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Abstract: Characterisation of the physiologic equine amniotic fluid (AF) proteome is a prerequisite to study its changes during diseases and discover new biomarkers. The aim of this study was to identify by a proteomic approach the most abundant proteins of equine AF. AF samples were collected at parturition from 24 healthy mares that delivered healthy foals. All samples were subjected to sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) on 4-12% gels. A pool of the 24 samples, after SDS-PAGE, was cut in 25 slices, trypsin-digested and analysed by mass spectrometry (MS) for protein identification. Mean AF protein concentration was 1.96±1.12 g/L. Thirty-four proteins were successfully identified by MS and subsequently categorised according to Gene Ontology (GO). Twelve proteins (e.g. fibronectin, lumican, thrombospondin and fibulin) belonged to or interacted with the extracellular matrix (ECM) playing an important role in the development of foetal tissues. Most of the remaining proteins were classified as transport (e.g. albumin, major allergen Equ c1 and alpha-fetoprotein) delivering nutrients, ions and lipids essential for foetal growth and development. Among these proteins, major allergen Equ c1 is widely studied in human medicine because it induces Iq-E mediated type I allergic reaction. The absence of immunoglobulins in equine AF was also confirmed.

# **REVISION NOTE**

Animal Reproduction Science Ms. No. ANIREP-D-16-6583 Identification of the most abundant proteins in equine amniotic fluid by a proteomic approach.

# Dear Editor,

thank you for considering the present paper for publication in Animal Reproduction Science and for the opportunity to revise our manuscript.

As you can see in the following pages, we answered all the questions raised by the reviewer. Following the reviewer suggestions, some sentences with more details have been added improving the clarity and the quality of the manuscript. In particular, more details on the statistical analysis performed were added and the discussion section was partially modified.

As requested, the added paragraph/words are written in red. Moreover, we highlighted the deleted parts in yellow. We upload two version of the revised manuscript: one with the changes highlighted and a second version corrected and edited.

In addition, after an accurate revision of the manuscript, we did also other minor corrections to improve the editing according to Animal Reproduction Science author's guidelines.

# **RESPONSE TO REVIEWER 1**

• Reviewer #1: In general, this is a well-written manuscript. However, it is merely descriptive and does not follow any hypothesis. Nevertheless, it is based on a sufficient number of experimental animals and data appear to be reliable. It is therefore recommended for publication after moderate revision. The following points of criticism should be taken into account:

This research can be considered as pure, fundamental research, curiosity-driven and aimed to improve our knowledge on the complexity of equine amniotic fluid. The description of its proteome in healthy mares is a prerequisite for further investigations focused on specific diseases and/or pathologic conditions.

• Information in the abstract is very general, please add some data. As suggested by the reviewer, some more data and information have been added.

• In the introduction the authors should explain why they choose to analyze the protein of amniotic and not of allantoic fluid.

As suggested by the reviewer, a sentence explaining the motivation of analysing AF instead of allantoic fluid was added.

• *Line 63: "and discuss their function" should be deleted.* As suggested by the reviewer, the words have been deleted.

• Lines 74-75: It is stated that all mares were healthy based on clinical and ultrasonographic evaluation. Please add details on which parameters have been accessed at what times before and after parturition.

As suggested by the reviewer, more details on clinical and ultrasonographic evaluation have been added.

# • Information on statistical analysis is not detailed enough.

As suggested by the reviewer, more information have been added. Correlation analysis between AF total proteins and the other evaluated parameters was added. Nevertheless, to the author opinion, since the main objective of the present paper was the characterization/description of the AF proteome in healthy samples and it was not a comparison between different experimental groups, the statistics are mainly descriptive.

• The discussion does not start properly. It is always annoying for the reader if he/she has to deal with limitations of the study and has to wait for the more exciting points of the discussion. Please start with the most important results and shift the present beginning of the discussion to the end.

Following the reviewer suggestion, the discussion has been modified, moving the first paragraphs to the end of the section. In the author's opinion the non-depletion of albumin and of the other proteins is not a limitation, but a decision driven by the intention to maintain the integrity of the sample.

• The authors refer to the point that protein concentration of amniotic fluid analyzed in the present study differs from results of other studies and try to explain this discrepancy by the use of different methods. However, also in the present study wide variations were found. How can such an "inter-individual variability" be explained. What are the possible underlying causes?

As underlined by the reviewer, the inter-individual variability was quite high. However, to the authors knowledge, no publications are present in the literature that evaluate possible source of inter-individual variability in AF protein concentration. Therefore, to the authors opinion, the possible explanation for the variability reported in the present paper could rely on different pre-hospitalization factors, such as differences in feed and hydration status and in housing conditions. A sentence was added at lines 326-331.

- Lines 221 to 229: The publication by Ottsdottir et al. 2010 (Theriogenology 75 (2011) 1130-1138) on dynamics of MMP in equine fetal fluids has not been considered. As suggested by the reviewer, the citation has been added.
  - The conclusion has to be rewritten. At present it is another summary but does not present a "take home message" for the reader which would be preferable.

Following the suggestion of the reviewer the conclusion has been rewritten. However, as previously reported, since the study is mainly descriptive, also the "take home message" cannot have a direct practical/clinical application.

- *Table 1: please add bodyweight of mares and their fetuses* The requested data have been added.
  - All Tables and Figures should "stand alone" with complete descriptions of what is being presented as for example mean ± sem, numbers of animals, etc. so that the reader does not have to refer to the text.

As suggested by the reviewer, some more information have been added to the figures and tables legends. In legend of figure 1 the sample number of the analysed AF as reported in table 1 was also added. We also decided to remove table 2, since the data reported were already present in figure 2. However, if the reviewer considers the table as useful we will add it again.

Highlights

- Equine amniotic fluid was preliminary characterised for the first time.
- SDS-PAGE coupled to MS allowed the identification of the most abundant proteins.
- The three most abundant proteins were albumin, major allergen Eqc1 and fibronectin.
- The role of the extracellular matrix component in fetal maturation was highlighted.
- The importance of transport proteins like alfa-fetoprotein and PLTP was evidenced.

1	Identification of the most abundant proteins in equine amniotic fluid by a proteomic
2	approach
3	
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## 21 Abstract

22 Characterisation of the physiologic equine amniotic fluid (AF) proteome is a prerequisite to 23 study its changes during diseases and discover new biomarkers. The aim of this study was to 24 identify by a proteomic approach the most abundant proteins of equine AF. AF samples were 25 collected at parturition from 24 healthy mares that delivered healthy foals. All samples were 26 subjected to sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) on 4-12% gels. A pool of the 24 samples, after SDS-PAGE, was cut in 25 slices, trypsin-digested 27 28 and analysed by mass spectrometry (MS) for protein identification. Mean AF protein 29 concentration was 1.96±1.12 g/L. Thirty-four proteins were successfully identified by MS and subsequently categorised according to Gene Ontology (GO). Twelve proteins (e.g. 30 31 fibronectin, lumican, thrombospondin and fibulin) belonged to or interacted with the 32 extracellular matrix (ECM) playing an important role in the development of foetal tissues. 33 Most of the remaining proteins were classified as transport (e.g. albumin, major allergen Equ 34 c1 and alpha-fetoprotein) delivering nutrients, ions and lipids essential for foetal growth and 35 development. Among these proteins, major allergen Equ c1 is widely studied in human 36 medicine because it induces Ig-E mediated type I allergic reaction. The absence of 37 immunoglobulins in equine AF was also confirmed.

38

### 39 Keywords

- 40 Horse, pregnancy, electrophoresis, proteome.
- 41

42

#### 44 Introduction

45 Amniotic fluid (AF) is a complex dynamic milieu that changes as pregnancy progresses. AF contains nutrients and growth factors that facilitate foetal growth, provides 46 47 mechanical cushioning and antimicrobial effectors to protect the foetus and allows assessment of foetal maturity and disease (Underwood et al. 2005). In comparison to humans, the 48 49 physiology and pathophysiology of foetal fluids in domestic mammals are poorly understood 50 (Canisso et al. 2015). In horses, some studies have investigated biochemical composition, 51 particularly enzymes and electrolytes, in AF collected by ultrasound-guided transabdominal 52 amniocentesis, at delivery or at slaughter (Holdstock et al. 1995; Lyle et al. 2006; Williams et 53 al. 1993; Zanella et al. 2014). AF was also studied for evaluation of foal's lung maturity at 54 birth through lecithin/sphingomyelin ratio and lamellar body count (Castagnetti et al. 2007). More recently, significantly higher levels of lactate were found in AF collected during 55 56 parturition in mares delivering healthy foals (Pirrone et al. 2012).

57 Unlike the allantoic fluid, equine AF can be easily collected during parturition without 58 stressing the animal and avoiding any contamination (Castagnetti et al. 2007, Pirrone et al. 59 2012). As reported in women, the biochemical composition of AF, including proteins, is 60 primarily representative of the foetal profile and reflects its physiological status (Tong 2013), 61 thus it could be potentially useful to evaluate the high-risk foal born attended.

Proteomics is a powerful analytical approach providing a profile of proteins present in a biological sample at a given time. The high potential of this approach has been recently found to have a major role in different areas of veterinary medicine, from farm (Almeida et al. 2015) to companion (Ferlizza et al. 2015; Miller et al. 2014) animals. Proteomic techniques have recently been applied to the characterisation of horse amniotic membrane (Galera et al. 2015) and bovine conceptus fluids (Riding et al. 2008), whereas the equine AF proteome remains uncharacterised. Therefore, the aims of this study were to identify the most abundant 69 proteins in equine AF by SDS-PAGE separation followed by mass spectrometry70 identification.

71

## 72 Materials and Methods

#### 73 Animal selection and data collection

Twenty-four mares admitted for assisted delivery during three breeding seasons at the S. Belluzzi Equine Perinatology Unit of the Department of Veterinary Medical Sciences, University of Bologna, were included. The mares were hospitalised at about 310 days of pregnancy because the owners requested an attended parturition, and remained under observation for at least 7 days *postpartum*. They were housed in wide straw bedding boxes and fed with hay *ad libitum* and concentrates twice a day. During the day, the mares were allowed to go to pasture.

81 All the mares included in the study were healthy based on clinical and 82 ultrasonographic evaluation. At admission, a complete clinical evaluation, including complete 83 blood count, serum biochemistry and transrectal ultrasonography, was performed. Severe 84 maternal illness, uterine discharge, premature lactation, twinning, abnormal foetal presentation, placenta oedema, and signs of foetal distress were ruled out. During the course 85 of hospitalisation, mares were clinically evaluated twice a day and by transrectal 86 87 ultrasonography every 10 days until parturition. The following ultrasonographic parameters were evaluated: combined thickness of the uterus and placenta, foetal fluids echogenicity, 88 89 foetal activity, and foetal orbital area. Foals were born between 320 and 365 days of 90 pregnancy by normal delivery, had an Apgar score  $\geq 8$  recorded within 5 minutes from birth 91 (Vaala et al. 2002) and had a normal clinical evaluation during the course of hospitalisation, 92 including a complete blood count and serum biochemistry at birth and an immunoglobulin G 93 (IgG) serum concentration  $\geq$ 800 mg/dL at 18-24 hours of life.

For each mare, the following data were recorded: breed, age, parity, days of
pregnancy, body weight, length of stage II labour (minutes), and foal's body weight and
Apgar score. All procedures on the animals were carried out with the approval of the Ethical
Committee, in accordance with DL 116/92, approved by the Ministry of Health (approval
number: n.18/64/11; date of approval 22/02/2011). Oral informed consent was given by the
owners.

100

101 Sample collection

At foaling, a sample of AF was collected from each mare with a 50 mL syringe by needle puncture of the amniotic sac within few minutes of its appearance through the vulva during stage II of labour. The AF was then immediately transferred to 5 mL test tubes and stored at -80°C until SDS-PAGE and protein identification were performed. AF protein concentration was determined by the Biuret method using bovine serum albumin as standard.

107

108 SDS-PAGE

109 To optimise protein separation, different protocols were tested including 4-12% and 110 12% polyacrylamide gels (NuPage/Thermo Fisher Scientific, Waltham, Massachusetts, USA) in 2-(N-morpholino) propanesulfonic acid buffer (MOPS; NuPage/Thermo Fisher Scientific) 111 112 or 2-(N-morpholino) ethanesulfonic acid buffer (MES; NuPage/Thermo Fisher Scientific) 113 with sodium dodecyl sulphate (SDS). Each AF sample (n=24) was analysed at least twice 114 with the protocol assuring the best protein separation in our experimental conditions (4-12%) 115 gels, in MOPS buffer). Twenty µg of proteins were loaded for each sample and the gels were 116 stained with Coomassie G250 compatible with mass spectrometry analysis. After staining, each gel was digitalised by ChemiDoc<sup>TM</sup>MP (BioRad, Hercules, California, USA) and its 117 pherogram was obtained using ImageLab 5.2.1 software (BioRad). The software determines 118 119 the volume of each protein band through the analysis of the pixel values in the digital image, meaning as volume the sum of all the pixels intensities within the band boundaries. The band volumes are subsequently compared to the entire volume of the lane and the relative abundances reported in percentage. A pool was prepared by collecting and mixing 50  $\mu$ g of protein from each AF (n=24) and analysed twice with the same protocol used for each sample.

124

#### 125

# Protein identification by mass spectrometry

126 The pool lanes were divided manually into 25 slices and subjected to in-gel tryptic 127 digestion as previously described (Bellei et al. 2013). Digested dried samples were then re-128 suspended in 97% Water/3% ACN added of 1% formic acid (Buffer A) and analysed by a Nano LC-CHIP-MS system (ESI-Q-TOF 6520; Agilent Technologies, Santa Clara, 129 130 California, USA). Four microliters of each sample were loaded into the system and transported to the Chip enrichment column (Zorbax C18, 4 mm x 5 µm i.d., Agilent 131 132 Technologies) by a capillary pump, with a loading flow of 4 µL/min, using 95% 133 ACN/5% water added of 0.1% formic acid (buffer B) as mobile phase. Nitrogen was used as 134 the nebulising gas. A separation column (Zorbax C18, 43 mm x 75 µm i.d., Agilent 135 Technologies), at flow rate of  $0.4 \,\mu$ L, was used for peptide separation.

136 Since the horse protein database is not well annotated, a broader taxonomy, namely "all mammals", was selected for identification to be based on sequence homology. Protein-137 138 identification peak lists were generated using the Mascot search engine 139 (http://mascot.cigs.unimo.it/mascot) against the UniProt database (UniProt.org) specifying the 140 following parameters: mammalian taxonomy, parent ion tolerance ±20 ppm, MS/MS error 141 tolerance  $\pm 0.12$  Da, alkylated cysteine as fixed modification and oxidised methionine as 142 variable modification, and two potential missed trypsin cleavages, as previously described 143 (Bertoldi et al., 2013). Proteins with a score >80 or identified with at least two or more 144 significant sequences were selected. The significant threshold in Mascot searches was set to

obtain a false discovery rate <5% (5% probability of false match for each protein with a score</li>
above 80).

147

148 Statistical analysis

Data (AF total proteins, mare's age, mare's and foal's body weight, parity, days of pregnancy, length of stage II labour (minutes), foal's Apgar score and number of bands) were analysed with statistical software (R version 2.15.1) and reported as mean  $\pm$  standard deviation (SD). Shapiro-Wilk normality test was performed to evaluate data normal distribution. Pearson coefficient of correlation was calculated between AF total proteins and the other data recorded for each mare (mare's age, mare's and foal's body weight, parity, days of pregnancy, length of stage II labour (minutes), foal's Apgar score and number of bands).

The identified proteins were categorised by biological process, molecular function and
cellular component with Gene Ontology terms according to Gene Ontology (GO) and Human
Protein Reference Database (HPRD).

159

160 **Results** 

161 *Clinical data* 

162 Clinical data collected from the 24 mares included in the study are shown in Table 1. Mean 163 AF total protein concentration was  $1.96\pm1.12$  g/L, ranging from 0.36 to 4.16 g/L. No 164 significant correlation was found between AF protein concentration and the other data 165 recorded.

166

167 SDS-PAGE and protein identification by mass spectrometry

168 Representative gel and pherogram of AF are reported in Figure 1A and 1B. The mean 169 number of bands was 23±1.5. All samples presented a similar pattern characterised by two 170 clusters of bands: the first with molecular weights (MW) higher than 62 kDa and the second with MW lower than 34 kDa. In the middle, very few faint bands were present. Figure 2 reports the relative abundance in percentage of AF protein bands. Out of the 25 slices cut from the gel (Figure 3), 20 yielded significant results leading to the unambiguous identification of 34 proteins (Table 2). Serum albumin and major allergen Equ c1 (ALL1) were the two most abundant proteins, followed by fibronectin, transferrin and haemoglobin; these five proteins represented >60% of the equine AF proteome. Fibronectin, versican and albumin were also identified in bands characterised by different MW.

The identified proteins categorised by their molecular function and biological process according to Gene Ontology (GO) and Human Protein Reference Database (HPRD) are shown in Table 3 and Figure 4. Most of the proteins were involved in cellular growth and/or maintenance (38%), transport (26%) and protein metabolism (9%). The vast majority of the identified proteins were classified as extracellular (79%).

183

## 184 **Discussion**

The present paper aimed to explore the complexity of equine AF proteome and to identify its most abundant proteins. The study was carried out on 24 mares of different breed, age and parity referred to the Equine Perinatology Unit, and they can be considered representative of a typical equine hospital population. Therefore, the proteomic profile described can be considered a useful starting point for further applied studies on the equine AF proteins.

Most of the 34 proteins identified were involved in cellular growth and maintenance, transport and protein metabolism reflecting the dynamic biological functions of AF. Regarding cellular growth and/or maintenance, 12 of the proteins identified belonged to or interacted with the extracellular matrix (ECM) that plays an important role in the development of foetal tissues. All these proteins, with the exception of versican and proteoglycan 4, were also identified in human AF (Cho et al. 2007; Michaels et al. 2007) and/or in equine amniotic

membrane (Galera et al. 2015). Among the ECM structural proteins, fibronectin is a 197 198 multifunctional glycoprotein known to participate in the organisation of ECM binding to 199 integrins. During pregnancy, fibronectin is expressed in the junction between maternal and 200 foetal membranes as well as in the uterus and placenta (Mogami et al. 2013). Lumican, a 201 member of the family of small leucine-rich proteoglycans, is the major keratan sulphate 202 proteoglycan of the cornea and is also present in the ECM throughout the body, including 203 human and equine amniotic membrane (Galera et al. 2015; Kao et al. 2006; Mavrou et al. 204 2008). Based on its interaction with fibrillar collagen and its ability to modulate cell 205 proliferation and migration, lumican could play a role in the maturation of foetal tissues 206 (Mavrou et al. 2008). Regarding the non-structural proteins involved in ECM development 207 and organisation, thrombospondin and fibulin are regulatory proteins belonging to the group 208 of the matricellular proteins. These proteins represent a bridge between matrix proteins and 209 cell surface receptors, or other molecules such as cytokines that can interact with the cell 210 surface (Bornstein, 1995). They are typically expressed at low levels in adult tissues, but are 211 strongly expressed during development or following injury or pathology (Morris and 212 Kyriakides, 2014). Gelsolin is a multifunctional actin regulatory protein involved in cytoskeleton dynamics and structure. In addition to its role in aiding chemotaxis and 213 214 movement of intracellular structures, plasma gelsolin binds to a variety of proinflammatory 215 and bioactive molecules including fibronectin, platelet activating factor and the bacterial 216 surface lipids lipoteichoic acid and lipopolysaccharide (Peddada et al. 2012). The role of 217 gelsolin in AF is still unknown, but it has been suggested to modulate inflammation and 218 bacterial infections in human AF (Sezen et al. 2009). In association with gelsolin, vinculin is 219 also a component of the actine cytoskeleton and is involved in integrin-mediated focal 220 adhesion, cell motility and other cellular functions such as migration, proliferation and 221 differentiation (Wu et al. 2014). Other proteins interacting with ECM that could play 222 important roles in the development of foetal tissue are type IV collagenase (MMP2) and

223 metalloproteinase inhibitor 1 (TIMP1) belonging to the matrix metalloproteinases (MMPs) 224 and tissue inhibitors respectively. The MMPs are a family of over 20 enzymes acting on the 225 ECM components, regulated at different levels via their activators, inhibitors and localization 226 on the cell surface (Sternlicht and Werb, 2001). The biological functions of these enzymes 227 and their inhibitors have been widely studied, in particular MMP2 is important for bone 228 development and angiogenesis regulation and has been identified and studied in plasma and 229 AF of pregnant women (Anumba et al. 2010; Turner et al. 2014). MMPs activity was also 230 studied in amniotic and allantoic fluid from mares that delivered live term foals and from 231 mares with preterm delivery, suggesting that MMPs may have a role as markers for high risk 232 pregnancy in the mare (Oddsdóttir et al. 2011).

233 Among transport proteins, albumin, transferrin, alpha-fetoprotein, apolipoprotein A1 234 and phospholipid transfer protein (PLTP) transport nutrients, ions and lipids essential for 235 foetal growth and development and have been identified as common components of AF also 236 in humans (Cho et al. 2007; Michaels et al. 2007). Alpha-fetoprotein is member of the albuminoid superfamily and is present in the allantoic and amniotic fluids of domestic 237 238 animals (Luft et al. 1984; Smith et al. 1979). In mammalian foetuses alpha-fetoprotein is 239 associated with oestrogen-binding, anti-oxidative properties and immunoregulation (DeMees 240 et al. 2006; Mizejewski 2001) and it is highly expressed during early pregnancy by the 241 equine conceptus (Simpson et al. 2000). In women, AF alpha-fetoprotein is actively 242 investigated for pathologies such as Down syndrome, trisomies 13 and 18, intra-amniotic 243 infection, preterm delivery, pre-eclampsia, membrane rupture, and foetoplacental hypoxia 244 (Cho et al. 2007). Recently, Canisso et al. (2015) confirmed the presence of alpha-fetoprotein in equine foetal fluids during the third trimester of pregnancy and found increased maternal 245 246 plasma concentrations of the protein in mares with experimentally induced placentitis. The 247 presence of ALL1 in AF is challenging. ALL1 is a glycoprotein of 21.7 kDa belonging to the 248 family of lipocalins, whose function is to carry small hydrophobic molecules such as

249 odorants, steroids and pheromones. This protein is expressed in salivary glands and in the 250 liver and is highly concentrated in secretory fluids such as saliva and urine as well as in hair 251 and dander (Botros et al. 2001). ALL1 is widely studied in human medicine because it 252 induces an IgE-mediated type I allergic reaction in the majority of patients allergic to horses 253 (Lascombe et al. 2000). The physiological role of this protein is still unknown and to our 254 knowledge, this is the first study reporting the presence of ALL1 in AF. PLTP is a monomeric 255 glycoprotein involved in lipid transport, lipoprotein metabolism and lipopolysaccharide 256 binding. It is ubiquitously expressed in human tissues and is secreted into the plasma, where 257 its central role has been well established (Albers et al. 2012). PLTP is highly expressed in 258 lung epithelial cells, and may play a role in surfactant metabolism during foetus lung 259 development (Brehm et al. 2014).

260 The proteomic approach applied in the present study led to the successful 261 identification of the most abundant proteins, even though a few additional points should be 262 taken into consideration. The first one regards the choice of non-depleting albumin and other 263 major proteins before electrophoresis and MS identification. Complex biological samples 264 contain thousands of different protein species, few of them characterised by high abundance 265 and many others by low or very low abundance. The presence of very high abundance 266 proteins like albumin and immunoglobulins often hampers the separation and characterisation 267 of serum and AF proteomes, therefore the depletion of these major components has been 268 applied in human proteomics (Cho et al. 2007; Michaels et al. 2007). However, this approach 269 can lead to the loss of some low abundance proteins due to the "sponge effect" of albumin 270 that can bind a variety of other proteins or peptides (Bellei et al. 2011). From this point of 271 view, equine AF is a preferential sample due to the absence of immunoglobulins. In domestic 272 animals, the passage of immunoglobulins is influenced by the placental structure: in horses, 273 pigs and ruminants the placenta is epitheliochorial, thus impermeable to immunoglobulins 274 (Furukawa et al. 2014), whereas in dogs, the endotheliochorial placenta allows only 5% to

10% transfer of maternal antibodies to the foetus (Dall'Ara et al. 2015). From an analytical
point of view, the absence of immunoglobulins in equine AF, as confirmed by this study, can
be considered an advantage, allowing to perform SDS-PAGE and MS identification without
affecting proteome integrity and complexity.

The second point is related to the sample collection. This study collected AF only at parturition because mares were client-owned and transabdominal amniocentesis is still not recommended for clinical use. Recently, Canisso et al. (2014) described a safe technique to perform multiple ultrasound-guided foetal fluid samplings during the last trimester of gestation in mares. On this basis, abdominal amniocentesis will probably be preferred more frequently also in a clinical setting and to evaluate gestational changes in the AF proteome as reported in women (Michaels et al. 2007).

286 The last point regards AF total protein concentrations, which were similar to those 287 reported by Williams et al. (1993) (3.1±2.6 g/L) and Paccamonti et al. (1995) (1-2 g/L), but 288 lower than those reported by Kochhar et al. (1997) (9.1±2 g/L), and higher than those of 289 Zanella et al. (2014) (0.3±0.1 g/L). The reported discrepancies could be related to the use of 290 quantification methods characterised by different analytical performances; also the influence 291 of wide inter-individual variability cannot be excluded. Many environmental and 292 physiological factors can contribute to this variability; in particular, since the mares were 293 client-owned, pre-hospitalisation conditions, such as housing, feeding, nutrition and hydration 294 status, might have affected AF total protein concentration.

295

#### 296 Conclusions

Applying a qualitative proteomic approach, this study identified the 34 most abundant proteins of the AF proteome from healthy mares that delivered live term foals. GO categorisation demonstrated that these proteins are involved in different biological processes and molecular functions including cell growth/maintenance and transport. Some of these 301 proteins belonged to or interacted with the extracellular matrix, highlighting the role of its 302 components in foetal maturation. The study confirmed also the importance of transport 303 proteins like alpha-fetoprotein and PLTP, and reported for the first time the presence of ALL1 304 in AF. Though entirely descriptive, these findings can be considered valuable context for 305 further investigations to gain insights into the function of the proteins identified and to 306 discover potential biomarkers of foetal disease at birth or during pregnancy.

307

# 308 Acknowledgements

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310 Castagnetti C.

311

# 312 **Conflict of interest**

313 The authors have no conflict of interests to declare.

315	Table 1. Clinical	data collected from	the 24 mares i	ncluded in the s	tudy. Data are reported as
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316 mean  $\pm$  standard deviation. AF amniotic fluid; TP total proteins.

Sample	Breed	Age	Mare weight	Foal weight	Parity	Length of pregnancy	Length of stage II	Foal's Apgar score	AF TP
		years	Kg	Kg		days	minutes		g/L
1	Saddlebred	11	660	51.7	1	355	25	10	2.74
2	Thoroughbred	6	645	51	1	344	20	10	1.66
3	Standardbred	5	565	50	2	335	14	8	1.51
4	Standardbred	6	585	44	1	329	8	8	3.07
5	Saddlebred	7	660	53	2	330	16	9	2.64
6	Saddlebred	12	500	40	1	360	20	9	1.01
7	Saddlebred	17	650	58	3	330	13	9	0.55
8	Quarter Horse	11	560	42.6	5	333	14	9	0.58
9	Saddlebred	14	660	53.5	1	354	20	10	0.36
10	Arabian	5	450	42	2	335	15	10	0.90
11	Standardbred	7	565	45.3	1	328	11	10	1.39
12	Standardbred	19	578	43.5	3	349	12	8	1.25
13	Standardbred	12	546	45	2	352	15	8	1.30
14	Standardbred	16	590	42.5	11	347	9	8	1.73
15	Standardbred	10	535	50	6	343	8	10	0.58
16	Standardbred	14	610	45	6	336	21	9	2.50
17	Arabian	12	430	50	6	326	20	10	2.29
18	Standardbred	6	620	59	1	341	18	10	1.66
19	Standardbred	20	606	45	12	338	12	10	4.01
20	Saddlebred	16	650	47	2	360	5	8	3.01
21	Standardbred	18	680	50	4	332	9	9	4.16
22	Thoroughbred	13	580	58	4	354	12	8	2.14
23	Quarter Horse	16	425	41	3	357	17	9	4.16
24	Standardbred	19	660	56	3	345	25	8	1.75
		12 ± 9.5	583.8 ± 74.1	48.7 ± 11.4	3 ± 3	342 ± 10.7	15 ± 5.7	9 ± 1	1.96 ± 1.12

**Table 2.** Proteins identified in equine amniotic fluid by mass spectrometry. Identified proteins

are listed according to the number of the band as marked in Figure 3.

Band	<sup>a</sup> Entry name <sup>b</sup>	Protein full name	MW (kDa) <sup>c</sup>	Score <sup>d</sup>	Pept. <sup>e</sup>	Sign. Pept. <sup>f</sup>	Seq. <sup>g</sup>	Sign Seq. <sup>h</sup>	% id. <sup>i</sup>
1	CSPG2_BOVIN	Versican core protein	371.8	257	76	21	15	6	77.2
2	FINC_HORSE	Fibronectin	58.1	1451	155	89	19	18	100
3	FINC_HORSE	Fibronectin	58.1	440	76	34	15	10	100
	FINC_BOVIN	Fibronectin	275.5	155	46	16	16	7	95.4
4		VCollagen alpha-1(VI) chain	109.6	168	41	16	10	7	89
5	VINC_HUMAN	Vinculin	124.3	148	53	18	26	10	99.5
	ACE2_PAGLA	Angiotensin-converting enzyme 2	93.0	380	43	18	10	6	86
6	PLMN_HORSE	Plasminogen	38.1	288	36	18	8	5	100
U	FINC_CANFA	Fibronectin	58.2	148	18	11	7	5	96.4
	GELS_HORSE	Gelsolin	81.1	1911	193	122	27	21	100
	FBLN1_HUMAN		81.3	434	80	47	11	9	91.2
7	FINC_HORSE	Fibronectin	58.1	360	36	23	11	7	100
	LUM_MOUSE	Lumican	38.6	159	27	14	6	4	87.4
	TRFE_HORSE	Serotransferrin	80.3	2201	263	153	43	32	100
8		Lactotransferrin	77.9	268	203 61	30	43 19	52 11	100
9	TRFL_HORSE			1642					
9	TRFE_HORSE	Serotransferrin	80.3		194	106	34	24	100
	ALBU_HORSE	Serum albumin	70.5	2863	244	148	44	33	100
		Extracellular matrix protein 1	62.2	188	53	22	4	4	78.4
	MMP2_BOVIN	72 kDa type IV collagenase	74.8	468	50	27	14	9	95.4
10	PLTP_HUMAN	Phospholipid transfer protein	54.9	400	24	19	3	3	89.1
	LUM_MOUSE	Lumican	38.6	153	23	12	6	4	87.4
	FETA_HORSE	Alpha-fetoprotein	70.1	145	17	10	7	4	100
		Versican core protein	96.8	101	13	7	4	3	82
	ALBU_HORSE	Serum albumin	70.5	17663	1154	849	70	56	100
	FETA_HORSE	Alpha-fetoprotein	70.1	407	59	34	23	14	100
11	CSPG2_MACNE	Versican core protein	96.8	347	13	10	5	4	82
	BGH3_HUMAN	Transforming growth factor-beta- induced protein ig-h3	75.3	53	12	6	4	3	92.9
12	A1AT2_HORSE	Alpha-1-antiproteinase 2	47.1	195	11	8	4	2	100
	ALBU_EQUAS	Serum albumin	70.5	160	53	13	23	9	98.5
	CLUS_HORSE	Clusterin	52.7	453	48	28	15	10	100
	ACTB_BOVIN	Actin cytoplasmic 1	42.1	118	36	11	14	6	100
	GELS_HORSE	Gelsolin	81.1	40	12	4	7	4	100
13	IBP3_BOVIN	Insulin-like growth factor-binding protein	32.6	79	11	9	5	5	86.7
	FETUA BOVIN	Alpha-2-HS-glycoprotein	39.2	65	11	6	2	2	71.3
		VCollagen alpha-3(VI) chain	345.2	93	5	4	3	2	88.5
	HPT_BOVIN	Haptoglobin	45.6	52	3	2	3	2	78.9
	CLUS_HORSE	Clusterin	52.7	42	18	4	7	3	100
14	TSP1_BOVIN	Thrombospondin-1	133.4	29	13	3	9	3	96.8
14	SFTPA_HORSE	Pulmonary surfactant-associated protein A	26.5	71	4	3	3	2	100
	TSP1_HUMAN	Thrombospondin-1	133.3	751	60	49	12	11	98
15	ALL1_HORSE	Major allergen Equ c 1	21.9	642	58	33	10	8	100
	TIMP1_HORSE	Metalloproteinase inhibitor 1	23.7	189	15	7	5	3	100
	ALL1_HORSE	Major allergen Equ c 1	21.9	1620	193	107	17	11	100
16	TSP1_HUMAN	Thrombospondin-1	133.3	351	58	26	13	9	98
10		Apolipoprotein A-1	30.2	124	24	11	4	3	80.8
	HBB_HORSE	Haemoglobin sub. beta	16.1	2153	159	114	14	12	100
17	HBA_HORSE	Haemoglobin sub. alpha	15.3	824	87	53	8	7	100
18	PIP_HYLSY	Prolactin-inducible protein homolog	16.9	122	9	8	1	1	64.3
10	111_111L01	r rotacuit-inductore protein noniolog	10.7	122	7	0	1	1	04.3

19	THIO_HORSE	Thioredoxin	12.0	80	12	2	4	1	100
20	PRG4_HUMAN ALBU_EQUAS	Proteoglycan 4	152.2	130	18	8	3	2	88.6
20	ALBU_EQUAS	Serum albumin	70.5	75	18	10	4	3	98.5

- <sup>a</sup> Number of the identified band as marked in Figure 3.
- <sup>b</sup> Protein entry name from UniProt knowledge database.
- 323 <sup>c</sup> Theoretical protein molecular weight.
- <sup>d</sup> The highest scores obtained with Mascot search engine.
- <sup>e</sup> Peptides: total number of peptides matching the identified proteins.
- <sup>f</sup> Significant peptides: total number of significant peptides matching the identified proteins.
- <sup>g</sup> Sequence: total number of distinct sequences matching the identified proteins.
- 328 <sup>h</sup> Significant sequences: total number of significant distinct sequences matching the identified
- 329 proteins.
- <sup>330</sup> <sup>i</sup> Percentage of identical amino acids between the identified protein and the respective horse
- 331 protein.

**Table 3.** Biological and functional classification of the proteins identified in equine amniotic

Entry name <sup>a</sup>	Protein full name	MW (kDa)	Biol. Proc. <sup>c</sup>	Mol. Funct. <sup>d</sup>	Cell. Comp.
HPT_BOVIN	Haptoglobin	45.6	Acute-phase response	HB binding	Extracellular
BGH3_HUMAN	Transforming growth factor-beta-induced protein ig-h3	75.3	Cell communication/ signal transduction	Receptor binding	Extracellular
CO6A1_HUMAN	Collagen alpha-1(VI) chain	109.6	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
CO6A3_HUMAN	Collagen alpha-3(VI) chain	345.2	Cell growth/ maintenance	peptidase inhibitor	Extracellular
CSPG2_BOV	Versican core protein	371.8	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
ECM1_HUMAN	Extracellular matrix protein 1	62.2	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
FBLN1_HUMAN	1	81.3	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
FINC_HORSE	Fibronectin	58.1	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
GELS_HORSE	Gelsolin	81.1	Cell growth/ maintenance	Structural constituent of cytoskeleton	Extracellular
IBP3_BOVIN	Insulin-like growth factor- binding protein	32.6	Cell growth/ maintenance	protein binding	Extracellular
LUM_MOUSE	Lumican	38.6	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
PRG4_HUMAN	Proteoglycan 4	152.2	Cell growth/ maintenance	Binding/Cell adhesion molecule	Extracellular
FIMP1_HORSE	Metalloproteinase inhibitor 1	23.7	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
TSP1_HUMAN	Thrombospondin-1	133.3	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
VINC_HUMAN	Vinculin	124.3	Cell growth/ maintenance	Cytoskeletal protein binding	Cytoplasm
CLUS_HORSE	Clusterin	52.7	Cell morphogenesis/ cell death	protein binding-chaperone	eCytoplasm
THIO_HORSE	Thioredoxin	12.0	Metabolism/ energy pathways	Catalytic activity	Cytoplasm
FETUA_BOVIN	Alpha-2-HS-glycoprotein	39.2	Mineral balance	protein binding	Extracellular
ACTB_BOVIN	Actin cytoplasmic 1	42.1	Protein folding	protein binding	Cytoskeletor
ACE2_PAGLA	Angiotensin-converting enzyme 2	93.0	Protein metabolism	Carboxylpeptidase	Plasma membrane
MMP2_BOVIN	72 kDa type IV collagenase		Protein metabolism	Metallopeptidase	Extracellular
PLMN_HORSE	Plasminogen	38.1		Peptidase	Extracellular
SFTPA_HORSE	Pulmonary surfactant- associated protein A	26.5	Respiratory gaseous exchange	carbohydrate/metal ion binding	Extracellular
ALBU_HORSE ALL1_HORSE	Serum albumin Major allergen Equ c 1	70.5 21.9	Transport Transport	Transporter Transporter	Extracellular Extracellular
	Apolipoprotein A-I	30.2	Transport	Binding	Extracellular HDL
FETA_HORSE	Alpha-fetoprotein	70.1	Transport	Transporter	Extracellula
HBA_HORSE	Haemoglobin subunit alpha	15.3	Transport	Transporter	Extracellular
HBB_HORSE	Haemoglobin subunit beta	16.1	Transport	Transporter	Extracellular
PLTP_HUMAN	Phospholipid transfer protein	54.9	Transport	Transporter	Extracellular
TRFE_HORSE TRFL_HORSE	Serotransferrin Lactotransferrin	80.3 77.9	Transport Transport	Transporter Transporter	Extracellular Secretory
					4 5

fluid. Identified proteins are listed according to the Biological Process category.

					granule
A1AT2_HORSE	Alpha-1-antiproteinase 2	47.1	Unknown	Protease inhibitor	NA
PIP_HYLSY	Prolactin-inducible protein homolog	16.9	Unknown	Binding	Extracellular

334

- <sup>a</sup> Protein entry name from UniProt knowledge database.
- 336 <sup>b</sup> Theoretical protein molecular weight.
- <sup>c</sup> Biological Process according to Gene Ontology and Human Protein Reference Database.
- <sup>d</sup> Molecular Function according to Gene Ontology and Human Protein Reference Database.
- <sup>e</sup> Cellular Component according to Gene Ontology and Human Protein Reference Database.
- 340

## 342 Figure Legends

Figure 1. Representative SDS-PAGE of 6 out of the 24 amniotic fluid samples on 4-12% gel in MOPS buffer. Twenty micrograms of proteins were loaded for each lane and the gel was stained with Coomassie Blue. A) Representative AF samples (lane 1, molecular weight marker; lanes 2-7, AF collected from six different healthy mares [samples 16-21]); B) representative pherogram obtained from lane 3

348

Figure 2. Relative abundance of each protein band compared to the entire volume of the lane; data are expressed as percentage (%) and reported as mean  $\pm$  standard deviation (n=24).

351

**Figure 3.** SDS-PAGE of the amniotic fluid pool on 4-12% gel in MOPS buffer. The pool was prepared by collecting and mixing 50  $\mu$ g of proteins from each AF sample (n=24). Two replicates are reported. Arrows and numbers indicate the slices that have been excised and analysed by ESI-Q-TOF as listed in Table 2. Asterisk (\*) indicates bands that did not give significant results by MS identification.

357

Figure 4. Distribution of amniotic fluid proteins in the Biological process category according
to Gene Ontology (GO) and the Human Protein Reference Database (HPRD) as reported in
Table 3.

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# 1 Identification of the most abundant proteins in equine amniotic fluid by a proteomic

- 2 approach
- 3
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- 20

## 21 Abstract

22 Characterisation of the physiologic equine amniotic fluid (AF) proteome is a prerequisite to 23 study its changes during diseases and discover new biomarkers. The aim of this study was to 24 identify by a proteomic approach the most abundant proteins of equine AF. AF samples were 25 collected at parturition from 24 healthy mares that delivered healthy foals. All samples were 26 subjected to sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) on 4-12% gels. A pool of the 24 samples, after SDS-PAGE, was cut in 25 slices, trypsin-digested 27 28 and analysed by mass spectrometry (MS) for protein identification. Mean AF protein 29 concentration was 1.96±1.12 g/L. Thirty-four proteins were successfully identified by MS and subsequently categorised according to Gene Ontology (GO). Twelve proteins (e.g. 30 31 fibronectin, lumican, thrombospondin and fibulin) belonged to or interacted with the 32 extracellular matrix (ECM) playing an important role in the development of foetal tissues. 33 Most of the remaining proteins were classified as transport (e.g. albumin, major allergen Equ 34 c1 and alpha-fetoprotein) delivering nutrients, ions and lipids essential for foetal growth and 35 development. Among these proteins, major allergen Equ c1 is widely studied in human 36 medicine because it induces Ig-E mediated type I allergic reaction. The absence of 37 immunoglobulins in equine AF was also confirmed. The present study successfully applied SDS-PAGE coupled to MS identifying the most abundant proteins of equine AF, highlighting 38 39 the importance of ECM and transport proteins. 40

41 Keywords

42 Horse, pregnancy, electrophoresis, proteome.

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44

### 46 Introduction

47 Amniotic fluid (AF) is a complex dynamic milieu that changes as pregnancy progresses. AF contains nutrients and growth factors that facilitate foetal growth, provides 48 49 mechanical cushioning and antimicrobial effectors to protect the foetus and allows assessment of foetal maturity and disease (Underwood et al. 2005). In comparison to humans, the 50 51 physiology and pathophysiology of foetal fluids in domestic mammals are poorly understood 52 (Canisso et al. 2015). In horses, some studies have investigated biochemical composition, particularly enzymes and electrolytes, in AF collected by ultrasound-guided transabdominal 53 54 amniocentesis, at delivery or at slaughter (Holdstock et al. 1995; Lyle et al. 2006; Williams et 55 al. 1993; Zanella et al. 2014). AF was also studied for evaluation of foal's lung maturity at birth through lecithin/sphingomyelin ratio and lamellar body count (Castagnetti et al. 2007). 56 57 More rRecently, significantly higher levels of lactate were found in AF collected during 58 parturition in mares delivering healthy foals (Pirrone et al. 2012).

Unlike the allantoic fluid, equine AF can be easily collected during parturition without
stressing the animal and avoiding any contamination (Castagnetti et al. 2007, Pirrone et al.
2012). As reported in women, the biochemical composition of AF, including proteins, is
primarily representative of the foetal profile and reflects its physiological status (Tong 2013),
thus it could be potentially useful to evaluate the high-risk foal born attended.

Proteomics is a powerful analytical approach providing a profile of proteins present in a biological sample at a given time. The high potential of this approach has been recently found to have a major role in different areas of veterinary medicine, from farm (Almeida et al. 2015) to companion (Ferlizza et al. 2015; Miller et al. 2014) animals. Proteomic techniques have recently been applied to the characterisation of horse amniotic membrane (Galera et al. 2015) and bovine conceptus fluids (Riding et al. 2008), whereas the equine AF proteome remains uncharacterised. Therefore, the aims of this study were to identify the most abundant

- proteins in equine AF by SDS-PAGE separation followed by mass spectrometry identification
   and discuss their functions.
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# 74 Materials and Methods

#### 75 Animal selection and data collection

Twenty-four mares admitted for assisted delivery during three breeding seasons at the S. Belluzzi Equine Perinatology Unit of the Department of Veterinary Medical Sciences, University of Bologna, were included. The mares were hospitalised at about 310 days of pregnancy because the owners requested an attended parturition, and remained under observation for at least 7 days *postpartum*. They were housed in wide straw bedding boxes and fed with hay *ad libitum* and concentrates twice a day. During the day, the mares were allowed to go to pasture.

83 All the mares included in the study were healthy based on clinical and 84 ultrasonographic evaluation. At admission, a complete clinical evaluation, including complete 85 blood count, serum biochemistry and transrectal ultrasonography, was performed. Severe 86 maternal illness, uterine discharge, premature lactation, twinning, abnormal foetal presentation, placenta oedema, and signs of foetal distress were ruled out. During the course 87 of hospitalisation, mares were clinically evaluated twice a day and by transrectal 88 89 ultrasonography every 10 days until parturition. The following ultrasonographic parameters 90 were evaluated: combined thickness of the uterus and placenta, foetal fluids echogenicity, 91 foetal activity, and foetal orbital area. Foals were born between 320 and 365 days of pregnancy, by normal delivery, had an Apgar score  $\geq 8$  recorded within 5 minutes from birth 92 93 (Vaala et al. 2002) and had a normal clinical evaluation during the course of hospitalisation, 94 including a complete blood count and serum biochemistry at birth and an immunoglobulin G 95 (IgG) serum concentration  $\geq$ 800 mg/dL at 18-24 hours of life.

96 For each mare, the following data were recorded: breed, age, parity, days of

97 pregnancy, body weight, length of stage II labour (minutes), and foal's body weight and

98 Apgar score. All procedures on the animals were carried out with the approval of the Ethical

99 Committee, in accordance with DL 116/92, approved by the Ministry of Health (approval

100 number: n.18/64/11; date of approval 22/02/2011). Oral informed consent was given by the

101 owners.

102

103 Sample collection

At foaling, a sample of AF was collected from each mare with a 50 mL syringe by needle puncture of the amniotic sac within few minutes of its appearance through the vulva during stage II of labour. The AF was then immediately transferred to 5 mL test tubes and stored at -80°C until SDS-PAGE and protein identification were performed. AF protein concentration was determined by the Biuret method using bovine serum albumin as standard.

109

110 SDS-PAGE

111 To optimise protein separation, different protocols were tested including 4-12% and 12% polyacrylamide gels (NuPage/Thermo Fisher Scientific, Waltham, Massachusetts, USA) 112 in 2-(N-morpholino) propanesulfonic acid buffer (MOPS; NuPage/Thermo Fisher Scientific) 113 114 or 2-(N-morpholino) ethanesulfonic acid buffer (MES; NuPage/Thermo Fisher Scientific) 115 with sodium dodecyl sulphate (SDS). Each AF sample (n=24) was analysed at least twice 116 with the protocol assuring the best protein separation in our experimental conditions (4-12%) 117 gels, in MOPS buffer). Twenty µg of proteins were loaded for each sample and the gels were 118 stained with Coomassie G250 compatible with mass spectrometry analysis. After staining, each gel was digitalised by ChemiDoc<sup>TM</sup>MP (BioRad, Hercules, California, USA) and its 119 120 pherogram was obtained using ImageLab 5.2.1 software (BioRad). The software determines 121 the volume of each protein band through the analysis of the pixel values in the digital image, meaning as volume the sum of all the pixels intensities within the band boundaries. The band volumes are subsequently compared to the entire volume of the lane and the relative abundances reported in percentage. A pool was prepared by collecting and mixing 50  $\mu$ g of protein from each AF (n=24) and analysed twice with the same protocol used for each sample.

126

#### 127

# Protein identification by mass spectrometry

128 The pool lanes were divided manually into 25 slices and subjected to in-gel tryptic 129 digestion as previously described (Bellei et al. 2013). Digested dried samples were then re-130 suspended in 97% Water/3% ACN added of 1% formic acid (Buffer A) and analysed by a Nano LC-CHIP-MS system (ESI-Q-TOF 6520; Agilent Technologies, Santa Clara, 131 132 California, USA). Four microliters of each sample were loaded into the system and transported to the Chip enrichment column (Zorbax C18, 4 mm x 5 µm i.d., Agilent 133 134 Technologies) by a capillary pump, with a loading flow of 4 µL/min, using 95% 135 ACN/5% water added of 0.1% formic acid (buffer B) as mobile phase. Nitrogen was used as 136 the nebulising gas. A separation column (Zorbax C18, 43 mm x 75 µm i.d., Agilent 137 Technologies), at flow rate of  $0.4 \,\mu$ L, was used for peptide separation.

Since the horse protein database is not well annotated, a broader taxonomy, namely 138 "all mammals", was selected for identification to be based on sequence homology. Protein-139 140 identification peak lists were generated using the Mascot search engine 141 (http://mascot.cigs.unimo.it/mascot) against the UniProt database (UniProt.org) specifying the 142 following parameters: mammalian taxonomy, parent ion tolerance ±20 ppm, MS/MS error 143 tolerance  $\pm 0.12$  Da, alkylated cysteine as fixed modification and oxidised methionine as 144 variable modification, and two potential missed trypsin cleavages, as previously described 145 (Bertoldi et al., 2013). Proteins with a score >80 or identified with at least two or more 146 significant sequences were selected. The significant threshold in Mascot searches was set to

obtain a false discovery rate <5% (5% probability of false match for each protein with a score</li>
above 80).

149

150 Statistical analysis

Data (AF total proteins, mare's age, mare's and foal's body weight, parity, days of pregnancy, length of stage II labour (minutes), foal's Apgar score and number of bands) were analysed with statistical software (R version 2.15.1) and reported as mean ± standard deviation (SD). Shapiro-Wilk normality test was performed to evaluate data normal distribution. Pearson coefficient of correlation was calculated between AF total proteins and the other data recorded for each mare (mare's age, mare's and foal's body weight, parity, days of pregnancy, length of stage II labour (minutes), foal's Apgar score and number of bands).

The identified proteins were categorised by biological process, molecular function and
cellular component with Gene Ontology terms according to Gene Ontology (GO) and Human
Protein Reference Database (HPRD).

161

162 **Results** 

163 *Clinical data* 

164 Clinical data collected from the 24 mares included in the study are shown in Table 1. Mean 165 AF total protein concentration was 1.96±1.12 g/L, ranging varied from 0.36 to 4.16 g/L. No 166 significant correlations were found between AF protein concentration and the other data 167 recorded.

168

169 SDS-PAGE and protein identification by mass spectrometry

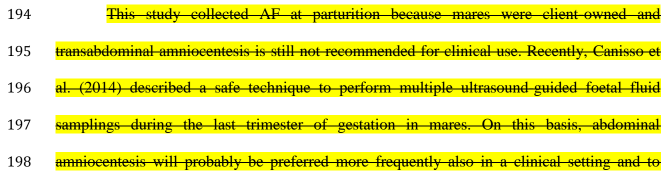
170 Representative gel and pherogram of AF are reported in Figure 1A and 1B. The mean 171 number of bands was 23±1.5. All samples presented a similar pattern characterised by two 172 clusters of bands: the first with molecular weights (MW) higher than 62 kDa and the second with MW lower than 34 kDa. In the middle, very few faint bands were present. Figure 2 and Table 2 reports the relative abundance in percentage of AF protein bands. Out of the 25 slices cut from the gel (Figure 3), 20 yielded significant results leading to the unambiguous identification of 34 proteins (Table 2 3). Serum albumin and major allergen Equ c1 (ALL1) were the two most abundant proteins, followed by fibronectin, transferrin and haemoglobin; these five proteins represented >60% of the equine AF proteome. Fibronectin, versican and albumin were also identified in bands characterised by different MW.

The identified proteins categorised by their molecular function and biological process according to Gene Ontology (GO) and Human Protein Reference Database (HPRD) are shown in Table 3 **4** and Figure 4. Most of the proteins were involved in cellular growth and/or maintenance (38%), transport (26%) and protein metabolism (9%). The vast majority of the identified proteins were classified as extracellular (79%).

185

#### 186 **Discussion**

187 The present paper aimed to explore the complexity of equine AF proteome and to 188 identify its most abundant proteins. Since this The study was carried out on 24 mares of 189 different breeds, age and parity referred to the Equine Perinatology Unit, the animals were of 190 different breed, age and parity, and they can be considered representative of a typical equine 191 hospital population reflecting the local equine population. Therefore, the proteomic profile 192 described can be considered a useful starting point for further applied studies on the equine 193 AF proteins.



- 199 evaluate gestational changes in the AF proteome as in women (Michaels et al. 2007).
- 200 Regarding total proteins, values found in the present research (1.96±1.12 g/L) are similar to
- 201 those reported by Williams et al. (1993) (3.1±2.6 g/L) and Paccamonti et al. (1995) (1-2 g/L),
- 202 but lower than those reported by Kochhar et al. (1997) (9.1±2 g/L) and higher than those of
- 203 Zanella et al. (2014) (0.3±0.1 g/L). The use of different methods for AF total proteins
- 204 quantification or the wide inter individual variability in concentration could be the cause of
- 205 the reported discrepancies.
- 206 Complex biological samples contain thousands of different protein species, few of them characterised by high abundance and many others by low or very low abundance. The 207 208 presence of very high abundance proteins like albumin or immunoglobulins often hampers the 209 separation and characterisation of serum and AF proteomes, therefore the depletion of these major components has been widely applied in human proteomics (Cho et al. 2007; Michaels 210 211 et al. 2007). However, this approach can lead to the loss of some low abundance proteins due 212 to the "sponge effect" of albumin that can bind a variety of other proteins or peptides (Bellei 213 et al. 2011). From this point of view, equine AF is a preferential sample due to the absence of 214 immunoglobulins. In domestic animals, the passage of immunoglobulins is influenced by the placental structure: in horses, pigs and ruminants the placenta is epitheliochorial, thus 215 216 impermeable to immunoglobulins (Furukawa et al. 2014), whereas in dogs, the 217 endotheliochorial placenta allows only 5% to 10% transfer of maternal antibodies to the 218 foetus (Dall'Ara et al. 2015). The absence of immunoglobulins in equine AF, as confirmed by 219 this study, was advantageous for its more reliable characterisation, as a consequence to 220 preserve proteome integrity and complexity the most abundant proteins were not depleted.

Most of the 34 proteins identified were involved in cellular growth and maintenance, transport and protein metabolism reflecting the dynamic biological functions of AF. Regarding cellular growth and/or maintenance, 12 of the proteins identified belonged to or interacted with the extracellular matrix (ECM) that plays an important role in the development 225 of foetal tissues. All these proteins, with the exception of versican and proteoglycan 4, were 226 also identified in human AF (Cho et al. 2007; Michaels et al. 2007) and/or in equine amniotic 227 membrane (Galera et al. 2015). Among the ECM structural proteins, fibronectin is a 228 multifunctional glycoprotein known to participate in the organisation of ECM binding to 229 integrins. During pregnancy, fibronectin is expressed in the junction between maternal and 230 foetal membranes as well as in the uterus and placenta (Mogami et al. 2013). Lumican, a 231 member of the family of small leucine-rich proteoglycans, is the major keratan sulphate 232 proteoglycan of the cornea and is also present in the ECM throughout the body, including 233 human and equine amniotic membrane (Galera et al. 2015; Kao et al. 2006; Mavrou et al. 234 2008). Based on its interaction with fibrillar collagen and its ability to modulate cell proliferation and migration, lumican could play a role in the maturation of foetal tissues 235 236 (Mavrou et al. 2008). Regarding the non-structural proteins involved in ECM development 237 and organisation, thrombospondin and fibulin are regulatory proteins belonging to the group 238 of the matricellular proteins. These proteins represent a bridge between matrix proteins and 239 cell surface receptors, or other molecules such as cytokines that can interact with the cell 240 surface (Bornstein, 1995). They are typically expressed at low levels in adult tissues, but are strongly expressed during development or following injury or pathology (Morris and 241 Kyriakides, 2014). Gelsolin is a multifunctional actin regulatory protein involved in 242 243 cytoskeleton dynamics and structure. In addition to its role in aiding chemotaxis and 244 movement of intracellular structures, plasma gelsolin binds to a variety of proinflammatory 245 and bioactive molecules including fibronectin, platelet activating factor and the bacterial 246 surface lipids lipoteichoic acid and lipopolysaccharide (Peddada et al. 2012). The role of 247 gelsolin in AF is still unknown, but it has been suggested to modulate inflammation and 248 bacterial infections in human AF (Sezen et al. 2009). In association with gelsolin, vinculin is 249 also a component of the actine cytoskeleton and is involved in integrin-mediated focal 250 adhesion, cell motility and other cellular functions such as migration, proliferation and

251 differentiation (Wu et al. 2014). Other proteins interacting with ECM that could play 252 important roles in the development of foetal tissue are type IV collagenase (MMP2) and 253 metalloproteinase inhibitor 1 (TIMP1) belonging to the matrix metalloproteinases (MMPs) 254 and tissue inhibitors respectively. The MMPs are a family of over 20 enzymes acting on the 255 ECM components, regulated at different levels via their activators, inhibitors and localization 256 on the cell surface (Sternlicht and Werb, 2001). The biological functions of these enzymes 257 and their inhibitors have been widely studied, in particular MMP2 is important for bone 258 development and angiogenesis regulation and has been identified and studied in plasma and 259 AF of pregnant women (Anumba et al. 2010; Turner et al. 2014). MMPs activity was also 260 studied in amniotic and allantoic fluid from mares that delivered live term foals and from 261 mares with preterm delivery, suggesting that MMPs may have a role as markers for high risk 262 pregnancy in the mare (Oddsdóttir et al. 2011).

263 Among transport proteins, albumin, transferrin, alpha-fetoprotein, apolipoprotein A1 264 and phospholipid transfer protein (PLTP) transport nutrients, ions and lipids essential for 265 foetal growth and development and have been identified as common components of AF also 266 in humans (Cho et al. 2007; Michaels et al. 2007). Alpha-fetoprotein is member of the 267 albuminoid superfamily, and is present in the allantoic  $\frac{\text{fluid}}{\text{fluid}}$  and  $\frac{\text{AF}}{\text{AF}}$  amniotic fluids of 268 domestic animals (Luft et al. 1984; Smith et al. 1979). In mammalian foetuses alpha-269 is associated with oestrogen-binding, anti-oxidative fetoprotein properties and 270 immunoregulation (DeMees et al. 2006; Mizejewski 2001) and In horses, alpha-fetoprotein it 271 is highly expressed during early pregnancy by the equine conceptus (Simpson et al. 2000) and 272 in mammalian foetuses, it is associated with estrogen binding, anti-oxidative properties and immunoregulation (DeMees et al. 2006; Mizejewski 2001). In women, AF alpha-fetoprotein 273 274 is actively investigated for pathologies such as Down syndrome, trisomies 13 and 18, intra-275 amniotic infection, preterm delivery, pre-eclampsia, membrane rupture, and foetoplacental 276 hypoxia (Cho et al. 2007). Recently, Canisso et al. (2015) confirmed the presence of alpha277 fetoprotein in equine foetal fluids during the third trimester of pregnancy and found increased 278 maternal plasma concentrations of the protein in mares with experimentally induced 279 placentitis. The presence of ALL1 in AF is challenging. ALL1 is a glycoprotein of 21.7 kDa 280 belonging to the family of lipocalins, whose function is to carry small hydrophobic molecules 281 such as odorants, steroids and pheromones. This protein is expressed in salivary glands and in 282 the liver and is highly concentrated in secretory fluids such as saliva and urine as well as in 283 hair and dander (Botros et al. 2001). ALL1 is widely studied in human medicine because it 284 induces an IgE-mediated type I allergic reaction in the majority of patients allergic to horses 285 (Lascombe et al. 2000). The physiological role of this protein is still unknown and to our 286 knowledge, this is the first study reporting the presence of ALL1 in AF. PLTP is a monomeric 287 glycoprotein involved in lipid transport, lipoprotein metabolism and lipopolysaccharide 288 binding. It is ubiquitously expressed in human tissues and is secreted into the plasma, where 289 its central role has been well established (Albers et al. 2012). PLTP is highly expressed in 290 lung epithelial cells, and may play a role in surfactant metabolism during foetus lung 291 development (Brehm et al. 2014).

292 The proteomic approach applied in the present study led to the successful 293 identification of the most abundant proteins, even though a few additional points should be 294 taken into consideration. The first one regards the choice of non-depleting albumin and other 295 major proteins before electrophoresis and MS identification. Complex biological samples 296 contain thousands of different protein species, few of them characterised by high abundance 297 and many others by low or very low abundance. The presence of very high abundance 298 proteins like albumin and immunoglobulins often hampers the separation and characterisation 299 of serum and AF proteomes, therefore the depletion of these major components has been 300 applied in human proteomics (Cho et al. 2007; Michaels et al. 2007). However, this approach 301 can lead to the loss of some low abundance proteins due to the "sponge effect" of albumin 302 that can bind a variety of other proteins or peptides (Bellei et al. 2011). From this point of

303 view, equine AF is a preferential sample due to the absence of immunoglobulins. In domestic 304 animals, the passage of immunoglobulins is influenced by the placental structure: in horses, 305 pigs and ruminants the placenta is epitheliochorial, thus impermeable to immunoglobulins 306 (Furukawa et al. 2014), whereas in dogs, the endotheliochorial placenta allows only 5% to 307 10% transfer of maternal antibodies to the foetus (Dall'Ara et al. 2015). From an analytical 308 point of view, the absence of immunoglobulins in equine AF, as confirmed by this study, was advantageous for its more reliable characterisation, as a consequence to preserve proteome 309 310 integrity and complexity the most abundant proteins were not depleted, can be considered an 311 advantage, allowing to perform SDS-PAGE and MS identification without affecting proteome 312 integrity and complexity.

The second point is related to the sample collection. This study collected AF only at parturition because mares were client-owned and transabdominal amniocentesis is still not recommended for clinical use. Recently, Canisso et al. (2014) described a safe technique to perform multiple ultrasound-guided foetal fluid samplings during the last trimester of gestation in mares. On this basis, abdominal amniocentesis will probably be preferred more frequently also in a clinical setting and to evaluate gestational changes in the AF proteome as reported in women (Michaels et al. 2007).

320 The last point regards **Regarding** AF total protein concentrations, which values found 321 in the present research (1.96±1.12 g/L) with a wide variability (range 0.36-4.16 g/L) are were 322 similar to those reported by Williams et al. (1993) (3.1±2.6 g/L) and Paccamonti et al. (1995) (1-2 g/L), but lower than those reported by Kochhar et al. (1997) (9.1 $\pm$ 2 g/L), and higher than 323 324 those of Zanella et al. (2014) (0.3±0.1 g/L). The use of different methods for AF total proteins quantification or the wide inter-individual variability in concentration could be the cause of 325 326 the reported discrepancies. The reported discrepancies could be related to the use of 327 quantification methods characterised by different analytical performances; also the influence 328 of wide inter-individual variability cannot be excluded. Many environmental and

physiological factors can contribute to this variability; in particular, since the mares were
client-owned, pre-hospitalisation conditions, such as housing, feeding, nutrition and hydration
status, might have affected AF total protein concentration.

332

333 Conclusions

334 Applying a qualitative proteomic approach, this study identified the 34 most abundant 335 proteins of the AF proteome from healthy mares that delivered live term foals The present 336 applyied a qualitative proteomic approach to characterise the AF proteome from healthy 337 mares. The 34 proteins identified GO categorisation demonstrated that these proteins are 338 involved in different biological processes and molecular functions including cell 339 growth/maintenance and transport. Some of these proteins belonged to or interacted with the 340 extracellular matrix, highlighting the role of its the ECM components in foetal maturation. 341 The study confirmed also the importance of transport proteins like alpha-fetoprotein and 342 PLTP, and reported for the first time the presence of ALL1 in AF. Further studies are needed 343 to define reference intervals for AF total protein in healthy mares and Though entirely 344 descriptive, these findings can be considered valuable context for further investigations to gain insights into the function of the proteins identified and to discover potential biomarkers 345 of foetal disease at birth or during pregnancy. In particular, 2DE could be applied to better 346 characterise the physiologic AF proteome and to evaluate differentially expressed proteins in 347 348 <del>case of disease.</del> 349

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353

354 **Conflict of interest** 

355 The authors have no conflict of interests to declare.

**Table 1.** Clinical data collected from the 24 mares included in the study. Data are reported as

358	mean $\pm$ standard deviation. AF amniotic fluid; TP total proteins.
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Sample	Breed	Age	Mare weight	Foal weight	Parity	Length of pregnancy	Length of stage II	Foal's Apgar score	AF TP
		years	Kg	Kg		days	minutes		g/L
1	Saddlebred	11	660	51.7	1	355	25	10	2.74
2	Thoroughbred	6	645	51	1	344	20	10	1.66
3	Standardbred	5	565	50	2	335	14	8	1.51
4	Standardbred	6	585	44	1	329	8	8	3.07
5	Saddlebred	7	660	53	2	330	16	9	2.64
6	Saddlebred	12	500	40	1	360	20	9	1.01
7	Saddlebred	17	650	58	3	330	13	9	0.55
8	Quarter Horse	11	560	42.6	5	333	14	9	0.58
9	Saddlebred	14	660	53.5	1	354	20	10	0.36
10	Arabian	5	450	42	2	335	15	10	0.90
11	Standardbred	7	565	45.3	1	328	11	10	1.39
12	Standardbred	19	578	43.5	3	349	12	8	1.25
13	Standardbred	12	546	45	2	352	15	8	1.30
14	Standardbred	16	590	42.5	11	347	9	8	1.73
15	Standardbred	10	535	50	6	343	8	10	0.58
16	Standardbred	14	610	45	6	336	21	9	2.50
17	Arabian	12	430	50	6	326	20	10	2.29
18	Standardbred	6	620	59	1	341	18	10	1.66
19	Standardbred	20	606	45	12	338	12	10	4.01
20	Saddlebred	16	650	47	2	360	5	8	3.01
21	Standardbred	18	680	50	4	332	9	9	4.16
22	Thoroughbred	13	580	58	4	354	12	8	2.14
23	Quarter Horse	16	425	41	3	357	17	9	4.16
24	Standardbred	19	660	56	3	345	25	8	1.75
		12 ± 9.5	583.8 ± 74.1	48.7 ± 11.4	3 ± 3	342 ± 10.7	15 ± 5.7	9 ± 1	1.96 ± 1.12

- 361 Table 2. Relative abundance in percentage of each protein band compared to the entire
- 362 volume of the lane (n=24).
- 363

<mark>MW</mark>	<mark>Mean</mark>	<mark>SÐ</mark>
<mark>(kDa)</mark>	<mark>(%)</mark>	<mark>-(%)</mark>
<mark>&gt;165</mark>	<mark>0.3</mark>	<mark>0.2</mark>
<mark>&gt;165</mark>	<mark>1.5</mark>	<mark>0.4</mark>
<mark>&gt;165</mark>	<mark>5.2</mark>	<mark>1.9</mark>
<mark>&gt;165</mark>	<mark>1.4</mark>	<mark>0.3</mark>
<mark>-145</mark>	<mark>0.3</mark>	<mark>0.2</mark>
<mark>131</mark>	<mark>0.1</mark>	<mark>0.1</mark>
<mark>117</mark>	<mark>0.6</mark>	<mark>0.1</mark>
<mark>102</mark>	<mark>0.6</mark>	<mark>0.2</mark>
<mark>91</mark>	<mark>0.7</mark>	<mark>0.2</mark>
<mark>83</mark>	<mark>3.6</mark>	<mark>0.7</mark>
<mark>73</mark>	<mark>2.0</mark>	<mark>0.9</mark>
<mark>64</mark>	<mark>25.7</mark>	<mark>2.8</mark>
<mark>53</mark>	<mark>1.4</mark>	<mark>0.5</mark>
<mark>41</mark>	<mark>1.4</mark>	<mark>0.5</mark>
<mark>35</mark>	<mark>0.7</mark>	<mark>0.2</mark>
<mark>29</mark>	<mark>1.6</mark>	<mark>0.5</mark>
<mark>24</mark>	<mark>19.2</mark>	<mark>3.6</mark>
<mark>21</mark>	<del>0.7</del>	<mark>0.2</mark>
<mark>18</mark>	<mark>4.2</mark>	<mark>1.2</mark>
<mark>14</mark>	<mark>3.0</mark>	<mark>2.9</mark>
<mark>&lt;13</mark>	<mark>3.4</mark>	<mark>1.0</mark>
<mark>&lt;13</mark>	<mark>2.8</mark>	<del>1.2</del>
<mark>&lt;13</mark>	<mark>4.0</mark>	<mark>0.8</mark>



ınd	<sup>a</sup> Entry name <sup>b</sup>	Protein full name	MW (kDa) <sup>c</sup>	Score <sup>d</sup>	Pept. <sup>e</sup>	Sign. Pept. <sup>f</sup>	Seq. <sup>g</sup>	Sign Seq. <sup>h</sup>	% i
1	CSPG2_BOVIN	Versican core protein	371.8	257	76	21	15	6	77.
2	FINC_HORSE	Fibronectin	58.1	1451	155	89	19	18	10
3	FINC_HORSE	Fibronectin	58.1	440	76	34	15	10	10
4	FINC_BOVIN	Fibronectin	275.5	155	46	16	16	7	95
4	CO6A1_HUMAN	Collagen alpha-1(VI) chain	109.6	168	41	16	10	7	8
5	VINC_HUMAN	Vinculin	124.3	148	53	18	26	10	99
	ACE2_PAGLA	Angiotensin-converting enzyme 2	93.0	380	43	18	10	6	8
6	PLMN_HORSE	Plasminogen	38.1	288	36	18	8	5	10
	FINC_CANFA	Fibronectin	58.2	148	18	11	7	5	96
	GELS_HORSE	Gelsolin	81.1	1911	193	122	27	21	1(
-	FBLN1_HUMAN	Fibulin-1	81.3	434	80	47	11	9	91
7	FINC_HORSE	Fibronectin	58.1	360	36	23	11	7	1(
	LUM_MOUSE	Lumican	38.6	159	27	14	6	4	87
_	TRFE_HORSE	Serotransferrin	80.3	2201	263	153	43	32	1(
8	TRFL_HORSE	Lactotransferrin	77.9	268	61	30	19	11	1(
9	TRFE HORSE	Serotransferrin	80.3	1642	194	106	34	24	1(
/	ALBU_HORSE	Serum albumin	70.5	2863	244	148	44	33	10
		Extracellular matrix protein 1	62.2	188	53	22	4	4	78
	MMP2 BOVIN	72 kDa type IV collagenase	74.8	468	50	27	- 14	9	95
10	PLTP HUMAN	Phospholipid transfer protein	54.9	400	24	19	3	3	89
10	LUM_MOUSE	Lumican	38.6	400 153	24	19	6	4	87
			58.0 70.1	135	23 17	12		4	10
	FETA_HORSE	Alpha-fetoprotein					7		
		Versican core protein	96.8	101	13	7	4	3	8
	ALBU_HORSE	Serum albumin	70.5	17663	1154	849	70	56	1(
1.1	FETA_HORSE	Alpha-fetoprotein	70.1	407	59	34	23	14	10
11	CSPG2_MACNE	Versican core protein	96.8	347	13	10	5	4	8
	BGH3_HUMAN	Transforming growth factor-beta- induced protein ig-h3	75.3	53	12	6	4	3	92
12	A1AT2_HORSE	Alpha-1-antiproteinase 2	47.1	195	11	8	4	2	10
	ALBU_EQUAS	Serum albumin	70.5	160	53	13	23	9	98
	CLUS_HORSE	Clusterin	52.7	453	48	28	15	10	10
	ACTB_BOVIN	Actin cytoplasmic 1	42.1	118	36	11	14	6	10
	GELS_HORSE	Gelsolin	81.1	40	12	4	7	4	10
13	IBP3_BOVIN	Insulin-like growth factor-binding protein	32.6	79	11	9	5	5	86
	FETUA BOVIN	Alpha-2-HS-glycoprotein	39.2	65	11	6	2	2	71
		Collagen alpha-3(VI) chain	345.2	93	5	4	3	2	88
	HPT_BOVIN	Haptoglobin	45.6	52	3	2	3	2	78
	CLUS_HORSE	Clusterin	52.7	42	18	4	7	3	10
14	TSP1_BOVIN	Thrombospondin-1	133.4	29	13	3	9	3	96
14	SFTPA_HORSE	Pulmonary surfactant-associated protein A	26.5	71	4	3	3	2	1(
	TSP1_HUMAN	Thrombospondin-1	133.3	751	60	49	12	11	9
15	ALL1_HORSE	Major allergen Equ c 1	21.9	642	58	33	10	8	1(
-	TIMP1_HORSE	Metalloproteinase inhibitor 1	23.7	189	15	7	5	3	10
	ALL1_HORSE	Major allergen Equ c 1	21.9	1620	193	107	17	11	1(
		Thrombospondin-1	133.3	351	58	26	13	9	9
16	TSP1 HUMAN		100.0				4		80
16	TSP1_HUMAN APOA1 CANFA	-	30.2	124	24	11	4	<b>)</b>	
	APOA1_CANFA	Apolipoprotein A-1	30.2	124 2153	24	11 114		3	
16 17	APOA1_CANFA HBB_HORSE	Apolipoprotein A-1 Haemoglobin sub. beta	16.1	2153	159	114	14	12	1(
	APOA1_CANFA	Apolipoprotein A-1							10 10 64

Table 2 3. Proteins identified in equine amniotic fluid by mass spectrometry. Identified
proteins are listed according to the number of the band as marked in Figure 3.

20	PRG4_HUMAN	Proteoglycan 4	152.2	130	18	8	3	2	88.6
20	ALBU_EQUAS	Serum albumin	70.5	75	18	10	4	3	98.5

- <sup>a</sup> Number of the identified band as marked in Figure 3.
- 369 <sup>b</sup> Protein entry name from UniProt knowledge database.
- 370 <sup>c</sup> Theoretical protein molecular weight.
- <sup>d</sup> The highest scores obtained with Mascot search engine.
- <sup>e</sup> Peptides: total number of peptides matching the identified proteins.
- <sup>f</sup> Significant peptides: total number of significant peptides matching the identified proteins.
- <sup>g</sup> Sequence: total number of distinct sequences matching the identified proteins.
- <sup>375</sup> <sup>h</sup> Significant sequences: total number of significant distinct sequences matching the identified
- 376 proteins.
- <sup>377</sup> <sup>i</sup> Percentage of identical amino acids between the identified protein and the respective horse
- 378 protein.

## **Table 3 4.** Biological and functional classification of the proteins identified in equine

Entry name <sup>a</sup>	Protein full name	MW (kDa)	Biol. Proc. <sup>c</sup>	Mol. Funct. <sup>d</sup>	Cell. Comp.
HPT_BOVIN	Haptoglobin	45.6	Acute-phase response	HB binding	Extracellular
BGH3_HUMAN	Transforming growth factor-beta-induced protein ig-h3	75.3	Cell communication/ signal transduction	Receptor binding	Extracellular
CO6A1_HUMAN	Collagen alpha-1(VI) chain	109.6	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
CO6A3_HUMAN	Collagen alpha-3(VI) chain	345.2	Cell growth/ maintenance	peptidase inhibitor	Extracellular
CSPG2_BOV	Versican core protein	371.8	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
ECM1_HUMAN	Extracellular matrix protein 1	62.2	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
FBLN1_HUMAN	Fibulin-1	81.3	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
FINC_HORSE	Fibronectin	58.1	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
GELS_HORSE	Gelsolin	81.1	Cell growth/ maintenance	Structural constituent of cytoskeleton	Extracellular
IBP3_BOVIN	Insulin-like growth factor- binding protein	32.6	Cell growth/ maintenance	protein binding	Extracellular
LUM_MOUSE	Lumican	38.6	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
PRG4_HUMAN	Proteoglycan 4	152.2	Cell growth/ maintenance	Binding/Cell adhesion molecule	Extracellular
TIMP1_HORSE	Metalloproteinase inhibitor 1	23.7	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
TSP1_HUMAN	Thrombospondin-1	133.3	Cell growth/ maintenance	Extracellular matrix structural constituent	Extracellular
VINC_HUMAN	Vinculin	124.3	Cell growth/ maintenance	Cytoskeletal protein binding	Cytoplasm
CLUS_HORSE	Clusterin	52.7	Cell morphogenesis/ cell death	protein binding-chaperon	eCytoplasm
THIO_HORSE	Thioredoxin	12.0	Metabolism/ energy pathways	Catalytic activity	Cytoplasm
FETUA_BOVIN	Alpha-2-HS-glycoprotein	39.2	Mineral balance	protein binding	Extracellular
ACTB_BOVIN	Actin cytoplasmic 1	42.1	Protein folding	protein binding	Cytoskeleton
ACE2_PAGLA	Angiotensin-converting enzyme 2	93.0	Protein metabolism	Carboxylpeptidase	Plasma membrane
MMP2_BOVIN	72 kDa type IV collagenase	74.8	Protein metabolism	Metallopeptidase	Extracellular
PLMN_HORSE	Plasminogen	38.1	Protein metabolism	Peptidase	Extracellular
SFTPA_HORSE	Pulmonary surfactant- associated protein A	26.5	Respiratory gaseous exchange	carbohydrate/metal ion binding	Extracellular
ALBU_HORSE ALL1_HORSE	Serum albumin Major allergen Equ c 1	70.5 21.9	Transport Transport	Transporter Transporter	Extracellular Extracellular
APOA1_CANFA	Apolipoprotein A-I	30.2	Transport	Binding	Extracellular HDL
FETA_HORSE	Alpha-fetoprotein	70.1	Transport	Transporter	Extracellular
HBA_HORSE	Haemoglobin subunit alpha	15.3	Transport	Transporter	Extracellular
HBB_HORSE	Haemoglobin subunit beta	16.1	Transport	Transporter	Extracellular
PLTP_HUMAN	Phospholipid transfer protein	54.9	Transport	Transporter	Extracellular
TRFE_HORSE	Serotransferrin	80.3	Transport	Transporter	Extracellular
TRFL_HORSE	Lactotransferrin	77.9	Transport	Transporter	Secretory

# amniotic fluid. Identified proteins are listed according to the Biological Process category.

					granule
A1AT2_HORSE	Alpha-1-antiproteinase 2	47.1	Unknown	Protease inhibitor	NA
PIP_HYLSY	Prolactin-inducible protein homolog	16.9	Unknown	Binding	Extracellular

381

- <sup>a</sup> Protein entry name from UniProt knowledge database.
- <sup>b</sup> Theoretical protein molecular weight.
- <sup>c</sup> Biological Process according to Gene Ontology and Human Protein Reference Database.
- <sup>d</sup> Molecular Function according to Gene Ontology and Human Protein Reference Database.
- <sup>e</sup> Cellular Component according to Gene Ontology and Human Protein Reference Database.

387

389 Figure Legends

Figure 1. Representative SDS-PAGE of 6 out of the 24 amniotic fluid samples on 4-12% gel in MOPS buffer. Twenty micrograms of proteins were loaded for each lane and the gel was stained with Coomassie Blue. A) Representative AF samples (lane 1, molecular weight marker; lanes 2-7, AF collected from six different healthy mares [samples 16-21]); B) representative pherogram obtained from of lane 3

395

Figure 2. Relative abundance in percentage of each protein band compared to the entire
volume of the lane; data are expressed as percentage (%) and reported as mean ± standard
deviation (n=24).

399

Figure 3. SDS-PAGE of the amniotic fluid pool on 4-12% gel in MOPS buffer. Two replicates of the pool was prepared by collecting and mixing 50  $\mu$ g of proteins from each AF sample (n=24). Two replicates are reported. Arrows and numbers indicate the slices that have been excised and analysed by ESI-Q-TOF as listed in Table 2  $\frac{3}{2}$ . Asterisk (\*) indicates bands that did not give significant results by MS identification.

405

Figure 4. Distribution of amniotic fluid proteins in the Biological process category according
to Gene Ontology (GO) and the Human Protein Reference Database (HPRD) as reported in
Table 3.

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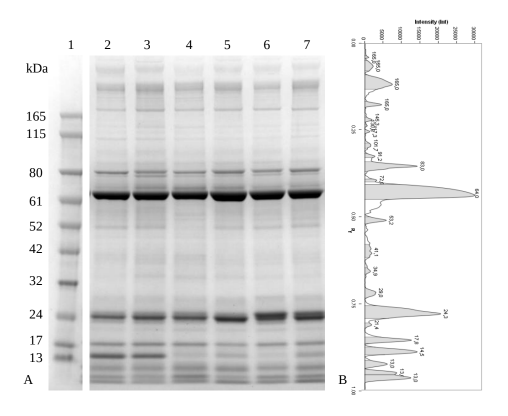
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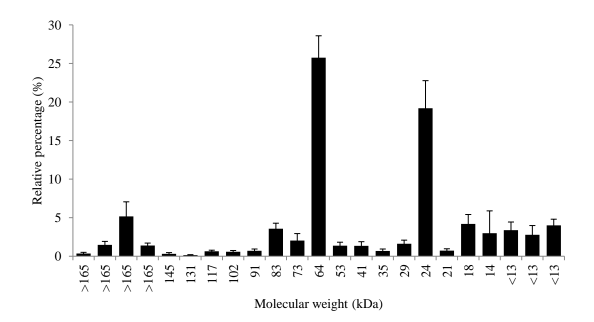
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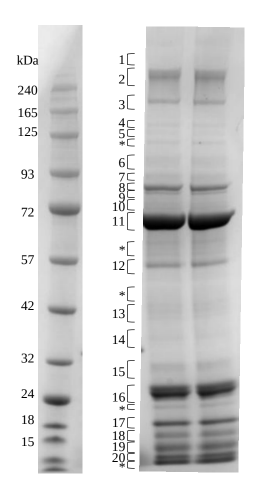
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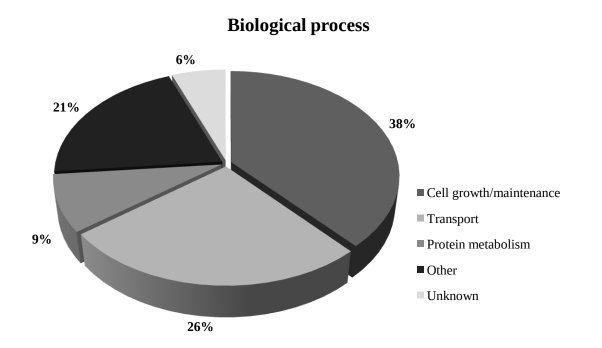












### **Conflict of interest: none**

The authors have no conflict of interests to declare.