## 1 Actin proteolysis during ripening of dry fermented sausages at different pH values

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## 26 ABSTRACT

27 In dry fermented sausages, myofibrillar proteins undergo intense proteolysis generating small peptides and free amino acids that play a role in flavour generation. This study aimed to identify 28 29 small peptides arising from actin proteolysis, as influenced by the type of processing. Therefore, two acidification profiles were imposed by adding a different dose of dextrose, in order to mimic 30 the pH normally obtained in southern-type ("high pH"; pH 5.2 after fermentation and pH 6.0 after 31 ripening) and northern-type ("low pH"; pH 4.9 after fermentation and pH 5.0 after ripening) dry 32 fermented sausages. The identification of peptides within the raw meat was done by liquid 33 chromatography coupled to mass spectrometry in a data-independent positive mode of acquisition 34 (LCMS<sup>E</sup>), both after fermentation (4 days) and at the end of ripening (28 days). During 35 manufacturing of the dry fermented sausages, actin was highly proteolysed, especially in nine 36 regions of the sequence. After fermentation, 52 and 42 actin-derived peptides were identified at 37 38 high and low pH, respectively, which further increased to 66 and 144 peptides, respectively, at the end of ripening. Looking at the cleavage sites, cathepsins B and D likely played an important role. 39 40 The activity of exopeptidases was also evident since many peptides differed by a single amino acid.

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42 Keywords: actin, proteolysis, dry fermented sausages, mass spectrometry.

#### 44 1. INTRODUCTION

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Dry fermented sausages are processed meat products that undergo a fermentation and drying period 46 before consumption. During processing, quality-affecting physico-chemical changes occur of which 47 several are driven by proteolysis (Spaziani et al., 2009; Ordóñez et al., 1999). Proteolysis is carried 48 out by enzymes that are endogenous to the meat as well as by enzymes from microbial origin 49 50 (Molly et al., 1997). This process is of fundamental importance because it influences the flavour of dry fermented sausages (Ordóñez et al., 1999). The resulting small peptides and amino acids not 51 only directly influence taste but also serve as substrates for microorganisms that further convert 52 53 them into flavour compounds, as is the case for coagulase-negative staphylococci (Stavropoulou et al., 2015; Sánchez Mainar et al., 2016). 54

Proteolysis in dry fermented sausages is usually studied by determining protein and non-55 56 protein nitrogen (Defernando at al., 1991) and free  $\alpha$ -NH<sub>2</sub>-N (Verplaetse et al., 1992), by electrophoresis techniques (Hughes et al., 2002), and by measuring the release of amino acids 57 during ripening (Defernando at al., 1991). In addition, the use of mass spectrometry-based 58 proteomic techniques now also allows for the identification of peptides generated during ripening 59 60 (Mora et al., 2015). As a result, cleavage sites of proteolytic enzymes can be determined to better 61 unravel the proteolytic mechanisms at play during ripening. Proteomic approaches also have the potential to generate a more accurate view on the effect of processing factors on the dynamics of 62 proteolysis. It is for instance known that proteolytic enzymes have different pH optima (Hughes et 63 al., 1999, 2000). Therefore, the prevailing acidification profile during processing is expected to 64 affect proteolysis and thus flavour formation. Dry fermented sausages can be roughly divided in 65 two groups with different acidity levels, namely northern-type and southern-type products (Ravyts 66 et al., 2012). In the northern-type, the pH drops below 5.0 during fermentation and stays more or 67 less at that level throughout ripening. On the contrary, in the southern-type, the pH drops only 68

moderately during fermentation and increases during the drying phase, resulting in a final pH
between 5.5 and 6.0 (Demeyer et al., 2000).

Since the pH drop during meat fermentation not only influences flavour by creating an acid taste but also through proteolysis, it is important to better understand the details of this mechanism. Therefore, the aim of this study was to identify the peptides originated from actin degradation, which is one of the most abundant proteins in meat, during ripening of dry fermented sausages at two different pH values.

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## 77 2. MATERIAL AND METHODS

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## 79 2.1. Dry fermented sausage preparation

Dry fermented sausages were prepared by mixing lean pork (70.5 %), pork backfat (27.0 %), 80 81 sodium chloride (2.5 %), sodium nitrate (0.015 %, m/m), sodium ascorbate (0.05 %, m/m), and a starter culture strain (Lactobacillus sakei CTC 494) originating from the culture collection of the 82 Research Group of Industrial Microbiology and Food Biotechnology (Vrije Universiteit Brussel, 83 Brussels, Belgium). Two pH treatments were installed by adding either 0.25 %, m/m (high-pH 84 treatment) or 0.7 %, m/m (low-pH treatment) of dextrose. The batter was stuffed into collagen 85 86 casings of 50 mm diameter (Naturin, Weinheim, Germany) and ripened for 28 days in a climate chamber. During the first four days, fermentation was performed at a temperature of 24 °C and a 87 relative humidity of 94 %. For the drying process, the temperature was dropped to 12 °C and 88 relative humidity was set at 94 % for the first 14 days and at 80 % for the last 10 days. Samples 89 were taken at days 0, 4 (end of fermentation), and 28 (end of ripening). The manufacturing 90 91 processes and sampling were repeated in triplicate, resulting in three independent replicates.

#### 93 2.2 pH and weight loss

In each manufacturing process, three randomly selected sausages per treatment were weighed and the pH was recorded after their preparation and during ripening. The pH was measured directly in the sausages [ISO 2917 (1999)] and the pH meter was calibrated in buffers of pH 4.0 and 7.0. Weight loss was expressed as a percentage of the initial weight and the mean of the three records was calculated.

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## 100 *2.3 Peptide extraction*

101 The peptide extraction was carried out as described by Mora et al. (2015). Briefly, peptides were 102 extracted in 0.01 N HCl and proteins were precipitated by addition of EtOH. Finally, the peptide 103 extract was dried in a rotary evaporator and the peptides were dissolved in 25 ml 0.01 N HCl.

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## 105 *2.4 Size-exclusion chromatography*

Size-exclusion chromatography was carried out to select peptides between 500 and 4000 Da. Peptides were fractionated using an Akta Purifier (GE Healthcare Life Sciences, Uppsala, Sweden) on a Sephadex G25 Fine column ( $2.6 \times 60$  cm) and 0.01 N HCl was used as mobile phase at a flow rate of 1 mL/min. The fraction corresponding to elution volumes from 80 to 220 mL was collected and aliquots of 100 µL were lyophilised.

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# 112 $2.5 LCMS^E$ analysis

Lyophilised peptides were suspended in 40  $\mu$ l of 100 mM NH<sub>4</sub>HCO<sub>2</sub> (pH 10). Samples were filtered (0.22  $\mu$ m) and centrifuged. The peptide identification was done according to Devos et al. (2015) by liquid chromatography coupled to mass spectrometry in a data-independent positive mode of acquisition (LCMS<sup>E</sup>). Briefly, 2  $\mu$ l sample was injected on a NanoAcquity UPLC<sup>®</sup> system (Waters Corporation, Milford, MA, USA) for peptide separation. Solvent A1 and B1 were composed of 20 mM ammonium formate in water and acetonitrile (pH 10), respectively. Solvent A2 and B2 were composed of 0.1 % formic acid in water and 0.1 % formic acid in acetonitrile, respectively. The sample was loaded onto an Xbridge<sup>TM</sup> BEH130 C18 column (300  $\mu$ m × 50 mm, 5 µm; Waters) at 50 % solvent B1 at 2  $\mu$ L/min. Peptides were eluted and trapped on a Symmetry<sup>®</sup> C18 trapping column (180  $\mu$ m × 20 mm, 5  $\mu$ m; Waters Corporation) and finally separated on a HSS T3 C18 analytical column (75  $\mu$ m × 250 mm, 1.8  $\mu$ m; Waters Corporation) at 40°C at 250 nL/min by increasing the acetonitrile concentration from 5 to 50 % B2 over 60 min.

The outlet of the column was directly connected to a PicoTip Emitter (uncoated SilicaTip<sup>TM</sup> 10 ± 1 µm, New Objective, Woburn, MA, US) mounted on a Nanolockspray source of a SYNAPT<sup>TM</sup> G1 HDMS mass spectrometer (Waters Corporation). Accurate mass data were collected by alternating between low (5 V) and high (ramping from 15 to 35 V) energy scan functions (Geromanos et al., 2009). The selected m/z range was 125–2000 Da. The capillary voltage was set to 3.0 kV, the sampling cone voltage was 26 V, and the extraction cone voltage was 2.65 V. The source temperature was set at 80°C.

The LCMS<sup>E</sup> data were processed using the ProteinLynx Global SERVER<sup>™</sup> v2.5 (PLGS, 132 Waters Corporation) (Geromanos et al., 2009). The identification of actin peptides was done by 133 using a Uniprot database containing 753 protein entries from actin of the organism Sus scrofa 134 (downloaded from the Uniprot website, March 2016). The primary digest reagent was set as "none". 135 136 The precursor and fragment ion tolerance were determined automatically. The default protein identification criteria used included a detection of minimal three fragment ions per peptide, and 137 minimal three fragment ions per protein. Methionine oxidation was selected as a variable 138 modification. A false positive rate of 4 % was allowed. 139

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## 141 2.6 Identification of peptides cleaved by cathepsins B and D

To identify the actin-derived peptides that were likely generated by the action of cathepsins B and D
during meat fermentation, the previously determined cleavage sites of these enzymes on bovine Factin, were used (Hughes et al., 1999, 2000).

## 146 **3. RESULTS**

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## 148 *3.1 pH and weight loss*

The initial pH in the raw meat was 5.69 and 5.68 for the high-pH and low-pH treatments respectively. During the fermentation phase, the pH decreased to 5.23 ( $\pm$  0.02) and 4.89 ( $\pm$  0.01) in the high-pH and low-pH treatments, respectively. In the high-pH treatment, the pH increased throughout the drying phase to a final pH of 5.98 ( $\pm$  0.07) on day 28. On the contrary, in the low-pH treatment, the pH decreased to a minimum of 4.76 ( $\pm$  0.01) on day 15 and subsequently increased to a final pH of 4.96 ( $\pm$  0.01) at the end of ripening on day 28.

Weight loss showed similar trends in both treatments. Indeed, all samples lost about 2 % and 30 % of the initial weight at the end of fermentation and ripening, respectively.

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## 158 *3.2 Peptides identified from actin degradation*

The extracted peptides from the raw meat and the sausages at the end of fermentation and ripening 159 were subjected to identification by LCMS<sup>E</sup>. In the raw meat, no peptides arising from actin were 160 found. At the end of fermentation, however, 52 and 42 peptides were identified in the high-pH and 161 162 low-pH sausages, respectively (Tables 1 and 2). At this stage, both sausage types showed 37 peptides in common (highlighted in bold in the tables). At the end of ripening, the number of 163 peptides identified increased to 66 in the high-pH sausages (Table 3). In the low-pH sausages, 144 164 165 peptides were identified, of which 47 peptides were also found in the high-pH variant (Table 4). The peptides identified covered more than 50 % of the entire actin sequence and were generated 166 from all four actin subdomains, in which nine actin regions containing the majority of the identified 167 peptides are highlighted in Figure 1. Most peptides were identified in the regions 1 and 9, which are 168 near the N- and C-terminus and are located in the first subdomain together with the regions 3, 4 and 169 5 (region 1 is also partially lying in the second subdomain). Regions 2 and 7 were located in the 170

second and fourth subdomain, respectively. Finally, in the third subdomain, peptides were mainly
released from regions 6 and 8. On average, more peptides were identified at the end of ripening.
However, in the regions 6 and 9, similar amounts of peptides were identified at both days.

The identified peptides were mapped on to the 3D actin structure (Figure 2). Although after the fermentation phase the number of peptides identified in the two pH treatments was similar, some peptides (in red) were identified only at high pH (Figure 2A). On the contrary, at the end of ripening, the higher number of peptides identified in the low-pH compared to the high-pH treatment is reflected in the higher number of blue zones in the 3D structure (Figure 2B). However, the majority of peptides were identified in both treatments and are highlighted in yellow.

180 Based on the known specific cleavage sites of cathepsins B and D (Figure 1), an analysis of 181 the proteolytic breakdown by the latter enzymes was done. At the end of fermentation, 19 and 17 peptides (on a total of 52 and 42) were cleaved off either from the N or C-terminus, at high and low 182 183 pH, respectively (Tables 1 and 2). Throughout ripening, cleavage site analysis indicated that cathepsins B and D remained active leading to 29 and 67 identified peptides (on a total of 66 and 184 144) at high and low pH, respectively. The peptides Thr<sub>108</sub>-Lys<sub>120</sub> and Ile<sub>359</sub>-Lys<sub>375</sub> that were present 185 in the low-pH samples were likely generated by cathepsin D. In addition, cathepsin D was probably 186 responsible for the following cleavage sites: Cys12-Asp13, Thr79-Asn80, Thr91-Phe92, Ile153-Val154, 187 188 Leu155-Asp156, Lys240-Ser241, Trp358-Ile359, Thr91-Phe92, His103-Pro104, and Lys120-Met121. According to the known cleavage sites for cathepsin B, this enzyme likely cleaved actin at the following sites: 189 Ala24-Gly25, Thr68-Leu69, Met49-Gly50, His75-Gly76, Gly76-Ile77, Lys86-Ile87, Thr105-Leu106, Thr251-190 191 Ile<sub>252</sub>, Lys<sub>330</sub>-Ile<sub>331</sub>, and Glu<sub>363</sub>-Tyr<sub>364</sub>. Because of the occurrence of some common cleavage sites by both cathepsins, it was not possible to denote which of both enzymes mainly cleaved at the sites 192 193 Gly22-Phe23, Arg30-Ala31, Phe92-Tyr93, Arg97-Val98, and Leu107-Thr108.

A lot of identified peptides were truncated differing only by a single amino acid. In particular for the peptides  $Ile_{153}$ -Gly<sub>170</sub>, Tyr<sub>242</sub>-Arg<sub>258</sub>, and Trp<sub>358</sub>-His<sub>373</sub>, single amino acids were cleaved after already four days of fermentation and smaller peptides were generated in both high-

pH and low-pH sausages. In addition to the peptides reported above for the fermentation phase, the 197 198 peptides Met<sub>49</sub>-Gly<sub>65</sub> and Ser<sub>237</sub>-Arg<sub>256</sub> (at high and low pH) as well as Asp<sub>13</sub>-Ala<sub>31</sub> and Phe<sub>23</sub>-Gly<sub>44</sub> (at low pH only) were intensively hydrolysed at the end of the ripening phase. Similarly, several 199 peptides were truncated differing by two or three amino acids, indicating the release of dipeptides 200 and tripeptides. In the final products, the dipeptides Thr-Lys, Tyr-Ala, and Met-Gln were released 201 in the high-pH sausages. The dipeptides and tripeptides Gln-Gly, Arg-His, Val-Phe, Met-Gln, Gly-202 203 Ser-Gly, Leu-Tyr-Arg, Ile-Val-Gly, Ile-Val-His, and Ile-Leu-Thr were released in the low-pH 204 sausages.

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#### 206 4. DISCUSSION

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Since actin is one of the most abundant proteins in muscle (Dominguez and Holmes, 2011), its 208 209 degradation during ripening of dry fermented sausages is of interest for flavour formation. Breakdown of actin during ripening has been demonstrated previously, usually via electrophoresis 210 techniques (Hughes et al., 2002). Yet, more detailed information on its degradation patterns requires 211 state-of-the-art analysis via proteomic analysis. In the present paper, the small peptides generated 212 from actin hydrolysis were identified by LCMS<sup>E</sup>. The results showed that a high degree of 213 214 proteolysis took place since the identified peptides covered more than 50 % of the actin sequence and peptides were identified from all four subdomains. In a recent study, Lopez et al. (2015) 215 identified some peptides arising from actin in fermented sausage models. Several peptides identified 216 217 from that group originated from the regions 1, 3, and 7 that were also highlighted in the present study, confirming that those regions are susceptible to proteolysis. 218

It is evident that actin proteolysis started already during the fermentation phase: whereas no actin peptides were found in the raw meat, a more or less comparable peptide generation became already clear after four days of fermentation for both types of sausages. After 28 days of ripening, however, the number of peptides identified was more than twofold higher in the low-pH sausages.

In contrast to the rather brief fermentation step, ripening lasted for a longer time and led to more 223 224 pronounced pH differences (pH 4.9 versus pH 6.0 at the end of ripening). Demeyer et al. (2000) 225 also reported lower actin degradation in sausages with higher pH and this might be due to the low optimum pH of cathepsin D like muscle enzymes, which play a major role in actin hydrolysis 226 (Molly et al., 1997). In dry fermented sausages, proteolysis is generally divided in two steps: firstly, 227 endopeptidases break down intact proteins generating small peptides; secondly, the generated 228 229 peptides are further degraded by exopeptidases which release single amino acids, dipeptides, and tripeptides. The lysosomal cathepsins B and D are believed to be the main endopeptidases 230 responsible for the first protein breakdown in dry fermented sausages (Molly et al., 1997). The 231 232 identification of the cleavage sites of cathepsins B and D on bovine actin, which has the same sequence of pig actin, by Hughes et al. (1999, 2000) allowed to understand which peptides were 233 likely generated by these enzymes. Indeed, some of these cleavage sites were also the starting point 234 235 of several peptides identified at day 4 in the present study, supporting the finding that cathepsins are already likely active during the first days of fermentation (Demeyer et al., 1992). The pH at the end 236 of fermentation was in the activity range of cathepsin B and D in both treatments. Indeed, Schwartz 237 and Bird (1977) reported that rabbit actin was degraded by cathepsins B and D at pH 5.0. These 238 239 enzymes remain stable and active for several months during processing of dry cured meat products 240 (Toldrà et al., 1993) and it is not surprising that the number of peptides likely generated by cathepsin B and D increased at the end of ripening in the present study. This was particularly 241 evident in the low-pH treatment were the number of identified peptides greatly increased at the end 242 243 of ripening. In the high-pH sausages, the contribution of these enzymes was more limited during the drying phase since their activity is very low at pH 6.0 (Schwartz and Bird, 1977). Although for 244 245 some peptides the contribution of cathepsins B and D seems indisputable, this might be doubtful for other peptides. Indeed, in some cases the same peptide but with an additional residue after the 246 cathepsin cleavage site was also identified. At this point, it is impossible to know whether the 247

shorter peptide was the result of endopeptidase activity or if it was generated from the longerpeptide by hydrolysis of the additional residue made by an exopeptidase.

The further degradation of peptides exerted by exopeptidases releases amino acids, 250 dipeptides, and tripeptides from the N- and C-terminus. These enzymes, in dry fermented sausages, 251 are either of muscle or microbial origin (Mora et al., 2015). In the present study, the activity of 252 aminopeptidases and carboxypeptidases was evident since a lot of identified peptides were 253 254 truncated differing by a single amino acid. There are five aminopeptidases known to be active postmortem: leucyl, arginyl, alanyl, pyroglutamyl, and methionyl aminopeptidases (Toldrá, 2006). Ile, 255 Leu, Tyr, and Gly were probably released by alanyl aminopeptidase (Flores et al., 1996). Methionyl 256 257 aminopeptidase mainly cleaves Lys, Ala and Leu (Flores et al., 2000) and arginyl aminopeptidase cleaves basic amino acids (Flores et al. 1993). These aminopeptidases have optimum activities at 258 neutral pH and only retain some activity at pH 5.0 (Toldrá, 2006). Nevertheless, the action of 259 260 exopeptidases generated more unique peptides at low pH. We suggest that some endopeptidases like cathepsins, which have optimum activity at very acidic pH, provided more substrates to 261 exopeptidases at low pH. From the C-terminus, the activity of carboxypeptidases was also evident, 262 although knowledge about these enzymes in meat products is still limited. There are two lysosomal 263 carboxypeptidases (A and B) known to have optimal activity at acidic pH, with the former cleaving 264 265 hydrophobic amino acids and the latter having a wider activity (Toldrá, 2006). Several peptides were also truncated differing by two or three amino acids, indicating that dipeptidyl, such as DPP I 266 and DPP II (Sentandreu and Toldrá, 2000, 2001), and tripeptidyl peptidases were active during 267 268 ripening.

In addition to the above, the contribution of microbial enzymes cannot be ignored. In the present study, all sausages were inoculated with *L. sakei*. This species, which has often been isolated from spontaneously fermented dry fermented sausages, is often used as starter culture for its high competitiveness and ability to produce antimicrobial compounds with strong antilisterial activity (Leroy and De Vuyst, 2005; Ravyts et al., 2012). Moreover, the background microbiota that is usually constituted of coagulase-negative microbiota may also be relevant for proteolytic activity,
as well as for the further conversion of amino acids in aroma compounds (Sánchez Mainar et al.,
2016). It is generally believed that meat endogenous enzymes initiate proteolysis and the
contribution of microbial enzymes is relevant only in a later stage of ripening (Hughes et al., 2002;
Molly et al., 1997). The species *L. sakei* has been shown to possess some proteolytic activity
(Candogan and Acton, 2004), which was in particular reported to release Leu and Ala from peptides
(Sanz and Toldrá, 1997).

Peptides, especially the ones with a low molecular mass < 5000 Da, are potential flavour 281 compounds and flavour precursors. The taste of peptides depends on the conformational 282 283 characteristics and on the amino acid composition. Charges on the side chains or on the terminals can give salty or sour taste, hydrophobic side chains can give bitter taste and umami taste was 284 reported for some small peptides in meat products (Temussi, 2011). Nevertheless, there are only 285 286 few studies that investigated the influence of peptides on the taste of meat products. Henriksen and Stahnke (1997) evaluated the effect of fractions of small peptides and free amino acids extracted 287 from dry fermented sausages on taste. The results suggested that bitterness and sourness were 288 correlated with hydrophobic amino acids and glutamic acid, respectively, and bouillon taste was 289 290 provided by a mixture of different peptides and amino acids. The results of the present study 291 revealed that numerous peptides of low molecular weight are generated by actin degradation, especially at low pH conditions. Therefore, actin might be a key protein in determining the final 292 taste characteristics in dry fermented sausages considering the high abundance of this protein. 293

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#### 295 5. CONCLUSION

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In the present study, the generation of peptides from actin, which is one of the most abundant proteins in muscle, was studied during ripening of dry fermented sausages through LCMS<sup>E</sup>. The understanding of proteolysis is of great importance since small peptides and amino acids influence the sensory characteristics of these products. The results showed that actin is already being hydrolysed during the fermentation phase, but proteolysis is most intense during the drying phase. In addition, a more acid pH profile, characteristic of northern-type fermented sausages, generated more unique peptides. The relevance of the findings of the present study will need to be confirmed and contrasted with the analysis of proteolytic products of other major muscle proteins in future research.

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## 313 **Conflict of interest**

314 The authors declare that no competing interests exist.

#### 315

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# 400 Figure Captions

- **Figure 1.** Actin sequence and 3D structure. The majority of the identified peptides were lying in the
- 402 underlined regions.
- **Figure 2.** Peptides were mapped on the 3D structure. Peptides identified for high-pH and low-pH
- fermentations are shown in red and blue, respectively. Common peptides are shown in yellow.

- Table 1. Peptides of actin identified by LCMS<sup>E</sup> in high-pH dry fermented sausages after 4 days of 1
- ripening (fermentation phase) 2

N.	P <sub>1</sub> <sup>a</sup>	Peptide sequence	P1,b	P. s. s. <sup>c</sup>	zd	m/z <sup>e</sup>	Info
1	E	DETTALVCDNGSGLVK	A	4	2	811.3924	
2	С	DNGSGLVKAGFAGDDAPR	А	12	3	582.9503	CD <sup>f</sup> (N-ter), CDB <sup>h</sup> (C-ter)
3	ĸ	AGFAGDDAPR	A	20	2	488.7295	(,, (,
4	G	FAGDDAPRAVFPSIVG	R	22	2	809.9159	CBD <sup>h</sup> (N-ter)
5	G	FAGDDAPRAVFPS	I	22	2	675.3301	CBD <sup>h</sup> (N-ter)
6	D	SYVGDEAQSKRG	I	53	2	648.816	(,
7	Y	VGDEAQSKRG	Ī	55	2	523.7644	
8	F	YNELRVAPEEHPTL	L	92	3	556.6154	CBD <sup>h</sup> (N-ter)
9	F	YNELRVAPEE	H	92	2	610.3024	CBD <sup>h</sup> (N-ter)
10	L	RVAPEEHPTL	L	96	2	574.8065	022 (11 002)
11	R	VAPEEHPTL	L	97	2	496.7555	
12	A	PEEHPTL	L	99	2	411.7003	
13	Т	GIVLDSGDGVTHNVPIYEG	Y	151	3	647.9892	
14	G	IVLDSGDGVTHNVPIYEG	Ŷ	152	2	942.9739	
15	G	IVLDSGDGVTHNVPIYE	G	152	2	914.4625	
16	I	VLDSGDGVTHNVPIYEG	Y	153	2	886.4331	CD <sup>f</sup> (N-ter)
17	I	VLDSGDGVTHNVPITEG	G	153	2	857.9189	CD <sup>f</sup> (N-ter)
18	I	VLDSGDGVTHNVPITE	E	153	2	793.3958	CD <sup>f</sup> (N-ter)
19	I	VLDSGDGVTHNVP	L	153	2	655.3221	CD <sup>f</sup> (N-ter)
20	V	LDSGDGVTHNVPIYEG	Ϋ́	154	2	836.8954	CD (N CEI)
20	L	DSGDGVTHNVPIYEG	Y	155	2	780.3534	CD <sup>f</sup> (N-ter)
22	L	DSGDGVTHNVPIYE	G	155	2		CD <sup>f</sup> (N-ter)
23	L			155	2	751.843	
		DSGDGVTHNVPIY SGDGVTHNVPIYEG	E Y		2	687.3205	CD <sup>f</sup> (N-ter)
24	D			156		722.8397	
25	S	DGVTHNVPIYEG	Y	158	2	650.8137	
26	L	EKSYELPDGQVITIGN	E	238	2	881.946	
27	S	YELPDGQVITIGNERFR	С	241	3	669.6759	
28	S	YELPDGQVITIGNERF	R	241	2	925.9726	
29	S	YELPDGQVITIGNER	F	241	2	852.4363	$ODG \left( O + z \right)$
30	S	YELPDGQVIT	I	241	1	1134.5723	CB <sup>g</sup> (C-ter)
31	Y	ELPDGQVITIGNERF	R	242	2	844.4377	
32	Y	ELPDGQVITIGNER	F	242	2	770.9025	
33	М	SGGTTMYPGIADRMQ	K	301	2	792.862	
34	M	SGGTTMYPGIADR	M	301	2	663.312	
35	S	TFQQMWITKQEY	D	352	3	534.9266	
36	M	WITKQEYDEAGPSIVH	R	357	3	624.9725	
37	M	WITKQEYDEAGPS	I	357	2	762.3543	
38	W	ITKQEYDEAGPSIVH	R	358	3	562.9473	CD <sup>f</sup> (N-ter)
39	W	ITKQEYDEAGPS	I	358	2	669.3152	CD <sup>f</sup> (N-ter)
40	I	TKQEYDEAGPSIVH	R	359	2	787.3807	
41	K	QEYDEAGPSIVH	R	361	2	672.8068	
42	Q	EYDEAGPSIVHRK	С	362	2	750.8767	
43	Q	EYDEAGPSIVHR	K	362	2	686.8291	
44	Q	EYDEAGPSIVH	R	362	2	608.781	
45	E	YDEAGPSIVHRK	С	363	3	457.9011	CBg (N-ter)
46	Ε	YDEAGPSIVHR	K	363	2	622.3057	CB <sup>g</sup> (N-ter)
47	Ε	YDEAGPSIVH	R	363	2	544.2557	CB <sup>g</sup> (N-ter)
48	Y	DEAGPSIVHRK	С	364	2	604.824	
49	Y	DEAGPSIVHR	K	364	2	540.7746	
50	Y	DEAGPSIVH	R	364	2	462.7249	
51	Ε	AGPSIVHR	K	366	2	418.7418	
52	Ε	AGPSIVH	R	366	1	680.3765	

<sup>a</sup> Position of the amino acid residue preceding the peptide sequence (N-terminus). <sup>b</sup> Position of the amino acid residue following the peptide sequence (C-terminus).

<sup>c</sup> Peptide sequence start in actin.

5 6 <sup>d</sup> Charge (+).

- 7 <sup>e</sup> Mass to charge ratio.
- 8 <sup>f</sup> Cathepsin D cleavage site.
- <sup>g</sup> Cathepsin B cleavage site.
- 9 10 <sup>h</sup> Cathepsin B and D common cleavage site.

Peptides that were also found in low pH dry fermented sausages are indicted in bold.

- Table 2. Peptides of actin identified by LCMS<sup>E</sup> in low-pH dry fermented sausages after 4 days of 13
- ripening (fermentation phase) 14

N.	P <sub>1</sub> <sup>a</sup>	Peptide sequence	<b>P</b> 1, <sup>b</sup>	P. s. s. <sup>c</sup>	z <sup>d</sup>	m/z <sup>e</sup>	Info
1	G	FAGDDAPRAVFPSIVG	R	22	2	809.9204	CBD <sup>h</sup> (N-ter)
2	G	FAGDDAPRAVFPS	I	22	2	675.3318	
3	Н	GIITNWDDMEK	I	75	2	661.3103	
4	F	YNELRVAPEEHPTL	L	92	3	556.6168	CBD <sup>h</sup> (N-ter)
5	F	YNELRVAPEEHPT	L	92	3	518.9217	CBD <sup>h</sup> (N-ter)
6	F	YNELRVAPEE	Н	92	2	610.3063	CBD <sup>h</sup> (N-ter)
7	L	RVAPEEHPTL	L	96	2	574.8041	
8	A	PEEHPTL	L	99	2	411.7017	
9	G	IVLDSGDGVTHNVPIYEG	Y	152	2	942.9772	
10	I	VLDSGDGVTHNVPIYEG	Y	153	2	886.4335	CD <sup>f</sup> (N-ter)
11	I	VLDSGDGVTHNVPIYE	G	153	2	857.9228	CD <sup>f</sup> (N-ter)
12	I	VLDSGDGVTHNVPIY	Ε	153	2	793.3984	CD <sup>f</sup> (N-ter)
13	V	LDSGDGVTHNVPIYEG	Y	154	2	836.8956	
14	L	DSGDGVTHNVPIYEG	Y	155	2	780.3545	CD <sup>f</sup> (N-ter)
15	L	DSGDGVTHNVPIYE	G	155	2	751.8453	CD <sup>f</sup> (N-ter)
16	L	DSGDGVTHNVPIY	Ε	155	2	687.3256	CD <sup>f</sup> (N-ter)
17	D	SGDGVTHNVPIYEG	Y	156	2	722.8389	
18	G	DGVTHNVPIYEG	Y	158	2	650.8106	
19	G	YALPHAIM	R	170	2	458.2395	
20	S	YELPDGQVITIGNERFR	С	241	3	669.6783	
21	S	YELPDGQVITIGNERF	R	241	2	925.9741	
22	S	YELPDGQVITIGNER	F	241	2	852.4399	
23	Y	ELPDGQVITIGNERF	R	242	2	844.4383	
24	Y	ELPDGQVITIGNER	F	242	2	770.9013	
25	М	WITKQEYDEAGPSIVHRK	С	357	4	540.0309	
26	М	WITKQEYDEAGPSIVH	R	357	3	624.9791	
27	М	WITKQEYDEAGPS	I	357	2	762.3554	
28	W	ITKQEYDEAGPSIVH	R	358	3	562.9483	CD <sup>f</sup> (N-ter)
29	W	ITKQEYDEAGPS	I	358	2	669.3148	CD <sup>f</sup> (N-ter)
30	I	TKQEYDEAGPSIVH	R	359	2	787.3782	
31	K	QEYDEAGPSIVH	R	361	2	672.8107	
32	Q	EYDEAGPSIVHRK	С	362	2	750.8771	
33	Q	EYDEAGPSIVH	R	362	2	599.7724	
34	Ε	YDEAGPSIVHRK	С	363	3	457.9007	
35	Ε	YDEAGPSIVHR	K	363	2	622.3088	CB <sup>g</sup> (N-ter)
36	Ε	YDEAGPSIVH	R	363	2	544.2598	CB <sup>g</sup> (N-ter)
37	Y	DEAGPSIVHRK	С	364	2	604.8242	
38	Y	DEAGPSIVHR	Κ	364	2	540.7753	
39	Y	DEAGPSIVH	R	364	2	462.724	
40	D	EAGPSIVHRK	С	365	2	547.3041	
41	Ε	AGPSIVHR	K	366	2	418.7422	
42	Е	AGPSIVH	R	366	1	680.3748	

15 <sup>a</sup> Position of the amino acid residue preceding the peptide sequence (N-terminus).

16 <sup>b</sup> Position of the amino acid residue following the peptide sequence (C-terminus).

17 <sup>c</sup> Peptide sequence start in actin.

18 <sup>d</sup> Charge (+).

19 <sup>e</sup> Mass to charge ratio.

20 <sup>f</sup> Cathepsin D cleavage site.

21 <sup>g</sup> Cathepsin B cleavage site.

22 23 <sup>h</sup> Cathepsin B and D common cleavage site.

Peptides that were also found in high pH dry fermented sausages are indicted in bold.

- **Table 3.** Peptides of actin protein identified by LCMS<sup>E</sup> in high-pH dry fermented sausages after 28
- 25 days of ripening

. P1 <sup>a</sup> Peptide sequence	P1,b	P. s. s. <sup>c</sup>	zď	m/z <sup>e</sup>	Info
1 C DNGSGLVKAGFAGDD	А	12	2	711.8276	CD <sup>f</sup> (N-ter)
2 K AGFAGDDAPRAVFPSIVG	R	20	2	873.9467	
3 A GFAGDDAPRAVFPSIVG	R	21	2	838.4287	
4 G FAGDDAPRAVFPSIVG	R	22	2	809.9159	CBD <sup>h</sup> (N-ter)
5 F AGDDAPRAVFPSIVGRP	R	23	3	575.6369	
6 F AGDDAPRAVFPSIVGR	Р	23	3	543.2865	
7 F AGDDAPRAVFPSIVG	R	23	2	736.3813	
G MGQKDSYVGDEAQSKRG	I	48	3	619.2906	
9 G MGQKDSYVGDEAQ	S	48	2	714.3076	
M GQKDSYVGDEAQSKRG	I	49	3	575.6074	CB <sup>g</sup> (N-ter)
M GQKDSYVGDEAQSK	R	49	2	756.3522	CBg (N-ter)
2 M GQKDSYVGDEAQS	K	49	2	692.3051	CB <sup>g</sup> (N-ter)
3 M GQKDSYVGDEAQ	S	49	2	648.7893	CB <sup>g</sup> (N-ter)
4 Q KDSYVGDEAQSKR	G	51	3	494.9124	
5 H GIITNWDDMEK	I	75	2	661.3104	CB <sup>g</sup> (N-ter), CB <sup>g</sup> (C-ter)
6 G IITNWDDMEK	I	76	2	632.7996	CB <sup>g</sup> (N-ter), CB <sup>g</sup> (C-ter)
7 I ITNWDDMEK	I	77	2	576.2543	CB <sup>g</sup> (C-ter)
3 I TNWDDMEKIWHHT	F	78	3	571.587	CD <sup>f</sup> (N-ter), CD <sup>f</sup> (C-ter)
9 W DDMEKIWHHT	F	81	2	656.292	CD <sup>f</sup> (C-ter)
D F YNELRVAPEEHPTL	L	92	3	556.614	CBD <sup>h</sup> (N-ter)
1 F YNELRVAPEEHPT	L	92	3	518.9213	CBD <sup>h</sup> (N-ter), CB <sup>g</sup> (C-ter)
2 F YNELRVAPEE	Н	92	2	610.3028	
3 Y NELRVAPEEHPT	L	93	2	696.3501	CB <sup>g</sup> (C-ter)
4 L RVAPEEHPTL	L	96	2	574.8061	02 (0 001)
5 R VAPEEHPTLL	T	97	2	553.2978	CBD <sup>h</sup> (N-ter), CBD <sup>h</sup> (C-ter
6 L YASGRTTGIVLDSGDGVTHNVPIYEG	Ŷ	144	3	893.4371	CD <sup>f</sup> (N-ter)
7 T GIVLDSGDGVTHNVPIYEG	Y	151	2	971.4831	CD (IN CEL)
G IVLDSGDGVTHNVPIYEG	Y	151	2	942.9721	
9 I VLDSGDGVTHNVPIYEGYA	L	153	2	1003.4841	CD <sup>f</sup> (N-ter)
0 I VLDSGDGVTHNVPTTEGTA	Ч	153	2	886.4301	CD <sup>f</sup> (N-ter)
	G		2		CD <sup>f</sup> (N-ter)
		153		857.9226	
	I	153	2	655.323	CD <sup>f</sup> (N-ter)
3 V LDSGDGVTHNVPIYEG	Y	154	2	836.8967	ODf(N, h, r)
4 L DSGDGVTHNVPIYEG	Y	155	2	780.3542	CD <sup>f</sup> (N-ter)
5 L DSGDGVTHNVP	I	155	2	549.2497	CD <sup>f</sup> (N-ter)
6 D SGDGVTHNVPIYEG	Y	156	2	722.843	
7 S GDGVTHNVPIYEG	Y	157	2	679.3209	
B G DGVTHNVPIYEG	Y	158	2	650.8119	
9 S SLEKSYELPDGQVITIGNER	F	236	3	750.0474	
) S <b>SLEKSYELPDGQVIT</b>	I	236	2	839.9317	CB <sup>g</sup> (C-ter)
1 L EKSYELPDGQVITIGNER	F	238	3	683.345	
2 L EKSYELPDGQVITIGN	E	238	2	881.951	
3 E KSYELPDGQVITIGNER	F	239	2	959.9952	
4 E KSYELPDGQVITIGN	E	239	2	817.4255	
5 K SYELPDGQVITIGNERF	R	240	2	969.4918	CDf (N-ter)
6 K SYELPDGQVITIGNER	F	240	2	895.9536	CD <sup>f</sup> (N-ter)
7 Y ELPDGQVITIGNERF	R	242	2	844.4362	
BY ELPDGQVITIGNER	F	242	2	770.905	
9 I GMESAGIHETTYNS	I	269	2	748.8201	
) R KDLYANNVMSGGTTM	Y	292	2	801.3673	
1 M SGGTTMYPGIADRMQ	K	301	2	792.8641	
2 M SGGTTMYPGIADR	М	301	2	663.3132	
3 M SGGTTMYPGIAD	R	301	1	1169.5151	
4 W ITKQEYDEAGPSIVH	R	358	3	562.9504	CD <sup>f</sup> (N-ter)
5 W ITKQEYDEAGPS	I	358	2	669.3137	CD <sup>f</sup> (N-ter)
6 I TKQEYDEAGPSIVHRK	С	359	3	619.9816	
7 I TKQEYDEAGPSIVHR	K	359	3	577.2846	
B I TKQEYDEAGPSIVH	R	359	2	787.3806	
9 I TKQEYDEAGPS	I	359	2	612.7735	
) K QEYDEAGPSIVH	R	361	2	672.808	
1 Q EYDEAGPSIVHR	K	362	3	452.213	
2					
~					CB <sup>g</sup> (N-ter)
~			<b>EAGPSIVH</b> R 362 <b>AGPSIVHR</b> K 363		

65	Y	DEAGPSIVHR	K	364	2	540.7759
66	Ε	AGPSIVHR	K	366	2	418.7395

- <sup>a</sup> Position of the amino acid residue preceding the peptide sequence (N-terminus). <sup>b</sup> Position of the amino acid residue following the peptide sequence (C-terminus).
- 26 27 28 29 30 31 32 <sup>c</sup> Peptide sequence start in actin.
- <sup>d</sup> Charge (+).
- <sup>e</sup> Mass to charge ratio.
- <sup>f</sup> Cathepsin D cleavage site.
- <sup>g</sup> Cathepsin B cleavage site.
- 33 Peptides that were also found in low pH dry fermented sausages after 28 days of ripening are indicted in bold.

**Table 4.** Peptides of actin protein identified by LCMS<sup>E</sup> in low-pH dry fermented sausages after 28

# 35 days of ripening

N.	P <sub>1</sub> <sup>a</sup>	Peptide sequence	P1, <sup>b</sup>	P. s. s. <sup>c</sup>	zd	m/z <sup>e</sup>	Info
1	С	DNGSGLVKAGFAGDDAPRA	V	12	3	606.6272	CD <sup>f</sup> (N-ter)
2	С	DNGSGLVKAGFAGDDAPR	A	12	3	582.952	CD <sup>f</sup> (N-ter), CBD <sup>h</sup> (C-ter)
3	D	NGSGLVKAGFAGDDAPRAVFPSIVG	R	13	3	801.4185	
4	D	NGSGLVKAGFAGDDAPRA	V	13	3	568.287	
5	D	NGSGLVKAGFAGDDAPR	А	13	3	544.6087	CBD <sup>h</sup> (C-ter)
6	Ν	GSGLVKAGFAGDDAPRA	V	14	3	530.2726	
7	Ν	GSGLVKAGFAGDDAPR	А	14	3	500.5875	CBD <sup>h</sup> (C-ter)
8	G	LVKAGFAGDDAPRA	V	17	3	463.2476	
9	G	LVKAGFAGDDAPR	А	17	2	658.8565	CBD <sup>h</sup> (C-ter)
10	L	VKAGFAGDDAPR	А	18	2	602.3102	
11	V	KAGFAGDDAPRA	V	19	2	588.2925	
12	K	AGFAGDDAPRAVFPS	I	20	2	739.3563	
13	K	AGFAGDDAPRA	v	20	2	524.2473	
14	K	AGFAGDDAPR	Ā	20	2	488.7296	CBD <sup>h</sup> (C-ter)
15	A	GFAGDDAPRAVFPS	I	21	2	703.8424	022 (0 001)
16	A	GFAGDDAPRA	V	21	2	488.7285	
17	G	FAGDDAPRAVFPSIVGRPRHQG	v	22	4	588.3061	CBD <sup>h</sup> (N-ter)
18	G	FAGDDAPRAVFPSIVGRPRHQG	Q	22	4 3	722.3819	
10 19	G			22	3	624.6606	
		FAGDDAPRAVFPSIVGRP	R				
20 21	G	FAGDDAPRAVFPSIVGR	P	22 22	3 2	592.3103	
	G	FAGDDAPRAVFPSIVG	R			809.9169	
22	G	FAGDDAPRAVFPS	I	22	2	675.34	. ,
23	G	FAGDDAPRAVFP	S	22	2	631.8156	
24	G	FAGDDAPRAVF	Р	22	2	583.2892	CBD <sup>h</sup> (N-ter)
25	F	AGDDAPRAVFPSIVGRP	R	23	3	575.6375	
26	F	AGDDAPRAVFPSIVGR	Р	23	3	543.2854	
27	F	AGDDAPRAVFPSIVG	R	23	2	736.3833	
28	A	GDDAPRA	V	24	1	683.3168	CB <sup>g</sup> (N-ter)
29	D	DAPRAVFPS	I	26	2	480.251	
30	A	VFPSIVGRPRHQG	V	31	3	483.9377	
31	F	PSIVGRPRHQG	V	33	2	602.3381	
32	R	HQGVMVGMGQK	D	41	2	586.286	
33	G	VMVGMGQKDS	Y	44	2	526.2501	
34	V	GMGQKDSYVGDEAQSKRG	I	47	3	638.2976	
35	G	MGQKDSYVGDEAQSKRGILT	L	48	3	728.3632	CB <sup>g</sup> (C-ter)
36	G	MGOKDSYVGDEAQSKRG	I	48	3	619.2863	. ,
37	G	MGQKDSYVGDEAQSKR	G	48	3	600.2819	
38	G	MGQKDSYVGDEAQSK	R	48	3	548.2491	
39	G	MGQKDSYVGDEAQ	S	48	2	714.3086	
40	M	GOKDSYVGDEAQSKRG	I	49	3	575.6082	CB <sup>g</sup> (N-ter)
41	M	GQKDSYVGDEAQSKR	G	49	3	556.5999	CB <sup>g</sup> (N-ter)
42	M	GQKDSYVGDEAQSK	R	49	2	756.3502	CB <sup>g</sup> (N-ter)
43	D	SYVGDEAQSKR	G	53	2	620.3017	CD <sup>3</sup> (IN CCT)
44	S	YVGDEAQSKRGILT	L	54	2	768.9041	CB <sup>g</sup> (C-ter)
44 45	s S	YVGDEAQSKRGILI	Г	54 54	2	605.297	
45 46	s S	YVGDEAQSKR	L G	54 54	2	576.7844	
46 47	S Y		G I	54 55	2		
		VGDEAQSKRG				523.7643	$CD^{q}$ (C tor)
48	V	GDEAQSKRGILT	L	56	2	637.842	CB <sup>g</sup> (C-ter)
49	L	TLKYPIE	H	67	2	432.2482	
50	H	GIITNWDDMEK	I	75	2	661.3095	· · · · · ·
51	G	IITNWDDMEKIWHHTF	Y	76	3	695.999	
52	G	IITNWDDMEKIWHHT	F	76	3	646.9729	
53	G	IITNWDDMEK	I	76	2	632.8001	
54	I	ITNWDDMEKIWHHTF	Y	77	3	658.303	CBD <sup>h</sup> (C-ter), CD <sup>f</sup> (C-ter)
55	I	ITNWDDMEKIWHH	Т	77	3	575.5944	
56	I	ITNWDDMEK	I	77	2	576.2577	
57	I	TNWDDMEKIWHHTF	Y	78	3	620.6089	CD <sup>f</sup> (N-ter), CBD <sup>h</sup> (C-ter)
58	I	TNWDDMEKIWHHT	F	78	3	571.5958	
59	I	TNWDDMEKIWHH	Т	78	3	537.9025	
60	I	TNWDDMEK	I	78	2	519.7155	. ,
61	Ť	NWDDMEKIWHHT	F	79	3	537.9053	
62	Ň	WDDMEKIWHHTF	Y	80	3	548.9097	
52			T	80	3	466.203	022 (0 001)
63	Ν	WDDMEKIWHH					

64	W	DDMEKIWHHT	F	81	2	656.2975	CD <sup>f</sup> (C-ter)
65	F	YNELRVAPEEHPTL	L	92	3	556.6167	
66	F	YNELRVAPEEHPT	L	92	3	518.9219	CBD <sup>h</sup> (N-ter)
67	F	YNELRVAPEE	Н	92	2	610.3025	CBD <sup>h</sup> (N-ter)
68	Ε	LRVAPEEHPTL	L	95	2	631.3515	
69	E	LRVAPEEH	P	95	2	475.7531	CD <sup>f</sup> (C-ter)
					2		CD (C LEI)
70	L	RVAPEEHPTL	L	96		574.805	
71	L	RVAPEEHPT	L	96	2	518.2621	CB <sup>g</sup> (C-ter)
72	R	VAPEEHPTLL	Т	97	2	553.2994	CBD <sup>h</sup> (N-ter), CBD <sup>h</sup> (C-ter)
73	R	VAPEEHPT	L	97	2	440.2114	CBD <sup>h</sup> (N-ter), CB <sup>g</sup> (C-ter)
74	A	PEEHPTL	L	99	2	411.7021	(,, (,
75	L	LTEAPLNPKAN	R	106	2	584.3263	
76	L	TEAPLNPKANREKM	Т	107	3	533.6141	CBD <sup>h</sup> (N-ter)
77	L	TEAPLNPKANREK	М	107	3	489.9309	CBD <sup>h</sup> (N-ter), CD <sup>f</sup> (C-ter)
78	L	TEAPLNPKANRE	K	107	2	670.3511	CBD <sup>h</sup> (N-ter)
79	L	TEAPLNPKAN	R	107	2	527.7844	CBD <sup>h</sup> (N-ter)
80	G	IVLDSGDGVTHNVPIYEG	Y	152	2	942.9775	
81	G	IVLDSGDGVTHNVPIYE	G	152	2	914.462	_
82	I	VLDSGDGVTHNVPIYEG	Y	153	2	886.4333	
83	I	VLDSGDGVTHNVPIYE	G	153	2	857.9201	CD <sup>f</sup> (N-ter)
84	I	VLDSGDGVTHNVPIY	E	153	2	793.3949	CD <sup>f</sup> (N-ter)
85	I	VLDSGDGVTHNVP	I	153	2	655.3256	CD <sup>f</sup> (N-ter)
86	L	DSGDGVTHNVPIYEG	Y	155	2	780.3532	CD <sup>f</sup> (N-ter)
87	L	DSGDGVTHNVPIYE	G	155	2	751.8449	CD <sup>f</sup> (N-ter)
88	L	DSGDGVTHNVPIY	E	155	2	687.3255	CD <sup>f</sup> (N-ter)
89	L	DSGDGVTHNVP	I	155	2	549.2509	CD <sup>f</sup> (N-ter)
90	D	SGDGVTHNVPIYEG	Y	156	2	722.8416	(
91	D	SGDGVTHNVP	I	156	2	491.7312	
92	S	GDGVTHNVPIYEG	Y	157	2	679.3201	
93	G	DGVTHNVPIYEG	Y	158	2	650.814	
94	Y	ALPHAIMRL	D	171	2	511.2992	
95	T	AASSSSLEKSYELPDGQVITIGN	Ē	231	2	1183.5982	
96	A	ASSSSLEKSYELPDGQVIT	I	232	2	1005.9979	CB <sup>g</sup> (C-ter)
97	S	SLEKSYELPDGQVITIGNERF	R	236	3	799.0719	
98	S	SLEKSYELPDGQVITIGN	E	236	2	982.0108	
99	S	SLEKSYELPDGQVIT	I	236	2	839.9343	CB <sup>g</sup> (C-ter)
100	S	SLEKSYELPDGQVI	T	236	2	789.4068	()
101	S	LEKSYELPDGQVITIGNER	F	237	3	721.0399	
102	S	LEKSYELPDGQVIT	I	237	2	796.4186	CB <sup>g</sup> (C-ter)
103	L	EKSYELPDGQVITIGN	Ε	238	2	881.9505	
104	L	EKSYELPDGQVIT	I	238	2	739.8722	
105	K	SYELPDGQVITIGNERF	R	240	2	969.4866	CD <sup>f</sup> (N-ter)
		-			2		CD <sup>f</sup> (N-ter)
106	K	SYELPDGQVITIGNER	F	240		895.9547	CD <sup>2</sup> (N-ter)
107	S	YELPDGQVITIGNERFR	С	241			
108	S	YELPDGQVITIGNERF	R	241	2	925.9733	
109	S	YELPDGQVITIGNER	F	241	2	852.4358	
110	S	YELPDGQVIT	I	241	1	1134.5734	
111	Y		R	241	2		
		ELPDGQVITIGNERF				844.4392	
112	Y	ELPDGQVITIGNER	F	242	2	770.9043	
113	Y	ELPDGQVI	Т	242	1	870.4615	
114	E	TLFQPSF	I	261	1	839.3847	
115	Y	NSIMKCDIDIRK	D	281	2	718.3576	
116	D	LYANNVMSGGTTMYPGIADRMOKE	I	294	3	883.0785	
		~					
117	A	NNVMSGGTTMYPGIADRMQKE	I	297	3	767.3495	
118	A	NNVMSGGTTMYPGIADR	М	297	2	892.4089	
119	М	SGGTTMYPGIADRMQKE	I	301	3	614.62	
120	М	SGGTTMYPGIADRMQK	E	301	3	571.6075	
121	М	SGGTTMYPGIADRMQ	K	301	2	792.8625	
121	M		M	301	2		
		SGGTTMYPGIADR				663.3142	
123	М	SGGTTMYPGIAD	R	301	1	1169.5235	
124	K	IIAPPERKYS	V	330	2	587.3361	CBg (N-ter)
125	М	WITKQEYDEAGPSIVHRK	С	357	4	540.0306	
126	М	WITKQEYDEAGPSIVHR	K	357	3	677.0105	
120	M	WITKOEYDEAGPS	I	357	2	762.3605	
		~					
128	W	ITKQEYDEAGPSIVHRK	С	358	4	493.5112	
129	W	ITKQEYDEAGPSIVHR	K	358	3	614.984	CDf (N-ter)
130	W	ITKQEYDEAGPSIVH	R	358	3	562.9486	CDf (N-ter)
131	W	ITKQEYDEAGPS	I	358	2	669.3159	
132	I	TKQEYDEAGPSIVHRK	Ċ	359	3	619.9871	/
TJC	1	INZEIDERGESIVIER	C	222	J	1106.610	
				6			

133	I	TKQEYDEAGPSIVHR	K	359	3	577.2883		
134	I	TKQEYDEAGPSIVH	R	359	2	787.38		
135	I	TKQEYDEAGPS	I	359	2	612.7782		
136	Q	EYDEAGPSIVHRK	С	362	2	750.8781		
137	Q	EYDEAGPSIVHR	K	362	2	686.8298		
138	Q	EYDEAGPSIVH	R	362	2	608.7845		
139	Ε	YDEAGPSIVHRK	С	363	2	686.3556	CBa	(N-ter)
140	Ε	YDEAGPSIVHR	K	363	2	622.3098	CBg	(N-ter)
141	Ε	YDEAGPSIVH	R	363	2	544.2598	CBg	(N-ter)
142	Y	DEAGPSIVHRK	С	364	2	604.8217		
143	Y	DEAGPSIVHR	K	364	2	540.7767		
144	Ε	AGPSIVH	R	366	1	680.3776		

36 <sup>a</sup> Position of the amino acid residue preceding the peptide sequence (N-terminus).

37 <sup>b</sup> Position of the amino acid residue following the peptide sequence (C-terminus).

38 <sup>c</sup> Peptide sequence start in actin.

39 <sup>d</sup> Charge (+).

<sup>e</sup> Mass to charge ratio.

40 41

<sup>f</sup> Cathepsin D cleavage site. <sup>g</sup> Cathepsin B cleavage site. Peptides that were also found in high pH dry fermented sausages after 28 days of ripening are indicted in bold. 42 43

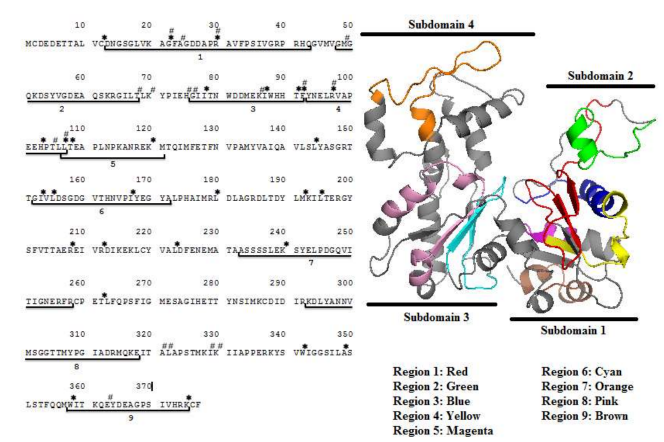


Figure 1.

- \* Cleavage sites of cathepsin D according to Hughes et al. (2000)
- <sup>5</sup> <sup>#</sup> Cleavage sites of cathepsin B according to Hughes et al. (1999)

7 Figure 2.

