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Analysis of pig serum proteins based on shotgun liquid chromatography-tandem mass spectrometry

Keshan Zhang, Yongjie Liu, Youjun Shang, Haixue Zheng, Jianhong Guo, Hong Tian, Ye Jin, Jijun He and Xiangtao Liu*

State Key Laboratory of Veterinary Etiological Biology, National Foot and Mouth Disease Reference Laboratory, Lanzhou Veterinary Research Institute, Chinese Academy of Agricultural Science, Xujiaping No.1, Lanzhou, Gansu, 730046, PR China.

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Recent advances in proteomics technologies have opened up significant opportunities for future applications. We used shotgun liquid chromatography, coupled with tandem mass spectrometry (LC-MS/MS) to determine the proteome profile of healthy pig serum. Samples of venous blood were collected and subjected to sodium dodecyl sulfate-polyacrylamide gel electrophoresis separation and in-gel trypsin digestion. The peptides were then processed using shotgun LC-MS/MS. Serum proteins were subjected to protein identification and bioinformatics analysis. A total of 392 proteins were identified, and 179 were annotated according to their molecular functions and biological processes, excluding 142 hypothetical proteins and 71 immune globulins. To the best of our knowledge, this represents the first porcine serum proteomics analysis based on shotgun LC-MS/MS. This method and the resulting proteomics information may prove valuable for ensuring good animal welfare practice and for monitoring swine health and disease status.

Key words: Analysis, pig serum, shotgun coupled with tandem mass spectrometry (LC-MS/MS).

INTRODUCTION

Serum is a major body fluid. Serum composition thus reflects the overall health status of the individual animal and is often used to monitor health and disease in farm animals (Bendixen et al., 2011; Eckersall et al., 1996). Studies of protein distribution characteristics in serum may provide significant information to help unravel the mechanisms of disease and for the identification of biomarkers associated with new drug targets and early diagnosis (Issaq et al., 2007; Wan et al., 2006). Human serum protein maps have already been established (Millioni et al., 2012). Detailed serum protein two-

Abbreviations: LC-MS/MS, Liquid chromatography-tandem mass spectrometry; 2-D PAGE, two-dimensional polyacrylamide gel electrophoresis; MWs, molecular weights; pl, isoelectric point; PRRSV, porcine reproductive and respiratory syndrome virus; PCV-2, porcine circovirus-2; PRV, pseudorabies virus; HCV, hog cholera virus; FMDV, foot-andmouth disease virus. dimensional gel electrophoresis (2-DE) identification maps have been described for healthy pigs, and 27 highto-medium-abundance plasma proteins, including some examples of infection/inflammation-regulated proteins in healthy Landrace × Large white pigs (Miller et al., 2009).

Proteome analysis is most commonly accomplished using a combination of 2-DE to separate and visualize proteins and mass spectrometry (MS) for protein identification (Gygi et al., 2000a). However, the disadvantages of this technique include extensive sample handling, a limited dynamic range, and difficulties in resolving lowabundance proteins with extreme isoelectric points (pls) and molecular weights (MWs), as well as hydrophobic proteins such as membrane proteins (Corthals et al., 2000; Gygi et al., 2000b; Oh-Ishi et al., 2000). Liquid chromatography, coupled with tandem mass spectrometry (LC-MS/MS), represents a powerful technique for the proteomic analysis of complex samples, where peptide masses may still overlap, even with a highresolution mass spectrometer (Adams and Zubarev, 2005; Wysocki et al., 2005). LC-MS/MS has been increasingly used for the accurate detection of changes in

^{*}Corresponding author. E-mail: hnxiangtao@hotmail.com.

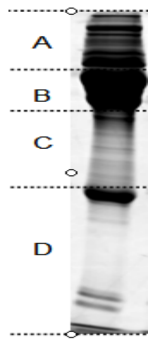


Figure 1. Separation of serum proteins by onedimensional SDS-PAGE. Samples of 150 μ g of proteins were separated on 12% bis-Tris gels and stained to allow protein identification. Four sections were excised and subsequently used for digestion.

protein profiles and to infer biological function (Aebersold and Mann, 2003; Crockett et al., 2005). The shotgun LC-MS/MS proteomics method has been used to identify thousands of proteins in human body fluids, including blood, seminal plasma and tear fluid.

In the present study, gel-LC-MS/MS and bioinformatics analysis methods were used to develop a pig serum protein profile. These results are expected to provide valuable information to assist in the practice of good animal welfare, and for monitoring swine health and disease status.

MATERIALS AND METHODS

Animals

Four Landrace femal pigs, aged about six months, were bought from a local farm and bred in separated rooms. All the pigs were free of the following pathogenic agents: porcine reproductive and respiratory syndrome virus (PRRSV), porcine circovirus-2 (PCV-2), pseudorabies virus (PRV), hog cholera virus (HCV) and foot-andmouth disease virus (FMDV), which were detected by polymerase chain reaction or reverse transcription-polymerase chain reaction (datas not shown). The animal experiments were conducted in accordance with the International Guiding Principles for Biomedical Research Involving Animals, issued by the Council for the International Organizations of Medical Sciences.

Sample preparation and sodium dodecyl sulfatepolyacrylamide gel electrophoresis separation

Samples of intravenous (iv) blood were collected, they were incubated at 37°C for 2 h, then 4°C for 6 h; at last serum were separated at 5000 rpm, for 5 min. The serum protein concentration was determined by quantitative kit (GE) according to the instructions and stored at -80°C until use (Hsieh et al., 2006). Four serum samples (150 µg) were pooled and denatured at 100°C for 5 min in an equal volume of 2x protein loading buffer [0.1 M Tris buffer, pH 6.8, 4% sodium dodecyl sulfate (SDS), 0.2% mercaptoethanol, 40% glycerol, and 0.002% bromophenol blue], and subjected to SDS-polyacrylamide gel electrophoresis. Samples were separated using 12% homogeneous SDS polyacrylamide slab gels and Tris-glycine-SDS buffer (10 mM Tris, 50 mM glycine, 0.1% SDS, pH 8.0) using a Bio-Rad mini-protean tera system (Bio-Rad). Electrophoresis was carried out at a constant current of 15 mA/gel followed by 30 mA for about 1.5 h until the bromophenol blue reached the bottom of the gel. The gels were then stained with Coomassie brilliant blue G250 (Sigma, USA). Images were acquired using a GS-800 densitometer (Bio-Rad, Hercules, CA).

In-gel digestion

The in-gel trypsin digestion of proteins was conducted according to Wilm et al. (1996). The protein lane of the stained gel was cut into four slices (A, B, C, and D), depending on protein molecular weight (MW) (Figure 1). Each slice was diced into 1 x 1 mm pieces and subjected to in-gel tryptic digestion. The gel pieces were rinsed three times using Milli-Q water and destained with 0.2 ml of 100 mM NH₄HCO₃ in 50% acetonitrile for 45 min at 37°C, until complete depigmentation. The gel pieces were then dried in a vacuum centrifuge. 10 µl of 10 mM dithiothreitol in 100 mM NH₄HCO₃. sufficient to cover the gel pieces, was added to the proteins at 56°C for 1 h. After cooling to room temperature, the dithiothreitol solution was replaced with the same volume of 55 mM iodoacetamide in 100 mM NH₄HCO₃. After 45 min incubation at room temperature in the dark, the gel pieces were washed with 100 µL of 100 mM NH₄HCO₃ for 10 min, dehydrated in 100 µL of acetonitrile, swollen by rehydration in 100 μL of 100 mM NH₄HCO₃, and shrunk again by adding the same volume of acetonitrile. The proteins were subsequently digested with 20 ng/ μ L porcine trypsin (modified proteomics grade, Sigma) overnight at 37°C. Peptides were extracted by one change of 20 mM NH₄HCO₃ and three changes of 5% formic acid in 50% acetonitrile (20 min for each change) at room temperature (Li et al., 2009; Zhang et al., 2007).

Shotgun LC-MS/MS analysis

The extracted peptides from each gel piece were analyzed using an Ettan MDLC system (GE Healthcare, USA). In this system, samples were desalted on RP trap columns (Zorbax 300 SB C18, Agilent Technologies, USA), and then separated on an RP column (150 μ m internal diameter, 100 mm long, Column Technology Inc., Fremont, CA). Mobile phase A (0.1% formic acid in HPLC-grade water) and mobile phase B (0.1% formic acid in acetonitrile) were selected. 20 μ g of tryptic peptide mixture was loaded onto the columns and separation was carried out at a flow rate of 2 μ L/min using a linear gradient of 4 to 50% B for 120 min. A Finnigan LTQ linear ion trap MS (Thermo Electron, USA), equipped with an electrospray

Parameter	Number of protein	Percentage (%)
Total peptides	5390	100
Total proteins	848	15.7
Protein groups	392	46.2
immune globulin	71	17.1
annotated proteins	179	45.7
Hypothetical proteins	142	36.2

Table 1. Numbers of peptides and proteins identified in porcine serum.

interface, was connected to the LC setup to detect the eluted peptides. Data-dependent MS/MS spectra were obtained simultaneously. Each scan cycle consisted of one full MS scan in profile mode, followed by five MS/MS scans in centroid mode with the following Dynamic exclusion settings: repeat count 2, repeat duration 30 s and exclusion duration 90 s. Each sample was analyzed in triplicate.

Protein identification and bioinformatics analysis

Peptides and proteins were identified using Biowork 3.2 software (Thermo Finnigan, San Jose, CA), which uses the MS and MS/MS spectra of peptide ions to search against the Suina protein database. MASCOT protein scores (based on combined MS and MS/MS spectra) > 72 were considered statistically significant ($p \le$ 0.05). We accepted individual MS/MS spectra with a statistically significant (confidence interval ≥ 95%) ion score (based on MS/MS spectra). The protein identification and annotation criteria were based on Delta CN (\geq 0.1) and Xcorr (one charge \geq 1.9, two charges \geq 2.2 and three charges \geq 3.75). Protein classification was performed using Gene Ontology Annotation (GOA; http://www.ebi.ac.uk/goa/), according to their molecular functions and biological processes. The subcellular locations of different proteins were predicated with PSORT (http://psort.hgc.jp/).

RESULTS

Serum protein SDS-PAGE separation

Serum proteins were separated by one dimensional SDS-PAGE and the gel was cut into four pieces, according to MW, for shotgun LC-MS/MS analysis (Figure 1).

Identification of proteins

A total of 5390 peptides were detected by shotgun LC-MS/MS. A total of 848 proteins were identified, some of which belonged to the same peptides groups. Thus, 392 proteins were finally identified (Table 1).

Characterization of protein profile

The molecular mass and pl value distributions of the 392 identified serum proteins were analyzed. Their molecular masses ranged from 3.2 to 360 kDa, with most being

between 10 and 100 kDa (Figure 2A). The pls of the proteins ranged from 4 to 11.6 (Figure 2B), with most being between 5 and 7. The predicted 2-DE distribution (Figure 3) showed that the pls of about 95% of the identified proteins were between 4 and 10, representing proteins that are usually difficult to separate by 2-DE. About 40% of the identified proteins fell outside the typical limits of protein resolution obtained by 2-DE. Furthermore, about 14 proteins had higher pls (> 10), which are also usually difficult to separate by 2-DE, but these proteins were also identified successfully by shotgun LC-MS/MS.

Bioinformatics analysis

A total of 189 peptides were annotated according to the GOA database and were classified on the basis of molecular function or biological process. They could be divided into about five functional molecular groups (Figure 4A): the classical protein group (101, 32.6% of 189 annotated peptides) and the cellular protein group (88, 31.7%) were the most common. The classical serum protein group can be further sub-classified into five subgroups, based on their specific functions (Figure 4B); most proteins were proteases or other enzymes (46, 47.9%), common circulating blood proteins (19, 19.0%), or coagulation and complement factors (18, 18.8%) which are important categories of classical serum proteins. The cellular protein group is also sub-classified into five subgroups according to their function or biological process (Figure 4C): signaling channels, hormone regulation, the cytoskeleton, the nucleus, and cellular metabolic secretions.

DISCUSSION

Serum contains many high-abundance proteins that perform various housekeeping functions, as well as numerous secreted or shed low-abundance proteins that are critical for signal transduction and regulatory events. During necrosis, apoptosis, and hemolysis, cell contents may be released into the serum. In a certain time period, the presence, absence or concentration of a specific protein from serum may be related with the

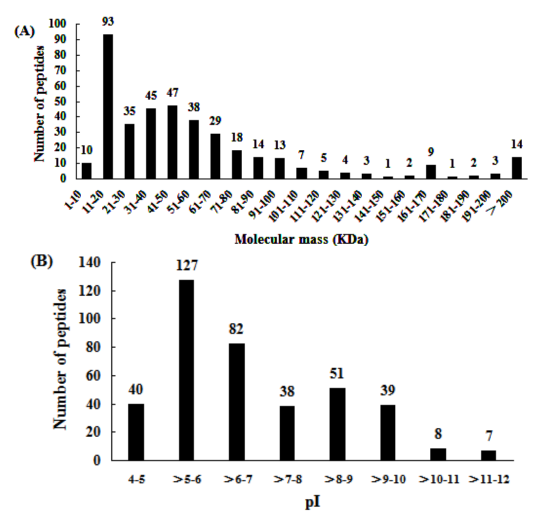


Figure 2. Distributions of molecular weights and pl values for proteins identified by LC-MS/MS, (A) Distribution of molecular weights. (B) Distribution of pl values.

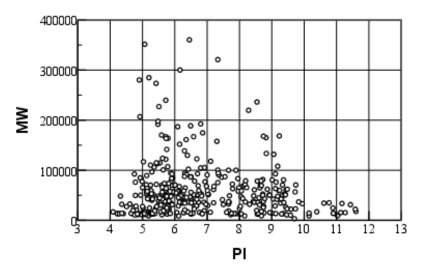


Figure 3. Theoretical 2-DE distribution of proteins from porcine serum. The theoretical pIs and MWs of the proteins were calculated using compute pI/Mw tools according to protein amino acid sequence or ID.

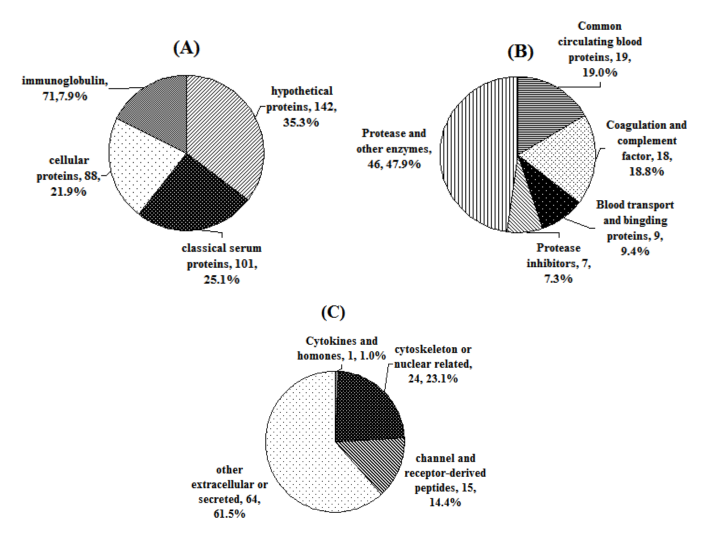


Figure 4. Categorization of 189 serum proteins by molecular functions or biological processes. (A) All identified proteins. (B) Classical serum proteins. (C) cellular proteins.

pathophysiological performance of body, and the presence of these components in blood reinforces the significance of a proteomic approach to identifying biomarkers for disease status. Previous proteomic characterizations of pig serum have used twodimensional PAGE (Miller et al., 2009). Result shows (Figure 3) 40% proteins that are usually difficult to separate by 2-DE. This study used the shotgun LC-MS/MS proteomics technique combined with informatics analysis to determine the proteome profile of pig serum. This technique represents an efficient strategy for swine serum proteomics research, and overcame the disadvantage of 2DE that cannot separate polarity protein. To reduce the individual differences, serum from four pigs were merged together and used for gel electrophoresis separation. In order to identify proteins according to molecular weight, brand A and B mixed, and brand C and D mixed, respectively (Figure 1).

In this study, given profile of serum protein from porcine, we identified a total of 392 proteins (Table 1), of which 189 (Table 2) were annotated and classified based on their molecular function or biological process (Figure 4). As we expected, besides immune globulin, albumin, apolipoprotein, hemoglobin and actin, which are both ubiquitous in the red blood cells were successfully identified. These represented the main serum proteins, and are involved in the combination and transportation of small molecules (Alaupovic, 1996; Bondarenko et al., 2002). Few coagulation or complement factors associated with whole process of blood coagulation were identified successfully as well. Sodium channel protein, transmembrane channel-like protein related to signaling pathways and receptors were identified, with important functions in signal transduction (Naren et al., 1997). The identified proteins also included 46 kinds of proteases or enzymes related to many important biological processes, such as biosynthesis, metabolic regulation, nucleotide replication, damage repair, transcription and posttranslational modification. Fibronectins involved in cell adhesion, cell motility, opsonization, wound healing, and

Table 2. List of 189 annotated serum proteins.

Protein name	PepCount	Unique PepCount	Cover percent	MW kD	p/	Accession number
Common circulating blood protein 15						
Albumin	787	52	74.1	69.4	5.92	gi 833798
Alpha-2-macroglobulin	91	37	32.2	167	5.58	gi 31125621
Ceruloplasmin	71	26	33.7	121.8	5.72	gi 31126951
Apolipoprotein B	31	22	12.3	300	6.19	gi 951375
Apolipoprotein A-I	63	15	52.1	30.3	5.48	gi 461519
Apolipoprotein A-II	8	3	26	11.1	7.73	gi 29774730
Apolipoprotein C-III	4	2	28.1	107	4.76	gi 416627
Hemoglobin subunit beta	33	9	72.1	16.2	7.1	gi 3041678
Porcine hemoglobin	30	9	67.8	16	6.76	gi 5542425
Haptoglobin	22	9	30.8	38.5	6.51	gi 41019122
Hemopexin precursor	37	12	45.3	51.3	6.59	gi 47522736
Apolipoprotein E	12	7	24.6	36.6	5.62	gi 461527
Apolipoprotein D	3	2	13.7	21.5	4.76	gi 31126982
Angiotensinogen-like	3	2	6.3	37.8	8.74	gi 31127118
Spectrin alpha chain	2	2	1.7	284.9	5.2	gi 31124655
Coagulation and complement factor 19						
Complement C3	272	55	46.8	186.8	6.09	gi 47522844
Coagulation factor X protein	1	1	2.3	53.1	5.28	gi 11320581
Coagulation factor IX	1	1	2.7	45.5	5.19	gi 6039224
Complement component C3	70	12	60.5	33.4	5.69	gi 29565664
Complement factor B	36	11	22.6	85.9	7.45	gi 16213824
Complement component 4	65	20	19.5	192.5	6.8	gi 15853775
Complement C2	1	1	3	83.3	7.95	gi 15612013
Complement component C5	11	6	5.9	188.6	6.49	gi 3767794
Complement component C6	5	3	5.1	105.3	6.92	gi 14822653
Complement component C7 precursor	1	1	1.5	93.1	6.7	gi 47523630
Complement component C8A	6	4	13.1	66	5.61	gi 14790521
Complement component C8B	7	3	7.4	69.2	8.14	gi 14823541
Component C8G	2	2	10.9	22.3	5.59	gi 14822322
Complement component C9	5	1	3.3	62.3	5.92	gi 14823369
Complement factor I	11	4	11.8	67.1	8.06	gi 31126268
Complement C1	5	2	15.8	26.5	9.43	gi 51491906
Blood coagulation factor XIV	9	4	13.9	51.8	6.23	gi 571399
Coagulation factor XII	1	1	3.6	68	6.98	gi 3503907
Galectin-8	1	1	6.3	36.3	7.86	gi 21866446
Protease inhibitors 7						
Serpin A3-1	89	17	36.9	60.9	8.46	gi 31126151
Inter-alpha-trypsin inhibitor	40	15	22.8	102.1	6.42	gi 48374067
Inhibitor of carbonic anhydrase	34	13	26.9	77.6	5.88	gi 4752316
Alpha-1protease inhibitor	16	6	26.8	47.2	5.54	gi 1703026
Clusterin precursor	12	6	20.4	51.7	5.62	gi 4752277(
Plasma protease C1 inhibitor	7	2	6.5	54.6	6.77	gi 17805671
Plasminogen activator inhibitor	2	1	4.7	44.8	8.6	gi 31125919

Table 2. Count'd.

Blood transport and binding proteins 9						
Serotransferrin	258	46	74.7	77	6.93	gi 136192
Vitamin D-binding protein	12	6	24.3	24.5	5.02	gi 5186337
Hemoglobin subunit alpha	10	6	57.4	150.3	8.76	gi 122465
Transthyretin	7	5	48	16.1	6.29	gi 1717817
C4b-binding protein alpha chain-like	4	3	8.3	67.4	6.14	gi 31126515
Transgelin-2-like	3	2	12.6	54.7	6.04	gi 31125401
Polyadenylate-binding protein 4-like	2	1	3.8	70.5	9.33	gi 31125894
Polypyrimidine tract-binding protein 1	1	1	5	59.9	9.24	gi 47523538
Telethonin binding protein	1	1	9.6	18.9	5.38	gi 22480955
Channel and receptor derived proteins 15						
Voltage-dependent anion-selective						
Channel protein 1	3	1	6.7	30.7	8.62	gi 7505046
Signal sequence receptor, alpha	2	1	5.2	32	4.36	gi 29763242
Calreticulin	2	1	7	48.3	4.32	gi 29075000
Signal recognition particle 68 kda protein	1	1	2.4	70.4	8.65	gi 31126675
Lycine receptor subunit alpha-1	1	1	1.4	50.2	8.93	gi 31127408
Sodium channel protein	1	1	0.8	206.8	4.92	gi 31126695
Transmembrane channel-like protein	1	1	1.8	92.5	5.91	gi 31124591
Phosphoinositide 3-kinase adapter protein 1	1	1	1.6	100.7	5.64	gi 19404178
Insulin receptor substrate 4	1	1	2.3	53.1	8.83	gi 25859076
Transient receptor potential cation channel	1	1	0.9	236.2	8.54	gi 31124591
Sodium channel and clathrin linker 1	1		0.9 6.4	230.2	4.98	gi 3112459
	-	1				•
Mitochondrial import receptor subunit TOM34	1	1	3.1	50.9	9.42	gi 31127490
Calmodulin-like Syntaxin-3-like	1 1	1 1	14.8 3.9	16.8 48.9	4.09 8.44	gi 31125267 gi 31124761
Protease or other enzymes 44						
Alpha-1-antichymotrypsin 2	32	9	40.2	46.7	6.28	gi 4752327
Plasminogen	18	9	15.8	90.6	7	gi 14634548
Fumarate hydratase	20	8	11.8	13.9	6.3	
Antithrombin-III						gi 4752363
	28	7	24.8	52.4	5.84	gi 19401866
Prothrombin precursor	19	7	18.8	70.1	5.62	gi 17207265
Membrane primary amine oxidase-like	12	6	15.7	78.3	6.61	gi 31126715
Glyceraldehyde 3-phosphate dehydrogenase	18	4	29.4	35.8	8.57	gi 2407184
Serum paraoxonase/arylesterase 1	8	4	18.8	39.9	5.29	gi 16762141
Kininogen-1 isoform 2	5	4	11	43.8	6.64	gi 31126976
Plasma kallikrein	8	3	7.9	72.3	7.78	gi 4752296
L-lactate dehydrogenase B chain	5	3	17.7	36.6	5.57	gi 1107387
Beta-enolase	19	2	9.22	47.1	8.05	gi 11320594
Pyruvate kinase isozymes	7	2	5.4	64.9	7.98	gi 31126085
ATP synthase subunit alpha, mitochondrial	6	2	7.8	59.7	9.21	gi 29759197
ATP-dependent RNA helicase A	1	1	4.5	44.6	5.53	gi 31126494
Alpha-1-antichymotrypsin 1	5	2	13.8	24.7	5.22	gi 9968809
Transketolase	4	2	9.3	67.8	7.21	gi 16295205
Carbonic anhydrase 1	3	2	12.7	29	6.67	gi 19403709
Pig muscle 3-phosphoglycerate kinase	3	2	8.5	43.4	8.78	gi 1339964
ADP/ATP translocase 1-like isoform 1	3	1	6.7	24.7	10.89	gi 31125441
Phosphoglycerate mutase 1-like isoform 2	3	1	10.5	28.9	6.51	gi 19404179
Ribose-phosphate pyrophosphokinase	3	1	4.7	34.8	8	gi31127676
Carboxypeptidase B2	2	1	4.3	48.6	6.83	gi 19404062
Transmembrane protease serine 4-like	2	1	2.5	64.5	8.22	gi 31126400

Table 2. Count'd.

Bifunctional aminoacyl-trna synthetase	2	1	1	161.1	1.35	gi 311265228
Glucosaminefructose-6-phosphate	2	1	2.7	59.5	7.01	gi 311249541
Aminotransferase	2	1	2.7	59.5	7.01	gi 311249541
Polypeptide N-acetylgalactosaminyltransferase	2	1	4.8	64.2	8.63	gi 194042623
Threonyl-trna synthetase	1	1	4	37.7	6.28	gi 311273548
Nicotinamide N-methyltransferase	1	1	9	29.5	5.63	gi 118573081
Phosphoinositide 3-kinase adapter protein	1	1	1.5	107.1	5.78	gi 28860138
Mismatch repair endonuclease PMS2	1	1	1.9	94	6.31	gi 311250873
Alanyl-trna synthetase, cytoplasmic-like	1	1	23.5	8.6	6.12	gi 311257020
Adenylosuccinate synthetase	1	1	2	50.1	8.72	gi 189031714
Inorganic pyrophosphatase	1	1	6.6	27.5	5.44	gi 311271315
Ubiquitin carboxyl-terminal hydrolase 22-A	1	1	2.2	94.4	7.34	gi 311276293
Tyrosine-protein kinase	1	1	1	122.6	6.68	gi 311249266
Ubiquitin-conjugating enzyme E2 L3	1	1	16.2	17.9	8.68	gi 297591969
Serum paraoxonase/arylesterase 1	7	4	14.3	45.6	6.36	gi 118403912
Cholinephosphotransferase	1	1	2.2	42.3	9.05	gi 311262709
Serine/threonine-protein kinase 25-like	1	1	3.5	48.7	6.18	gi 311273415
Serine/threonine-protein kinase Nek5	1	1	0.1	79	8.67	gi 311266294
Rho gtpase-activating protein 23-like	1	1	1	131.8	9.06	gi 311268532
Carbonyl reductase [NADPH] 3-like	1	1	6.1	30.7	5.57	gi 311270205
Cytokines or homones 1						
Interleukin enhancer-binding factor 2	3	2	13.1	43.1	5.19	gi 311254260
Other extracellular or secreted 55						
Spreading factor	6	4	10.7	52.6	5.6	gi 1351418
Heparin cofactor 2	6	4	11.2	55.8	6.5	gi 194043402
Matrin-3-like isoform 1	2	2	5.1	94.7	5.87	gi 311250254
Elongation factor 1-alpha 2	6	1	6.3	50.2	9.33	gi 311263706
60S ribosomal protein L15	2	1	7.8	17.7	11.6	gi 6174950
40S ribosomal protein S5-like isoform 1	2	1	9.8	22.9	9.73	gi 311259613
Vitamin K-dependent protein S	2	1	5.9	27.6	5	gi 311270126
Leucine-rich alpha-2-glycoprotein	2	1	3.8	29.7	7.02	gi 311248408
Macrosialin-like isoform 1	1	1	4.3	42.8	9.29	gi 311261974
Fibronectin isoform 3	27	18	13.9	239.7	5.72	gi 311273025
Histidine-rich glycoprotein	56	12	25.3	61.5	7.2	gi 311269757
Gelsolin	30	11	25.7	84.8	5.93	gi 121118
Alpha-1B-glycoprotein	42	10	30.8	54.4	5.99	gi 311259609
Alpha-2-HS-glycoprotein	59	8	36.3	38.8	5.5	gi 311269753
60 kda heat shock protein	21	8	25.1	60.9	5.7	gi 194044029
Actin	50	6	31.2	41.7	5.29	gi 311250866
Heat shock cognate 71kda protein	26	5	25.9	50.4	5.41	gi 311264120
Heat shock 70 kda protein 1B	18	5	17.5	70.1	5.6	gi 56748897
Heat shock cognate protein HSP 90-beta	13	5	12.3	83.2	4.96	gi 31160516
Heat shock protein HSP 90-alpha	8	3	7.4	84.7	4.93	gi 47522774
T-complex protein 1 subunit alpha-like	13	4	15.4	60.3	5.71	gi 194033404
Vimentin-like	13	4	11.5	70.3	6.01	gi 257096532
Complex of Bdellastasin With Porcine Trypsin	27	3	22.4	23.4	8	gi 257472074
Clathrin heavy chain	7	3	3.8	191.6	5.48	gi 224492556
Fetuin-B-like	4	3	15.4	41.2	7.4	gi 31126975
Zinc-alpha-2-glycoprotein-like	3	3	17.1	34.4	5.88	gi 311250971
Eukaryotic translation initiation factor 3	1	1	16.8	12.6	4.84	gi 311253491
Lumican-like	7	2	11.4	38.8	5.82	gi 194037683

Table 2. Count'd.

Ubiquitin-like modifier-activating enzyme 1	5	2	3.4	114.6	5.54	gi 31127623
Elongation factor 1-gamma	5	2	11.4	50	6.15	gi 31124748
Fascin	3	2	8.5	54.7	6.07	gi 22637295
40S ribosomal protein S15	3	2	28.3	17	10.39	gi 51338618
Zinc finger protein 7	1	1	1.9	76.3	9.13	gi 31125323
Zinc finger protein AEBP2-like	1	1.4	1.4	54.2	5.13	gi 31125067
Zinc finger protein 425-like	1	1	2.6	70.6	9.74	gi 31126476
Polyubiquitin-C-like	1	1	1.2	80.8	9.36	gi 3112700
Adiponectin	1	1	10.3	15.5	8.94	gi 33694199
Troponin T	1	1	6.11	31.2	5.92	gi 66773803
Transcription activator BRG1	1	1	1.4	165	8.82	gi 31124865
Mitogen-activated protein kinase 9	1	1	4	48.4	5.5	gi 31124953
Cytotoxic T-lymphocyte protein 4	1	1	4.5	24.4	5.42	gi 1264450
Centromere protein F-like	1	1	0.5	351.5	5.07	gi 31126500
Transcription factor AP-2 gamma	1	1	4.9	49.1	7.69	gi 17805653
Cell division control protein 42 homolog	1	1	6.8	21.3	5.76	gi12206330
94 kda glucose-regulated protein	1	1	2.1	92.5	4.75	gi 1786569
Ubiquilin-4-like	1	1	3.3	63.9	5.14	gi 31125413
Annexin A5-like	1	1	3.9	42.9	5.16	gi 31126260
Apoptosis regulator protein 1-like	1	1	2.2	71.8	5.43	gi 31127128
C-reactive protein	1	1	11.3	24.9	5.75	gi 62899989
Myosin-9	1	1	1	227	5.51	gi 31125516
ADP-ribosylation factor 1-like	1	1	15	20.7	6.31	gi 31124948
Golgi membrane protein 1-like	1	1	5.9	27.2	4.51	gi 31126550
40S ribosomal protein S28	1	1	17.5	9.1	11.03	gi 4526896
78 kda glucose-regulated protein	4	4	9.5	73.8	5.68	gi 19403359
Leucine-rich PPR motif-containing protein	1	1	2.1	87	7.55	gi31125271
Sytokeleton or nuclear related 24						0
Heterogeneous nuclear ribonucleoprotein A1	10	4	20.9	34.2	9.27	gi 11617525
Eukaryotic initiation factor 4A-I	10	3	14.5	46.1	5.32	gi 15414766
Tubulin beta chain isoform 1	6		14.5	40.1	4.78	
	0 7	3				gi 19404012
Histone H2A type 2-C-like		2	37.2	14	10.9	gi 3112544(
Plastin-2 isoform 1	3	2	6.4	70.2	5.25	gi 19404062
Collagen alpha-3(VI)	1	1	0.8	32.1	7.33	gi 1940437
Microtubule-associated protein 4-like	3	1	1.87	116.9	5.03	gi 31126880
Heterogeneous nuclear ribonucleoprotein A/B	3	1	6.3	32	8.31	gi 16295182
Histone H3.1-like	3	1	23.5	15.4	11.13	gi 31125987
RNA-binding protein FUS-like	3	1	6.2	52.5	9.4	gi 31125125
Plastin-3 isoform 1	2	1	2.4	63.9	5.73	gi 31127682
Heterogeneous nuclear ribonucleoprotein F	2	1	4.1	45.7	5.32	gi 31127122
	2	1	1.7	123.9	5.62	gi 5040367
Heterogeneous nuclear ribonucleoprotein Q	1	1	2.9	69.6	8.68	gi 19403529
Actin related protein	1	1	8.3	19.7	8.53	gi 1955622
	50	6	31.2	41.7	5.29	gi 31125086
Nucleolysin TIAR isoform 2	1	1	4.6	43.4	8.1	gi 31127191
Heterogeneous nuclear ribonucleoprotein D0	1	1	9.5	22.9	9.47	gi 31126290
Small nuclear ribonucleoprotein E	1	1	27.2	10.8	9.46	gi 14790320
Histone H3.2	1	1	23.5	15.4	11.27	gi 31125441
Histone h1t-like	1	1	5.2	22.2	11.58	gi19403983
Nucleophosmin-like isoform 1	1	1	4.8	32.6	4.61	gi 31127393
Nuclear envelope pore membrane protein POM	1	1	7.3	31.2	11.43	gi 31126561
Sister chromatid cohesion protein PDS5	1	1	3.6	37.1	7.66	gi 31126202

maintenance of cell shape were identified (Hakkinen et al., 2010). Some organellar proteins were found (including 40S ribosomal protein S15 and 40S ribosomal protein S28), as well as a few eukaryotic translation initiation factors. Heat-shock proteins (HSPs) are specific proteins that can protect cells and play an important role in growth, development, differentiation and other physiological activities (Arrigo and Simon, 2010; Burel et al., 1992). In a word, this overview map of pig serum protein provided a large number of reference information.

According to function information of proteins, we can make a particular study of partially serum protein in some aspect of disease. Additionally, as can be seen from the sub-cellular localization of identified proteins, these proteins distributed mainly in nucleus, cytoplasm, extracellular matrix, mitochondrion cytoskeleton, and perform their functions in these areas. Therefore, subcellular localization of protein from serum has potential values in research on diseases.

Due to the limited number of pig proteins available in the public databases, protein annotation for some of the proteins was impossible and a number of peptide mass fingerprinting was unmatched effectively. The identification of total proteins in pig serum will be achievable as soon as the complete and accurately annotated genome and protein sequence databases for pig become available.

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