC-UV detection	Identification of pork in meatball products	HPLC-UV in combination with PCA could classify meatballs containing pork and beef in the products using variable of hydrolysis of Triacylglycerols (TAGs). However, the authors did not mention which TAG markers contribute to this classification	[30]
HPLC-Fluorescence detector	Identification of pork through amino acid composition	HPLC using fluorescence detector has been successfully applied for differentiation of pork and other animal meats based on analysis of derivatized amino acids with ortho-phthalaldehyde. The amino acid VAL can be identified as marker for differentiating pork from the other meats studied (beef, chicken mutton, and chevon)	[31]
HPLC-Fluorescence detector	Detection of pig collagen using D, L-amino acids	Pre column derivatization using R (-)-4-(3-isothiocyanato pyrrolidin-1-yl)-7-(N, N-dimethylamino sulfonyl)-2,1,3-benzoxadiazole [R (-)-DBD-PyNCS] could be used to determine D, L-amino acids in pork collagen. Three amino acids of D-Asp, D-Pro, and D-Hyp were first detected in pork collagen samples	[32]
LC-MS/MS with multiple reaction monitoring (MRM)	Detection of Horse and Pork in Halal Beef	Biomarker peptides were successfully identified by a shotgun proteomic approach using tryptic digests of protein extracts. Pork was identified by peptide markers: TLAFLFAER (from myosin-4) and SALAHAVQSSR (from myosin-1 and myosin-4). The detection limit is 0.55% horse or pork in a beef matrix	[33]
HPLC-MS/MS with MRM	Detection of pork in highly processed food by analysis of specific tryptic marker peptides	HPLC-MS/MS using MRM has been successfully applied for analysis of pork in some processed food products (cooking, frying and baking) based on peptide markers which are specific for pork. The peptide markers of pork identified based on MRM experiment were: marker 1 (YDIINLR) markers 2 (TLAFLFAER) and 3 (SALAHAVQSSR)	[34]
LC-MS/MS	Differentiation of porcine gelatine and bovine gelatine	LC-MS/MS in combination with exploratory data analysis of PCA could discriminate porcine and bovine gelatines. Based on loading plot PCA, peptides appearing in retention time (t_R) 32 min could be identified as peptide markers	[35]
Nano UPLC-Q-TOF-MS	Differentiation of porcine and bovine gelatin in food products	Marker peptide of bovine and porcine gelatin could be detected using nano UPLC-Q-TOF-MS based data dependent technique in yogurt, cheese, and ice cream. The method could be used to detect bovine and porcine gelatin in the mixtures	[36]
Nano UPLC-Q-TOF-MS	Differentiation of porcine and bovine gelatin in food products	Marker peptide of bovine and porcine gelatin could be detected using nano UPLC-Q-TOF-MS based data dependent technique in yogurt, cheese, and ice cream. The method could be used to detect bovine and porcine gelatin in the mixtures	[36]
LC-MS QTRAP	Gelatin speciation (bovine, porcine, and fish)	LC-MS in combination with PCA could differentiate bovine, porcine, and fish gelatin. PLS-DA could be used for classification of pure gelatin and adulterated gelatin (fish and bovine) with porcine gelatin using several concentration levels of porcine gelatin	[37]
LC-MS/MS	Discrimination of raw beef, pork, poultry and their mixtures	Protein of troponin I (TnI), enolase 3, L-lactate dehydrogenase (LDH), triose-phosphate isomerase (TPI), Tropomyosin 1, and carbonic anhydrase 3 could be used as potential markers to distinguish mammals and poultry	[38]

(Continued)

Table 1. (Continued).

Methods	Issues	Results	References
LC-Q-TOF-MS	Differentiation between dead-on arrival and normally slaughtered of poultry meat	LC-Q-TOF-MS could be used to differentiate between normally slaughtered and dead-on arrival poultry meat based on metabolic profiles analyzed using multivariate analysis. Using METLIN and analysis of chemical standards, metabolite of sphingosine was found to be potential marker for dead-on arrival poultry meat	[39]
UPLC-TOF-MS	Metabolite's differentiation of broiler chicken slaughtered using different techniques	UPLC-TOF-MS could be used to distinguish between halal slaughtering method and non-halal slaughtering method of broiler chicken based on their metabolite profiles. Non-halal slaughtered method demonstrated high amino acid and high glucose breakdown	[40]
LC-HRMS	Analysis of pork meat in meat mixtures using PRM	Five peptides of myosin were screened and used for PRM analysis using LC-Orbitrap HRMS. Peptide of KLETDISIQGEMEDIVQEAR was found to be the most sensitive peptide with LOD value of 0.5% in meat mixtures	[41]
UPLC-MS	Detection of pork adulteration in beef using metabolomics approach	PLS-DA using metabolomics data obtained from untargeted measurement could classify pure and adulterated beef samples with pork. There was a significant difference in the metabolism of inositol, glutathione, and sphingolipid between beef and pork	[42]
LC-MS/MS	Detection of pork adulteration in meat samples using carbonic anhydrase 3 as a marker	Three peptides from carbonic anhydrase 3 were found as marker of pork (EPITVSSDQMAK, GGPLTAAYR, HDPSLLPWTASYDPGSAK). Quantification analysis could be performed using those three peptides with perfect quantitative ability and provided good correlation and recovery results	[43]

were identified through shot-gun proteomics. Potential peptide marker identified for raw pork is myosin-2 having sequence of peptide marker of (F)DFNSLE(Q). OPLS-DA using variable of identified peptides could separate halal and non-halal meats.^[48] 215

Targeted proteomic analysis using LC-MS has been developed to investigate the heat stable protein in pork meat. Five heat treatments were applied such as (1) water bath heating at 78°C for 30 min; (2) boiling at 100°C for 30 min; (3) sterilizing at 121°C for 30 min; (4) frying using oil until golden brown color; and (5) baking at 200°C for 30 min. Protein extraction from samples was performed using buffer solution containing 2 M thiourea, 7 M urea and 50 mM Tris-HCl (pH 8.0). Proteins were digested using proteomic grade trypsin added with DTT to reduce disulfide bonds and IAA for alkylation. Incubation was carried out for at least 12 h at 37°C. Result showed that seven heat-stable specific peptides of pork were found such as DQLIHNLK from l-lactate dehydrogenase A chain, HDPSLLPWTASYDPGSAK from carbonic anhydrase 3, EPITVSSDQMAK from carbonic anhydrase 3, VNVDEVGGEALGR from hemoglobin subunit beta, HPGDFGADAQGAMSK from myoglobin, SLYSSAENEPVPLVR from carbonic anhydrase 3 and YLEFISEAIQVLQSK from myoglobin. Commercial samples such as Iberian dried ham, Pasteur dry sausage, import dried ham, lunch meat canned, sandwich sausage and Thuringia flavor sausage were used to identify the presence one or more pig heat-stable peptides. Results showed that the heat-stable peptides of pig could be found in various types of food products with different cooking process methods. It suggested that targeted proteomics analysis using seven heat stable peptides of pig could be used for halal authentication of food products especially meat-based food products containing pork.^[49] 220 225 230

Analysis using LC-MS employing MRM (multiple reaction monitoring) technique was successfully used to detect heat-stable peptides in cooked meats including pork meat. Thermal treatment applied was boiling at 100°C, grilling at 150°C and grilling at 180°C. After the protein was extracted, digestion process was performed using proteomic grade trypsin. Identification of homologous protein and 235

potential biomarkers of pork peptide was carried out using UPLC Triple TOF-MS equipped with a C-18 column (2.1 × 100 mm, 1.7 μm; Waters Corporation, Taunton, MA, USA and Wexford, Ireland). The mobile phase used was water containing 0.1% formic acid (A) and acetonitrile containing 0.1% formic acid (B) with flow rate of 0.3 mL/min. On the other hand, MRM analysis was performed using a SCIEX ExionLC AD system (AB SCIEX, Framingham, MA, USA) and an AB SCIEX QTRAP 4500 mass spectrometry system (AB SCIEX PTE. LTD., Marsiling, Singapore) equipped with a column of Waters ACQUITY UPLC BEH C18 (2.1 × 50 mm, 1.7 μm). Results showed that the potential peptide biomarkers in raw pork meat found were GHHEAELTPLAQSHATK from myoglobin, FAGGNLDVLK; ADMVIEAVFEELSILK; TVLGAPEVLLGILPGAGGTQR from trifunctional enzyme subunit alpha, mitochondrial and WGDAGATYVVESTGVFTTMEK from glyceraldehyde-3-phosphate dehydrogenase. Meanwhile, the heat-stable peptide biomarkers of pork were FAGGNLDVLK and TVLGAPEVLLGILPGAGGTQR from trifunctional enzyme subunit alpha, mitochondrial as well as WGDAGATYVVESTGVFTTMEK from glyceraldehyde-3-phosphate dehydrogenase. The MRM analysis confirmed the heat-stable peptide of pork in meat product samples. It suggested that LC-MS employing MRM method could be used as promising analytical technique for halal authentication of meat products.^[50]

Application of gas chromatography for analysis of non-halal components

Table 2 listed the application of gas chromatography for analysis of halal components in the food products. GC-MS combined with chemometrics has been proposed as tools for detection of lard as adulterant in olive oil using metabolomic approach. GC separation of fatty acid methyl esters (FAME) was achieved using HP-5 MS nonpolar capillary column. The identification of metabolites of FAMES was carried out using standard FAMES and mass spectrometer detector using the WILEY 2007 library. Some FAMES are specific, i.e., methyl behenate was only present in olive oil and methyl myristate was only detected in lard. PCA using identified FAMES was successful for separating lard, olive oil and olive oil adulterated with lard for halal authentication study.^[66]

Two-dimensional GC combined with time-of-flight mass spectrometer (GC x GC-TOF/MS) is successfully used for analysis of lard as adulterant in virgin coconut oil (VCO) through analysis of sterols. GC x GC system could perform the complete baseline separation of sterol trimethylsilyl ethers derived from cholesterol and cholestanol, which facilitate the detection of lard in VCO. Using GC x GC-TOF/MS, cholestanol trimethylsilyl ether (Cha-TME) and cholesterol trimethylsilyl ether (Che-TME) were separated into some peaks, identified as CHE₁, CHE_{bI}, CHE_{bII}, CHE₂ (Che-TME), and CHa₁, CHa_{bI}, CHa_{bII} and CHa₂ for Cha-TME. Quantification of these compounds could be used as tools for quantification of adulteration levels of lard in VCO.^[20]

GC-MS coupled with headspace solid-phase microextraction (HS-SPME) is successful for the analysis of volatile compounds in pork. The profiles of volatile compounds from different meats are different; therefore, the volatile compounds analyzed by GC-HS-SPME/MS could be used as fingerprinting tools for specific meats.^[67] In addition, VOCs also contribute to the aroma which can be used for the discrimination tools among animal meats.^[68] Analysis of VOCs is very challenging because of different factors, including the high number of volatile compounds, differences in volatility degree and the great amount of functional groups.^[69] Chen et al.^[70] have identified the key volatile compounds for differentiation of pork from different pig breeding. The volatile compounds contributing to the pork flavor identified during this study were 3-methyl-1-butanol, 1-nonanal, octanal, hexanal, 2-pentyl-furan, 1-penten-3-one, N-morpholinomethyl-isopropyl-sulfide, methyl butyrate, and (E,E)-2, 4-decadienal. Kosowska et al.^[71] reported that some volatile compounds namely octanal, nonanal, (E,E)-2,4-decadienal, methanethiol, methional, 2-furfurylthiol, 2-methyl-3-furanthiol, 3-mercapto-2-pentanone and 4-hydroxy-2,5-dimethyl-3-(2 H)-furanone are key features in cooked pork. Thus, the identification of marker volatile compounds in pork can be meaningful for pork identification during halal authentication analysis of products. GC-HS-SPME/MS and GC-MS using simultaneous distillation and extraction (SDE) are

Table 2. The application of gas chromatography (GC-FID and GC-MS) for analysis of halal components in the food and pharmaceutical products.

Methods	Issues	Results	References
GC-FID for analysis of alcohol	Determination of ethanol contents in vinegar	The maximum contents of ethanol in vinegar is 1.0%. GC-FID could determine the levels of ethanol (alcohol) in the marketed vinegar samples. The detection level of ethanol was about 0.4 mg%	[51]
GC-FID for analysis of ethanol in foods	Determination of ethanol in different processed foods and beverages	Extraction technique using aqueous extraction assisted magnetic-stirring could be used to extract ethanol from different foods and beverages. GC-FID successfully used to determine ethanol with good validity. The validated method was successfully used to determine ethanol in 108 food and beverage products	[52]
GC-MS for analysis of alcohol	Determination of alcohol in fermented black tape ketan using GC-MS	GC-MS could be used for quantitative analysis of alcohol content in fermented black tape ketan with good recovery (89%). The alcohol concentrations determined at 3, 10, 17, 24, and 31 days were 4.295, 4.23, 5.005, 4.747, and 5.344% v/v, respectively	[53]
GC-FID for analysis of lard	Differentiation of lard from other edible fats using GC-FID and chemometrics	Lard contains high amount of C18:2 <i>cis</i> and low amount of C16:0. Chemometrics of PCA and K-mean cluster analysis could differentiate lard adulteration on chicken fat and beef tallow at low concentrations (0.5%-10%)	[54]
GC-MS for analysis of pork	Analysis of fatty acids a fatty acid methyl esters of pork (non-halal meats) in sausages compared with beef sausages (halal meat)	The dominant fatty acids in pork sausage are palmitic, myristic, oleic acid, and lauric acids. While fatty acids dominating in beef sausage are palmitic, oleic, stearic and myristic acids. The chemometrics of PCA could classify sausages according to meat sources (beef and pork)	[55]
GC-MS for analysis of rat meat	Analysis of rat meat (non-halal meat) and its classification with other meats using chemometrics of PCA	Six fatty acids, i.e. myristic, palmitoleic, palmitic, linoleic, oleic and stearic acids combined with PCA could classify rat meat and other meats	[56]
Headspace GC-MS for analysis of pork	Differentiation of pork (non-halal meat) and pork sausages from beef, mutton and chicken meats	The samples were introduced into GC instrument using headspace, and volatile compounds present in the evaluated samples were separated using GC and detected by MS. The chemometrics of PCA provided good separation between pork-based sausages and halal meat-based sausages	[57]
GC-MS for analysis of lard	Analysis of lard (non-halal fat derived from adipose tissue of pig) in chocolate products	The fatty acid of 11,14-eicosadienoic acid is used as fatty acid marker for identification of lard	[58]
GC-MS-SPME for analysis of wild boar	Volatilomics analysis of non-halal (wild boar) meat ball using GC-MS-SPME and chemometrics	PLS-DA could be used to differentiate volatile compounds of halal meatball and non-halal meatball. Compounds of β -cymene, 3-methyl-butanal, and 2-pentanol were found to be potential markers for chicken meatball. Compounds of 5-ethyl-m-xylene, benzaldehyde, and 3-ethyl-2-methyl-1,3-hexadiene were associated to the potential markers of beef meatball. Compounds of pentanal, 2,6-dimethylcyclohexanone, 1-undecanol, cyclobutanol, 2,4,5-trimethyl-thiazole, and 5-ethyl-3-(3-methyl-5-phenyl pyrazol-1-yl)-1,2,4-triazol-4-amine could be used as potential markers as wild boar meatball	[59]

(Continued)

Table 2. (Continued).

Methods	Issues	Results	References
HS-SPME-GC-MS for analysis of minced beef and pork meat	Volatilomics analysis using HS-SPME-GC-MS combined with multivariate analysis to differentiate minced beef and pork meat	GC-MS based on volatilomics analysis and chemometrics of PCA and PLS-DA could be used to differentiate minced beef and pork meat. Heptanal, octanal, butanol, pentanol, hexanol, 1-penten-3-ol, 2-octen-1-ol, 3-hydroxy-2-butanone were associated to the potential markers of beef whereas pentanal, hexanal, decanal, nonanal, benzaldehyde, trans-2-hexenal, trans-2-heptenal could be used as potential volatile compound markers of pork meat	[60]
GC-MS for analysis of pork	Detection of pork in beef meatball using GC-MS and chemometrics	PCA using fatty acid compositions of pure beef meatball and adulterated beef meatball using pork as the variables successfully differentiate pure and adulterated beef meatball. The ratio of SFA: MUFA of pork meatball was 1.0	[61]
GC-MS for analysis of house rat	Detection of rat house in beef meatball by analysis of fat using G-CMS	The fatty acids composition of house rats were myristate ($0.19 \pm 0.03\%$), palmitoleate ($2.40 \pm 0.29\%$), methyl palmitate ($27.65 \pm 0.32\%$), oleate ($45.81 \pm 3.25\%$), and stearate ($4.65 \pm 0.28\%$). Analysis using PCA could differentiate beef meatball and beef meatball containing rat house meat. Further analysis using PCA demonstrated that fatty acids of house rats have high similarity to chicken fatty acids	[62]
GC-MS for analysis of lard	Detection of lard in wheat biscuits using GC-MS and chemometrics	PCA using fatty acids composition could differentiate lard, wheat biscuits, and adulterated wheat biscuits with lard. PLS-DA could be used to find potential marker for differentiation. Fatty acid of C18:3n6 is suggested as potential marker to distinguish pure wheat biscuits and adulterated wheat biscuits with lard	[63]
GC-MS for analysis of dog fat	Detection of dog fat from other animal fats using GC-MS and chemometrics	Nine types of fatty acids in dog fat were found such as lauric, myristate, pentadecanoate, palmitoleate, palmitate, margarate, oleat, stearic, and arachidonic. Analysis PCA showed that dog fat is close to lard	[64]
GC-MS for analysis of rat fat	Detection of Sprague Dawley rat fat in meatball using GC-MS and chemometrics	PCA could differentiate meatball and adulterated meatball with Sprague Dawley rat meats. Further analysis revealed that the Sprague Dawley rat fat is close to beef fat	[65]

also successful for identification of volatile compounds used for the identification of cooking braised pork. There are 109 aroma compounds identified, in which aldehydes were the most predominant in number, followed by alcohols, oxygen-containing heterocyclic compounds, acids and ketones. Methanethiol was the most abundant aroma substance in SPME, while anethole was the most abundant in SDE.^[72] 290

GC-HS-SPME/MS has been developed and validated as reliable analytical method for analysis of volatile organic compounds (VOCs) of minced pork meat during storage. The origin of aromatic hydrocarbons in pork was verified using migration test. Two chemometrics techniques, namely, PCA and OPLS-DA were employed for characterizing and profiling VOCs in pork meat and for identifying the marker VOCs associated with the spoilage of pork. There are 41 VOCs (consisting of 10 alcohols, 7 aldehydes, 7 ketones, 6 aromatic hydrocarbons, 6 linear hydrocarbons, 2 terpenes, 1 acid, 1 ester, 1 furan) were identified during this study. The major VOCs of minced pork are aromatic hydrocarbons, alcohols, 295

aldehydes, linear hydrocarbons, and ketones). From loading plot study, three VOCs namely ethanol, 2,3-butanediol and 2-ethyl-1-hexanol were selected and considered as important variables in the projection values, because these VOCs contribute to the discrimination of pork with different storage times.^[73] 300

Analysis of volatile organic compounds (VOCs) as fingerprinting profiles for identification of dried pork slices from different processing stages have been done using GC coupled with ion mobility spectrometry (GC-IMS). Using LAV software, 54 peaks were selected. During this study, thirty seven VOCs were detected in the evaluated samples, in which aldehydes and alcohols accounted for the largest proportion. 1-octene-3-ol has the flavor of cooked mushroom, is important compound contributing to the VOCs of pork. This compound is considered as the autoxidation product of linoleic acid.^[74] GC-MS has been employed for identification of key aroma in pork broth. The multivariate calibration of PLS is used for screening the relatively better flavor of pork broth among different stewing time and applied for assisting the quantitative analysis of VOCs using standard internal of 1,2-dichlorobenzene. From this study, the key odorants of the aroma profile of pork broth were identified namely 4-hydroxy-2,5-dimethyl-3(2 H)- furanone, hexanal, 1-octen-3-ol, (E)-2-octenal, (E)-2-decenal, (E)-2-undecanal, (E, E)-2,4-decadienal, nonanoic acid, decanoic acid, 2-heptanone, 3-hydroxy-2- butanone, δ -decanolactone and 2-acetylpyrrole.^[75] 305 310

GC-MS coupled with olfactometry (GC-MS/O) and in combination with chemometrics of PCA and PLS-DA was reported to differentiate Chinese marinated pork hocks from four different local brands. The results of PCA and PLS-DA indicated that both chemometrics using variable of VOCs could clearly separate marinated pork hocks according to its groups. There are nine odor-active compounds having the high loading capability for discrimination, namely, heptanal, nonanal, 3-carene, D-limonene, β -phellandrene, p-cymene, eugenol, 2-ethylfuran and 2-pentylfuran. This study concluded that the validated GC-MS/O offered an alternative tools for the differentiation of VOCs profile in different brands of marinated pork hocks.^[76] 315 320

Analysis alcoholic compounds in products using chromatographic techniques

GC-MS is an excellent method for analysis of alcoholic compounds in foods. Park et al. have validated and reported GC-MS for the simultaneous analysis of five alcohols (methanol, ethanol, propanol, butanol and pentanol) in fermented Korean foods. The separation of alcohols was carried out using silica-based INNOWAX column (film thickness 0.25 μm , i.d. 250 μm , length 30 m) coated with poly-ethylene glycol and applying mass selective detector set to determine the specific selected ions for each alcohol. The LoD and LoQ values ranged from 0.25 to 1.16 mg/kg. The precision and accuracy of GC-MS are acceptable as indicated by intra-day and inter-day RSDs for individual alcohols of below 7%, with recovery values of 90.79–101.50%. The method is valid; therefore, the developed method is suitable for analysis of alcohols in food samples intended in Halal food authentication supporting the certification processes.^[77] 325 330

Mahama et al. have applied GC with flame ionization detector (GC-FID) for analysis of alcohol (ethanol) in marketed post samples (Fruit and vegetable juices from concentrate, syrups, sauce samples, etc.) in Thailand for identification of non-halal components suspected to be present in the products. The internal standard used is n-propanol. Ethanol, internal standard and others were separated using capillary columns DB-WAXTER (Agilent Technologies, 30 m by 0.32 mm by 1.00 μm) with temperature of FID was set at 250°C. Some certification bodies have different regulation related to the maximum limits of ethanol, and the majority allowed the maximum limit is 1%. The surveillance results indicated that 1 of 24 sauce samples showed an ethanol concentration of 1.0%. Furthermore, an about of 4% of all the concentrated syrup samples exhibited a higher percentage of ethanol than that permitted for Halal products. GC-FID method using a column HP-5 (5% Phenyl 95% Methyl Siloxane) is also valid for analysis of vinegar samples from Indonesia and Saudi Arabia offering reliable technique for alcohol determination.^[51] 335 340

Šorgić et al. developed gas chromatography coupled with the flame ionization detector and head-space autosampler (HSS-GC/FID) method for analyzing volatile compounds in the wine samples. The HSS-GC/FID method was developed, validated and verified for determining content of methanol, 345

higher alcohols and esters. The developed method was met the validation requirement for linearity, range, sensitivity, accuracy and precision parameters. Two grape varieties namely Merlot and Cabernet Sauvignon were analyzed. It was found that contents of the methanol were 198.0 mg/L and 150.5 mg/L, higher alcohols were 398.5 mg/L and 335.8 mg/L, ethyl acetate were 42.0 mg/L and 55.6 mg/L, and acetaldehyde were 23.3 mg/L and 16.1 mg/L for Merlot and Cabernet Sauvignon varieties, respectively. This study revealed that the higher content of methanol was influenced by type of grape used for preparation as well as maceration duration. Further evaluation was carried out using PCA to assess the effect of genotypes variation and extraction methods on wine samples.^[78]

Gas chromatography combined with PCA and cluster analysis (CA) were successfully applied in determining content of alcoholic compound in Chinese beverages. According to the study, 21 aroma components were found to be important in the aroma profiles of Chinese liquor. Among all the compounds, seven alcoholic compound including methanol, 2-butanol, 1-propanol, isobutanol, *n*-butanol, isoamylol and phenylethanol were detected by validated GC analysis method. Isoamylol, isobutanol and 1-propanol were found as the dominant alcoholic compound with the content of 800.53, 637.67 and 338.84 mg/L, respectively. The dimensionality reduction of PCA was employed in this study to statistically separated young liquor (fresh) and aged liquors. Individual plot was generated as two-dimensional visualization constructed by PC1 and PC2 with total variance of 98.27%. Further separation using CA was built using the Euclidean distance. All liquor samples were clustered into two big groups of young liquor and aged liquors. This results proved the ability of PCA and CA to successfully separate and classify the different ages Chinese liquor samples.^[79]

In Indonesia, a majority Muslim country, it was stated by the government that the alcohol content (in percentage) of alcohol-containing drugs, traditional medicines, and supplements have to be declared on the label. Halal evaluation of alcohol content in noni (*Morinda citrifolia* L.) can be performed using gas chromatography method. The GC instrumentation was set as the inlet injection mode split of 2.5:1, injection temperature of 140°C, oven initial temperature FID detector of 40°C and hold for 5 min. The sample of noni herbal medicines was collected from herbal drug stores or online shops in Jakarta, Indonesia. Twenty samples were evaluated and categorized as beverages (18 samples) and herbal medicines (2 samples). It was found that 13 out of 20 samples contained alcohol in the range of 0.04%–1.07%. Unfortunately, none of them were labeled properly according to the regulation.^[80]

GC-FID has been used for analysis of ethanol in foods and beverages such as tea-based, fruit-based, cheese-based, milk-based, seaweed-based, instant dried noodle, etc. Ethanol stock solution was prepared (1 mg/mL) and internal standard of 0.1% v/v 1-propanol was used for sample preparation. Sample preparation was carried out using magnetic stirring aqueous extraction. Analysis was performed out using an HP-Innowax (Agilent Technologies) column (30 m x 0.25 mm x 0.25 µm). The sample injection volume was 1 µL using split ratio of 13:1. The developed method was validated according to the requirements of ISO/IEC 17025:2017. Validation result showed that the method had good linearity ($R^2 > 0.999$), good accuracy (recoveries of 96–105%) and good precision (RSD < 5%). The detection limit was low (0.006 mg/g). The determination of ethanol concentration was successfully applied in 108 samples of processed foods and beverages. Therefore, this method could be used as valid method for halal authentication of processed foods and beverages.^[52]

GC-MS using static headspace has been applied for determination of ethanol in solid and semi-solid consumer goods such as cakes, ice creams, sauces and powders. Sample preparation was carried out using mechanical homogenization and aqueous dilution of the products. Subsequently, the sample was analyzed using headspace GC-MS. Separation of analytes was performed using a capillary column DB-624 (30 m x 0.25 mm x 1.4 µm) and sample was injected in split mode employing ratio of 1:200. Identification and quantification of ethanol and ethanol-d6 was performed at scan range of 29–250 m/z with a rate of 6.1 scans/s. Result showed that the developed method was specific to detect ethanol and ethanol-d6 at the retention time of 2.65 and 2.61, respectively. The method demonstrated good linearity at the concentration range of 0.1%–2.0% v/v showed by its high R^2 value (>0.998). Additionally, good accuracy as well as good precision was obtained. The accuracy was represented

by recoveries value (average recoveries of 99.7%). The precision was demonstrated by its lower RSD value (<1.5%). From the above results, it suggested that headspace GC-MS could be used for identification and quantification of ethanol in a various solid and semi solid food products for halal authentication.^[81] 400

Identification of ethanol using headspace GC-MS has also been applied in Kombucha products. Kombucha is one of fermented beverages consist of sugar, tea, a symbiotic of bacteria and yeast which is commonly known as nonalcoholic beverage. The United States and Canada state that the content of alcoholic compounds in product must be <0.5% and <1.1% alcohol by volume, respectively, to be categorized as nonalcoholic drink. Propan-1-ol was used as internal standard for ethanol quantification. The condition of headspace was incubation temperature at 70°C, syringe temperature at 70°C, incubation time of 300 s, agitator speed at 500 rpm, injection volume of 500 µL and split ratio of 10:1. Analysis was performed using an Agilent J&W DB-624 UI (30 m x 0.25 mm x 1.4 µm) applying flow rate of 1.4 mL/min (constant flow). The developed method was linear ($R^2 > 0.995$) obtained at a concentration range of 0.025%-2.47%. The accuracy result was good demonstrated by its recovery value (102%) and good precision was also obtained (RSD<4%). The LOD and LOQ values were 0.0002% and 0.002%, respectively. It can be concluded that the method is suitable for identification and quantification of ethanol in Kombucha product. It indicated a rapid and easy integration of analytical method for halal authentication of Kombucha.^[82] 405 410 415

The development of GC-MS coupled with headspace and multidimensional (heart-cut) chromatography has been successfully applied to determine ethanol content in medicinal syrups. The aim was to ensure and guarantee the safety of the syrups. Samples used for analysis consist of adult and pediatric syrups. Monitoring and quality control of ethanol content in the products were important due to the efforts of industry to reduce the ethanol content in the food and medicinal products. Sample preparation was directly performed using headspace with condition as follows: heating syringe temperature of 90°C, incubator temperature of 100°C, incubation time of 15 min at 500 rpm, sample volume of 500 µL with split mode using ratio of 1:20. Two dimensional GC analysis was carried out using GC-MS equipped with analytical column of RTX-5 capillary column (Crossbond® 5% diphenyl/95% dimethyl polysiloxane, 30 m × 0.25 mm × 0.25 µm) for the first dimension then for the second dimension used an NST 100 MS column (Carbowax polyethylene glycol, 30 m × 0.25 mm × 2.00 µm). The method was validated according to National Agency of Sanitary Surveillance (ANVISA) with validation parameters of selectivity, linearity, precision, accuracy, LOD, LOQ and robustness. Selectivity test found that isopropyl alcohol was an interfering compound of ethanol determination in syrups. Linearity assay demonstrated linear model at concentration range of 0.25% to 10.00% v/v ($R^2 > 0.999$). The developed method was sensitive enough as shown by its LOD value (0.03% v/v) and LOQ value (0.06% v/v). The precision was measured for repeatability (CV = 3.04%) and intermediate precision (CV = 3.03%). The recoveries value obtained ranged from 97.28% to 101.38% indicating good accuracy. The robustness test showed that the method remains unchanged with the small changes of several parameters. This developed method could be used as rapid and easy analytical technique for halal authentication of syrups by determining of the ethanol content.^[83] 420 425 430 435

CONCLUSION

Chromatography-based method consist of liquid chromatography and gas chromatography using various detectors has been widely applied for food products authentication including halal analysis due to its advantages. The combination of chromatographic methods with chemometrics of multivariate analysis, a powerful statistical analysis to manage huge data generated from analytical measurement, could be used to identify potential markers to differentiate halal and non-halal samples. It will be very useful for the institutions which have responsibility for halal quality assurance. Chromatogram and peak separation profiles resulted as the instrument responses can be further evaluated for determination as well as quantification for halal and non-halal components in food products. Chromatographic-based methods were successfully carried out to analyze products 440 445

containing non-halal material such as pork and alcoholic compound. Combination of chromatographic-based method and chemometrics techniques with some scenarios can be applied for halal research on food products.

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Author contribution

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