Proteomic analysis of *Rickettsia akari* proposes a 44 kDa-OMP as a potential biomarker for Rickettsialpox diagnosis

František Csicsay 1, Gabriela Flores-Ramírez 1, Fernando Zuñiga-Navarrete 1, Mária Bartošová 1, Alena Fučíková 2, Petr Pajer 3, Jiří Dresler 3, Ludovít Škultétý 1,4* and Marco Quevedo-Díaz 1*

**Abstract**

**Background:** Rickettsialpox is a febrile illness caused by the mite-borne pathogen *Rickettsia akari*. Several cases of this disease are reported worldwide annually. Nevertheless, the relationship between the immunogenicity of *R. akari* and disease development is still poorly understood. Thus, misdiagnosis is frequent. Our study is aiming to identify immunogenic proteins that may improve disease recognition and enhance subsequent treatment. To achieve this goal, two proteomics methodologies were applied, followed by immunoblot confirmation.

**Results:** Three hundred and sixteen unique proteins were identified in the whole-cell extract of *R. akari*. The most represented protein groups were found to be those involved in translation, post-translational modifications, energy production, and cell wall development. A significant number of proteins belonged to amino acid transport and intracellular trafficking. Also, some proteins affecting the virulence were detected. In silico analysis of membrane enriched proteins revealed 25 putative outer membrane proteins containing beta-barrel structure and 11 proteins having a secretion signal peptide sequence. Using rabbit and human sera, various immunoreactive proteins were identified from which the 44 kDa uncharacterized protein (A8GP63) has demonstrated a unique detection capability. It positively distinguished the sera of patients with Rickettsialpox from other rickettsiae positive human sera.

**Conclusion:** Our proteomic analysis certainly contributed to the lack of knowledge of *R. akari* pathogenesis. The result obtained may also serve as a guideline for a more accurate diagnosis of rickettsial diseases. The identified 44 kDa uncharacterized protein can be certainly used as a unique marker of rickettsialpox or as a target molecule for the development of more effective treatment.

**Keywords:** *Rickettsia akari*, Rickettsialpox, Proteome, Surface-exposed proteins, Outer membrane proteins

---

**Background**

From the clinical and antigenic perspective, *Rickettsia* species (Order *Rickettsiales*, Family *Rickettssiaceae*) are obligate intracellular gram-negative alpha-proteobacteria that diverged into three major phylogenetic groups [1–3]. This includes the typhus group (TG), spotted fever group (SFG), and the transitional group of *Rickettsia* (TRG). *Rickettsia akari*, the causative agent of Rickettsialpox, was originally classified into the SFG. However, current genomic studies suggested placing this organism together with *R. australis* and *R. felis* into the TRG [4, 5]. This pathogen is usually transmitted to humans and animals by the rodent mites *Liponyssoides sanguineus* [6, 7]. Nevertheless, it was also detected in the mite *Leptotrombidium scutellare* [8] and Korean voles *Microtus fortis pelliceus* [9]. Rickettsialpox was first described in New York City in

© The Author(s). 2020 Open Access. This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated in a credit line to the data.
1946 [10] and has been since reported in diverse parts of Europe, Asia and North America [11-16]. Patients suffering from this illness describe fever, headache, lymphadenopathy, myalgia, and eschar at the site of the mite bite. Early in the febrile course of the disease, a maculopapular eruption with intraepidermal vesicles usually appears, sparing the palms and soles of the feet [17]. But due to similar symptoms, it is often confused with cutaneous anthrax or smallpox [18, 19]. Thus, it was recommended to confirm the clinical observations with serological testing. Although a high level of cross-reactivity in antibody responses is noted between R. akari and other rickettsiae from the Spotted fever group [20].

Rickettsia species possess a relatively small genome (1.1 to 1.3 Mb) compared to those of their free-living relatives. Particularly, the complete genome sequence of R. akari comprises 1.23 megabase pairs containing 1013 protein-coding genes, 274 pseudogenes, and 39 RNA genes (gene bank accession No. CP000847). This feature is a consequence of invariable genome reduction caused by specialization to a restricted set of hosts during adaptation to the parasitic lifestyle [21]. Further analysis of rickettsial genomes, including R. akari, showed a number of split genes and palindromic elements inserted into genes [2]. Some data are also available from proteomic investigations of Rickettsia species [22-27]. The majority of identified proteins play a crucial role in the mechanism of pathogenesis and virulence of the bacteria. Proteins, however, may also act in antibiotic resistance [28] and host-specific immune response. Recent investigations on rickettsia-host interactions have also identified several important proteins involved in rickettsial adhesion and/or invasion as well as activation of host-cell signaling [29].

In this study, we investigated the antigenic potential of R. akari proteins using gel-free and gel-based proteomic approaches coupled to Liquid Chromatography-Mass Spectrometry (LC-MS/MS) experiments. Particular interest was paid on immuno-dominant cell envelope associated proteins. These key antigens might represent targets for novel diagnostics or vaccine development.

Results

Identification of rickettsial proteins using gel-free and gel-based proteomic approaches

Using two independent proteomics approaches, we identified 288 R. akari proteins in the whole bacterial lysate, from which 39 were identified as uncharacterized proteins. The identified proteins were ranging from 5.2 to 214 kDa in molecular mass and from 4.4 to 13.0 in isoelectric points. Out of these 288 proteins, 41 proteins were identified with predicted molecular masses higher than 70 kDa, 127 proteins with predicted molecular masses between 30 to 70 kDa, and 120 proteins with predicted molecular masses lower than 30 kDa. The sequence coverage of the identified proteins ranged from 1.2% (A8GLW4 – cell surface antigen) to 76.8% (A8GPB6-60 kDa chaperonin GroEL), and the abundance values expressed in label-free quantification (LFQ) ranged from 35.8 (A8GPB6-60 kDa chaperonin GroEL) to 21.5 (A8GPP7 - aspartokinase), with an average value of 26.4 (Additional file 1).

The detected R. akari proteins were grouped into 25 distinct Clusters of Orthologous Groups (COGs), using the database EggNOG v5.0 (http://eggnog5.embl.de/#/app/home). According to this classification, 27.4% of proteins are involved in translation, ribosomal structure, and biogenesis (COG: J); 9% in energy production and conversion (COG: C); 8.3% in cell wall/membrane/envelope biogenesis (COG: M); 6.9% in posttranslational modification, protein turnover, and chaperones (COG: O); 5.6% in intracellular trafficking, secretion, and vesicular transport (COG: U); 4.5% in amino acid transport and metabolism (COG: E); 4.2% in transcription (COG: K) 9.4% with unknown function (COG: S) and 3.5% non-belong to orthologous group (NOG). The remaining 21.2% of proteins belong to 17 other COGs (A, CO, D, F, FG, FP, G, H, I, IQ, L, MU, OU, P, PQ, T, V) in the share of 0.3 to 2.8% (Fig. 1).

Surface-exposed and envelope associated proteins (SEPs)

SEPs of R. akari represent an essential interface in pathogen-host interactions. To recognize these proteins, two membrane-enriched rickettsial extracts were prepared as detailed in the materials and method section. In the first, the cell-impermeant reagent, Sulfo-NHS-LC-Biotin, was used to label SEPs. The biotinylated proteins were then isolated by streptavidin agarose affinity purification. On the other hand, the second protocol was based on the Triton X-114 phase partitioning. This non-ionic detergent substitutes the lipid molecules interacting with the hydrophobic domain of integral membrane proteins and forms soluble protein–detergent micelles. After temperature changes, phase separation occurs. Hydrophobic proteins aggregate in the detergent phase, while the hydrophilic remain in the aqueous layer [30].

All the obtained fractions were then separated by 12% SDS-PAGE and analyzed with LC-MS/MS. Using these approaches, we identified 83 unique R. akari proteins with molecular masses range from 5.5 to 168.0 kDa and pi from 4.6 to 12.1. Fourteen of these proteins were recognized with both approaches, including OmpB (A8GPL7), chaperone protein DnA (A8GMF9), 60 kDa chaperonin GroEL (A8GPB6), ATP synthase subunit alpha (A8GPZ6), a 44 kDa uncharacterized protein (A8GP63), superoxide dismutase (A8GNP0), putative adhesin A1C_06425 (A8GQ33), thioredoxin peroxidase 1 (A8GN15), inorganic pyrophosphatase (A8GP57),
stress-induced DNA-binding protein (A8GPZ9), nucleoside diphosphate kinase (A8GLZ8), 50S ribosomal protein L7/L12 (A8GMA6), 10 kDa chaperonin (A8GPB7) and 7 kDa uncharacterized protein (A8GNR0).

Out of the 83 identified proteins, 15 were predicted using the PSORTb prediction tool [31] as a membrane or secreted extracellular proteins (Fig. 2a, Table 1), 1 protein as periplasmic, 48 proteins as cytoplasmic, and 17 proteins with unknown followed by 2 with multiple localizations. Using the SOSUIgramN program [32], 20 proteins were anticipated as a membrane or extracellular (Fig. 2a, Table 1), 2 proteins as periplasmic, 57 proteins as cytoplasmic and 4 proteins with unknown locations. In addition, the PRED-TMBB online tool [33] recognized 25 proteins with beta-barrel structure (Fig. 2a, Table 1), and 11 proteins were predicted with the Signal P–5.0 server to possess a signal peptide at the N-terminus (Fig. 2a, Table 1). Interestingly, nine of these proteins apparently carry a Type I secretory signal peptides that are usually transported by the Sec or twin-arginine translocon (Tat). These proteins include the OmpB (A8GPL7), 44 kDa uncharacterized protein (A8GP63), putative adhesin A1C_06425 (A8GQ33), putative surface antigen (A8GM15), 19 kDa uncharacterized protein (A8GP34), 17 kDa uncharacterized protein (A8GP34), tail-specific protease (A8GMM2), protein export protein PrsA (A8GP43), and the 18 kDa uncharacterized protein (A8GNE2). Additional two proteins, namely 9 kDa uncharacterized protein (A8GNC4) and a peptidoglycan-associated lipoprotein (A8GPW0), were...
predicted to have a lipoprotein signal peptides suggesting Type II secretion.

Using the four bioinformatics algorithms, we suggested 33 proteins with predicted outer membrane localization or possessing signal peptides or beta-barrel structure as R. akari SEPs (Table 1). These proteins were then grouped into 8 distinct COGs, using the database EggNOG v5.0. According to this classification, 18.2% of proteins are involved in posttranslational modification, protein turnover, and chaperones (COG: O); 12.1% with unknown function (COG: S); 12.1% in cell wall/envelope biogenesis (COG: M); 12.1% in energy production and conversion (COG: C); 12.1% in amino acid transport and metabolism (COG: E); 9.1% of proteins are involved in translation, ribosomal structure, and biogenesis (COG: J). The remaining 6% of proteins belong to 2 other COGs (D and P) in the share of 3% each (Fig. 2b, Additional file 2). Five proteins were not classified into any clusters of orthologous groups due to unknown functions.

### Table 1: Surface exposed proteins of R. akari

| Uniprot Accession Number | Description | mW (KDa) | pI | Identified by: | Subcellular localization | Predicted Motifs | Immunogenic |
|--------------------------|-------------|---------|----|---------------|--------------------------|----------------|-------------|
| A8GQ33                   | Putative adhesin A1C_06425 | 24.1 | 9.7 | B/T/S         | outer membrane            | outer membrane  | 2.8190^a Sec/SPI (0.9982) | –            |
| A8GMA6                   | 50S ribosomal protein L7/L12 | 13.0 | 5.0 | B/T/S         | multiple                  | cytoplasmic    | 2.851^b no | –            |
| A8GPL7                   | Outer membrane protein B     | 167.9| 5.0 | B/T/S         | outer membrane            | extracellular   | 2.861^b Sec/SPI (0.3494) Tat/SPI (0.5171) yes |
| A8GBP8                   | 60 kDa chaperonin            | 58.7 | 5.3 | B/T/S         | cytoplasmic               | cytoplasmic    | 2.886^b no yes |
| A8GM15                   | Putative surface antigen     | 48.1 | 9.6 | B             | inner membrane            | extracellular   | 2.898^b Sec/SPI (0.9979) | –            |
| A8GN82                   | Malate dehydrogenase         | 33.6 | 5.4 | T/S           | cytoplasmic               | cytoplasmic    | 2.901^b no | –            |
| A8GN16                   | Membrane protease subunits   | 33.9 | 5.5 | T/S           | unknown                   | inner membrane  | 2.902^b no | –            |
| A8GP78                   | Rod shape-determining protein MreB | 37.4 | 7.0 | T/S           | cytoplasmic               | cytoplasmic    | 2.912^b no | –            |
| A8GM29                   | ATP-dependent protease subunit HslV | 19.8 | 6.7 | T             | cytoplasmic               | cytoplasmic    | 2.932^b no | –            |
| A8GM33                   | Elongation factor Ts         | 33.7 | 5.2 | T/S           | cytoplasmic               | cytoplasmic    | 2.932^b no | –            |
| A8GM25                   | Preprotein translocase subunit SecG | 6.5   | 9.5 | B             | unknown                   | unknown        | 2.936^b no | –            |
| A8GM80                   | Probable cytosol aminopeptidase | 53.3 | 6.0 | T/S           | cytoplasmic               | cytoplasmic    | 2.939^b no | –            |
| A8GP24                   | ATP synthase subunit beta    | 51.0 | 4.6 | T/S           | multiple                  | cytoplasmic    | 2.940^b no | –            |
| A8GP69                   | Phospho-N-acetyluramoyl-pentapeptide-transferase | 39.7 | 8.4 | B             | inner membrane            | inner membrane  | 2.946^b no | –            |
| A8GP26                   | ATP synthase subunit alpha   | 56.2 | 6.4 | B/T/S         | cytoplasmic               | cytoplasmic    | 2.947^b no | –            |
| A8GC4                    | Uncharacterized protein      | 37.4 | 7.0 | T/S           | cytoplasmic               | cytoplasmic    | 2.948^b Sec/SPI (0.9985) | –            |
| A8GH0                    | 2,3,4,5-tetrahydropyridine-2,6-dicarboxylate N-succinyltransferase | 30.0 | 7.0 | T/S           | cytoplasmic               | cytoplasmic    | 2.950^b no | –            |
| A8GM59                   | Heat shock protein           | 18.6 | 9.2 | T/S           | unknown                   | outer membrane  | 2.954^b no | –            |
| A8GP63                   | Uncharacterized protein      | 44.6 | 8.5 | B/T/S         | unknown                   | extracellular   | 2.957^b Sec/SPI (0.9980) yes |
| A8GFM9                   | Chaperone protein DnaK       | 67.7 | 4.8 | B/T/S         | cytoplasmic               | cytoplasmic    | 2.960^b no yes | –            |
| A8GN11                   | Aminotransfer_5 domain-containing protein | 40.7 | 6.7 | T             | cytoplasmic               | cytoplasmic    | 2.961^b no | –            |
| A8GL8                    | ATP synthase subunit a       | 27.3 | 9.0 | T             | inner membrane            | inner membrane  | 2.962^b no | –            |
| A8GNF1                   | 4-hydroxy-tetrahydrodipicolinate synthase | 32.4 | 7.2 | T             | cytoplasmic               | cytoplasmic    | 2.962^b no | –            |
| A8GP29                   | Stress induced DNA-binding protein | 16.1 | 4.9 | B/T/S         | cytoplasmic               | cytoplasmic    | 2.963^b no | –            |
| A8GLX8                   | 50S ribosomal protein L9     | 19.4 | 7.7 | B/S           | cytoplasmic               | cytoplasmic    | 2.964^b no | –            |
| A8GLM3                   | Uncharacterized protein      | 61.3 | 5.5 | T/S           | outer membrane            | inner membrane  | 2.986 no | –            |
| A8GP43                   | Protein export protein prsA  | 31.4 | 9.4 | T/S           | outer membrane            | cytoplasmic    | 2.989 Sec/SPI (0.9958) – |
| A8GP34                   | Uncharacterized protein      | 19.5 | 5.7 | B/S           | unknown                   | extracellular   | 2.990 Sec/SPI (0.9796) – |
| A8GM4                    | Tail-specific protease       | 50.1 | 7.3 | T/S           | inner membrane            | outer membrane  | 2.990 Sec/SPI (0.8945) – |
| A8GE2                    | Uncharacterized protein      | 18.8 | 9.0 | T             | unknown                   | periplasmic     | 2.993 Sec/SPI (0.9866) – |
| A8GP22                   | Uncharacterized protein      | 17.0 | 6.2 | B/S           | unknown                   | extracellular   | 3.006 Sec/SPI (0.9979) – |
| A8GP67                   | Actin polymerization protein RickA | 59.6 | 10.1| T/S           | outer membrane            | extracellular   | 3.035 no | –            |
| A8GPW0                   | Peptidoglycan-associated lipoprotein | 17.5 | 8.9 | T/S           | outer membrane            | unknown        | 3.106 Sec/SPI (0.9996) yes |

SEPs detected using, B biotinylation, T Triton X-114, and/or S Shotgun proteomic approaches. ^a values below the threshold 2.965 indicate the presence of beta-barrel structure.
Immunogenic SEPs

In order to identify antigenic proteins, the 2-dimensional electrophoresis (2-DE, Fig. 3a) of the whole R. akari cell extract were probed against the anti-R. akari polyclonal rabbit serum (Fig. 3b) and the serum of infected patient clinically diagnosed with rickettsialpox (Fig. 3c). From the identified immunoreactive proteins, five were recognized in this study as SEPs, namely a 60 kDa chaperonin GroEL (A8GPB6), OmpB (A8GPL7), DnaK (A8GMF9), Peptidoglycan-associated lipoprotein (A8GPW0), and a 44 kDa uncharacterized (A8GP63) protein.

The 44 kDa uncharacterized protein (A8GP63) together with the highly abundant 60 kDa (GroEL) (A8GPB6) and the DnaK (A8GMF9) were prepared by ectopic expression in E.coli BL21(DE3). Purified recombinant proteins were then separated by SDS-PAGE (Fig. 3d) and probed against five sera from patients with confirmed rickettsial infection (2 sera of patients with SFG rickettsiosis and 3 with Rickettsialpox) (Fig. 3e). Interestingly, the proteins GroEL and DnaK reacted with all the patient’s sera. However, the 44 kDa uncharacterized protein (A8GP63) conducted a response only with the sera of patients suffering from Rickettsialpox (Fig. 3e, lanes 3, 4, 5). No reactivity was observed with the sera of patients with SFG rickettsiosis. In this protein preparation, two bands (about 44 kDa and 22 kDa) were seen in the SDS-PAGE and the immunoblots. However, both correspond to the 44 kDa uncharacterized protein as determined by LC-MS/MS analysis. The lower and upper ones represent the truncated and full-length protein, respectively (Fig. 3c).

Characterization of the 44 kDa uncharacterized protein (A8GP63)

The homology searches for the R. akari putative 44 kDa uncharacterized protein against the NCBI database showed high identity (91.8%) to uncharacterized protein from R. australis and 83.2% to unknown protein (RF_0375) from R. felis (accession number AAY61226.1). However, only a low level of sequence similarity and protein size was noted in the comparison to other SFG rickettsiae (e.g., R. conorii (AAL0344). R. parkeri (WP_146709264.1), R. slovaca (WP_041472097.1), R. rickettsii (WP_041472549.1) (Fig. 4a). Such a drift usually occurs due to gene loss which is a major source of genome diversification within Rickettsia species and may reflect host specialization to distinct arthropods [16].

Based on precise bioinformatics predictions, we could suggest that this 44 kDa protein is having a Sec signal peptide (Sec/SPI) and secondary transmembrane...
structure (Fig. 4b) that is represented by beta-barrels typical for outer membrane proteins. The protein structure also showed a high percentage of total accessible surface area that could be easily available for antibody binding. Using the B-cell type epitope predictor, we revealed the candidate epitopes for the 44 kDa uncharacterized protein (A8GP63). We found 10 potential peptides that showed a score higher than the cut-off. Six of them were exposed on the protein surface and probably well available for antibody binding (Additional file 3). Thus, this observation may suggest that the *R. akari* protein can be involved in the specific interaction with the host cell.

To confirm surface localization of this protein we performed a dual immunofluorescence antibody assay. For this purpose, we developed a mouse polyclonal antibody against the 44 kDa uncharacterized protein and conjugated it with Alexa 488 which provides a green fluorescence signal. In order to significantly reduce cross-reactivity and to increase the specificity of protein recognition, we pre-absorbed the mouse serum with *E. coli* cells prior to immunofluorescence assay (IFA). As seen in Fig. 4c, the green fluorescence signal is apparently associated with the outer membrane of *R. akari*. We observed a robust protein accumulation clustering near the pole of rickettsial cells. Such a feature was also recorded for surface cell antigens (Sca) of *R. typhi*, including Sca2, Sca3, and Sca4, as seen in immunoelectron microscopy where clustering of Sca proteins on the surface of intact rickettsia was clearly demonstrated [34].

**Discussion**

In this work, a comprehensive proteomic study of *R. akari* was performed using gel-based and gel-free approaches with the aim to identify proteins having a key role in pathogen-host interactions. In the first step, proteins from the whole-cell extract of the bacterium were separated by 2-DE. In the gel (Fig. 3a), 356 protein spots were detected. The spots were excised and identified by ESI-LC-MS/MS mass spectrometry. Obtained data were then combined with protein identification from a gel-free (shotgun) proteomic approach that was based on LC-MS/MS analyses. These proteomic analyses resulted in the identification of 288 unique proteins that correspond to 28.4% of 1013 protein-coding genes annotated in the *R. akari* genome. Similar numbers of proteins

---

**Fig. 4** The 44 kDa protein (A8GP63) of *R. akari* as an outer membrane protein. (a) Homology search to 44 kDa protein using blast analysis (b) In silico prediction of the protein secondary structure (c) IFA using a “pre-absorbed” mouse polyclonal antibody against the 44 kDa protein
were identified in previous proteomic studies in which R. prowazekii, R. conori, R. raoultii, R. massiliae, R. slovaca, and R. parkeri were evaluated [22, 24–26, 35].

The largest group of identified proteins was associated with translation, ribosomal structure, and biogenesis. This group was tightly followed by enzymes involved in the processes of central metabolic pathways such as glycolysis, pentose phosphate pathway, tricarboxylic acid (TCA) cycle, oxidative phosphorylation, fatty acid metabolism, and amino acid metabolism. Four key enzymes of TCA cycle, which have been shown to be complete in Rickettsiae (KEGG, MAP 00020) were detected in this study, namely malate dehydrogenase (A8GN82) citrate synthase (A8GQ56), acetyltransferase component of pyruvate dehydrogenase complex (A8GNQ3), and 2 subunits (alpha and beta) of pyruvate dehydrogenase E1 component (A8GMR3 and A8GMR4 respectively). Despite the evidence of a functional presence of pyruvate dehydrogenase complex and TCA cycle, it was shown that Rickettsia spp cannot metabolize (glycolysis) or generate (gluconeogenesis) glucose [36]. Neither pyruvate can be synthesized due to the lack of metabolic pathways necessary for pyruvate synthesis [37]. Therefore, glucose and pyruvate should be uptaken directly or in the form of phosphoenolpyruvate from the host cells cytoplasm to support ATP generation via the TCA cycle [23]. In this case, phosphoenolpyruvate is converted to pyruvate by pyruvate phosphate dikinase (A8GNN4), which was detected in this study. Nevertheless, ATP can be synthesized by oxidative phosphorylation. The presence of four ATP synthase subunits (A8GPZ3, A8GPZ4, A8GPZ6, A8GPZ7) illustrates the importance of this pathway in Rickettsiae that is necessary to compensate for glycolysis defect [23].

In addition to exogenous uptake of major energy sources including glucose, ATP and NAD⁺ [36], Rickettsiae as intracellular organisms possess refined secretion mechanisms to translocate various effector proteins into the host cell in order to sustain survival. They are equipped with various types of secretion systems, including the Sec or Tat translocation pathways, type I secretion system (T1SS), type IV secretion system (T4SS), and type V autotransporter pathways (T5SS) [38]. As in earlier rickettsial proteome studies [22, 25], we identified several proteins of Sec-mediated translocation system, namely protein translocase subunit SecD (A8GP54), Sec translocon accessory complex subunit YajC (A8GP53), signal recognition particle receptor FtsY (A8GPW4), preprotein translocase subunit SecG (A8GM25), and a protein-export membrane protein SecE (A8GM27), that is unique to all rickettsiae [23, 25]. We also identified two out of the three major structural features of the T1SS, namely outer membrane efflux protein (A8GMM0) belonging to the TolC family [38] and a putative ABC transporter ATP-binding component (A8GM03). This protein is a constituent of ATP-binding cassette transporters that play a role in multidrug resistance [23, 39]. Moreover, we detected three components of T4SS, namely TrbL/VirB6 plasmid conjugal transfer proteins (A8GM63–67), VirB4 family protein (A8GM62), and VirD4 protein (A8GMV1). The last interact with a VirB4 protein and the ATPases VirB11 independently of other subunits and forms the ATPase complex that dock to the inner membrane. The VirD4 also acts as a receptor that may transport substrates to the translocation channel [40, 41]. Meanwhile, VirB6 is the key polytopic membrane protein forming the inner membrane channel that is essential for substrate secretion (Additional file 1).

Secreted proteins and SEPs play a key role in pathogen-host interactions by allowing infection of the host cell and promoting bacterial replication in the cytosol [34]; hence identification of these important molecules is essential. Obviously, the surface cell antigen (Sca) family proteins are undoubtedly included among the most dominant SEPs which play an important role in rickettsial pathogenesis [42–44]. Previous studies indicated that two major Sca proteins, outer membrane protein A (Sca0 or OmpA) and OmpB (Sca5), might play key roles in the adhesion of the Rickettsia cells to the host tissue [42, 44–46]. These molecules are likely major antigenic determinants of Rickettsia spp and represent good candidates for vaccine development. Nucleotide sequences of these proteins were even used for phylogenetic determination. While OmpB (A8GPL7) has been identified in several Rickettsia spp, including R. akari (in this study), the presence of OmpA was reported only in the SFG [47]. The latest protein is mostly not encoded in the genomes of TG Rickettsia [34]. Further evaluation of the identified proteins has revealed many similar SEPs as in other Rickettsiae, including the OmpB (A8GPL7), putative adhesin (A8GQ33), heat shock protein (A8GMS9), and chaperone proteins DnaK (A8GMF9) and GroEL (A8GPB6). Using enrichment protocols, we also detected the cell division coordinator CpoB (A8GMM6) surface protein, the TolB protein (A8GXM9), and the actin polymerization protein RickA (A8GP67). The CpoB was previously described as a YbgF protective antigen [27]. Together with TolB, these periplasmic proteins are essential for the Tol-Pal system, which has a role in the maintenance of cell envelope integrity. These proteins may also participate in translocation of virulence factors [48]. On the other hand, the identification of actin polymerization protein RickA (A8GP67), an activator of the host Arp2/3 complex, may indicate the actin-based motility of R. akari [49, 50]. Thus this protein could be proposed as an important factor of virulence in SFG Rickettsia. It promotes
spreading among host cells, unlike the Typhus group members [51, 52].

However, the main aim of this study was to investigate the antigenic potential of these important R. akari proteins. For this propose, the whole-cell extract was analyzed using a two-dimensional immunoblot technique coupled to LC-MS/MS identification. Under our experimental conditions, a strong immunological response of R. akari lipopolysaccharide against the polyclonal sera of infected rabbit and positive patient sera was evident. Nevertheless, among the proteins with predicted beta-barrel structure, which were proposed to be R. akari SEPs, we detected only four immunoreactive antigens, namely chaperone proteins GroEL (A8GPB6) and DnaK (A8GMF9), outer membrane protein B (A8GPL7), and an uncharacterized protein (A8GP63). Heat shock proteins GroEL and DnaK are the most frequently detected immunoreactive proteins with high abundance in Rickettsiae, as it was indicated in many previous studies on R. felis, R. conorii, R. helvetica, and R. rickettsii [24, 25, 53].

The upregulation of groEL and dnaK was also described in R. prowazekii as a response to heat shock exposure [54]. According to our shotgun label-free proteomic quantitation (LFQ), proteins GroEL and DnaK belong to the most abundant proteins of R. akari, with LFQ values 35.8 and 32.2, respectively, whereas the average value of all identified proteins achieved 26.4. Interestingly, one of the uncharacterized proteins, the 44 kDa uncharacterized protein (A8GP63) also seems to be abundant as it reached the LFQ value of 30.1.

So far, no evidence was published concerning this uncharacterized protein (A8GP63). The protein homology search showed very high sequence similarity with two unknown orthologous proteins from R. australis (H8K6W8) and R. felis (Q4UM12). However, only a very low similarity was noted to other SEF rickettsiae (Fig. 4a). This is in line with the classification to TRG of Rickettsia since R. akari is phylogenetically related the most closely to R. australis and R. felis [4]. To better describe this unique protein, it was prepared by ectopic expression in E. coli. At the same time, we also produced the highly abundant 60 kDa (GroEL) and 70 kDa (DnaK) of R. akari for comparative purposes. Then, these recombinant proteins were tested against 3 positive sera from Rickettsialpox patients and 2 positive patient sera with spotted fever group rickettsiosis. As we predicted, all positive patient sera distinguished the recombinant proteins GroEL and DnaK. However, the 44 kDa unknown protein was recognized exclusively with the positive sera from Rickettsialpox patients. Moreover, IFA using specific mouse serum against this 44 kDa uncharacterized protein (A8GP63) together with precise bioinformatics suggested membrane association. Thus, the 44 kDa uncharacterized protein (A8GP63) appears to be a valuable biomarker of R. akari.

Conclusions

This paper represents the first comprehensive proteomic study of R. akari that was accomplished using gel-based and gel-free proteomics approaches. Altogether we identified 316 unique proteins that correspond to 31.2% of 1013 protein-coding genes annotated in the R. akari genome. From them, 33 proteins found in the cell envelope enrichment fractions were predicted as outer membrane proteins or proteins possessing a signal peptide or a beta-barrel structure indicating membrane association. These proteins were proposed as R. akari SEPs that may play a key role in pathogen-host interactions. However, antigenic protein seems to be the most important in terms of effective vaccine development and the discovery of new biomarkers for clinical diagnosis. Among these proteins, we recognized an immunodominant 44 kDa uncharacterized protein (A8GP63) that was not yet described in any Rickettsia. It seems this protein has beta-barrel structures indicating outer membrane character which was proved by IFA. In addition, the protein shows significant immunoreactivity against the sera of patients with Rickettsialpox. Thus, the 44 kDa putative outer membrane protein (A8GP63) represents a good candidate for improved differential diagnosis of rickettsial diseases especially those caused by TRG of Rickettsia.

Methods

Bacterial growth and purification of R. akari

R. akari reference strain Hartford was propagated in egg yolk sacs of pathogen-free chicken embryos in BSL-3 containment as described previously [55] with some modifications. Briefly, 6 days old chicken embryos were inoculated and incubated at 35°C. The bacterial inoculum was adjusted; therefore, most embryos died between 8 to 9 days after inoculation. Yolk sacs were collected, homogenized in 2 M NaCl solution and the suspension was centrifuged at 22000 x g at 4°C. Pellet was resuspended in PBS and centrifuged again at low speed (200 x g for 10 min) to remove debris. The supernatant was then overlaid on 25% w/w sucrose cushion and centrifuged at 22000 x g at 4°C. Resulted pellet was re-suspended in sucrose phosphate glutamate (SPG) buffer and purified by two rounds of Ultravist 370 (Bayer, Germany) discontinuous gradient centrifugation by using 32, 36, and 42% layers at 9000×g at 4°C for 50 min. Light bands containing intact bacteria were collected, and centrifuged at 22000 x g for 30 min [55]. Finally, the pellet was re-suspended in PBS and stored at −80°C until analysis. Giménez staining technique was employed to evaluate the purity of rickettsial cells.

Gel-free proteomics

The bacterial cells of R. akari (4 mg/mL) were pelleted by centrifugation (18,000 x g; 20 min; 4°C) and washed...
with 300 μL PBS. The resulting pellets were re-suspended in 100 μL of 50 mM Tris pH 7.5 containing 0.1% RapigestTM SF (Waters, UK) [56] and then incubated for 10 min at 95 °C. After cooling, 200 μL of 0.1% RapigestTM SF in 8 M guanidinium chloride (Sigma-Aldrich, USA) was added and incubated for another 20 min. Then, filter aided sample preparation – (FASP) [57] protocol was applied. Briefly, inactivated samples were transferred onto Amicon® Ultra – 10 kDa filters (Millipore) and washed twice with 100 mM ammonium bicarbonate (Sigma-Aldrich). Subsequently, proteins were quantified by bicinchoninic acid assay (QuantiPro™ BCA Assay Kit, Sigma-Aldrich) [58]. The samples were then reduced with 100 mM Tris (2-carboxyethyl) phosphine hydrochloride (TCEP, Sigma-Aldrich) and alkylated with 300 mM iodoacetamide (Sigma-Aldrich). Finally, the samples were digested with 2 μg of sequencing grade trypsin (Promega) overnight at 37 °C. Empore® SPE Cartridges, C18, standard density, bed I.D. 4 mm (Sigma-Aldrich) were used to desalt peptide mixtures before drying to completion in a speed-vac. Prior to mass spectrometry analysis, the samples were re-suspended in 30 μL of 2% acetonitrile (ACN)/0.1% trifluoroacetic acid. The samples were further analyzed by Ultimate 3000 RSLCnano system controlled by Chromeleon software (Dionex, USA), involving targeted mass spectrometry and LFQ as described earlier [59] with some modifications. Briefly, extracted peptide mixtures were loaded onto a PepMap100 C18, 3 μm, 100 Å, 0.075 × 20 mm trap column (Dionex) at 5 μL/min for 5 min. Separation was performed on a PepMap RSLC C18 column (0.075 × 150 mm, particle size 2 μm; Dionex) using 68 min gradient of 4–34% mobile phase B (80% ACN, 0.1% FA) with mobile phase A (0.1% formic acid, FA) and 21 min gradient of 34–55% mobile phase B at flow rate of 0.3 μL/min. Eluted peptides were electrosprayed into a Q-Exactive mass spectrometer using a Nanospray Flex ion source (Thermo Scientific, Bremen, Germany) to obtain positive ion full-scan MS spectra in the range 350–1650 m/z.

Acquired raw files were further processed in MaxQuant (version 1.6.7.0) [60]. Andromeda search engine [61] software was applied to identify proteins against the *Rickettsia akari* strain Hartford databases downloaded from Uniprot (September 21st, 2019). Identifications were accepted if at least two distinct reliable peptides matched the protein sequence, or the sequence coverage achieved at least 15%. Relative quantification was performed using the default parameters of the MaxLFQ algorithm [62], with the minimum ratio count set to 2.

**Protein preparation and 2-D electrophoresis (2-DE)**

The rickettsial pellet (2 mg wet-weight) was re-suspended in 4 mL of lysis buffer [28 mM Tris-HCl, 22 mM Tris-base, 200 mM dithiothreitol (DTT, Promega, USA), 2% SDS, protease inhibitor] and incubated for 30 min at 4 °C under shaking. After incubation sample was boiled for 5 min at 100 °C and cooled down on the ice for 5 min, followed by incubation with benzonase nuclelease (Sigma-Aldrich, USA) 1ul/mL during 1 h at room temperature. The supernatant was collected after centrifugation at 14000 x g for 20 min at 4 °C and proteins precipitated with an equal volume of chloroform, 5 volumes of methanol, and 3 volumes of MilliQ water was added prior centrifugation at 14000 x g for 5 min at 4 °C. After phase separation, the upper phase was discarded, the same volume of methanol was added and vortexed. The protein pellet was collected by centrifugation at 14000 x g for 20 min at 4 °C, dried with N2 and stored at −80 °C. Samples were dissolved in lysis buffer (8 M urea, 2 M thiourea, 85 mM DTT, 2.5% Triton X-100) prior to rehydration.

The IPG strip (18 cm, pH 3–10 NL; GE Healthcare, USA) was rehydrated overnight with approx. 140 μg of total protein with 1.0% v/v IPG Buffer (GE Healthcare) and 0.001% bromophenol blue in buffer No.7. The isoelectric focusing (IEF) was carried out in the Ettan IEP-phor 3 IEF System apparatus (GE Healthcare, USA). After IEF each IPG strip was incubated in equilibration buffer composed of 6 M urea, 2% SDS, 30% glycerol, 375 mM Tris-HCl (ph 8.8) in the presence of 1.0% DTT for 15 min, followed by equilibration for 15 min in 3.75% iodoacetamide (IAA, Sigma-Aldrich, USA) solution in the same equilibration buffer. Equilibrated strips were transferred onto 13% polyacrylamide gels to perform second dimension SDS-PAGE. After electrophoresis, gels were stained with Coomassie brilliant blue R-250 (Serva, Germany) and stored at 4 °C until spot excision and trypsin digestion.

**In-gel peptide digestion and MS analysis**

Protein spots were excised, transferred into 1.5 mL Eppendorf tubes and distained in 50 mM ammonium bicarbonate buffer containing 50% acetonitrile (ACN, Merck, Germany), reduced with 10 mM DTT in 100 mM ammonium bicarbonate, followed with alkylation in 50 mM IAA in 100 mM ammonium bicarbonate buffer [63]. Proteins were digested with porcine trypsin in 10 mM ammonium bicarbonate with 10% acetonitrile at 37 °C overnight. The enzymatic reaction was stopped with a 70% acetonitrile solution containing 1.0% trifluoroacetic acid (TFA, Merck, Germany), and peptides were subsequently extracted. Supernatants were concentrated on SpeedVac (Eppendorf, Germany) to a final volume of 20 μL and analyzed by automated nanoflow reverse-phase (RP) liquid chromatography coupled to a Q-TOF (quadrupole time-of-flight) Premier Electrospray Ionisation (ESI) tandem mass spectrometer (Waters, USA).
Separation of peptides was carried out on RP column BEH 130 C18 (200 mm × 75 μm, particle size 1.7 μm; Waters) using a 60 min gradient elution of 5–40% acetonitrile with 0.1% (w/w) formic acid at a flow rate of 0.3 μl/min. Samples were nanosprayed at 3.4 kV capillary voltage to Q-TOF detector and spectra were recorded from alternate scans at low (4 eV) and high (20–40 eV ramp) collision energies to obtain full-scan mass in ion range 50–1950 m/z. Finally, the data were processed by ProteinLynx Global Server 3.0.3 (Waters) searching the UniProt database (https://www.uniprot.org/proteomes/UP000291740). For peak picking the following thresholds were applied: low energy 140 counts and high energy 30 counts. Precursors and fragment ions were coupled, using correlations of chromatographic elution profiles in low/high energy traces. Spectra were searched against the assembled Rickettsiaceae proteome database (downloaded on January 20th, 2019). Workload parameters for the protein identification queries were: i) maximum one possible trypsin mis cleavage; ii) a fixed carbamidomethyl cysteine, variable oxidized methionine and deamidated asparagine/glutamine; iii) the precursors and fragments mass tolerance was automatically determined by the software; iv) peptide matching was limited to less than 4% false discovery rate against the randomized database. Identifications were accepted if at least two distinct reliable peptides matched the protein sequence or the sequence coverage achieved at least 15%.

**Membrane protein enrichment methods**

The first method was based on the Triton X-114 phase separation [30]. Briefly, the purified *R. akari* cells were resuspended in Triton X-114 extraction solution (1% Triton X-114, 10 mM Tris pH 7.5, 5 mM EDTA) and incubated on rotating platform at 4 °C for 4 h. After the extraction, sample was centrifuged at 15000 x g at 4 °C for 15 min, resulting in cell debris containing pellet and supernatant. This supernatant was layered over a chilled sucrose cushion, incubated at 37 °C for 30 min, and centrifuged at 500 x g for 20 min at 30 °C to separate the lower detergent and the upper aqueous phase. The proteins from the Triton X-114 and aqueous fractions were precipitated in acetone at −20 °C prior to separation.

As a second membrane enrichment technique, the cell surface biotinylation method was applied. Briefly, purified *R. akari* bacteria were labeled with Sulfo-NHS-LC-Biotin (Thermo Scientific, USA) and the biotinylated proteins were captured on streptavidin agarose resin (Thermo Scientific) as described previously [64]. The captured proteins were eluted from the streptavidin resin with a 5% solution of 2-mercaptoethanol-PBS at 30 °C for 30 min, repeated 3 times, and the eluted proteins were precipitated in acetone at −20 °C overnight. All protein fractions were then dissolved in Laemmli sample buffer and subjected to SDS-PAGE on 12% polyacrylamide gel. The whole lane was cut and processed by in-gel trypsin digestion and analyzed by LC-MS/MS as described above.

**Preparation of recombinant proteins**

The genes *groEL*, *dnaK* and 44 kDa uncharacterized protein (locus: A1C_04610) were amplified from the genomic sequence of *R. akari* (GenBank accession number: CP000847) with *groEL* primer pairs, 5′- TATA-CCATGG (NcoI)-CAACAAAAACTTATTAAACCG – 3′ and 5′- TATA-AGATCT (BglII)-GGAAATCCAT ACCGCCCAT - 3′ and *dnaK* primer pairs, 5′- TATA-CCATGG (NcoI)-GAAAAATGTAATGATT TGACCTTG - 3′ and 5′- TATA-AGATCT (BglII) - TCTTCTTGCTACATCTGAAAATCG-3′ and 44 kDa uncharacterized protein (locus: A1C_04610) primer pairs, 5′- AG-CCATGG (NcoI)-GTAAATTTAATAAAA TTAAATTTAATTTGC – 3′ and 5′- CTGCAG (PstI)-CTAAATCTAATTTAACCCTGCTCTAA – 3′. Each PCR amplified products was inserted into the pEcoli-Cterm6xHN vector (Clontech, Takara Bio USA) and then competent *E. coli* BL21(DE3) (Novagen, Merck Bioscience) cells were transformed with each recombinant plasmid and cultivated in LB broth (Amresco, USA) with addition of carbenicillin (Sigma-Aldrich). Expression of recombinant proteins were induced with 0.1 mM isopropyl-β-D-thiogalactopyranoside (IPTG, Bioline, UK) at OD$_{600}$ = 0.5–0.6, overnight 25 °C under shaking.

Recombinant proteins were purified by a combination of two methods: in first-round *E. coli* pellets (200 mg wet-weight per each clone) were lysed and purified according to the manufacturer's instruction with HisTalon buffer set (Clontech, USA) and using Talon Metal affinity resin (Clontech, USA). Additional purification of protein eluates was achieved with AKTA pure protein purification system (GE Healthcare, USA), using HisTrap FF crude (GE Healthcare, Sweden) 1 mL columns according to the manufacturer’s instruction. Elution was done with a gradient of 20 mM to 200 mM imidazole in a buffer of 20 mM sodium phosphate pH 7.4 containing 0.5 M NaCl. Concentrations of dialyzed protein eluates were determined by Pierce bicinchoninic acid assay kit (BCA, Thermo Scientific, USA) [58].

**Human and animal sera**

Passive rabbit and BALB/c mice sera were prepared as described previously [27] with some small modifications. Briefly, 10⁵ live *Rickettsia akari* cells were intraperitoneally administered into rabbit in two consequent steps 28 days away. On the other hand, 30 μg of purified 44 kDa recombinant protein mixed with complete Freund’s adjuvant were intraperitoneally administered into the mice, followed by booster immunization with the same dose of...
protein combined with incomplete Freund’s adjuvant on
the 28th-day after priming dose. Blood samples from the
both experiments were collected at the 7th day after the
booster by cardiac puncture under general anesthesia
with sodium thiopental (for mice 50 mg/kg and 20 mg/
kg for rabbit). Mice were sacrificed by cervical dislocation
and the rabbit was euthanized by intravenous overdose
(100 mg/kg) of sodium thiopental (VUAB pharma, Czech
republic). All animal experiments were conducted in the
experimental animal facility at Biomedical Research
Center of the Slovak Academy of Sciences (BMC, SAS) in
accordance with the European Directive 2010/63/EU on
the protection of animals used for scientific purposes
and were approved by the ethical committee of BMC,
SAS. The IgG titers of animal sera were determined by
ELISA using the corresponding antigen.

The sera of rickettsia positive patients, diagnosed base
on clinical signs and ELISA, were obtained from the Na-
ional Reference Centre for Rickettsioses (established by
Regional Authority of Public Health in Banska Bystrica,
Slovakia). Before the experiments, sera were verified by
IFA using a panel of rickettsial antigens, including SFG
rickettsiae (R. slovaca, R. conorii, R. helvetica, R. ricket-
sii), R. typhi and R. akari. IFA test was considered posi-
tive when the antibody titers achieved a cutoff of 1/128
for IgG and 1/64 for IgM. A fourfold and greater in-
tensive when the antibody titers achieved a cutoff of 1/128

Serological analysis of R. akari antigen and recombinant
proteins using western blot

R. akari proteins resolved by 2-DE were transferred (at
100 V/90 min) onto 0.45 μm PVDF membrane ( Pall Life
Sciences, USA) and blocked in 5% non-fat dried milk
(Bio-Rad, USA) in PBS with 0.1% Tween-20 at 4 °C over-
night. Membranes were incubated with a serum of rabbit
infected with R. akari or serum from infected patients at
1:1000 dilutions. After incubation, the membranes were
probed with horseradish peroxidase-conjugated polyclo-
atal swine anti-rabbit (Dako Denmark A/S, 1:3000) or
polyclonal rabbit anti-human IgG (Dako Denmark A/S,
1:3000) secondary antibody, respectively. Visualization
was carried out with a method of enhanced chemilumi-
nescence (ECL) [65].

In order to characterize the immunogenic properties of
recombinant proteins, 16 μg of each purified proteins
were resolved by 12% SDS PAGE gels and transferred to
the PVDF membranes. Then, each membrane was sliced
into 8 strips (containing approx. 2 μg of protein) and in-
cubated separately with different sera from infected
patients at 1:1000 dilutions. After incubation and re-
peated washings, the membrane strips were probed and
visualized as described above.

Preparation of “pre-absorbed” serum against rickettsial
44 kDa protein

The absorption procedure was described by Zhang et al.
[66] with minor modifications. E. coli BL 21DE3 over-
night cultures in LB broth with a density of 10^8 cells/ml
were centrifuged, washed twice and re-suspended in
100 μl of rickettsial 44 kDa protein mouse serum, incu-
bated for 2 h at 37 °C and then overnight at 4 °C on a ro-
tator. E. coli cells were centrifuged at 10000 x g for 30
min; the supernatant was collected and used for Im-
munofluorescence assay.

Dual fluorescence staining of R. akari and 44 kDa
uncharacterized protein (A8GP63)

Purified R. akari cells on a coverslip were fixed and
permeabilized as described earlier [67]. Rickettsial cells
were washed three times in PBS (pH 7.2) containing 2%
bovine serum albumin (BSA; mixtures, PBSA) and then
blocked with 5% BSA in PBS for 1 h at 37 °C. After
washing, bacterial cells were incubated with a “pre-
absorbed” mouse anti-serum against 44 kDa hypothetical
protein (1:100) diluted in 2% PBSA for 1 h at 37 °C. After
washing, cells were incubated with goat anti-mouse IgG
secondary antibody conjugated with Alexa fluor 488
(Life Technologies, USA) diluted 1:1000 in PBS contain-
ing 2% PBSA. The cells were washed three times with
PBSA and blocked again with 5% BSA in PBS for 1 h at
37 °C. R. akari cells were then stained by using poly-
clonal rabbit antiserum against live R. akari diluted 1:200
and a goat anti-rabbit IgG conjugated with Rhodamine
(Life technologies, USA) diluted 1:2000. After five
times washing with PBS, the coverslip was dried and
mounted with Vectashield (Vector Laboratories) and
viewed with fluorescence microscopy (model Eclipse Ni,
Nikon Japan).

Database use and in silico analyses

Prediction of membrane proteins was performed using
four independent programs: signal peptide prediction of
identified peptides was completed by the program SignalP-5.0
(http://www.cbs.dtu.dk/services/SignalP/) [68]. Predictions of the subcellular localization of identified
proteins was performed by program PSORTb version
3.0.2 (https://www.psort.org/psortb/) [31] and
SOSUI-GramN (http://harrier.nagahama-i-bio.ac.jp/
sosui/sosuigramn/sosuigramn_submit.html) [32]. Predic-
tion of beta-barrel outer membrane proteins was made
by a hidden Markov model PRED-TMBB program
(http://bioinformatics.biol.uoa.gr/PRED-TMBB/input.
jsp) [33]. Homology of identified protein with other
rickettsial proteins was compared using the Basic Local Alignment Search Tool (BLAST) (https://blast.ncbi.nlm.nih.gov/Blast.cgi) [69]. Protein classification into COGs functional classes was carried out using public database EggNOG v5.0 (http://eggnog5.embl.de/#/app/home) [70]. Analysis of the 44 kDa protein was submitted to the bcell program (http://tools.immuneepitope.org/bcell/) which predicts flexible linear B-cell epitopes. The sequence was submitted with a signal peptide and a threshold 0.35 [71].

Supplementary information
Supplementary information accompanies this paper at https://doi.org/10.1186/s12866-020-01877-6.

Additional file 1. Identified R. akari proteins classified in the COGs functional category. * S - gel-free approach “Shotgun”; 2D - two dimensional electrophoresis; B - biotinylation; T - Triton X-114 phase partitioning.

Additional file 2. Rakari predicted surface-exposed proteins classified in the COGs functional category.

Additional file 3. Predicted B-cell epitopes of the 44 kDa uncharacterized protein (A8GP63). Threshold 0.35. The orientation of the peptide in the outer membrane is defined as out-faced to the external space, in-faced the internal lipid bilayer, and membrane-embedded in the outer membrane.

Abbreviations
2-DE: Two-dimensional gel electrophoresis; ACN: Acetonitrile; AG: Ancestral group; ATP: Adenosine triphosphate; BSL3: Biosafety level 3; COG: Clusters of Orthologous Groups; ESI-LC-MS/MS: Electrospray ionisation liquid chromatography coupled with tandem mass spectrometry; IPG: Immobilized pH gradient; kDa: Kilodaltons; MW: Molecular weight; Omp: Outer membrane protein; pl: Isoelectric point; PVD: Polyvinylidene fluoride; Sca: Surface cell antigen; SEP: Surface exposed protein; SFG: Spotted fever group; SPG: Sucrose phosphate glutaminate; TG: Typhus group; TRG: Transitional group.

Acknowledgments
The authors thank Jana Vierikova for devoted technical support and Radivojka Banova for precious advice and help.

Authors’ contributions
MQD and FC have designed and performed the biological experiments and wrote the manuscript. MB participated in recombinant protein preparation. AF and PP prepared proteomics samples. FZN and JD performed LC-MS/MS analyses; GFR interpreted MS data. LS coordinated the whole study. LS, MQD, and GFR revised the manuscript. All authors have read and approved the final version of the manuscript.

Funding
This study was supported by the grants from the Scientific Grant Agency of the Ministry of Education of the Slovak Republic and the Slovak Academy of Sciences (VEGA-2/0057/19, VEGA-2/0068/18, 2/0052/19) and the Slovak Research and Development Agency (APVV 19–0066 and APVV 19–0519). Gel free proteomics experiments were funded from the grant of Ministry of Interior, Czech Republic, [VH2017200012: Preparation of the collection of biologically significant toxins with the support of European biological European biodefence laboratory network], the grant of Ministry of Defence of the Czech Republic through a Long-term organization development plan 907930101413, and the grant of Ministry of Education, Youth and Sports of the Czech Republic (L01509).

Availability of data and materials
The LC-MS/MS proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE (Perez-Riverol et al., 2019) partner repository with the dataset identifier PXD017762 and PXD018060 (https://www.ebi.ac.uk/pride/archive/login).

Ethics approval and consent to participate
The usage of laboratory animals in the study was approved by the Ethical Committee of the Biomedical Research Center of the Slovak Academy of Sciences and the State Veterinary and Food Administration of the Slovak Republic under permission number: SK U 01016 292/16-22/1). The experiments were accomplished under regular conditions in the experimental animal facility of the Biomedical Research Center of the Slovak Academy of Sciences. Laboratory animals used in this study were purchased from accredited breeding facility of the Department of Toxicology and Laboratory animals breeding IEP, CEM, Slovak Academy of Science.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Author details
1Institute of Virology, Biomedical Research Center of the Slovak Academy of Sciences, Dúbravská cesta 9, 845 05 Bratislava, Slovak Republic. 2Department of Biology, Faculty of Science, University of Hradec Králové, Hradecká 1285, 500 03 Hradec Králové, Czech Republic. 3Military Health Institute, Military Medical Agency, Tychonova 1, 160 00 Prague 6, Czech Republic. 4Institute of Microbiology of the Czech Academy of Sciences, Vedienska 1083, 142 20 Prague 4, Czech Republic.

Received: 13 March 2020 Accepted: 25 June 2020
Published online: 08 July 2020

References
1. Gillespie JJ, Beter MS, Rahman MS, Amerman NC, Shollom JM, Purkayastha A, et al. Plasmids and Rickettsial evolution: insight from Rickettsia felis. PLoS One. 2007;2:1–7.
2. Merhej V, Angelakis E, Socolovici C, Raoult D. Genotyping, evolution and epidemiological findings of Rickettsia species. Infect Genet Evol. 2014:25:122–37. https://doi.org/10.1016/j.meegid.2014.03.014.
3. Merhej V, Raoult D. Rickettsial evolution in the light of comparative genomics. Biol Rev. 2011;86:379–405.
4. Gillespie JJ, Drisco TP, Verhoeve V, Utsuki T, Husseneder C, Chouljenko VN, et al. Genomic diversification in strains of Rickettsia felis isolated from different arthropods. Genome Biol Evol. 2014;7:35–56.
5. Gillespie JJ, Williams K, Shulak M, Snyder EE, Nordberg BK, Cerial SM, et al. Rickettsia phylogenomics: unwinding the intricacies of obligate intracellular life. PLoS One. 2008;3:21–7.
6. Huebner R, Jefferon W, Pomaranz C. Rickettsiosapox-a newly recognized rickettsial disease. IV. Isolation of a rickettsia apparently identical with the causative agent of rickettsiosapox from Allodermamythus sanguinarius. A rodent mite. Public Heal Rep. 1946;61:1677–82.
7. Huebner R, Jefferson W, CA. Rickettsiosapox-a newly recognized rickettsial disease. V. Recovery of Rickettsia akari from a house mouse (Mus musculus). Public Heal Rep. 1947;62:777–80.
8. Ogawa M, Takahashi M, Matsumot I, Takada N, Noda S, Sajio M. Obligate intracellular bacteria diversity in unfed Leptotrombidium scutellare larvae highlights novel bacterial endosymbionts of mites. Microbiol Immunol. 2020:1–9.
9. Jackson EB, Danaukas JX, Coale MC, Smadel JE. Recovery of Rickettsia akari from the korean vole Microtus fortis hystrix. Am J Epidemiol. 1957;66:301–8.
10. Huebner RJ, Stamps PAC. Rickettsiosapox; a newly recognized rickettsial disease; isolation of the etiological agent. Public Heal Rep. 1946;61(45): 1605–14.
11. Radulovic S, Feng HM, Murovic M, Djelalija B, Popov V, Crocquet-Valdes P, et al. Isolation of Rickettsia akari from a patient in a region where Mediterranean spotted fever is endemic. Clin Infect Dis. 1996;22:216–20.
12. Dzelalija B, Punda-Polic V, Medic A, Dobec M. Rickettsiosapox and rickettsial diseases in Croatia: implications for travel medicine. Travel Med Infect Dis. 2016;14:436–43.
65. Mruk DD, Cheng CV. Enhanced chemiluminescence (ECL) for routine immunoblotting. Spermatogenesis. 2011;1:121–2.
66. Zhang W, Liu G, Tang F, Shao J, Lu Y, Bao Y, et al. Pre-absorbed immunoproteomics: a novel method for the detection of Streptococcus suis surface proteins. PLoS One. 2011;6.
67. Válárková J, Sekeyová Z, Škultetý L, Bohácsová M, Quevedo-Díaz M. New way of purification of pathogenic Rickettsiae reducing health risks. Acta Virol. 2016;60:206–10.
68. Nielsen H, Tsirigos KD, Brunak S, von Heijne G. A brief history of protein sorting prediction. Protein J. 2019;38:200–16. https://doi.org/10.1007/s10930-019-09838-3.
69. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. Basic local alignment search tool. J Mol Biol. 1990;215:403–410.
70. Huerta-Cepas J, Szklarczyk D, Heller D, Hernández-Plaza A, Forslund SK, Cook H, et al. EggNOG 5.0: a hierarchical, functionally and phylogenetically annotated orthology resource based on 5090 organisms and 2502 viruses. Nucleic Acids Res. 2019;47:D309–14.
71. El-Manzalawy Y, Dobbs D, Honavar V. Predicting flexible length linear B-cell epitopes. Comput Syst Bioinformatics Conf. 2006;7:121–32.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.