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1 **Species specific marker peptides for meat authenticity assessment: a multispecies quantitative**
2 **approach applied to Bolognese sauce**

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15 **Abstract**

16 Food frauds are a critical issue in the field of food safety and quality. Given the high added value, and the
17 complexity of the matrix, processed meat products are among those most susceptible of adulteration.
18 Despite all the efforts made by the official control authorities and by the food industry to counteract these
19 frauds, the undeclared replacement of meat species with cheaper ones is still widespread. The meat
20 species allowed for food consumption are many, and their specific and accurate detection in highly
21 processed food products requires very sensitive and selective analytical methods. In this work, a LC-MS
22 method was developed to identify and quantify eight different meat species (duck, rabbit, chicken, turkey,
23 buffalo, equine, deer and sheep) in a complex food matrix, such as Bolognese sauce. After protein
24 extraction and trypsin digestion, a species-specific peptide marker for each species was chosen for
25 qualification and quantification. The method was validated on real Bolognese sauce samples prepared in an
26 industrial environment, showing a good sensitivity (LOD 0.2-0.8% on whole finished product) and the
27 possibility, using specifically defined calibration lines, to quantify the amount of meat present coming from
28 different species.

29 **Keywords**

30 Food fraud, meat authenticity, peptide markers, LC-MS, Bolognese sauce, species detection

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1. Introduction

The progress of globalization, together with the economic opportunity and, in some cases, the low severity and probability of the punishment, has led to a steady increase in food frauds (Pustjens, Weesepeel, & van Ruth, 2016). Thus, a growing attention is being given to food authenticity assessment. Consumers are getting more and more informed about food risks related to frauds, and they ask for valuable tools for food protection and control (Tiozzo, Mari, Ruzza, Crovato, & Ravarotto, 2017). In 2013, the EU Food Fraud Network (FFN) was created in response to the horse meat crisis, and since then meat has always been at the centre of the attention in food frauds cases. From a recent report of FFN, emerged that, of 176 total reported cases of cross-border food frauds, meat products were involved in 47 records (19 for the only poultry and 28 for the remaining meat products) (European Commission, 2016). The types of adulteration that can occur in meat are many and diverse: meat origin, meat substitution, meat processing treatment and non-meat ingredient addition (Ballin, 2010). After the horse meat scandal in 2013, the problem of meat species substitution has obviously gained more and more attention. The substitution of a species with a cheaper one in a meat product is a common fraud in the sector, and it is sometimes difficult to detect, especially in processed products. In a recent survey, the presence of an undeclared species was detected in the 68% of meat products analysed, with prevalence in sausages, burger patties and deli meats (Cawthorn, Steinman, & Hoffman, 2013). **The survey was carried out in South Africa, collecting samples from retail outlets and butcheries over a five months period (April 2012 - August 2012), and 95 of the 139 samples contained a meat species which was not declared in the product label.** Pork and chicken were the most commonly detected species, but also cases of donkey, goat and water buffalo were detected. Also among game meat, mislabelling cases due to economic gain are reported (18.5% of the samples analysed) (Quinto, Tinoco, & Hellberg, 2016). The substitution of the more expensive game meat with domestic species has also been reported (Amaral et al., 2015), for example the substitution of duck with chicken. These meat species substitutions have important implications on food safety, because the undeclared species is not subjected to any veterinary health check, microbiological and contamination control. In addition to this, some lifestyles exclude the use of particular meat species, for example pork meat is not allowed in Islamic and Jewish dietary rules. In a recent paper undeclared pork meat presence was found in a high percentage (54%) of meat samples, even if declared halal (Amaral, Santos, Oliveira, & Mafra, 2017).

Analytical tools for meat authenticity assessment are continuously evolving in response to the new challenging issues and they cover a wide variety of techniques. In meat species substitution, differences in the genetic materials can be detected by the use of genetic approaches, such as DNA-hybridization and polymerase chain reaction (Rahmati, Julkapli, Yehye, & Basirun, 2016). DNA barcoding is among the most promising genetic techniques for food authentication in processed meat, being able to recognize with 68.3% of success meat species through sequencing of the target gene (cytochrome c oxidase subunit I

66 gene) (Hellberg, Hernandez, & Hernandez, 2017). Another innovative technique is based on the
67 development of chemiluminescent optical fibre genosensors, which has already been applied with success
68 for the detection of pork in minced raw meat mixtures down to 1% (Torelli, Manzano, & Marks, 2017).
69 Despite being a powerful tool for species identification, DNA methods can easily fail in heavily processed
70 food matrices, due to DNA degradation, and also the exact quantitation of the different species detected
71 might be cumbersome and difficult.

72 Another common technique is based on the recognition of species-specific proteins via immunoassays. A
73 great number of enzyme linked immunosorbent assays (ELISA) have been developed for meat speciation,
74 due to their speed and ease of use (Perestam, Fujisaki, Nava, & Hellberg, 2017). These methods can also be
75 applied to processed products, for example pork can be detected in heat-processed meat products by
76 monoclonal antibody-based ELISA even at low percentages (0.5%). Anyway, also these methods suffer in
77 many cases of false positives and false negatives in heavily processed foodstuff, and their accuracy tends to
78 be very much product-specific.

79 As mentioned before, food frauds are most likely to occur in processed food products, where minced meat
80 is present (preventing visual recognition). In these thermally treated processed food, several detection
81 problems can arise due to interfering compounds, nucleic acids and proteins degradation (Bauer, Weller,
82 Hammes, & Hertel, 2003; Terry, Harris, & Parkes, 2002). Hydrolysis and denaturation phenomena are
83 particularly harsh in those foodstuffs that also have acid pH (for example lemon juice or vinegar) and/or
84 that undergo to severe heat treatments (for example prolonged cooking or sterilization). Thus, for the most
85 complex food matrices, having different ingredients mixed together or subjected to severe thermal
86 treatments, the need for robust, sensitive and selective methods is still an issue (Sentandreu & Sentandreu,
87 2014).

88 A new method for the identification of species-specific molecular markers has been lately gaining
89 momentum, based on the coupling two very performing techniques in the field of food analysis: liquid
90 chromatography (LC) coupled to mass spectrometry (MS), applied to the detection of peptides. Label free
91 relative quantification of meat species-specific proteins is achieved through the detection of marker
92 peptides, generated by enzymatic hydrolysis. **The combination of off-gel fractionation and LC-MS detection,**
93 **allowed the detection of chicken in meat mixtures at very low percentages (0.5% w/w) with high**
94 **confidence (Sentandreu, Fraser, Halket, Patel, & Bramley, 2010). In more recent researches, even small**
95 **amount (1% w/w) of beef, pork, chicken, duck and goose in cooked meat could be detected by this**
96 **methodology (Montowska, 2017; Montowska & Fornal, 2017). With the contribution of off gel**
97 **fractionation, specific peptides, deriving from myosin light chain 1 and 2, could be identified for buffalo and**
98 **sheep. 0.5% of buffalo meat could be detected both in raw and cooked mixture (Naveena et al., 2017).**
99 LC/MS detection of peptides was demonstrated to be more efficient than PCR and ELISA assays for the

100 determination of the origin in products undergone severe heat treatment or acid/alkali extraction, for
101 example in the case of gelatine, where porcine and bovine collagen peptides could be identified down to
102 0.4-1% of contamination (Grundy et al., 2016). This method has also been successfully applied to meat
103 mixtures of beef, pork, horse and lamb, exploiting myoglobin derived peptides, with a good sensibility (1%)
104 (Watson, Gunning, Rigby, Philo, & Kemsley, 2015). The good stability of peptides to thermal treatments,
105 rather than intact proteins or nucleic acids, is also demonstrated by the possibility to identify markers for
106 horse and pork also in processed foodstuff even in small amount (0.24%) (von Bargaen, Brockmeyer, &
107 Humpf, 2014). Several efforts have been made to make the mass spectrometry analysis much faster and
108 easier, and appreciable results were obtained using the LESA-MS methodology. This technique allowed
109 detecting pork, horse, turkey and chicken meat at amount ranging from 5 to 10% (Montowska, Alexander,
110 Tucker, & Barrett, 2014). Demonstrating the robustness of this approach in heavily processed food
111 matrices, in a previous work we developed a LC-MS method for the identification and quantification of beef
112 and pork meat in a challenging food matrix, Bolognese sauce, where these two types of meat are usually
113 employed as ingredient (Prandi et al., 2017).

114 In the present paper, with the aim of achieving a fast and robust method for multispecies determination
115 and quantification, we have extended the methodology to other eight species of food interest, covering
116 most of the European meat market. In order to demonstrate its relevance for the food industry in a real
117 environment, its applicability has been demonstrated on real samples prepared at the industrial level.

118 More specifically, we took into account, beside bovine and pork, poultry and rabbit meat, equine meat
119 (donkey and horse), buffalo, red deer and sheep. Horsemeat has been the focus of attention in 2013, but
120 also turkey was involved in several scandals, such as the substitution of halal lamb with turkey meat at the
121 beginning of 2017 (European Commission, 2017). The method developed and here reported, based on
122 LC/MS determination of peptides, is the first one able to detect and quantify simultaneously these eight
123 species in Bolognese sauce.

124 **Materials and methods**

125 **2.1 Samples.**

126 Bolognese sauces were prepared in a pilot plant following the standard industrial procedure. Traditional
127 Italian recipe was opportunely modified to have standard reference matrices of each meat species. Thus,
128 Bolognese sauces were prepared with 19% of pure meat of: duck, rabbit, chicken, turkey, donkey, buffalo,
129 horse and red deer. Other ingredients in the Italian traditional recipe are: tomato pulp 28%, water, tomato
130 concentrate 16.5%, onion, carrots 4%, celery 3%, sunflower oil, corn starch, salt, yeast extract, sugar, laurel
131 0.06%, sage extract, black pepper. Briefly, chopped onion is suffused with vegetables. Then, tomato sauce,

132 tomato concentrate and freshly minced meat are added, and the sauce is cooked for two hours (approx.
133 temperature 90-100°C). After cooking, Bolognese sauce is poured into the jar at 70°C. Finally, the
134 sterilization is performed into an autoclave: the temperature is brought from 30°C to 121°C, then the jars
135 are cooled again to 30°C (total sterilization time: 90 min).

136 Besides Bolognese sauce prepared with pure meat species, mixed Bolognese sauces were also prepared,
137 where the total amount of meat (19%) was kept constant. Three different sets of samples were prepared;
138 each consisting of 3 concentration levels (for calibration purpose) and blind samples (for cross validating
139 purpose) as reported in Table 1.

140 2.2 Reagents and solvents

141 Deionized water was obtained with Select water purification system (Suez water, Thame, UK).
142 Tris(hydroxymethyl)-aminomethane (99+%), urea (99-101%), hydrochloric acid (≥37%), formic acid (≥95%),
143 DL-dithiothreitol (≥98%), iodoacetamide (crystalline) and trypsin from porcine pancreas (1,000-2,000 BAEE
144 units/mg solid) were purchased by Sigma Aldrich (Saint Louis, MO, USA). Thiourea was purchased by Carlo
145 Erba (Cornaredo, MI, Italy). Acetonitrile (≥99.9%) was purchased by Honeywell-Riedel de Haën (Seelze,
146 Germany). Ammonium bicarbonate (≥99%) was purchased by Fluka Chemie (Buchs, Switzerland).

147 2.3 Protein extraction

148 One hundred grams of Bolognese sauce were homogenized with a fixed rod homogenizer (Ultra Turrax IKA
149 T50 digital, Staufen im Breisgau, Germany) for 2 min at a speed of 22,000 rpm. The homogenized samples
150 were then lyophilized for 44 hours in a Lio 5P freeze drier (5Pascal, Milan, Italy) at a vacuum level of 0.80
151 mbar and a temperature of -50°C. One gram of lyophilized Bolognese sauce was extracted with 10 mL of a
152 0.05 M TrisHCl, 6 M urea and 1 M thiourea (pH 8), using a fixed rod homogenizer (Ultra Turrax IKA T18
153 digital, Staufen im Breisgau, Germany), for 5 min at 10,000 rpm. Samples were centrifuged at 3,220g for 10
154 min at 4°C (Eppendorf 5810R, Hamburg, Germany) and supernatant was filtered through 1 µm glass fibre
155 syringe filters (Acrodisc, Waters, Milford, MA, USA). Salts and caotropic agents were removed by solid
156 phase extraction (SPE) using Sep-Pak C18 Plus short cartridges (Waters, Milford, MA, USA) according to the
157 manufacturer instructions. The cartridges have 360 mg sorbent per cartridge, with a 55-105 µm particle
158 size. The loaded volume of sample was around 5 mL of protein solution. Desalted protein extracts were
159 lyophilized for 6 hours in a Lio 5P freeze drier (5Pascal, Milan, Italy) at a vacuum level of 0.80 mbar and a
160 temperature of -50°C and stored at -20°C.

161 2.4 Protein digestion

162 Freeze dried samples were dissolved in 1 mL of ammonium bicarbonate (0.05 M in water). Disulphide
163 bridges were reduced by adding 5 µL of dithiothreitol (0.2 M in water) and incubating the samples at room

164 temperature for 1 hour. Then, sulfhydryl groups were alkylated by adding 4 μL of iodoacetamide (1 M in
165 water) and incubating in the dark for 1 hour at room temperature. Excess iodoacetamide was neutralized
166 by the addition of 20 μL dithiothreitol (0.2 M in water) and incubation at room temperature for 1 hour.
167 Then, protein content was measured with the Quant-it protein assay kit (Thermo Fisher Scientific,
168 Waltham, MA, USA) using the Qubit Fluorimeter (Thermo Fisher Scientific, Waltham, MA, USA), according
169 to the manufacturer instruction. Trypsin digestion was carried out using an enzyme to substrate ratio of
170 1:20 and incubating overnight at 37°C in an orbital shaker incubator (ES 20, Biosan, Riga, Latvia) with a
171 speed of 100 rpm. Digestion was stopped by the addition of 50 μL of formic acid (10% v/v in water).
172 Samples were stored at -20°C until UHPLC/ESI-MS/MS analysis. Just before injection, samples were
173 centrifuged at 15,093g for 10 min at 4°C and the supernatant was saved for the analysis.

174 2.5 UHPLC/ESI-MS/MS analysis

175 Peptides generated by enzymatic cleavage were analysed using reverse phase ultra-high performance liquid
176 chromatography (UHPLC) coupled to electrospray ionization tandem mass spectrometry (ESI-MS/MS).

177 Aeris PEPTIDE 1.7 μm XB-C18 column (100 \AA , 150 \times 2.1 mm; Phenomenex, Torrance, CA, USA) was used for
178 the chromatographic analysis, equipped with a Security Guard ULTRA Cartridge (C18-Peptide, ID 2.1 mm;
179 Phenomenex, Torrance, CA, USA). Chromatographic separation was run in a Dionex Ultimate 3000 UHPLC
180 (Sunnyvale, CA, USA). Flow was set at 0.2 mL/min, column temperature at 35°C and sample temperature at
181 18°C; eluent A was water with 0.1% (v/v) of formic acid and 0.2% (v/v) of acetonitrile, eluent B was
182 acetonitrile with 0.1% (v/v) formic acid and 0.2% of water. A gradient elution was performed, according to
183 the following parameters: 0–7 min 100% A, 7–50 min from 100% A to 50% A, 50–52.6 min 50% A, 52.6–53
184 min from 50% A to 0% A, 53–58.2 min 0% A, 58.2–59 min from 0% A to 100% A, 59–72 min 100% A (total
185 analysis time 72 min). **Injection volume was 2 μL for 100% pure species and 10 μL for calibration curves and**
186 **blind samples.**

187 Detection was achieved using a triple-stage quadrupole mass spectrometer (TSQ Vantage, Thermo Fisher
188 Scientific, Waltham, MA, USA) with the following parameters: solvent delay 0-7 min, acquisition 7-58.2 min,
189 ionization type positive ions; spray voltage 3,500 V, vaporizer temperature 250°C; sheath gas pressure 22
190 (arbitrary units); capillary temperature 250°C. For the Q1MS Scan mode, the acquisition range was set at
191 100-1,500 m/z. For the Product Scan mode, different collision energies were applied (20, 25, 30, 35 and 40
192 V) to fragment the selected ions, and the acquisition range for fragments detection was 100-1,500 m/z. For
193 the Selected Reaction Monitoring (SRM) method, the monitored transitions are reported in **Table 2**. The
194 first reported fragment was used as quantifier, the second one as qualifier. UHPLC/ESI-MS data were
195 elaborated using Xcalibur software (Thermo Fisher Scientific, Waltham, MA, USA). **The calibration curves**

196 were constructed using both the “TIC” trace (sum of the two monitored transitions), and the trace of the
197 first transition (the most intense), giving fully comparable results.

198 **2.6 μ HPLC-LTQ-OrbitRAP analysis**

199 For protein and peptide identification through tandem mass spectrometry analysis, samples were analyzed
200 with μ HPLC (Dionex Ultimate 3000, Sunnyvale, CA, USA) coupled to Orbitrap LTQ XL mass spectrometer
201 (Thermo Scientific, Waltham, MA, USA). Software employed for the analysis was Xcalibur 2.0.7 (Thermo
202 Scientific, Waltham, MA, USA). Eluent A was water + 0.2% formic acid, eluent B was acetonitrile with 0.2%
203 formic acid. Sample loading conditions were: enrichment cartridge: μ -Precolumn Cartridge, Acclaim
204 PepMap100 C18 5 μ m, 100 Å, 300 μ m \times 5 mm, loading flow: 30 μ L/min, 50% eluent A and 50% eluent B.
205 Sample elution condition were: column: Phenomenex Jupiter 4 μ m Proteo 90Å 150 mm \times 0.3 mm, column
206 temperature: 35°C, gradient: 0-4 min 10% B, 4-60 min linear from 10% B to 50% B, 60-62 min from 50 to
207 95% B, 62-72 min 95% B (column washing), 72-73 min from 95% B to 10% B, 73-82 min 10% B (column
208 equilibration). HRMS acquisition was performed through 5 subsequent events: event 1: full scan acquisition
209 from 250 to 2000 m/z in high resolution mode (resolution at 400 m/z = 30,000); events from 2 to 5: data
210 dependent scan, at each cycle the four most intense ions (with charge $z > 1$ and with a minimum signal of
211 500 counts) identified in event 1 are fragmented. The same ion (tolerance 10 ppm and isolation window 2
212 m/z) can be observed for a maximum of 2 cycles, and then it is automatically inserted in the exclusion list
213 for a maximum time of 20 seconds. Fragmentation is performed in the linear trap of the instrument in CID
214 mode with collision energy of 35.

215 Proteins were identified with Peaks Studio (Bioinformatics Solutions, Waterloo, ON, Canada). Parameters
216 were: precursor ion tolerance 5 ppm, fragment ion tolerance 0.8 Da, decoy database search: strict 0.01,
217 relaxed 0.05, fixed modifications: cysteine carbamidomethylation, variable modifications: methionine
218 oxidation, hydroxyproline and hydroxylysine. Searches were run both on the specific database of each
219 species (for example *Gallus gallus* or *Equus caballus*) and on the class database (for example Aves or
220 Mammals). Peptides which did not align with any protein present in the database were identified with *de*
221 *novo* sequencing using both high and low resolution mass spectrometry.

222 **3. Results and discussion**

223 **3.1 Marker peptides identification**

224 Bolognese sauces prepared with the meat of several single pure species were first analysed. The proteins
225 were extracted and digested with trypsin (details in the experimental section) and the peptide mixtures
226 were analysed by LC coupled to high resolution mass spectrometry. Samples were analysed in Data
227 Dependent Scan mode using an LTQ-OrbitRAP instrument. The obtained spectra were analysed with the

228 Peaks™ software to identify the main proteins and peptides present. The main proteins identified in the
229 Bolognese sauces made with different types of meat are reported in Table S1 (supplementary on line
230 material).

231 In Bolognese sauce made with poultry, rabbit, goat or sheep, the identified peptides were mostly deriving
232 from collagen, myosin and tropomyosin, demonstrating that stomatic (collagen) and myofibrillar (myosin
233 and tropomyosin) proteins are the most resistant to the thermal treatments applied to Bolognese sauce,
234 generating a consistent amount of peptides. On the opposite, low amount of myofibrillar proteins-derived
235 peptides were found in donkey Bolognese sauce, where collagen (stomatic protein) and haemoglobin
236 (sarcoplasmic protein) gave the highest number of peptides. Haemoglobin derived peptides are found also
237 in buffalo and red deer, together with the muscle counterpart myoglobin and with a good amount of
238 myofibrillar proteins derived peptides (alpha and beta actin). Most of the horse peptides derived instead
239 from myosin (myofibrillar protein). The amount and protein types of the peptides derive from a complex
240 combination of factors: the meat cut used for Bolognese sauce preparation (some parts of the carcasses are
241 richer in collagen), the prevalence of the single protein class in that meat species and the resistance to
242 thermal treatment of the single protein.

243 To select good candidate marker peptides, only peptides having length longer than six amino acids were
244 chosen, as previously performed by other authors (Watson, Gunning, Rigby, Philo, & Kemsley, 2015). In this
245 way, the peptide length ensures the species-specificity. Then, the candidate peptide ions were cross
246 checked in the chromatogram of the other meat species and only the peptides that were present in one
247 meat types were selected. Moreover, a database search was performed to ensure peptide specificity (Basic
248 Local Alignment Search Tool, BLAST). Among the species specific marker peptides identified, one peptide
249 for each meat species was finally chosen taking into account the following factors: high abundance, good
250 signal to noise ratio at low concentrations, high specificity, no missed cleavages and trypsin specific
251 cleavage sites at both ends. Identified marker peptides are reported in **Table 2, together with** the MS
252 parameters used for their detection. The SRM chromatograms of the peptide markers for each species are
253 reported in Figure 1. For each chromatogram, all the transitions for all the species were monitored, to
254 ensure the sensitivity of the method (each species having its peptide marker) and the selectivity of the
255 method (only the specific marker peptide being present in each species).

256 All the identified marker peptides were re-analysed in a triple quadrupole instrument in order to re-confirm
257 their sequence. **Some of the** marker peptides chosen for red meat samples analysed derived from the
258 sarcoplasmic protein myoglobin. The highest abundance of myoglobin in red meat is probably the basis of
259 the largest amount of myoglobin derived peptides in buffalo and horse. On the other side, marker peptides
260 chosen for chicken and turkey (white meat) derived from fructose 1,6-bisphosphate aldolase, and the

261 myofibrillar proteins troponin T and myosin, respectively. In some cases it was possible to identify a marker
262 peptide very specific for the target species, such in case of rabbit, chicken, turkey and buffalo, whose
263 marker peptides were found to be present only in *O. cuniculus*, *G. gallus*, *M. gallopavo* and *B. bubalis*,
264 respectively. The marker peptide for rabbit (PHSHPALTPEQK) derives from the N-term of the metabolic
265 enzyme fructose bisphosphate aldolase and was previously studied in literature in order to assess chemical
266 modifications to the enzyme fructose 1,6-bisphosphate aldolase (Hopkins, O'Connor, Allen, Costello, &
267 Tolan, 2002), or it has been used as control to evaluate the effect of drift gas polarizability for the
268 separation of tryptic peptides (Ruotolo, McLean, Gillig, & Russell, 2004). So, this work describes for the first
269 time its use for authenticity assessment in food products. The marker peptide for turkey ALGQNPTNAEMNK
270 was already proven to be resistant to heat treatment, as it was detected in cooked beef mixture spiked
271 with 10% turkey meat (Montowska, Alexander, Tucker, & Barrett, 2014). No information was available in
272 literature about the troponin T peptide SDTEEVEHGEAHEAEVHEEAH and the myoglobin peptide
273 VETDVAGHGQEVL, present only in chicken and buffalo meat (respectively), which have been thus identified
274 here for the first time. The chicken marker peptide shows a post translational modification, an acetylation
275 on the N-term serine residue. This is a common feature in troponin T, where the N-term residue is
276 acetylated and, subsequently, it can be phosphorylated by a protein kinase (Gusev, Dobrovol'skiĭ, &
277 Severin, 1978). For what concerns sheep, the marker peptide AGEVGPPPPGPAGEK was the most
278 abundant among those that were specie specific for *O. aries*; this peptide derives from collagen $\alpha 1(I)$ and it
279 has two hydroxylated prolines (underlined in the sequence). Proline hydroxylation is a typical feature of
280 collagen protein, since it confers the compact structure to collagen helices. This peptide is known in
281 literature because of its use for the identification of sheep in natural leather goods (Izuchi, Takashima, &
282 Hatano, 2016), but it has never been used for food authenticity assessment. **There are many species that**
283 **share a very similar sequence, which only differs for one amino acid: the N-term alanine of the peptide is**
284 **present, in fact, only in sheep collagen.** Unfortunately, no species-specific peptides for goat could be here
285 detected, since all the identified peptides present in goat were also present in sheep. In previous works, it
286 has been possible to discriminate between keratins from sheep and goat exploiting specific keratin-derived
287 peptide markers, with the aim to assess wool and cashmere authenticity (Paolella et al, 2013). However,
288 the amount of keratin in a meat preparation is negligible, thus these peptide markers could not be used
289 here.

290 The myoglobin peptide HPGDFGADAQGAMTK has been previously used to detect horse in fresh meat
291 (Watson, Gunning, Rigby, Philo, & Kemsley, 2015). Given the genetic proximity of donkey and horse, which
292 belong to the same genus (*Equus*), we found this peptide also in Bolognese sauce prepared with donkey
293 meat. Since the alignment in the Uniprot database gave as the only outputs horse and zebra, this is
294 probably the first time that the myoglobin peptide HPGDFGADAQGAMTK has been found in donkey.
295 Unfortunately, all the peptides identified in the Bolognese sauce prepared with horse meat, were present

296 also in that prepared with donkey meat. Thus, this myoglobin peptide was used as a marker peptide for
297 equine meat (horse and donkey). The amino acid sequence of the marker peptide for duck (*Anas*
298 *platyrhynchos*) was not present in the protein database, thus it was determined through *de novo*
299 sequencing using a triple quadrupole instrument. The amino acid sequence was determined on the basis of
300 the fragmentation pattern. The amino acid sequence (QELADLAR) has a certain homology with the
301 sequences QENANTLAR and QENANALAR, found in fructose-1,6-bisphosphate aldolase of *A. platyrhynchos*
302 (the same protein from which the marker peptide for rabbit derives). The failure to align the peptide
303 sequences QELADLAR with duck can be explained by the limited number of proteins available in its
304 database. Unfortunately, duck proteome, to date, has not been intensively studied as it happened for other
305 animal species, such as for examples piglet or cattle. However, we found this peptide only in the duck
306 sample, and it was not present in any of the other meat samples under investigation neither in beef or pork
307 meat. Thus, this cross check covered almost all the meat used for human consumption in Western
308 countries. For what concerns red deer, we found that the peptide LNFKPEEEYPDLSK is present in M-type
309 creatine kinase of red deer, and it is present in the protein database of thirteen line ground squirrel,
310 Tasmanian devil and gray short tailed opossum. However, these species are not of food interest and quite
311 of difficult availability, thus it is unlikely that they could be used for meat adulteration. So, the peptide
312 LNFKPEEEYPDLSK was used as a marker for red deer presence.

313 **3.2 Meat species quantification**

314 Once identified, marker peptides for each of the species under study were analysed in order to define the
315 limit of detection and quantification of the method, and also the linearity of the response at very low
316 percentages was studied. More specifically, classical Bolognese sauces were purposely prepared in an
317 industrial pilot plant, following traditional recipe (50% bovine + 50% swine meat), with the addition of 1, 2.5
318 and 5% (w/w_{meat}) of duck, rabbit, chicken, turkey, donkey, buffalo, horse and red deer.

319 The aim of this part of the work was to quantify the amount of each meat species (in % w/w) present in
320 Bolognese sauce samples. To do that, the areas of the marker peptides were integrated (sum of both the
321 transitions monitored). Also the area of a common peptide (present in all the meat species) was integrated.
322 The ratio between the area of each marker peptide and the area of the common peptide was made to
323 normalize the data, and to avoid differences due to possible variations in protein extraction efficiency,
324 enzyme activity, and other factors. The peptide used for the normalization of the data was DMIPAQK
325 (deriving from C-term of creatine kinase M-type), used here as a molecular marker for the total meat
326 amount. The obtained ratio was plotted against the amount of each meat species (% w/w). A very good
327 linearity was found for equine, followed by duck and rabbit (Table 4), while lower correlation coefficients
328 were found for turkey and buffalo.

329 Limits of detection and quantification were determined from the calibration curves, according to the IHC
330 guidelines (2005), using the formulas: $LOD = 3.3 \times \frac{S_y}{S}$ and $LOQ = 10 \times \frac{S_y}{S}$, where S_y is the standard
331 deviation of the response and S is the slope of the calibration curve. The limits of species detection in mixed
332 meat, calculated as percentage on total meat, were in the range 0.8-3.6% (equine-turkey, respectively),
333 while the limits of quantification, always calculated as percentage on total meat were in the range 2.8-
334 12.1%.

335 Considering that the amount of meat used in Bolognese sauce preparation was 19%, the LOD and the LOQ
336 calculated on the total sauce are 0.15-0.68% and 0.53-2.30% on whole finished product, respectively. Thus,
337 the developed method is able to detect even small contamination of these meat species in the final product
338 and, in the case of massive adulteration, it is able to quantify with a good accuracy the different
339 proportions of meat added. These results were very encouraging also considering the complexity of the
340 tested product (Bolognese sauce), which is composed of many ingredients and submitted to prolonged
341 cooking and a strong sterilization process.

342 Finally, several blind samples were analysed to check the reliability of the method from very low
343 percentages (such as 0.38% for turkey and buffalo, around the LOD and LOQ values) to 6% (w/w, on whole
344 product) for sheep. The SRM chromatograms of the analysed blind samples are reported in Figure 2.

345 Very good trueness values (in the range 80-120%) were found for duck, chicken, and sheep 1 (Table 5). It is
346 important to note that, while the amounts of duck and sheep meat added, were higher than LOQ, chicken
347 meat was added in a percentage slightly lower than the LOQ. However, the developed method was able to
348 detect it with a good accuracy. Considering the total amount of meat present in the Bolognese sauce (19%
349 w/w), in most cases the amount of each added species was very low: from 0.38% (w/w, on whole product)
350 for turkey and buffalo to 6% (w/w, on whole product) for sheep. Turkey and buffalo (the two species tested
351 in lower amount) were correctly detected in the samples, even if with less satisfactory accuracy: turkey was
352 overestimated in the blind sample (5.8% instead of 2%, trueness 290%), while buffalo was underestimated
353 (1% instead of 2%, trueness 50%). However, it has to be taken into account that, since the absolute values
354 are low (low % of meat species added), the error percentage becomes high. For higher values (higher % of
355 meat species added), the trueness improves. Intermediate accuracy values were obtained for the other
356 species. The developed method has good specificity and sensitivity for the meat species under
357 investigation. Its application to quantify the amount of meat using a calibration curve in processed food can
358 be improved for some species; however, considering that no analytical methods are available at the
359 moment for this purpose, the results here shown represent a step forward in the field of food
360 authentication, also considering the complexity of the product, and the strong thermal treatment occurred.

361 Once established the threshold of the method for each species, several samples at unknown meat
362 composition were subjected to the new developed analysis for fraudulent meat species checking. The SRM
363 chromatograms of the blind samples analysed are reported in Figure 3.

364 As shown in Table 6, the proposed UHPLC/ESI-MS/MS method was always able to detect the meat addition,
365 thus no false negatives were present, demonstrating a high sensitivity of the method. Moreover, the
366 developed method has showed a high specificity, because when a certain meat species was not present,
367 samples always resulted below the LOD, thus no false positives were reported. This is particularly valuable
368 because we took into account also closely related species that could cross-react, such as for example turkey
369 and chicken.

370 **3.3 Conclusions**

371 The here developed method allows the simultaneous detection and quantification of 8 different meat
372 species in a 72 minutes run in Bolognese sauce preparations. Selected peptide markers were used to build
373 calibration curves with a good linearity, allowing obtaining the quantification of the meat species present.
374 This comprehensive methodology took into account the most used species in the meat sector, and was
375 applied to real industrial products prepared by a sausage company according to industrial recipes. On the
376 basis of the obtained results, the method showed, on blind samples, a very good specificity (no false
377 positives) and sensitivity (no false negatives). This is of the utmost importance giving the application on a
378 complex food matrix like Bolognese sauce, which is one of the most susceptible to frauds. The quantitative
379 performance of this method will allow to discriminate between an accidental contamination (low amount
380 detected) and a voluntary adulteration (presumably higher detected amount). With the current work, we
381 provide a useful tool both to the control bodies (to assess product authenticity), and to the food industries
382 (to monitor supplies of raw materials). Moreover, the success of the application of this method to a very
383 difficult food matrix demonstrates how the same approach can likely be successfully applied to a broad
384 range of food preparations.

385 **4. Conflict of interest**

386 The authors declare that they have no conflict of interests.

387 **5. Acknowledgements**

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389 Grant Agreement 613688 (Ensuring the Integrity of the European food chain – Food Integrity).

390

391 **6. Captions to figures**

392 **Figure 1.** SRM chromatograms obtained for each species: A – duck, B – rabbit, C – chicken, D – turkey, E –
393 buffalo, F – horse, G – deer, H – sheep. For each chromatogram, all the transitions for all the species were
394 monitored.

395 **Figure 2.** Overlaid SRM chromatograms of the blind samples analysed for sample set 1 (A – duck, rabbit,
396 chicken and turkey) and for sample set 2 and 3 (B – buffalo, equine, deer and sheep).

397 **Figure 3.** Overlaid SRM chromatograms of the detected meat species in the unknown samples.

398

399

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Table 1. Experimental plan: set 1 comprises poultry and rabbit meat, set 2 contains buffalo, equine and red deer meat, and set 3 contains sheep and goat meat.

Set 1	Set 2	Set 3
Mix beef/pork 96% + 1% duck, 1% rabbit, 1% chicken, 1% turkey	Mix beef/pork 96% + 1% donkey, 1% buffalo, 1% horse, 1% red deer	Mix beef/pork 98% + 1% sheep, 1% goat
Mix beef/pork 90% + 2.5% duck, 2.5% rabbit, 2.5% chicken, 2.5% turkey	Mix beef/pork 90% + 2.5% donkey, 2.5% buffalo, 2.5% horse, 2.5% red deer	Mix beef/pork 94% + 3% sheep, 3% goat
Mix beef/pork 80% + 5% duck, 5% rabbit, 5% chicken, 5% turkey	Mix beef/pork 80% + 5% donkey, 5% buffalo, 5% horse, 5% red deer	Mix beef/pork 90% + 5% sheep, 5% goat
Blind_1	Blind_2	Blind3A + Blind 3B

Table 2. Identified marker peptides for each species are reported with their retention times and SRM parameters.

Peptide	Species	Retention time (min)	Parent ion (m/z)	Product ions (m/z)	Collision Energy
QELADLAR	Duck	24.4	458.1	449.3; 545.0	20
PHSHPALTPEQK	Rabbit	18.8	447.8	234.8; 314.0	30
SDTEEVEHGAEAEAEVHEEAH	Chicken	24.0	636.3	226.6; 155.8	30
ALGQNPTNAEMNK	Turkey	22.2	694.2	904.5; 185.2	35
VETDVAGHGQEV L	Buffalo	26.0	677.1	1123.5; 200.6	30
HPGDFGADAQGAMTK	Equine	23.1	751.8	234.6; 742.6	35
LNFKPEEEYPDLSK	Deer	27.7	570.1	722.7; 135.5	25
AGEVGPPGPPGPAGEK	Sheep	21.4	724.7	518.3; 1092.6	30
DMIPAQK	Meat	22.3	802.3	443.2; 297.1	30

Table 3. Tandem mass spectrometry identification of the species specific marker peptides chosen for each meat species. Columns from “a” to “z” refer to the types of fragment ions observed in an MS/MS spectrum of the peptides, with the m/z ratio of the detected fragments. Number in brackets indicate the position of the broken bond (count starts from the N-term for a, b, and c fragments; from the C-term for x, y, and z fragments). In the “BLAST” column, the species whose proteins align with 100% identity with the peptide marker are reported.

Sequence	a	b	c	x	y	z	Protein	Blast
QELADLAR (Duck)		257.5 (2) 370.9 (3) 556.7 (5) 897.6 (8)		384.7 (3)	174.7 (1) 359.0 (3) 474.2 (4) 658.3 (6)	229.1 (2)	<i>De novo</i> sequencing	-
PHSHPALTP EQK (Rabbit)	207.0 (2)	235.0 (2) 459.3 (4) 1069.1 (10)			501.2 (4) 602.1 (5) 883.3 (8) 1020.8 (9) 1109.1 (10) 1245.5 (11)	257.6 (2)	Fructose- bisphosphate aldolase	<i>O. cuniculus</i>
<u>S</u> DTEEVEHG EAHEAEEVH EEAH (Chicken)		346.0 (3) 604.6 (5) 703.3 (6) 970.6 (9)		510.3 (4)	155.4 (1) 226.4 (2) 357.3 (3) 484.2 (4) 721.3 (6) 1051.7 (9) 1149.5 (20)		Troponin T	<i>G. gallus</i>
ALGQNPTN AEMNK (Turkey)	157.0 (2)	185.2 (2) 1127.1 (11) 185.2 (4)	299.2 (6)		391.7 (3) 592.2 (5) 706.5 (6) 807.0 (7) 904.5 (8) 1018.7 (9) 1203.7 (11) 452.4 (8) 573.8 (10)		Myosin essential light chain isoform 1	<i>M.</i> <i>gallopavo</i>

	201.1 (2)	229.1 (2)							
	417.2 (4)	330.2 (3)				132.1 (1)			
VETDVAGH	516.3 (5)	445.2 (4)				739.4 (7)	1007.5		
GQEVL	781.4 (8)	866.4 (9)	561.3 (5)	708.3 (6)		810.4 (8)	(10)	Myoglobin	<i>B. bubalis</i>
(Buffalo)	838.4 (9)	994.5 (10)				909.5 (9)	1108.5		
	1194.6	1123.5 (11)				1024.5 (10)	(11)		
	(12)	1222.6 (12)				1254.6 (12)			
						146.7 (1)			
						247.6 (2)			
						379.0 (3)			
HPGDFGAD		234.6 (2)				634.7 (6)			<i>E. burchelli</i>
AQGAMTK	109.7 (1)	683.2 (7)		406.5 (3)		705.8 (7)		Myoglobin	<i>E. caballus</i>
(Horse)	526.7 (5)	1124.9 (12)				892.5 (9)			
						948.8 (10)			
						1097.0 (11)			
						1269.0 (13)			
						1366.0 (14)			
LNFKPEEEY						233.8 (2)			<i>C. elaphus</i>
PDSLK		858.7 (7)		372.7 (3)		558.9 (5)		Creatine	<i>hippelaphus</i>
(Deer)		987.3 (8)				722.0 (6)		kinase	<i>I.</i>
						1206.8 (10)			<i>tridecemline</i>
									<i>atus</i>
									<i>S. harrisi</i>
									<i>M.</i>
									<i>domestica</i>
AGEVGPPG		258.2 (3)	146.4 (2)			147.1 (1)			
		357.3 (4)	374.3 (4)			333.2 (3)			
PPGPAGEK	229.6 (3)	414.6 (5)	431.8 (5)	302.9 (2)		500.7 (5)			
(Sheep)	385.9 (5)	947.1 (11)	658.0 (7)	681.3 (7)		558.2 (6)	905.9 (10)	Collagen	<i>O. aries</i>
		1116.7 (13)	1134.1			1035.5 (11)			
			(13)			1092.5 (12)			

Table 4. Calibration curves and limits of detection and quantification for the different meat species. Table 4
Calibration curves and limits of detection and quantification for the different meat species.

Meat species	Line equation	R ²	LOD (% w/w)	LOQ (% w/w)
Duck	$y = 0.002573x - 0.00138$	R ² =0.962	1.3	4.0
Rabbit	$y = 0.001937x - 0.00046$	R ² =0.931	1.8	5.5
Chicken	$y = 0.000919x - 0.00013$	R ² =0.912	2.1	6.3
Turkey	$y = 0.000367x + 0.000472$	R ² =0.734	4.0	12.1
Buffalo	$y = 0.000285x + 0.000674$	R ² =0.848	2.8	8.6
Equine	$y = 0.001252x + 0.000559$	R ² =0.995	0.9	2.8
Red Deer	$y = 0.000357x - 0.000104$	R ² =0.904	2.2	6.6
Sheep	$y = 0.001188x + 0.00017$	R ² =0.786	3.4	10.4

Table 5. Results obtained for the blind samples, where the different meat species were added.

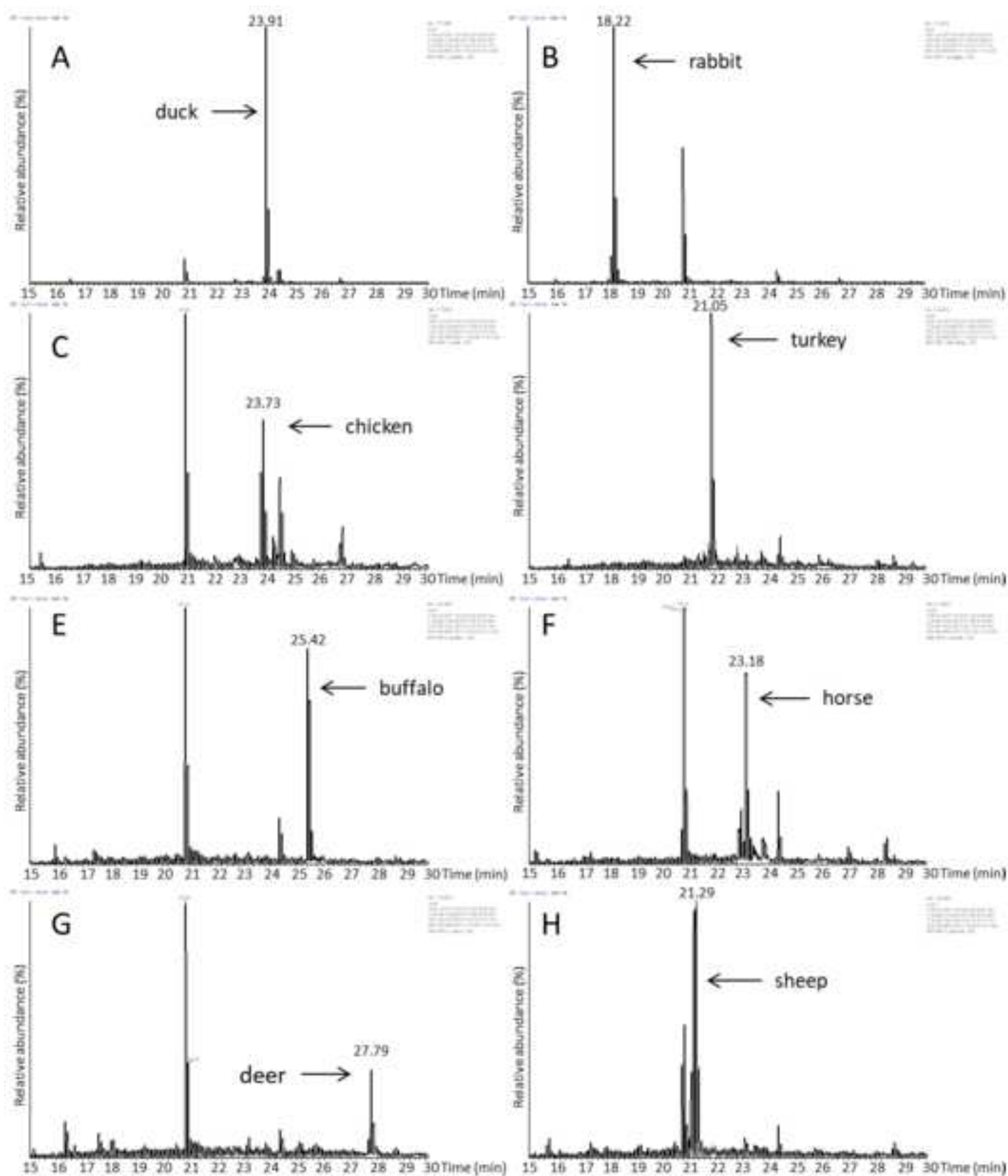
species	% (w/w) of meat added	% (w/w) of meat calculated
Duck	5.0	4.6±0.2
Rabbit	4.0	5.6±0.0
Chicken	6.0	5.0±0.5
Turkey	2.0	5.8±1.1
Buffalo	2.0	1.0±0.5
Equine	11.0	17.7±3.2
Red Deer	4.0	2.3±0.1
Sheep 1	31.6	30.8±1.4
Sheep 2	15.8	21.0±1.3

Table 6. Blind samples composition, expressed as % (w/w) of each species on the total amount of meat. Symbol √ means “detected above LOD”.

	Recipes (meat % on finished product)					Results				
	A	B	C	D	E	A	B	C	D	E
Duck	2.00		1.58			√	√			
Rabbit		2.00	1.58				√	√		
Chicken	2.00		1.58	2.00		√	√	√		
Turkey		2.00	1.58				√	√		
Buffalo		2.00	1.58				√	√		
Equine	2.00		1.58		2.00	√	√			√
Deer	2.00		1.58			√	√			
Sheep	2.00		1.58			√	√			

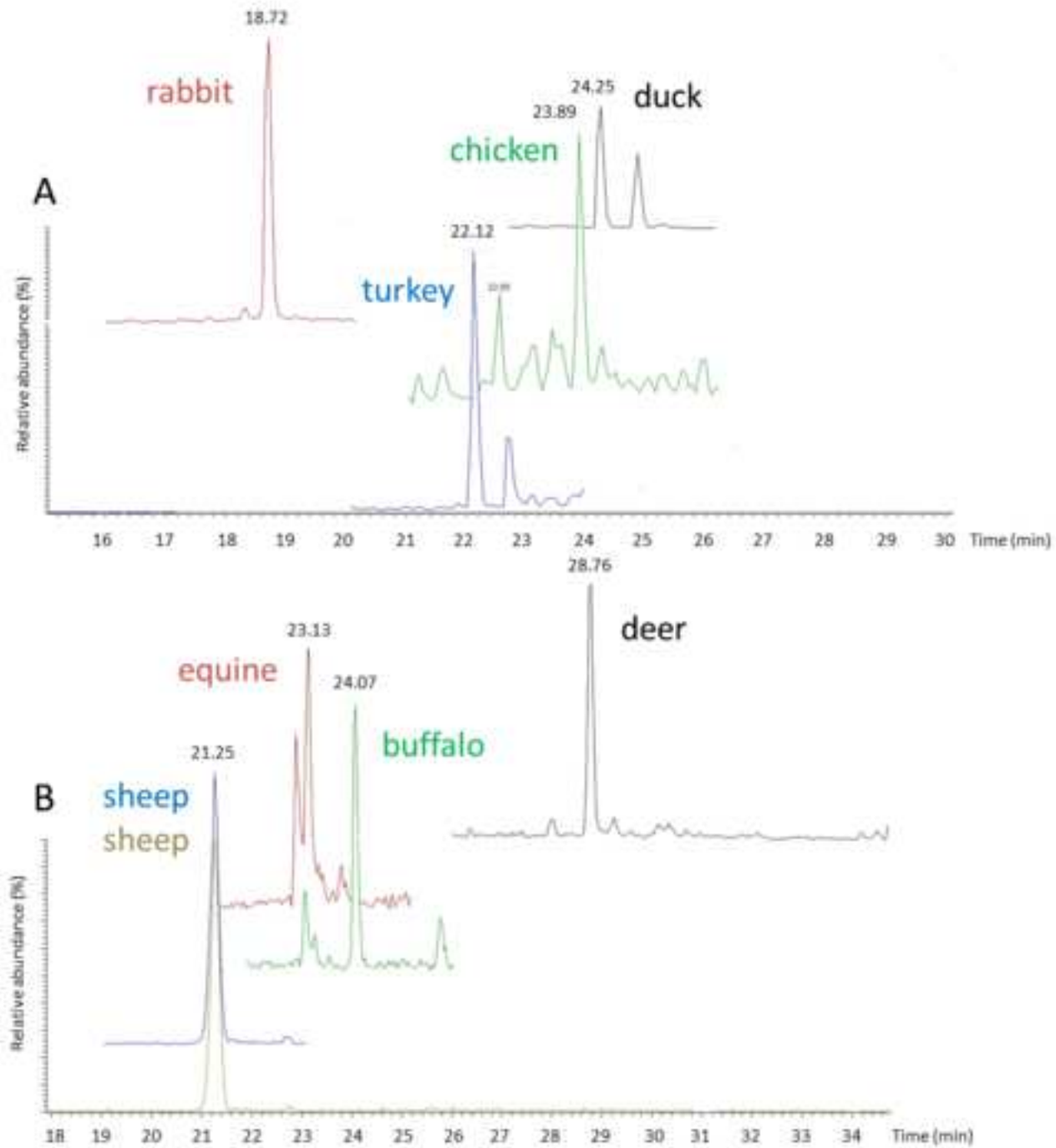
Figure

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Figure

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Figure

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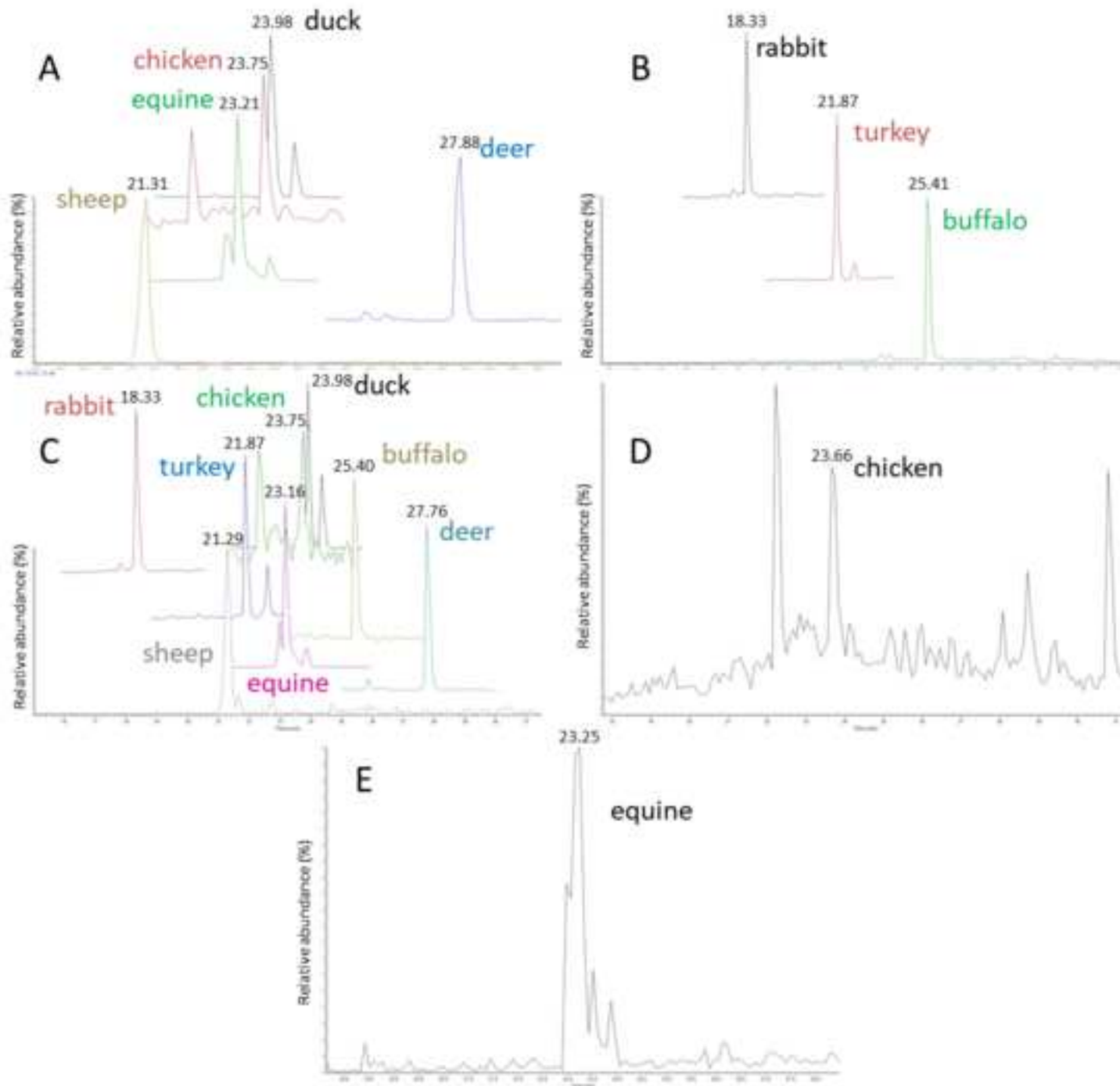
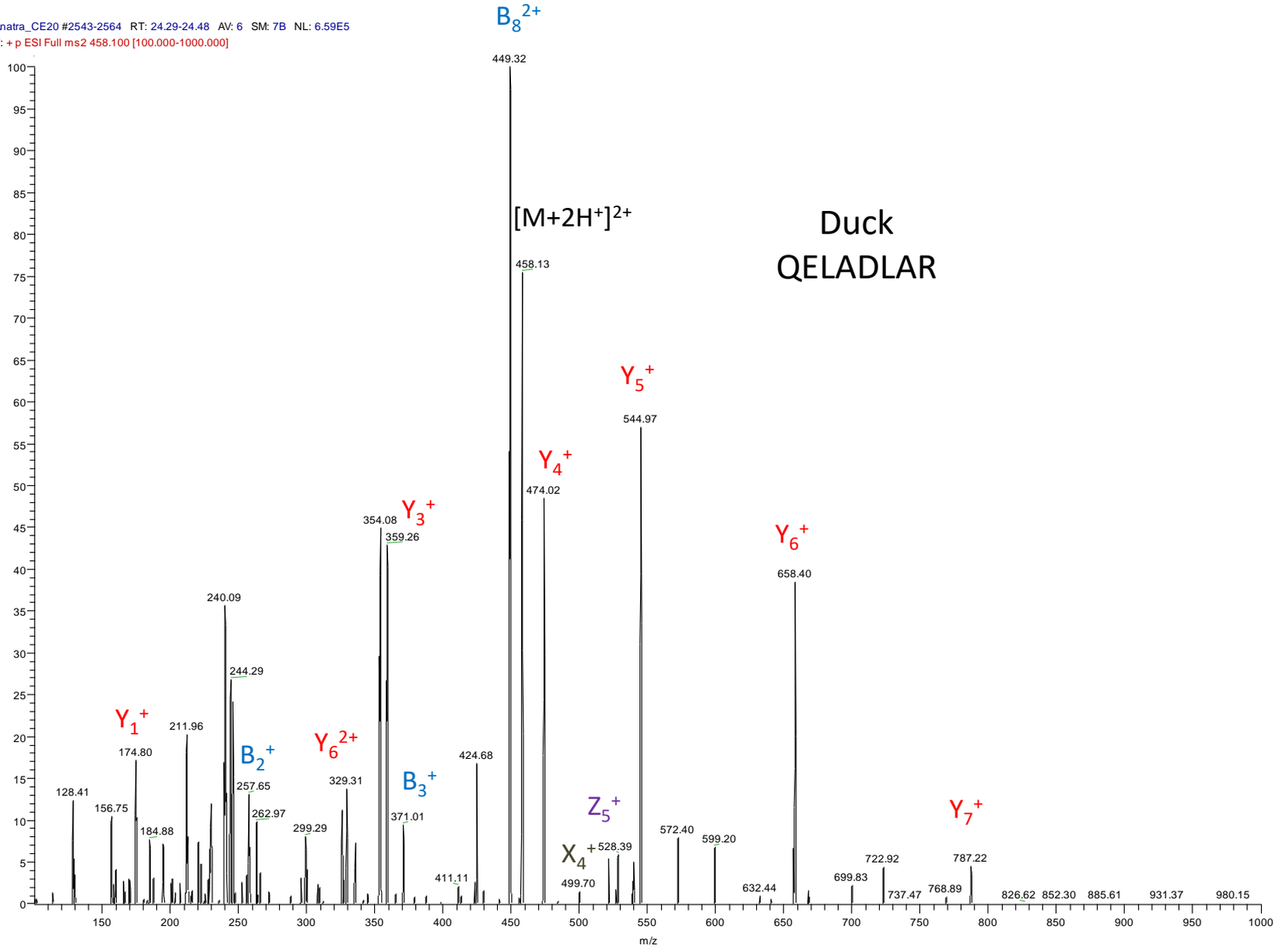


Table S1. Main proteins identified through bottom up proteomics in the different Bolognese sauce samples.

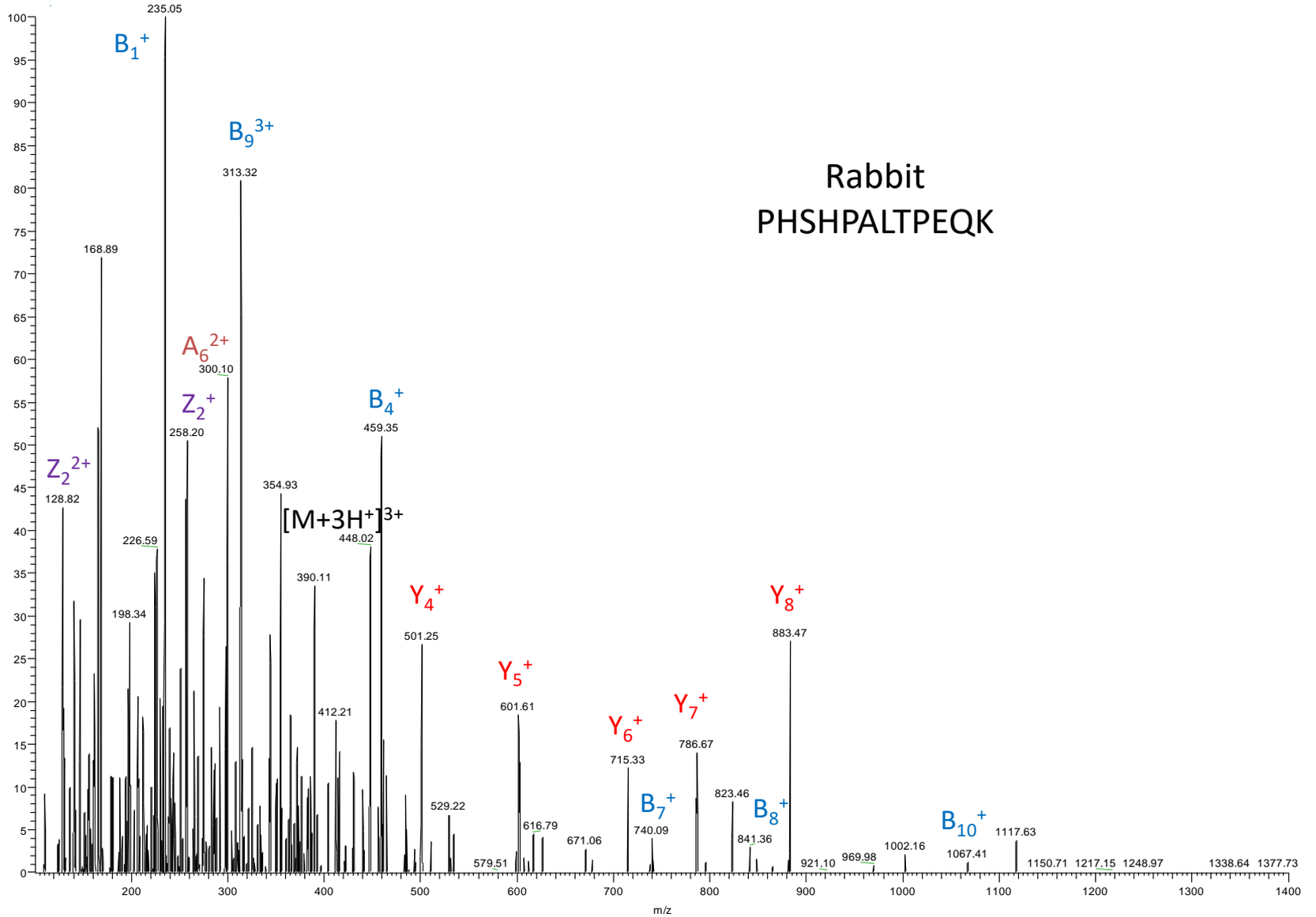
Accession	Description	Score	Coverage	MW [kDa]
<i>Duck (Anas platyrhynchos)</i>				
U3IW58	Uncharacterized protein	86	22	128.9
U3J7T0	Uncharacterized protein	81	10	216.0
U3J9U3	Uncharacterized protein	59	15	88.0
R0KZK3	Tropomyosin alpha-1 chain	50	13	23.6
<i>Rabbit (Oryctolagus cuniculus)</i>				
G1SJM7	Myosin-4	220	16	223.1
G1TKS9	Uncharacterized protein	196	13	223.5
L7N1T5	Uncharacterized protein	182	10	223.4
G1T4A5	Collagen alpha-1(I) chain	181	40	109.7
<i>Chicken (Gallus gallus)</i>				
P13538	Myosin heavy chain, skeletal muscle	111	13	223.0
P02457	Collagen alpha-1(I) chain	75	17	137.7
F1NM23	Tropomyosin alpha-1 chain	73	21	31.4
F1POH9	Collagen alpha-2(I) chain	48	21	128.8
<i>Turkey (Meleagris gallopavo)</i>				
F1P3X1	Uncharacterized protein	113	8	223.3
P02457	Collagen alpha-1(I) chain	101	23	137.7
G1NB83	Uncharacterized protein	64	16	128.0
F1NM23	Tropomyosin alpha-1 chain	64	23	31.4
<i>Donkey (Equus asinus)</i>				
B9VR88	Collagen alpha-1 type I chain	446	50	139.0
B9VR89	Collagen alpha-2 type I chain	350	50	128.7
P01959	Hemoglobin subunit alpha	131	29	15.2
D1MPT0	Hemoglobin beta chain	110	9	16.0
<i>Buffalo (Bubalus bubalis)</i>				
Q9TSN8	Hemoglobin subunit alpha-2	343	56	15.2
P84997	Myoglobin	312	53	17.2
Q0PEU1	Alpha 2 actin	250	35	20.1
A1XSX3	Beta-actin	230	36	41.7
<i>Horse (Equus caballus)</i>				
F6R839	Myosin-2	386	19	223.0
Q8MJV0	Myosin-1	386	19	222.9
Q8MJV1	Myosin-2	360	15	222.7
F6UP52	Uncharacterized protein	332	11	223.1
<i>Red deer (Cervus elaphus)</i>				
C0HJR0	Myoglobin	268	56	17.1
Q4TU70	Alpha hemoglobin chain	259	73	15.2
P01971	Hemoglobin subunit alpha	242	80	15.2
A0S012	Beta-actin	219	36	25.7
<i>Sheep (Ovis aries)</i>				
W5PPG6	Uncharacterized protein	257	12	222.8
W5PT09	Myosin heavy chain 2	240	10	222.4
W5NTT7	Collagen type I alpha 2 chain	223	19	129.0
A0A0H3V7A0	Myosin light chain 1 transcript variant 2	208	56	20.9
<i>Goat (Capra hircus)</i>				
F2X909	Collagen type I alpha 2 (Fragment)	228	84	13.6
F6KVT2	Troponin C type 1 slow	180	32	18.4
F6KVT3	Fast twitch skeletal muscle troponin C2	156	15	18.1
J9V335	Beta-actin (Fragment)	137	26	18.3

Supplementary Figure

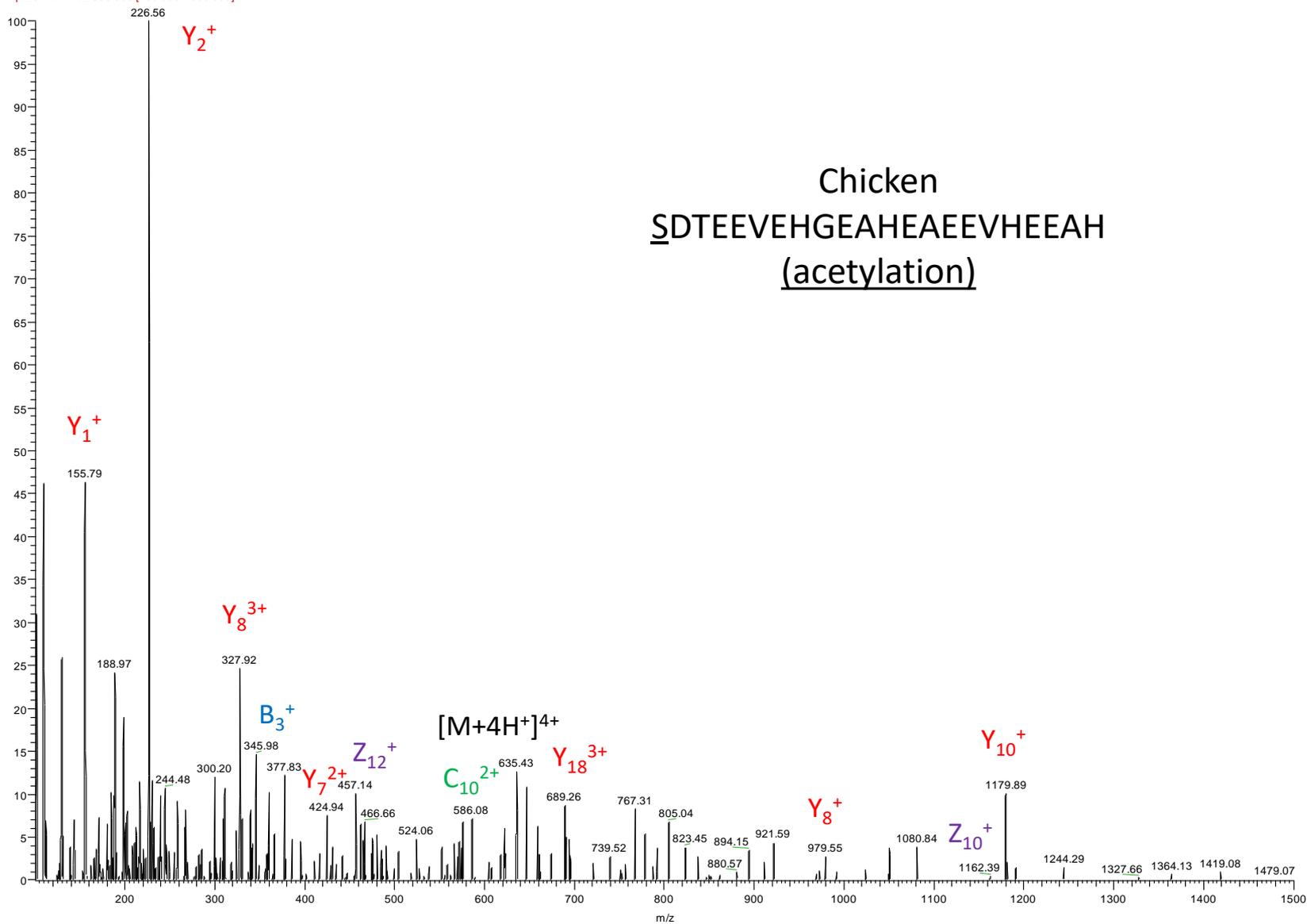
Anatra_CE20 #2543-2564 RT: 24.29-24.48 AV: 6 SM: 7B NL: 6.59E5
F: +p ESI Full ms2 458.100 [100.000-1000.000]



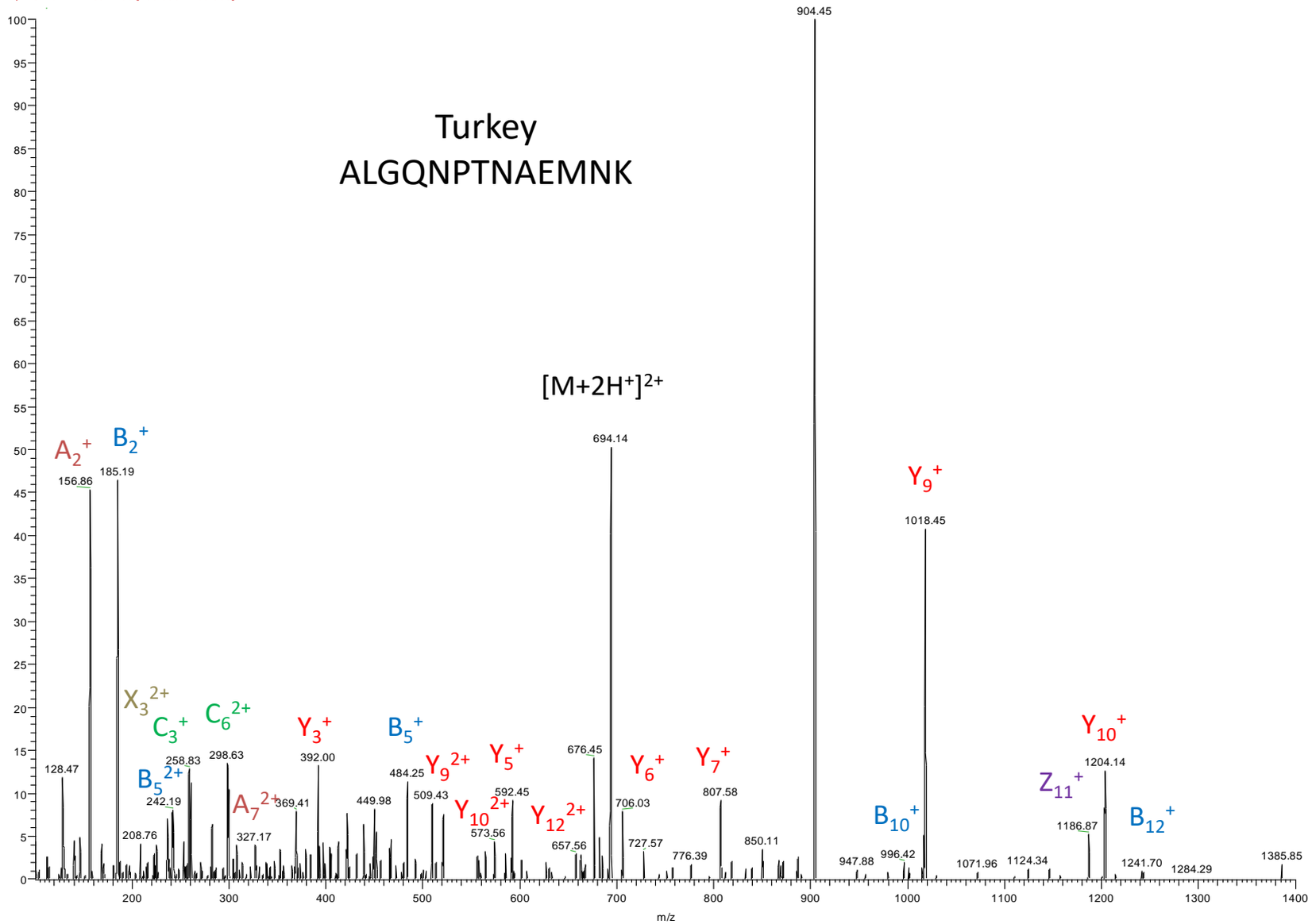
Rabbit PHSHPALTPEQK

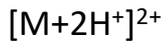


Chicken SDTEEVHGEAHEAEEVHEEAH (acetylation)

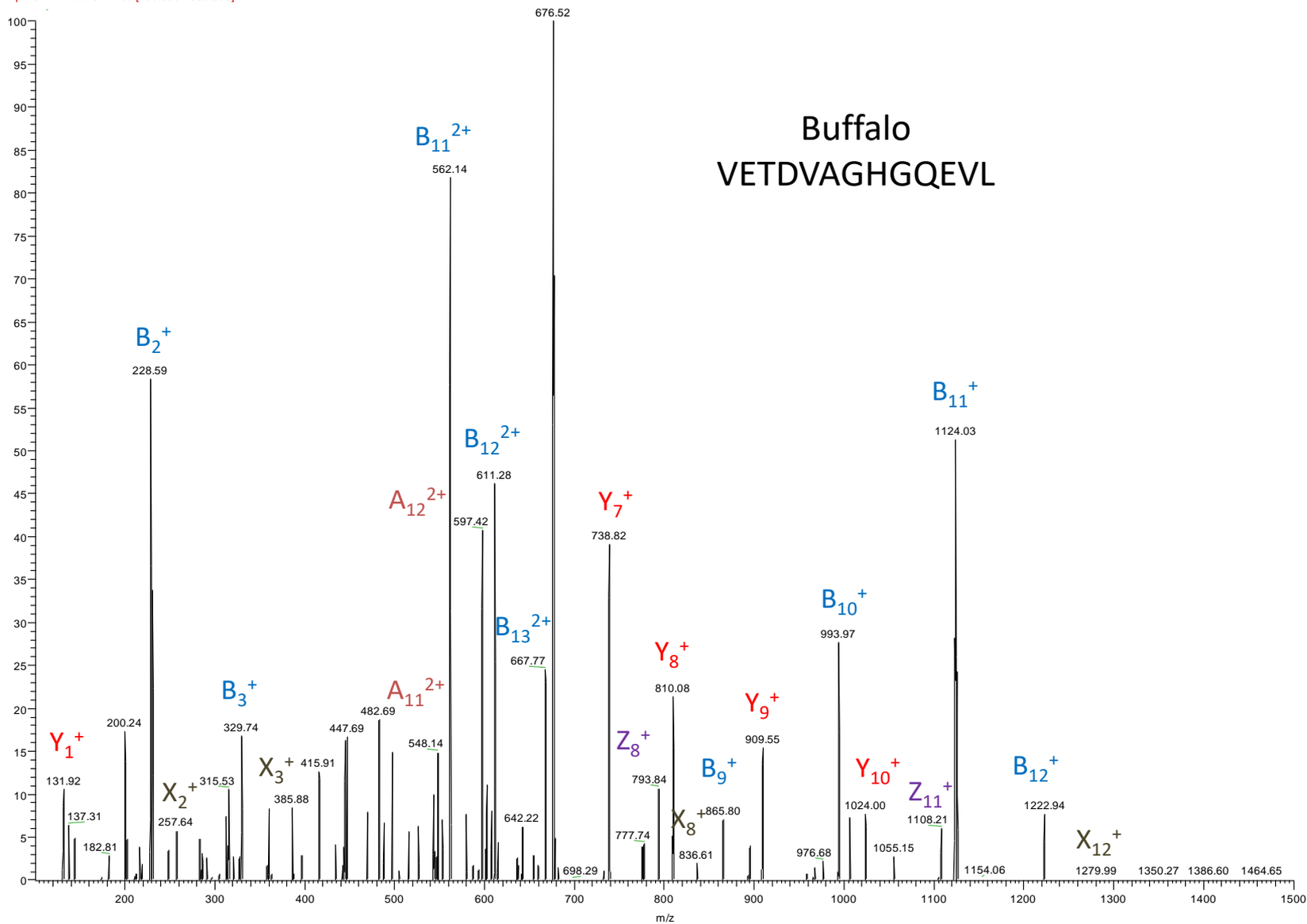


Turkey ALGQNPTNAEMNK

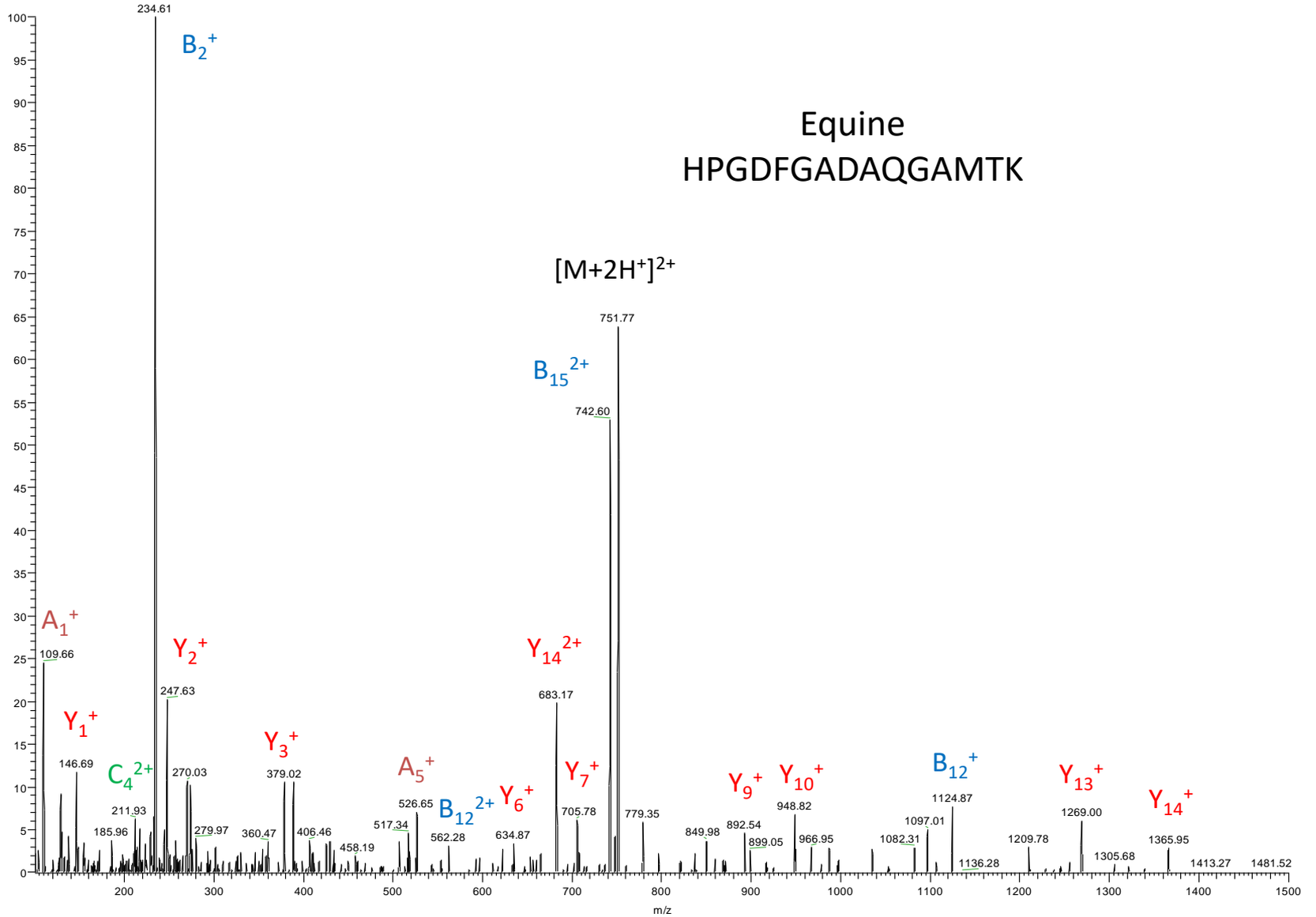




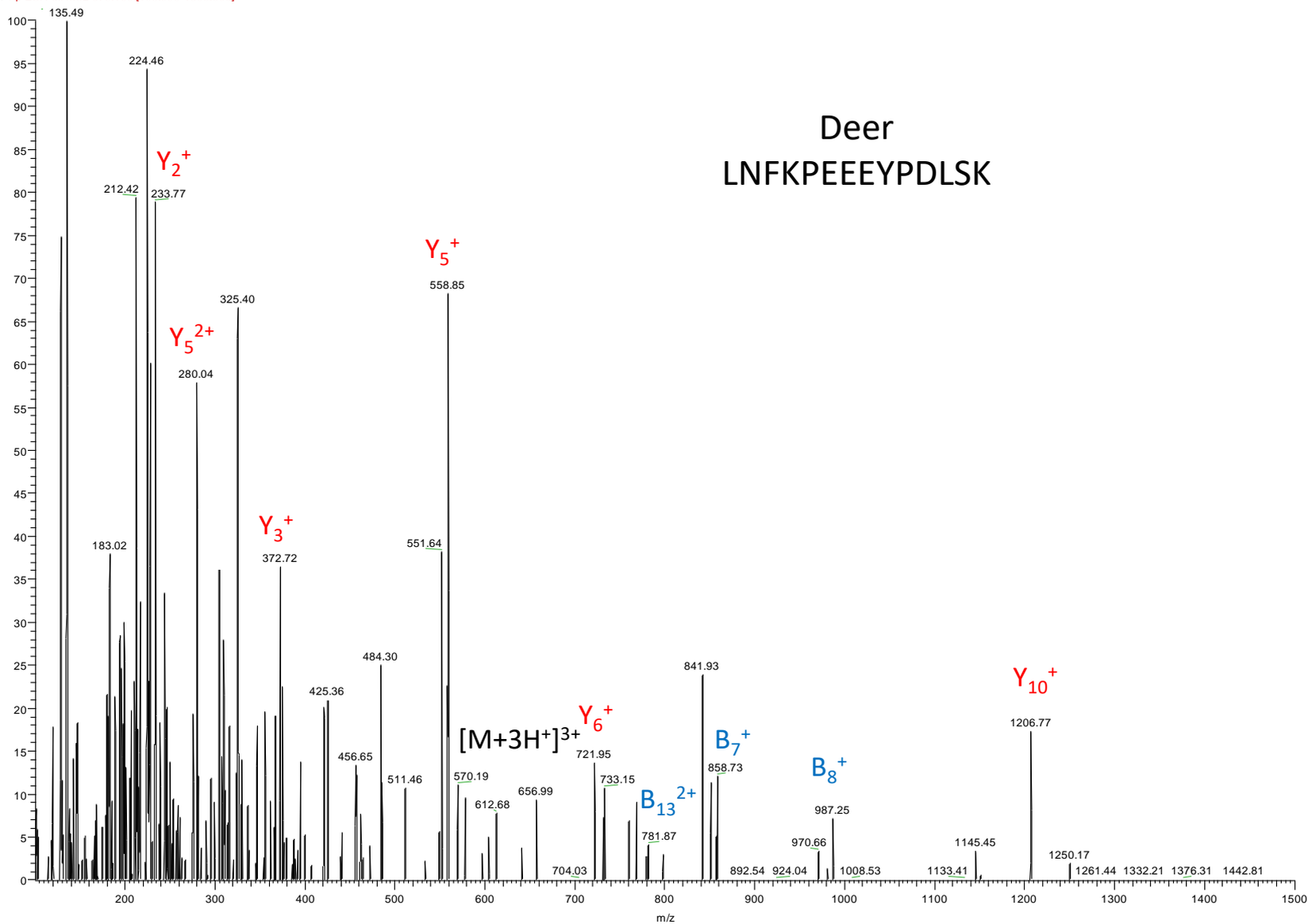
Buffalo VETDVAGHGQEVL



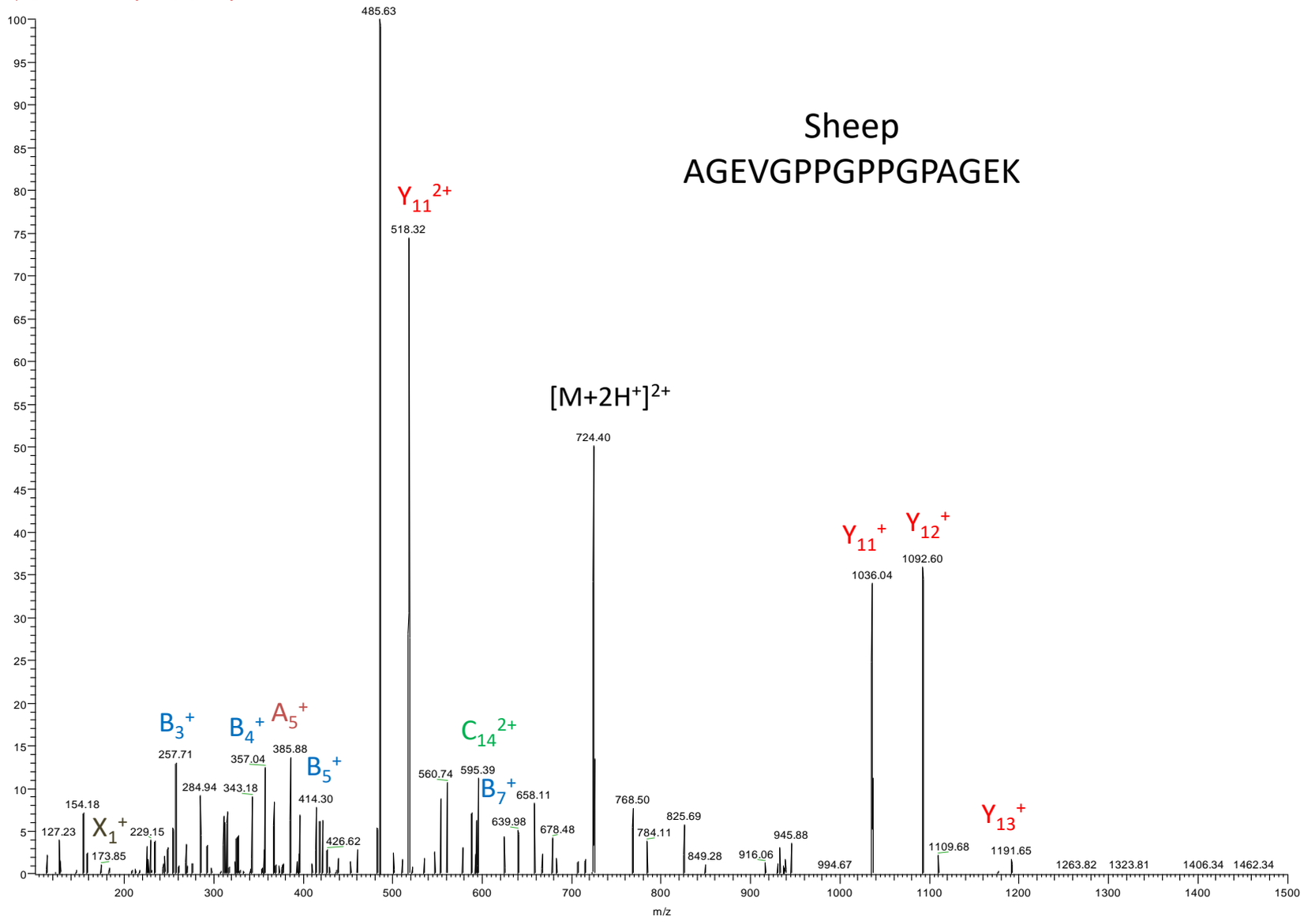
Equine HPGDFGADAQGAMTK



Deer LNFKPEEEYPDLISK



Sheep AGEVGPPGPPGPAGEK



HIGHLIGHTS

- Species-specific marker peptides were identified for eight meat species
- A LC-MS based method was developed for the detection of the marker peptides
- Bolognese sauces were prepared with increasing amount of each species
- Using proper calibration curves the method can quantify the meat species present
- A validation was performed analyzing blind Bolognese sauce samples