Profiling Antibody Responses to Infections by *Chlamydia abortus* Enables Identification of Potential Virulence Factors and Candidates for Serodiagnosis

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**Abstract**

Enzootic abortion of ewes (EAE) due to infection with the obligate intracellular pathogen *Chlamydia* (*C.*) *abortus* is an important zoonosis leading to considerable economic loss to agriculture worldwide. The pathogen can be transmitted to humans and may lead to serious infection in pregnant women. Knowledge about epidemiology, clinical course and transmission to humans is hampered by the lack of reliable diagnostic tools. Immunoreactive proteins, which are expressed in infected animals and humans, may serve as novel candidates for diagnostic marker proteins and represent putative virulence factors. In order to broaden the spectrum of immunogenic *C. abortus* proteins we applied 2D immunoblot analysis and screening of an expression library using human and animal sera. We have identified 48 immunoreactive proteins representing potential diagnostic markers and also putative virulence factors, such as CAB080 (homologue of the “macrophage infectivity potentiator”, MIP), CAB167 (homologue of the “translocated actin recruitment protein”, TARP), CAB712 (homologue of the “chlamydial protease-like activity factor”, CPAF), CAB776 (homologue of the “Polymorphic membrane protein D”, PmpD), and the “hypothetical proteins” CAB063, CAB408 and CAB821, which are predicted to be type III secreted. We selected two putative virulence factors for further characterization, i.e. CAB080 (cMIP) and CAB063, and studied their expression profiles at transcript and protein levels. Analysis of the subcellular localization of both proteins throughout the developmental cycle revealed CAB063 being the first *C. abortus* protein shown to be translocated to the host cell nucleus.

**Introduction**

*Chlamydia* (*C.*) *abortus* is an obligate intracellular bacterium with a biphasic developmental cycle involving infectious, spore-like elementary bodies (EBs) and metabolically active reticulate bodies (RBs), which reside and multiply within a non-fusogenic inclusion. The pathogen is the causative agent of enzootic abortion in ewes (EAE), a condition leading to considerable economic losses in sheep husbandry worldwide [1]. Transmission of the zoonotic chlamydiae to pregnant women can pose a life-threatening health risk. If not treated appropriately, the patients may develop severe septicemia with disseminated intravascular coagulation (DIC) resulting in spontaneous abortion of the fetus [2,3]. Despite general progress in chlamydial research, our knowledge about the epidemiology of *C. abortus* infection in humans, particularly the contact persons such as shepherds, agricultural workers and the rural population, is still poor. The fact that neither individual cases nor outbreaks of abortion are usually preceded by clinical signs renders prophylaxis and prevention of the disease particularly difficult. Alongside vaccination, screening flocks and animals using serological assays could be useful to control these infections. However, only a few chlamydial antigens have been used in standardized diagnostic assays so far [4], and serological tests for detection of *C. abortus*-specific antibodies in humans are not available yet. In these circumstances, the search for *C. abortus* proteins that are expressed in the natural host and recognized by the humoral immune response appears
relevant as the identified antigens broaden the spectrum of candidates for serodiagnosis and may imply identification of virulence factors, which become targets of the immune defence by interacting with host cell components. This is particularly important as tools for targeted genetic manipulation of chlamydiae are not available yet and the number of proven virulence factors of *Chlamydia* spp. is limited to a few. Virulence associated function could be assigned to the polymorphic membrane proteins (Pmps) of *C. abortus* [5], which have been identified in several pathogenic *Chlamydia* spp. and resemble autotransporters of the type V secretion system [6-8]. The members of the *pmp* gene family have been shown to be differentially expressed in the course of the developmental cycle leading to evasion of the immune response through antigenic diversity [5,6,8]. The PmpD homologue of *Chlamydia pneumoniae* has been described as an adhesion protein [9].

As the *C. abortus* genome is known to encode a type III secretion system (T3SS) [10], it seems reasonable to speculate that some of the chlamydial virulence factors are secreted into the host cell by use of this system. Therefore, our understanding of chlamydial pathogenesis could be improved through the identification of new T3SS effectors, the initial stage of which can be conducted *in silico* [11].

In order to identify immunoreactive *C. abortus*-specific antigens suitable for serodiagnosis, we have screened the pathogen’s proteome by serological 2D immunoproteomic analysis, using sera from naturally and experimentally infected sheep, as well as from women suffering from *C. abortus* infection during pregnancy. Analysis of reactive spots was performed by mass spectrometry-based assays. The search for immunoreactive proteins was complemented by screening a 2D immunoblot analysis and protein identification.

Materials and Methods

Ethics statement

The human and animal sera were taken from already-existing serum collections of the authors [3,12-14]. Serum samples were anonymized and their use in this study was approved by the local ethics committee of the University of Ulm (#96/09). The generation of polyclonal mice and rabbit sera against recombinant proteins was conducted in accordance with the Animal Welfare Act (TierSchG) and the institutional guidelines and was approved by the Animal Care Commission of the State government of Baden-Württemberg/Germany (#35/9185.82-2).

Organisms and culture

Strain *C. abortus* S26/3 [10] was used throughout the study and cultured on cycloheximide-treated HeLa 229 cell monolayers (American Type Culture Collection; CCL 2.1) in Quantum 101 (PAA Laboratories, Pasching, Austria) in six-well culture plates according to standard procedures [12]. At 48 to 72 h post infection (hpi), when ≥90% of cells contained inclusions, cells were harvested and homogenized with glass beads. HeLa cell debris was removed by centrifugation at 990 x g for 10 min at 4°C. Aliquots of the supernatant were frozen at -80°C. For protein preparation aliquots were thawed and EB were purified by density gradient centrifugation as described previously [15]. EB samples were solubilized in lysis buffer (7M urea, 2M thiourea, 2% w/v CHAPS, 1% w/v DTT, and 0.8% v/v Ampholyte 3/10 from Bio-Rad, Munich, Germany) and sonicated. Protein concentrations were determined using BIO-RAD Protein Assay, (Bio-Rad) according to manufacturer’s instructions.

Animal and human sera

Sera were taken from PCR-positive naturally infected and from experimentally infected animals at abortion [14]. In addition, two sera were taken from animals that were part of an infected flock and that were tested negative for *C. abortus* by PCR. Two human sera were obtained from pregnant women with severe *C. abortus* infections [3]. The control group comprised a serum sample of a sheep that was free of a *C. abortus* infection (“negative control”) and two samples from SPF lambs experimentally infected with *C. pecorum*, as well as two human sera collected from patients with PCR-confirmed *C. pneumoniae* or *C. trachomatis* infection, who had specific IgM and elevated IgG antibody titers, respectively (see Table 1). Sera were used after approval of the local ethical committee of the university of Ulm (#96/09).

Two-dimensional immunoblot analysis and protein identification

The analysis was conducted as described previously [13] using approximately 40 µg protein of purified *C. abortus* EBs for isoelectric focusing. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) gels were blotted onto Immobilon polyvinylidene difluoride (PVDF) membrane (Millipore, Eschborn, Germany) for immunoblot analysis or they were stained by use of the FireSilver Staining Kit (Proteome Factory, Berlin, Germany) for immunoblot analysis or they were stained by use of the FireSilver Staining Kit (Proteome Factory). For isolation and identification of single protein spots by mass spectrometry-based assays (see below). For immunoblot analysis human serum sample were diluted 1:200 and animal samples 1:800. Molecular weights were calculated by use of High-Range Rainbow Molecular Weight Markers” (GE Healthcare, Little Chalfont, UK).

Protein identification

Two 2D gels were stained using the FireSilver Staining Kit (Proteome Factory) according to manufacturer’s instructions and dried. Reactive spots detected by immunoblot analysis were correlated to silver-stained proteins using Delta2D software (DECODON, Greifswald, Germany). The spots were
Table 1. Animal and human sera.

| Serum ID in Table 3 | Original Serum ID | Donor | Causative Pathogen | Source / Reference |
|---------------------|-------------------|-------|--------------------|-------------------|
| A                   | SG SN             | naturally infected ewe (abortion) | C. abortus | Tierseuchenkasse Baden-Württemberg, Stuttgart, Germany |
| B                   | T167/2-4527       | naturally infected ewe (abortion) | C. abortus | Stiftung Tierärztliche Hochschule Hannover, Germany |
| C                   | SZ178/19-7423     | naturally infected ewe (infection is endemic in the flock) | C. abortus | Stiftung Tierärztliche Hochschule Hannover, Germany |
| D                   | SZ178/23-4207     | naturally infected ewe (infection is endemic in the flock) | C. abortus | Stiftung Tierärztliche Hochschule Hannover, Germany |
| E                   | 1291B             | experimentally infected ewe | C. abortus | Livingstone et al., 2005 [14] |
| F                   | 1871A             | experimentally infected ewe | C. abortus | Livingstone et al., 2005 [14] |
| G                   | 815B              | experimentally infected ewe | C. abortus | Livingstone et al., 2005 [14] |
| H                   | 966B              | experimentally infected ewe | C. abortus | Livingstone et al., 2005 [14] |
| I                   | H03-02831H        | human serum (infection during pregnancy) | C. abortus | Medical University of Innsbruck, Austria [3] |
| J                   | H06-12754W        | human serum (infection during pregnancy) | C. abortus | Medical University of Innsbruck, Austria |
| K                   | 1825A             | control serum of an uninfected sheep | no infection | Moredun Research Institute, Edinburgh, UK |
| L                   | 84-796            | serum from a sheep experimentally infected with C. pecorum ("subtype conjunctivitis") | C. pecorum | Moredun Research Institute, Edinburgh, UK |
| M                   | P787              | serum from a sheep experimentally infected with C. pecorum ("subtype arthritis") | C. pecorum | Moredun Research Institute, Edinburgh, UK |
| N                   | SE456             | human serum from a patient with C. pneumoniae infection | C. pneumoniae | University Hospital of Ulm, Germany |
| O                   | SE20353          | human serum from a patient with C. trachomatis infection | C. trachomatis | University Hospital of Ulm, Germany |

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Excised from dried gels and subjected to sequence analysis. Protein identification using nanoLC-ESI-MS/MS (nano Liquid-Chromatography coupled with Electrospray–Ionization tandem mass spectrometry) was performed by Proteome Factory AG, Berlin, Germany [13,16].

Construction and screening of an inducible genomic expression library of the strain C. abortus S26/3

Genomic DNA was purified from EBs using Qiamp DNA Mini Kit (QIAGEN, Hilden, Germany), amplified by use of REPL-I-g Mini Kit (QIAGEN), and partially digested with Sau3A under optimized conditions to yield fragments ranging from 0.5 to 2.0 kb. The expression vectors pTrcHisABC (Invitrogen, Berlin, Germany) were replicated onto nitrocellulose membranes, which were placed on LB plates containing Ampicillin (100 μg/ml). Finally, about 12,000 clones were stored in 96-well plates. The library was replicated onto nitrocellulose membranes, which were placed on LB plates containing 0.1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) to induce gene expression in cloned inserts and incubated at 30°C overnight. The colonies adhering to membranes were lysed using a solution of 10% SDS and 0.5 M sodium hydroxide. Membranes were air dried and blocked with a solution of 5% nonfat dried cow milk (GE Healthcare) overnight. Subsequent colony immunoblot analysis was conducted as described previously [12]. Reactive clones were related to the corresponding wells of microplates and streaked for single colonies again. Reactivity was confirmed by further immunoblot analysis. Finally, the chlamydial inserts of persistently reactive clones were identified by sequencing on an ABI Prism 310 automated DNA sequencer (Applied Biosystems, Darmstadt, Germany).

Cloning, expression, and purification of CAB063, CAB080 (cMIP), and CAB668 (cTUF)

Chlamydial genomic DNA was isolated from infected HeLa cells using the Qiamp DNA Mini Kit (Qiagen). Based on the published genome sequence of C. abortus S26/3, the sequences of CAB063, CAB080, and CAB668 were amplified by PCR using forward primers 5'-CGCGGATCCGGATTAAATCCAAAGC (CAB063), 5'-CGCGGATCCGATCGAAGTTC (CAB080), 5'-CGCGGATCCACGTACTAAAC (CAB668), and reverse primers 5'-CGCGGATCTTTAATCTCTGACACACTC (CAB063), CGCGGATCTTCTAGAAGCTGTTG (CAB080), 5'-CGCGGATCTTGAATCTGTTCCAG (CAB668), all of which contained engineered BamHI and EcoRI restriction sites.
(underlined) at the 5’- and 3’- ends, respectively. The resulting fragments were ligated in-frame into the multi-cloning site of the protein expression vector pTrcHisA (Invitrogen) and the ligated products were used to transform E. coli ER2566 competent cells. Recombinant plasmid DNA was purified with QIAprep Spin Miniprep kit (Qiagen), and the constructs were checked by sequencing on an ABI Prism 310 sequencer (Applied Biosystems). The expression of the recombinant proteins was induced with 0.1 mM IPTG and incubated for 5h at 37°C. The hexahistidine-tagged proteins were affinity purified using the Protino Ni-TED 1000 kit (Macherey-Nagel, Düren, Germany), according to the manufacturer’s instructions.

Production of antibodies

About 12µg each of the recombinant proteins CAB063, CAB080 (cMIP), and CAB668 (cTUF) were subjected to SDS-PAGE and transferred to nitrocellulose by semi-dry Western blotting. The membrane was stained with MemCode™ (Thermo SCIENTIFIC, Schwerte, Germany), and the main band of each protein at 54 kDa (CAB063), 28 kDa (cMIP), and 43 kDa (cTUF), respectively, was excised, solubilized in 150 µl DMSO and used for subcutaneous immunization of mice. BALB/c mice were primed subcutaneously on day 0, boosted on days 14 and 28 with the same formulations, and bled on day 42. In addition a rabbit was immunized subcutaneously with CAB063 according to the same immunization schedule.

Immunofluorescence microscopy

HeLa cells infected with C. abortus at a multiplicity of infection (MOI) of 1 were cultured on glass cover slips and fixed with either methanol or with 4% paraformaldehyde (PFA) dissolved in PBS for 10 minutes at room temperature. PFA-fixed cells were permeabilized with 1% Triton X-100 for an additional hour. After washing and blocking, the cell samples were subjected to antibody and chemical staining. Thus, 4′,6-diamidino-2-phenylindole (DAPI) stain was used for visualization of host cell nuclei and Evans blue for cytoplasmatic counterstaining. CAB063- and CAB080-specific mica sera were used as primary antibodies in 1:60 dilution and detected with Alexa Fluor 488-conjugated goat anti-mouse antibody (Invitrogen) diluted 1:400. Binding of CAB063-specific rabbit serum was detected with Alexa Fluor 488-conjugated goat anti-rabbit antibody (Invitrogen). Images were examined using a Zeiss EC “Plan-Neofluar” 63x/1,25 oil lens on an AXIO Imager M2 microscope (Carl Zeiss, Jena, Germany).

Quantitation of gene expression using real-time polymerase chain reaction (qPCR)

HeLa cells were infected with C. abortus and simultaneous purification of DNA and RNA from infected monolayers was performed with the AllPrep DNA/RNA Mini Kit (Qiagen) according to the manufacturer’s instructions. Reverse transcription (RT-PCR) was performed using a QuantiTect Reverse Transcription Kit (Qiagen). qPCR was conducted in a LightCycler Instrument (Roche Applied Science, Mannheim, Germany) using unique primers and probes for CAB063, CAB080, and ompA (CAB048) synthesized by TIB MOLBIOL (Berlin, Germany) (see Table 2).

| Gene       | Forward primer (5’-3’) | Reverse primer (5’-3’) | LC probe-3’-Fluorescein / 5’-LC Red640 probe |
|------------|------------------------|------------------------|---------------------------------------------|
| CAB048     | TGTAGGCATCACTCAAGGAATCG | CCGACTTAGTGTCAGTTGCCTG | AGGTGTCGTAGAAGTACAAGCTGATAAATT/CAGTACCGTATCGTAAAAGAAGGAACGG |
| CAB063     | AGCATAGACTCTAATGACTCAACCG | CCAACCTAGATGTTACTGTCTGTAA | AACGCTGGACAAATCACGAAGCACA/GAGCCAATAGCGTACAAACTTGCCTGTT |
| CAB080     | AAGAAAACCTCTCCCTAGCC | CTCAGAGGTTTCCCTGATATTG | AGGTGTCGTAGAAGTACAAGCTGATAAATT/CAGTACCGTATCGTAAAAGAAGGAACGG |

Table 2. LC PCR primers and probes.

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The amplification mixture consisted of 0.5 μM forward and reverse primer, respectively, and 0.15 μM LG and FL probes, respectively, as well as 10 μl Quantitect Probe PCR Master Mix (Qiagen) and 2 μl template cDNA, in a total volume of 20 μl. Samples were amplified using the following cycling profile: initial denaturation at 95 °C for 15 min, followed by 45 cycles of denaturation for 5 s at 95 °C, annealing for 30 s at 57°C (for CAB048 and CAB063) and at 54°C for CAB080 and elongation for 30 s at 72 °C.

Quantitation of gene expression was performed as described elsewhere [5]. Briefly, total numbers of genomes and transcripts were calculated using standard curves of the corresponding plasmid DNA (serial dilution with 10^2 - 10^8 copies). Transcripts were normalized against genomes for each time point.

Based on three independent experiments, statistical error was calculated using the standard error of the mean.

Subcellular fractionation of HeLa cells infected with C. abortus

The ProteoExtractTM subcellular proteome extraction kit (Cabiotech, Darmstadt, Germany) allows the successive isolation of cytosolic proteins, organelle-specific proteins, and nuclear proteins. As previously described, chlamydial inclusions of infected HeLa cells can be detected in the “organelle fraction” [17,18]. The extraction procedure was performed according to the manufacturer’s instructions, except that for the cytoplasmic extraction of HeLa cells “Buffer I” was supplemented with 1mM NaVO3. Proteins from each of the extracted fractions were separated by SDS-PAGE in a 10% polyacrylamide gel system. Gels were electroblotted onto Immobilon polyvinylidene difluoride membrane (Millipore, Billerica, MA, USA) for immunoblotting as described previously [12].

Transfection of HeLa cells

The sequences of CAB063 and CAB080 were amplified by PCR using the forward primers 5'-CCGGAAATTCATGGGAATTAATC (CAB063) and 5'-CCGGAAATTCATGAAAAAACAATGG (CAB080) as well as the reverse primers 5'-CTGGTGCAGCTTAACTCCTGACAC (CAB063) and 5'-CTGGTGCACTCATGAAGCTGTGTTTTG (CAB080) with engineered EcoRI and SalI restriction enzyme sites (underlined) at the 5' and 3' ends of the cloned sequence, respectively. The resulting fragment was ligated into the multi-cloning site of the pCI Mammalian Expression Vector (Promega, Mannheim, Germany). After transformation of E. coli ER2566 competent cells recombinant plasmid DNA of pCI-CAB063 was purified with EndoFree Plasmid Maxi Kit (Qiagen).

HeLa cells were plated on glass coverslips so that cells were approximately 60-80% confluent at the time of transfection. The transfection experiment was performed using Nanofectin (PAA) at a DNA (μg) to reagent (μl) ratio of 1:3.2 according to the manufacturer's instructions. After 24 h cells were washed once with PBS and fixed with PFA. For CAB063 transfection additional cell samples were fixed with methanol. Coverslips were stained using CAB063- and cMIP-specific sera, respectively as described above. The experiment was performed three times yielding transfection rates of about 50%.

Results

Identification of 42 immunoreactive C. abortus proteins by 2D immunoblots

In this study we used a proteomic approach to identify immunogenic proteins of C. abortus. Proteins prepared from purified elementary bodies were separated by 2D gel electrophoresis and immunoblot analysis was performed using an assortment of different sera including sheep sera from naturally and experimentally infected sheep as well as two human sera from women with severe C. abortus infection during pregnancy (see Figure 1).

Immunoblot images given in Figure 1 revealed largely similar patterns showing minor differences in the total number of stained spots and individual signal intensities (see Table S1). Some prominent spots have been identified as homologues of MOMP (major outer membrane protein), the heat shock proteins HSP60 and HSP70, elongation factor Tu (EF-Tu), and PMPD (see Figure 1).

In total 42 immunoreactive protein spots could be identified by mass spectrometry based methods. Two significant protein hits were identified for each of three spots (31, 35, and 39) (see Table S1), whereas three proteins were identified following the analysis of two individual spots (CAB080 from spots 15 and 31; CAB668 from spots 1 and 39; and CAB776 from spots 9 and 26).

Searching for C. abortus specific antigens, which show strong immunoblot signals with animal and human sera but no cross-reactivities using control sera, including a serum from an uninfected sheep and two sera from SPF lambs experimentally infected with C. pecorum, as well as two human sera from individuals with C. pneumoniae and C. trachomatis infection, respectively, revealed seven antigens: cElongation factor Ts (CAB046), cMIP (CAB080), cDNA-directed RNA polymerase subunit alpha (CAB113), cGlucose-6-phosphate isomerase (CAB703), cPMP18D (CAB776), and the two “hypothetical” proteins CAB031 and CAB821 (see Table S3).

Identification of additional immunoreactive proteins by screening of an expression library

Six further immunogenic proteins were identified by screening of a C. abortus expression library using the same serum samples from infected animals and humans as previously described for 2D immunoblot analysis: CAB201 (pmp2A), CAB279 (pomp90A, alternative name: pmp12G), CAB281 (pomp91A, alternative name: pmp13G), CAB596 (pomp91B, alternative name: pmp16G), CAB575 ("putative peptide ABC transport system permease protein"), and CAB712 (homologue of the serine-type peptidase CPAF). The antigens CAB048 (MOMP), CAB080 (cMIP), CAB200 (pmp1B), and CAB750 ("putative heat shock-related exported protease") were detected by both screening of an expression library and 2D immunoblot analysis. The unique chlamydial insert of the MOMP specific clone led to high signal intensity using C. abortus specific animal as well as human sera.
Figure 1. Representative examples of 2D immunoblot results. a, experimentally infected sheep, serum F. b, naturally infected sheep, infection is endemic in the flock, serum C. c, human sample, serum J. d, naturally infected sheep, abort, serum A. Serum samples refer to Table 1. Prominent spots identified as the homologues of MOMP (CAB048), HSP60 (CAB615), HSP70 (CAB237) and the elongation factor TU (CAB668) as well as the putative virulence associated proteins MIP (CAB080), PMP18D (CAB776), CAB063, and TARP (CAB167) are marked by circles.

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Table 3. Selection of immunoreactive *C. abortus*-proteins identified by 2D immunoblot analysis.

| Spot-no. | Gene symbol / Protein description | Locus tag | Calculated MW (kDa) | pI | A | B | C | D | E | F | G | H | I | J | K | L | M | N | O |
|----------|----------------------------------|-----------|---------------------|----|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|
| 13       | CAB031 / hypothetical protein    | CAB031    | 87.0                | 4.54 | +++ | - | + | - | + | + | ++ | +++ | + | - | - | - | - | - |
| 14       | tsf / elongation factor Ts       | CAB046    | 30.7                | 5.42 | + | + | ++ | - | +++ | (+) | ++ | +++ | ++ | - | - | - | - | - |
| 28       | CAB063 / hypoth. protein         | CAB063    | 34.0                | 6.75 | + | + | +++ | +++ | + | ++ | - | - | - | - | - | - | - | - |
| 15       | mip                              | CAB080    | 28.1                | 5.09 | (+) | + | + | - | +++ | +++ | ++ | +++ | ++ | - | - | - | - | - |
| 31       | mip / putative macrophage infectivity potentiator lipoprotein | CAB080 | 28.1 | 5.09 | (+) | + | + | - | +++ | +++ | ++ | +++ | +++ | - | - | - | - | - |
| 2        | rpoA                             | CAB113    | 43.5                | 5.62 | +++ | +++ | + | + | (+) | (+) | + | - | +++ | (+) | - | - | - | - |
| 36       | pgi                              | CAB703    | 57.9                | 5.4 | + | - | + | + | ++ | +++ | +++ | +++ | +++ | - | - | (+) | - | - |
| 9        | pmp18D                           | CAB776    | 163.0               | 5.27 | ++ | + | + | - | +++ | +++ | + | - | ++ | + | - | - | - | - |
| 26       | pmp18D / polymorphic outer membrane protein | CAB776 | 163.0 | 5.27 | +++ | ++ | - | ++ | +++ | +++ | ++ | +++ | +++ | - | - | - | - |
| 37       | CAB821 / hypothetical protein    | CAB821    | 64.0                | 5.98 | + | - | + | - | +++ | +++ | + | (+) | + | - | - | - | - | - |

For sera (A – O) see Table 1. The signal intensities of reactive protein spots are presented semiquantitatively: (-), negative; (+), very weak, corresponding to the intensity of MIP (spot 15) in Figure 1d; + weak, corresponding to the intensity of MIP in Figure 1b; ++ moderate corresponding to the intensity of MIP in Figure 1c; +++ strong, corresponding to the intensity of MIP in Figure 1a. Column 4 shows the calculated molecular mass and column 5 the isoelectric point (pI) of the identified proteins. A list of all identified immunoreactive *C. abortus*-proteins is presented in Table S1.

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analysis revealed that this clone codes for a short protein fragment of about 9 kDa, which comprises the variable sequence "VS1"[19], the following conserved sequence, and the beginning of "VS2" (data not shown). As a total of 43 reactive clones did not contain only unique chlamydial inserts representing parts of one specific gene but in addition at least small parts of further chlamydial gene sequences, subcloning will be needed to confirm the specific immunogenicity of the encoded antigens.

Identification of eight “unclassified” proteins by analysis of functional categories

We assigned all 48 positively identified immunoreactive proteins to functional categories essentially based on clusters of orthologous genes (COGs) as listed under http://www.ncbi.nlm.nih.gov/sutils/coxik.cgi?gi=646: Carbohydrate transport and metabolism: CAB062, CAB115, CAB703, CAB841, CAB932. Energy production and conversion: CAB407, CAB654, CAB656, CAB751, CAB752, CAB816, CAB937. Lipid transport and metabolism: CAB539. Cell wall / membrane biogenesis: CAB048, CAB467, CAB712. Nucleotide transport and metabolism: CAB840. Transcription: CAB113, CAB129, CAB452. Amino acid transport and metabolism: CAB531, CAB575. Translation, ribosomal structure and biogenesis: CAB046, CAB188, CAB453, CAB668, CAB785. Posttranslational modification, protein turnover, Chaperones: CAB080, CAB237, CAB615, CAB750, CAB948. Signal transduction mechanisms: CAB031. Intracellular trafficking and secretion: CAB888. Unclassified / Hypothetical proteins: CAB015, CAB063, CAB167, CAB393, CAB395, CAB408, CAB821, CAB929. Polymorphic outer membrane proteins: CAB200, CAB201, CAB279, CAB281, CAB596, CAB776 (see Figure 2). While most of the proteins are predicted to be involved in macromolecular processes the function of eight “hypothetical proteins” (category “Unclassified / Hypothetical proteins”) remained unclear. Four of the hypothetical proteins (CAB063, CAB167, CAB408 and CAB821) are predicted to be secreted by the Type III secretion system (http://www.chlamydiaedb.org/portal/web/chlamydiaedb). CAB167 belongs to a cluster of orthologs that comprises the Chlamydia trachomatis specific TARP (translocated actin recruitment protein), which has been already shown to be an effector translocated by T3SS [20]. The TARP orthologs of Chlamydia abortus and Chlamydia trachomatis show 35% “Identities” and 50% “Positives” over the length of 611 amino acids that comprise three putative actin binding domains (http://blast.ncbi.nlm.nih.gov/Blast.cgi, [21]). Up to now nothing is known about the “hypothetical proteins” CAB063, CAB408 and CAB821.

Figure 2. Assignment of the identified immunoreactive proteins to functional categories based on clusters of orthologous genes. For detailed information about the composition of the assigned categories, see results.

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Expression profiles of CAB063 and CAB080 during the C. abortus growth cycle

We selected the “hypothetical protein” CAB063 predicted to be secreted by the Type III secretion system and CAB080 (cMIP), another putative virulence factor, which is not supposed to be secreted [22] for comparison of their expression profiles at RNA and protein levels. Both factors led to efficient antibody production after immunizing mice (CAB080, CAB063) and a rabbit (CAB063) with the corresponding recombinant protein thereby permitting analysis of the expression and the subcellular localization of the chlamydial proteins during the developmental cycle by cell fractionation, immunofluorescence microscopy, and transfection experiments.

For transcriptional profiling total RNA was prepared from HeLa cells infected with C. abortus at 2, 20, 32, 44, 56 and 68 hpi. The number of transcripts was calculated against chlamydial genome copies for each time point as described by Wheelhouse et al. [5]. As an “internal control” we included the expression profile of CAB048 (ompA), and compared it to the profile previously described [5] (Figure 3c). Both profiles turned out to be similar. Transcript expression of ompA showed a significant rise between 2 and 20 hpi and exhibited peak expression at 32 hpi before decreasing again.

For CAB063 the results of the real-time PCR assays showed no significant gene expression up to 32 hpi. Subsequently an increase led to a maximum of gene expression at 44 hpi followed by decrease of transcripts to about half of maximum at 68 hpi (Figure 3a). The expression pattern of cMIP showed maximal expression between 32 and 44 hpi (Figure 3b).

In order to study the expression profile of CAB063 and cMIP at the protein level and to determine their subcellular localization during the developmental cycle C. abortus infected cells were harvested at 24 hpi, 36 hpi, 42 hpi, and 48 hpi. Subcellular fractionation was performed for the successive isolation of cytosolic proteins (fraction I), organelle specific proteins (fraction II), and nuclear proteins (fraction III). As already shown by others [17] we confirmed the enrichment of chlamydial inclusions of infected HeLa cells within the organelle specific fraction by use of polyclonal cTUF antibodies, which detected the chlamydial elongation factor solely in this fraction (see Figure 4a). Furthermore, selectivity of subcellular extraction could be demonstrated by immunoblotting with antibodies directed against fraction specific marker proteins such as HSP 90 (cytosolic fraction I), calnexin (organelle specific fraction II), and PARP (nuclear fraction III) (see Figure 4a).

At all time points analysed the full-length protein of cMIP was only found within the organelle/inclusion fraction II (see Figure 4b). This localization was confirmed by immunofluorescence microscopy of HeLa cells infected with C. abortus at 48 hpi (see Figure 5a). As expected the ectopic expression of cMIP in HeLa cells led to enrichment of the chlamydial protein in the cytoplasm of transfected HeLa cells (see Figure 6a).

The full-length protein of CAB063 was detected within the organelle fraction at 24 hpi and additionally in the nuclear fraction at 36 hpi, 42 hpi, and 48 hpi (see Figure 4c). The occurrence of smaller fragments might be due to biological protein processing in chlamydiae or might just reflect some artefactual phenomenon such as insufficient inhibition of protease activities. No signal was found in the “eukaryotic cytoplasmic fraction I”. Nuclear localization of CAB063 was confirmed by immunofluorescence microscopy of C. abortus infected HeLa cells at 48 hpi (see Figure 5b) and by transfection experiments. As shown in Figure 6b and Figure 6c ectopic expression of CAB063 led to strong accumulation of the chlamydial protein in the nucleus of all transfected HeLa cells.

Discussion

To identify novel candidates for serodiagnosis as well as putative virulence factors of C. abortus we performed a comprehensive analysis of immunoreactive proteins in animal and human C. abortus infection. Combining serological proteome analysis and screening of an expression library we identified 48 immunoreactive proteins and assigned them to functional categories that comprised energy metabolism, carbohydrate metabolism, lipid metabolism and several other macromolecular processes as well as membrane proteins and “hypothetical proteins” (see Figure 2). Prominent antigens in human as well as in animal infection included MOMP, HSP60, HSP70, and elongation factor TU, most of which have been previously described as immunogenic in other chlamydial species [13,23]. In addition further immunoreactive proteins could be identified such as the homologues of CPAF (CAB712), MIP (CAB080), TARP (CAB167), and PMPD (CAB776); all of them were also discussed as virulence associated factors in other chlamydial infections [6,7,20,22,24]. Eighteen % of the identified proteins could not be assigned to a functional category. Three of these so called hypothetical proteins have been predicted to be secreted by the T3SS (CAB063, CAB408, and CAB821). This study presents the first experimental evidence of their immunogenicity and hence their expression in vivo in humans and animals.

Our results show that both experimental procedures for detection of immunoreactive proteins may complement each other. 2D analysis detected full-length proteins in high resolution and led to the identification of 42 chlamydial antigens. Nevertheless, the sensitivity of this method is limited by the relative abundance of certain proteins within the analyzed EBs. Furthermore this approach can not detect proteins that are expressed only during growth within the host cell. So it was not surprising that a further six proteins were identified by screening of an expression library such as CAB712, the homologue of the C. trachomatis serine-type peptidase CPAF, which has been shown to be expressed only in RBs and which is described as a virulence factor that degrade host transcription factors required for major histocompatibility complex antigen expression in C. trachomatis [24,25]. Moreover, as most reactive library clones do not express full-length proteins but just fragments, this method also offers the chance for the identification of polypeptides comprising exclusively immunogenic domains, which subsequently may be used for further evaluation with greater numbers of sera in a diagnostic assay format. In fact, we
Figure 3. Transcriptional expression of CAB063 (a), CAB080 (b), and CAB048 (c) between 2 and 68 hpi. Transcript levels were normalised relative to the genome numbers for each time point. ± SEM of three independent experiments (see M&M).

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Figure 4. Subcellular localization of cMIP and CAB063 during the developmental cycle of HeLa cells infected with C. abortus. I, cytosolic proteins; II, organelle specific proteins comprising chlamydial inclusions; III, nuclear proteins. CL +, lysate of HeLa cells infected with C. abortus. CL -, lysate of HeLa cells not infected with C. abortus.

a. Selectivity of subcellular extraction was confirmed by immunoblotting with antibodies directed against the indicated marker proteins: HSP90 (cytosolic fraction, I); chlamydial elongation factor TUF and the endoplasmic reticulum protein calnexin (organelle fraction comprising chlamydial inclusions, II); nuclear Poly ADP ribose polymerase, PARP (nuclear fraction, III).

b. Detection of CAB080 (cMIP) solely in the organelle/chlamydial inclusion fraction II of HeLa cells infected with C. abortus.

c. Detection of CAB063 specific signals in fraction II of HeLa cells infected with C. abortus at all time points and additionally in the nuclear fraction at 36 hpi, 42 hpi, and 48 hpi. Arrows indicate the full-length protein of CAB063 (54 kDa).

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Figure 5. Expression and subcellular localization of cMIP and CAB063. Immunofluorescence microscopy of *C. abortus* infected HeLa cells fixed with PFA at 48 hpi shows localization of cMIP (Figure 5a) and CAB063 (Figure 5b) within the chlamydial inclusions and additional CAB063 specific signals in the host cell nucleus. (A) Evans blue, cytoplasmic counterstaining; (B) anti-CAB080 mouse antibodies (Figure 5a) and anti-CAB063 rabbit antibodies (Figure 5b), respectively; (C) DAPI, staining of nuclei. Scale bar = 5 µm.

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Figure 6. Ectopical expression of cMIP and CAB063 in HeLa cells. HeLa cells were transfected with pCI-CAB080 (a) and with pCI-CAB063 (b, c), respectively. The cells were analyzed by immunofluorescence microscopy using (A) Evans blue for cytoplasmic counterstaining, (B) anti-CAB080 mouse antibodies (a) and anti-CAB063 rabbit antibodies (b, c), respectively, and (C) DAPI to stain nuclei (c). Scale bar = 5 µm.

a, Enrichment of cMIP (CAB080) in the cytoplasm of a transfected HeLa cell located next to a not transfected cell. b, c: Ectopically expressed CAB063 localizes to the host cell nucleus showing an intensified rim around the nucleus. b and c show representative examples of transfected cells, fixed with PFA (b, showing a not transfected and two transfected cells) and methanol (c), respectively.

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CAB821 as well as CAB063, CAB167 and CAB408 are predicted to be secreted via T3SS (http://liferay.csb.univie.ac.at/portal/web/chlamydiaedb). CAB167 shows homology to the type III secreted effector TARP of C. trachomatis, which is implicated in the recruitment of actin at the site of chlamydial attachment to promote internalization into the host cell [20]. Nothing is known about the function of the “hypothetical” proteins CAB063, CAB408 and CAB821. As these potential T3SS effectors may play a role in chlamydial pathogenesis we started to analyse the expression and subcellular localization of one of these proteins, CAB063, in comparison to the MIP homologue CAB080 during the developmental cycle.

cMIP showed similar expression patterns at RNA and protein levels revealing maximum expression between 32 and 44 hpi. By cell fractionation, the full-length protein could be only detected within the “chlamydial inclusion fraction” (fraction II) throughout the developmental cycle. This subcellular localization was confirmed by immunofluorescence microscopy of HeLa cells infected with C. abortus. We could not observe any enrichment of the protein in the inclusion membrane as described for the MIP homologue of C. pneumoniae [30].

Transcripts of CAB063 were observed in the late stage of infection after 32 hpi showing only low level expression (less than 1 copy per genome). A similar low rate of expression showing less than 1 copy per genome has been already described for several polymorphic membrane proteins of C. abortus [5]. The expression pattern was confirmed at the protein level by subcellular fractionation of HeLa cells infected with C. abortus at different time points during the developmental cycle. The presence of CAB063 within the nuclear fraction corresponds to nucleus specific signals found by immunofluorescence microscopy and is consistent with the presence of a nuclear localization signal (NLS) "TRG_NLS_MonoExtN_4" (RPLKRKAP, aminoacid positions 214-221) that has been shown to be recognised by the importer protein importin-alpha [31]. Strikingly, nuclear localization of CAB063 was also observed when ectopically expressed by transfection of HeLa cells indicating that no further chlamydial proteins are necessary for translocation of CAB063 to the host cell nucleus and supporting the biological function of the identified NLS. CAB063 is the first C. abortus protein shown to be translocated to the host cell nucleus. To our knowledge only four chlamydial proteins with nuclear localization have been described until now [18,32,33]. While the C. trachomatis DUF582 proteins (CT620, CT621, CT711) are more abundant in the cytosol than in the nucleus, the "nuclear effector protein" NUE (CT737) shown to be a histone methyltransferase is localized predominantly in the nucleus. Interestingly, sequences of NUE as well as of CAB063 contain an internal nuclear localization signal.

Conclusion
We identified 48 immunoreactive proteins in ovine and human C. abortus infection including novel candidates for serodiagnosis as well as putative virulence factors. A systematic evaluation of the corresponding recombinant proteins with larger numbers of sera is on the way to confirm their diagnostic usefulness. Although the function of CAB063 remains to clarify this study presents the first experimental evidence for translocation of a C. abortus protein into the host cell nucleus at the end of the developmental cycle.

Supporting Information
Table S1. Immunoreactive C. abortus-proteins identified by 2D immunoblot analysis. The signal intensities of reactive protein spots are presented semiquantitatively: (-), negative; (+), very weak, corresponding to the intensity of MIP (spot 15) in Figure 1d; +, weak, corresponding to the intensity of MIP in Figure 1b; ++, moderate corresponding to the intensity of MIP in Figure 1c; ++++, strong, corresponding to the intensity of MIP in Figure 1a. Column 4 shows the calculated molecular mass and column 5 the isoelectric point (pl) of the identified proteins.

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Author Contributions
Conceived and designed the experiments: VFB AE. Performed the experiments: VFB CF US MW. Analyzed the data: VFB CF US MW AE. Contributed reagents/materials/analysis tools: DL GW CB MG KS. Wrote the manuscript: VFB AE KS.

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