Molecular cloning and characterization of NcROP2Fam-1, a member of the ROP2 family of rhoptry proteins in *Neospora caninum* that is targeted by antibodies neutralizing host cell invasion *in vitro*

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**SUMMARY**

Recent publications demonstrated that a fragment of a *Neospora caninum* ROP2 family member antigen represents a promising vaccine candidate. We here report on the cloning of the cDNA encoding this protein, *N. caninum* ROP2 family member 1 (NcROP2Fam-1), its molecular characterization and localization. The protein possesses the hallmarks of ROP2 family members and is apparently devoid of catalytic activity. NcROP2Fam-1 is synthesized as a pre-pro-protein that is matured to 2 proteins of 49 and 55 kDa that localize to rhoptry bulbs. Upon invasion the protein is associated with the nascent parasitophorous vacuole membrane (PVM), evaucules surrounding the host cell nucleus and, in some instances, the surface of intracellular parasites. Staining was also observed within the cyst wall of 'cysts' produced *in vitro*. Interestingly, NcROP2Fam-1 was also detected on the surface of extracellular parasites entering the host cells and antibodies directed against NcROP2Fam-1-specific peptides partially neutralized invasion *in vitro*. We conclude that, in spite of the general belief that ROP2 family proteins are intracellular antigens, NcROP2Fam-1 can also be considered as an extracellular antigen, a property that should be taken into account in further experiments employing ROP2 family proteins as vaccines.

Key words: *Neospora caninum*, rhoptries, ROP2 family, invasion.

**INTRODUCTION**

*Neospora caninum*, a member of the protozoan phylum Apicomplexa, is closely related to *Toxoplasma gondii*, with whom it shares a number of features, one of which is the intracellular lifestyle. To ensure host cell entry and intracellular survival and replication, most apicomplexan parasites possess specialized secretory organelles, namely micronemes (MIC), rhoptries and dense granules (DG). These organelles play a crucial role in invasion and modulation of the host cell (*Dubrelement et al. 1993; Carruthers and Sibley, 1997; Ngo et al. 2000; Joiner and Roos, 2002*). The contents of these 3 secretory organelles are sequentially released in order to establish invasion and to ensure the maintenance of the parasite–host cell interaction (*Fourniaux et al. 1996; Carruthers et al. 1999; Huynh et al. 2003; Carruthers and Tomley, 2008*). Initially, MIC proteins are secreted either prior to, or immediately upon, contact with the host cell membrane and they mediate adhesion and attachment to the host cell surface. The secretion of both rhoptry neck (RON) and MIC proteins contributes to the biogenesis of the moving junction complex (*Mordue et al. 1999; Besteiro et al. 2009*), a transient feature that associates with the host cell plasma membrane and propels the parasite into the developing parasitophorous vacuole (PV) (*Alexander et al. 2005; Lebrun et al. 2005*). The PV membrane (PVM) is formed from the host plasma membrane together with proteins secreted from the rhoptries (*Saffer et al. 1992; Ngo et al. 2004; Lebrun et al. 2005; Besteiro et al. 2009*). The DG proteins are exocytosed into the PV both during and after invasion and they either remain soluble in the lumen of the PV or become associated with the PVM or the tubulovesicular network (TVN), a membranous structure within the PV. DG proteins modify the environment within the PV, thereby performing essential functions related to intracellular survival and replication (*Mercier et al. 2005*).

Rhoptry proteins have been largely investigated in *T. gondii*. Subcellular fractionation and proteomic analysis allowed the identification of about 34 rhoptry proteins (*Bradley et al. 2005*), which were classified into 3 groups. The first group includes rhoptry bulb (ROP) proteins not exhibiting any homology in *Plasmodium falciparum* and homologous proteins were found only in genera closely related to *Toxoplasma* including *Neospora* and *Sarcocystis*, but...
not in *Eimeria*. The second group includes proteins restricted to the RON, which are conserved throughout the phylum Apicomplexa, including *Plasmodium* spp. The third group is formed of proteins encoding functions that are common to most eukaryotic cells, such as kinases, phosphatases and proteases (Bradley et al. 2005).

Proteins of the ROP2 family are among the best-studied rhoptry proteins in *Toxoplasma* and have been implicated in vaccine development (Dlugonska, 2008). This family possesses at least 12 members that share a number of features. All precursors of the ROP2 family have a signal peptide and most of them harbour a pro-region that is cleaved during maturation. The eventual maturation site is followed by an arginine-rich (R-rich) domain of 100–150 residues. Their C-termini consist of a kinase domain (El Hajj et al. 2006; Reese and Boothroyd, 2009), but then necessary catalytic residues for ATP-dependent phosphorylation are found only in some proteins of the family (such as ROP11, 17 and 18) but not in others (ROP2, 4, 5, 7 and 8). Rhoptry proteins have also been identified in *N. caninum* (Marugán-Hernández et al. 2011; Regidor-Cerrillo et al. 2011).

The function of most ROP2 family proteins is still largely unknown. Most members appear to play a role in PV biogenesis and in the modulation of the functional properties of the PVM. Anti-sense RNA-mediated depletion of TgROP2 resulted in several harmful effects, such as an impairment of rhoptry biogenesis and cytokinesis, a reduced capacity to invade and replicate *in vitro*, as well as an attenuation of virulence in mice (Nakaar et al. 2003). It was initially suggested that *T. gondii* ROP2 forms a tight interaction between the PV and the host cell mitochondria and endoplasmic reticulum (Sinai and Joiner, 2001; Nakaar et al. 2003; Boothroyd and Dubremetz, 2008); however, recent work by Pernas and Boothroyd (2010) showed this was not the case.

Several studies demonstrated essential roles for TgROP18 in parasite virulence (Saeij et al. 2006; Taylor et al. 2006; Melo et al. 2011; Niedelman et al. 2012). *In vivo*, overexpression of TgROP18 increased intracellular tachyzoite multiplication (El Hajj et al. 2007b) and TgROP18 was shown to phosphorylate other ROPs (ROP2, 4 and 8) on the PVM. In addition, TgROP18 binds to and phosphorylates immunity-related GTPases, preventing the clearance of intracellular parasites within inflammatory monocytes and IFN-γ-activated macrophages, thus conferring parasite survival *in vivo* and promoting virulence (Fentress et al. 2010; Steinfeld et al. 2010). In contrast to *T. gondii*, ROP18 in *N. caninum* is a pseudogene and the parasite is unable to phosphorylate the immunity-related GTPase 6 (Reid et al. 2012).

To date, no rhoptry protein has been described and characterized in *N. caninum* and only comparative genomics of *T. gondii* and *N. caninum* revealed that although both the genome and gene expression are remarkably conserved between species, some rhoptry genes are highly divergent (Reid et al. 2012). However, several studies from our group demonstrated that a recombinant, bacterially expressed C-terminal moiety of NcROP2Fam-1 (formerly designated recNcROP2) induced highly effective immunity against challenge infection with *N. caninum* tachyzoites in both cerebral and fetal infection mouse models (Debache et al. 2008, 2009, 2010).

In the present work, we report on the cloning of the cDNA encoding a full-length NcROP2Fam-1, the molecular characterization of the protein, its localization and its translocation during invasion of target cells by tachyzoites. Moreover, we show that NcROP2Fam-1 is expressed in both tachyzoite and bradyzoite stages. Lastly, using antisera produced against NcROP2Fam-1 antigenic peptides, potential immunoprotective regions of the protein have been identified.

**MATERIALS AND METHODS**

**Cell and parasite cultures**

Vero cells and human foreskin fibroblasts (HFF) were cultured at 37 °C under 5% CO2 in RPMI 1640 medium containing 10% fetal calf serum (Gibco-Invitrogen, Carlsbad, CA, USA), 2 mm L-glutamine, 50 IU mL−1 penicillin and 50 μg mL−1 streptomycin. *Neospora caninum* (Nc-Liverpool isolate) tachyzoites were grown in Vero cells in the same conditions but replacing fetal calf serum by 10% horse serum (Hemphill and Gottstein, 1996). Cultures of *N. caninum*-infected keratinocytes were prepared as described previously (Vonlaufen et al. 2002).

Free parasites were obtained by repeatedly passing infected cells through a 25G, 5/8 needle, washed in ice-cold medium and filtered through PD-10 columns (GE Healthcare, Piscataway, NJ, USA), as previously described (Hemphill and Gottstein, 1996).

Tachyzoite to bradyzoite *in vitro* stage conversion was performed using a published protocol (Guionaud et al. 2010), except that sodium nitroprusside (SNP; Sigma, St Louis, MO, USA) was used at 50 μM for 6 days instead of 25 μM for 9 days. BAG1, a classical bradyzoite-specific marker (McAllister et al. 1996; Vonlaufen et al. 2002, 2004), was used to monitor the conversion efficiency. For this purpose, parasites were purified from SNP-treated cells and the expression levels of BAG1 and *Bag1* mRNA were compared with those of tachyzoites by immunofluorescence (IF) and quantitative RT-PCR (qRT-PCR), respectively (Fig. S1 – in Online version only).
Oligonucleotides

Neospora caninum-specific primers were designed based on nucleotide sequences in the ToxoDB database ((Kissinger et al. 2003; Gajria et al. 2008); http://toxodb.org/toxo/) and on the formerly available ApiDots database ((Li et al. 2003); http://www.cbil.upenn.edu/apidots/). All primers were purchased from MWG (Ebersberg, Germany).

RNA isolation and first strand cDNA synthesis

Total RNA was isolated from N. caninum Liverpool tachyzoites and first-strand cDNA synthesis reactions were performed as described previously (Matz et al. 1999; Guionaud et al. 2010). Primers NcROP2Fam-1-R (Table 1) and T-primer 3 (5′-AAGCAGTGGTATCAACGCAGAGTAC(T)VN-3′, where V = A, C or G and N = any nucleotide) were used to prime the first-strand cDNA synthesis reaction for the 5′ RACE or conventional RT-PCR and 3′ RACE, respectively. A manual hot-start PCR was used in order to reduce background amplifications in conventional PCR and 3′-RACE reactions.

The cDNA corresponding to the 5′-end of NcROP2Fam-1 was amplified by RT-PCR in a 100 μL reaction volume containing 5 μL (1/10th) of the NcROP2Fam-1-R-primed first strand cDNA, 20 pmol of each NcROP2Fam-1-5′F (5′-TTCAGAACATGCCCCGCTAAC-3′) and NcROP2Fam-1-5′R (5′-TTGGAGAGGAAGAAGGCGGT-3′) primers, 0.2 mM of each dNTP and 2 U of Phusion High-Fidelity DNA polymerase (Finnzymes) in the supplied HF reaction buffer. Thirty-five PCR cycles with an annealing temperature of 58 °C were performed.

The remainder of the NcROP2Fam-1 cDNA was obtained by 3′-RACE. The 3′-RACE was carried out in a 100 μL reaction mixture containing 2 μL (1/10) of the first-strand cDNA prepared with T-primer 3, 4 pmol of Heel-carrier primer (5′-CTAATACGACTCACTATAGGGCAAGCAGTGATCAACGCAGAGTAC(T)_30 VN-3′), 20 pmol of each NcROP2Fam-1-3′F (5′-TTGGAGAGGAAGAAGGCGGT-3′) and NcROP2Fam-1-3′R (5′-TTGGAGAGGAAGAAGGCGGT-3′) primers, 0.2 mM of each dNTP and 2 U of Phusion High-Fidelity DNA polymerase (Finnzymes) in the supplied HF reaction buffer. The PCR reaction comprised 10 cycles with an annealing temperature (T_a) of 62 °C and 30 further cycles with a T_a of 55 °C.

Sequencing and sequence analyses

NcROP2Fam-1 5′-end and 3′-RACE amplicons were cloned into pCR Blunt II TOPO (Invitrogen) and sequenced on both strands using a primer walking approach. Five independent clones were sequenced.
for each PCR product. Sequencing reactions were carried out using BigDye v3.1 fluorescent dye terminators and run on an ABI PRISM 3100 Genetic Analyzer (Applied Biosystems, Foster City, CA, USA). Raw sequencing data were assembled and edited with the Staden package (http://staden.sourceforge.net/).

Homology searches were done using BLAST (http://www.ncbi.nlm.nih.gov/blast/) and the Conserved Domain Database CD-Search (http://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi) with default settings. Proteins were aligned on the Muscle server (http://www.drive5.com/muscle/), minimally edited and formatted with GeneDoc (http://www.nrbsc.org/gfx/genedoc/). Residue grouping and shading was according to the structurally derived matrix (SDM) 12 reduced alphabet (Prlic et al., 2000; Solis and Rackovsky, 2000; Peterson et al., 2009). Potential signal peptide cleavage sites were identified with SignalP 3.0 (http://www.cbs.dtu.dk/services/SignalP/).

Potential alpha helices in the RAH domain were searched using Jpred3 (http://www.compbio.dundee.ac.uk/www-jpred/), PSIPRED v3.0 (http://bioinf.cs.ucl.ac.uk/psipred/) and PSSpred (http://zhanglab.ccmb.med.umich.edu/PSSpred/). The properties of the helices (mean hydrophobicity $[H]$ and mean hydrophobic moment $[\mu H]$) were analysed on the HelixQuest server ([Gautier et al. 2008]; http://helixquest.ipmc.cnrs.fr) and helical wheel projections were plotted using a Java applet (http://rzlab.ucr.edu/scripts/wheel/wheel.cgi).

For Neospora (Liverpool) vs Toxoplasma (ME49) genomic sequence comparisons, 50 kb segments of genomic DNA centred on the loci of interest were extracted from the ToxoDB database and low complexity regions were identified with Repeat Masker (http://www.repeatmasker.org/cgi-bin/WEBRepeatMasker). Genomic dot plots were generated using Advanced PipMaker ([Schwartz et al. 2000]; http://pipmaker.bx.psu.edu/cgi-bin/pipmaker?advanced).

Quantitative reverse transcription-PCR

RNA transcript steady-state levels were determined by two-step qRT-PCR. Total RNA from free N. caninum tachyzoites or from free parasites cultured in bradyzoite conditions (up to 6 days in the presence of 50 μM SNP, as indicated) was prepared using Trizol reagent (Invitrogen) and RNA concentrations were adjusted to 200 ng μL$^{-1}$. Since some of the transcripts analysed derive from intron-less genes, RNA was digested with RNase-free DNaseI to degrade contaminating genomic DNA. Briefly, 5 μg RNA was incubated for 10 min at room temperature with ~7 Kunitz units of DNaseI in 100 μL of the 1× supplied buffer (Qiagen). DNaseI was heat-inactivated for 5 min at 95 °C and RNA cleaned-up using RNeasy total RNA purification kit columns (Qiagen). RNA was eluted from the columns with RNase-free water and its concentration measured with a spectrophotometer (NanoDrop ND-1000; Thermo Scientific). First-strand cDNA synthesis was carried out in 40 μL, using 3 μg RNA and Omniscript reverse transcriptase (RT) following the manufacturer’s instructions (Qiagen), except that 1 μg random hexamers (Promega) were used to prime RT reactions. We performed control reactions without RT to exclude the presence of remaining genomic DNA in the cDNA preparations. RT was inactivated by heating for 5 min at 95 °C. Quantitative PCR was carried out using the QuantiTect SYBR Green PCR Kit (Qiagen). The reaction mix contained 1 × SYBR Green Mix, 0.5 μM each primer (Table 1), cDNA (~ 40 ng) and water to a final volume of 20 μL. Quantitative PCR was performed on the Rotor-Gene 6000 (Corbett Research, Sydney, Australia). Gene expressions were determined (each sample run in triplicate) using the second derivative maximum method and a standard curve based on 4-fold dilutions (ranging from 1 : 4 up to 1 : 256, each in triplicate) of a positive cDNA sample. Gene expression values were normalized to the housekeeping Neospora alpha tubulin transcripts. For the procedure and associated calculations, we followed the ‘relative standard curve method’ (Singleplex) using an independent sample for a standard curve’ guidelines, as described in the Applied Biosystems online guide to real-time quantitative PCR (http://www3.appliedbiosystems.com/cms/groups/mcb_support/documents/generaldocuments/cms_042380.pdf).

After each run, a high-resolution melting analysis was performed to confirm the identity of the amplified products.

Anti (α-) NcROP2Fam-1 polyclonal antibodies (Ab)

Regions of NcROP2Fam-1 towards which affinity-purified Ab are directed are presented in Fig. S2A and B – in Online version only.

A fragment of NcROP2Fam-1 (aa 238–594), encompassing the whole kinase-like domain and referred to here as recNcROP2Fam-1, was expressed in Escherichia coli as a recombinant (His)$_6$-tagged fusion protein. Details on the construction of the recombinant protein, on the production and affinity-purification of rabbit α-NcROP2Fam-1 polyclonal Abs and use thereof to neutralize invasion in vitro can be found elsewhere (Debache et al. 2008). The recombinant protein, previously dubbed recNcROP2, was used in a number of vaccination trials in mice (Debache et al. 2008, 2009, 2010).

In addition to α-recNcROP2Fam-1 (kinase-like domain) polyclonal Abs, 2 rabbit polyclonal
anti-peptide antisera were produced. During the first round of peptide selection we only retained peptides with a high potential antigenicity, as predicted by the consensus epitope search algorithm of the BEPTOPE software (Odorico and Pellequer, 2003). Peptides with significant identity to mammalian cell antigens or to other ROP2 family members were eliminated.

The 2 selected NcROP2Fam-1 peptides, P1 (aa 459–472; CH₃CO-CSRQGRSYPDRKHKK-R-CONH₂) and P2 (aa 581–594; CH₃CO-CQGDDGVDDETDLR-CO₂H), with an N-terminal cysteine residue added for coupling (underlined), were synthesized and used to immunize rabbits (Thermo Scientific). Anti-peptide polyclonal α-P1 and α-P2 Abs were affinity-purified using the Thio-Link Gel Kit (Severn Biotech, Kidderminster, UK).

Protein extracts, SDS-polyacrylamide gel electrophoresis (SDS-PAGE) and Western blots

Total protein extracts from free parasites or from infected or uninfected Vero cells were prepared as previously described (Peixoto et al. 2010). We used 10 μL mL⁻¹ Halt protease inhibitor cocktail (Pierce-Thermo Scientific) in the lysis buffer. Extracts were clarified by centrifugation and the protein concentration in the supernatant was measured using the DC Protein Assay Kit (Bio-Rad, Hercules, CA, USA).

Protein samples were mixed with reducing, denaturing sample buffer, boiled for 5 min and resolved by SDS-PAGE. Proteins were transferred onto nitrocellulose or PVDF membranes (Bio-Rad) for 1 h at 100 V using a wet transfer apparatus (Bio-Rad). Membranes were blocked by incubation in TBST containing 5% non-fat dry milk. Blots were then incubated for 2 h at room temperature with the following primary antibodies (Abs), in PBS/0.3% BSA, were applied for 1 h: (a) affinity-purified rabbit polyclonal Abs α-P1, α-P2 (this work) or α-recNcROP2Fam-1 (Debache et al. 2008 and this work), each at a 1 : 2000 dilution; (b) a-whole N. caninum extract rabbit antisemum (Hemphill et al. 1996), diluted 1 : 2500; (c) CC2, a rat monoclonal Ab reacting with a yet uncharacterized cyst wall antigen (Gross et al. 1995), diluted 1 : 300; (d) a-BAG1, a polyclonal rabbit Ab directed against BAG1, a classical bradyzoite-specific antigen (McAllister et al. 1996; Vonlaufen et al. 2002, 2004), diluted 1 : 300; (e) a-SAG1, a rat monoclonal Ab directed against the major immunodominant tachyzoite surface antigen (Bjorkman and Hemphill, 1998), diluted 1 : 300; and (f) E7, a mouse monoclonal Ab directed against alpha-tubulin (Developmental Studies Hybridoma Bank), diluted 1 : 2000.

Incubations with primary antibodies were followed by 3 washes in PBS for 5 min each. Alexa Fluor (AF)-labelled secondary antibodies (Invitrogen) at a dilution of 1 : 3000 (AF-488, AF-568) or 1 : 300 (AF-350) in PBS/0.3% BSA were then applied. Finally, the preparations were washed 3 times in PBS for 5 min each and samples were mounted in ProLong Gold anti-fade reagent (Invitrogen). Nuclei were eventually counterstained with 4′,6-diamidino-2-phenylindole (DAPI; Sigma) at 1 μg mL⁻¹. Images were acquired on an Axioskop 2 fluorescence microscope (Carl Zeiss, Oberkochen, Germany). Confocal microscopy was performed on a FluoView FV1000 laser scanning microscope (Olympus, Tokyo, Japan). Images were processed with Image J 1.38 (http://rsb.info.nih.gov/ij/).

Immunogold-labelling and transmission electron microscopy

LR-White embedding and on-section labelling of N. caninum cultures were performed as previously described (Hemphill et al. 1997). Sections were loaded onto Formvar-carbon coated grids and non-specific binding sites were blocked for 2 h in PBS/1% BSA. They were then incubated with rabbit affinity-purified α-P1 or α-P2 antibodies diluted at 1:100 in glutaraldehyde in PBS), distributed onto poly-L-lysine coated coverslips, left for 15 min at room temperature in the fixative and subsequently kept in pre-cooled acetone/methanol (ν:ν) for 30 min at −20 °C for permeabilization.

Unless otherwise stated (time-course of infection in vitro, see below), cells were fixed and simultaneously permeabilized with acetone/methanol. Briefly, coverslips in 24-well plates were rinsed once in PBS, immersed into pre-cooled (−20 °C) acetone/methanol (ν:ν) and incubated at −20 °C for 2 h.

For IF staining, coverslips were subsequently washed 3 times with PBS, 5 min each, incubated in blocking buffer (PBS/3% BSA) for 2 h and the following primary antibodies (Abs), in PBS/0.3% BSA, were applied for 1 h: (a) affinity-purified rabbit polyclonal Abs α-P1, α-P2 (this work) or α-recNcROP2Fam-1 (Debache et al. 2008 and this work), each used at a 1 : 2000 dilution; (b) a-whole N. caninum extract rabbit antisemum (Hemphill et al. 1996), diluted 1 : 2500; (c) CC2, a rat monoclonal Ab reacting with a yet uncharacterized cyst wall antigen (Gross et al. 1995), diluted 1 : 300; (d) a-BAG1, a polyclonal rabbit Ab directed against BAG1, a classical bradyzoite-specific antigen (McAllister et al. 1996; Vonlaufen et al. 2002, 2004), diluted 1 : 300; (e) a-SAG1, a rat monoclonal Ab directed against the major immunodominant tachyzoite surface antigen (Bjorkman and Hemphill, 1998), diluted 1 : 300; and (f) E7, a mouse monoclonal Ab directed against alpha-tubulin (Developmental Studies Hybridoma Bank), diluted 1 : 2000.

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PBS/0.1% BSA for 1 h. After washing in 5 changes of PBS (2 min each), goat α-rabbit secondary Ab was conjugated to 10 nm diameter gold particles (Amersham) and applied at a dilution of 1:5 in PBS/0.1% BSA. After extensive washing in PBS, grids were air-dried and stained with lead citrate and uranyl acetate. Specimens were viewed on a Philips 400 TEM (Philips Electronics, Eindhoven, the Netherlands) operating at 80 kV.

In order to determine the subcellular localization of NcROP2Fam-1, we analysed a series of transmission electron microscopy (TEM) photographs labelled using α-P2 Ab. We only took into consideration rhoptries/portions of rhoptries (N = 34, 23 longitudinal/oblique sections and 11 transversal sections) with (i) sufficient contrast to check for the eventual localization of gold nanoparticles within electron-dense rhoptries and (ii) for which the rhoptry surface was clearly delineated.

**Immunolocalization of NcROP2Fam-1 during a time-course of infection in vitro**

The fate of NcROP2Fam-1 during infection in vitro was assessed by IF after large numbers of parasites were left to adhere to cells for a short duration. NcROP2Fam-1 was then chased for up to 24 h. Briefly, Vero cells (5×10⁴) were seeded onto glass coverslips in 24-well tissue culture plates and cultured for 24 h until a subconfluent monolayer (~10⁵ cells) was obtained. The culture medium was aspirated from the tissue culture plates and 2×10⁶ freshly purified parasites in 200 μL of RPMI 1640 supplemented with horse serum (complete RPMI medium) were added to each well (parasites : cells ratio = 20). After 3 min at room temperature, free non-adherent parasites were washed away twice with PBS and once with RPMI 1640. Pre-warmed (37°C) complete RPMI medium (1 mL) was then added to the wells and plates were returned to the 37°C/5% CO₂ incubator for further short (5–90 min) or longer (24 h) periods of incubation. Infected cultures were then washed with PBS and either (i) fixed with 4% PFA in PBS for 20 min at room temperature and processed immediately for IF analysis or (ii) fixed with 4% PFA in PBS for 20 min at room temperature and permeabilized with 0.1% saponin (Sigma) in PBS for 10 min at room temperature.

**Invasion inhibition assays**

Invasion inhibition assays were carried out in 6-well plates, as previously described (Debache et al. 2008). Wells were inoculated with 2×10⁵ HFF and grown to confluency. Briefly, freshly purified tachyzoites (1×10⁵) were pre-incubated for 30 min at room temperature in 200 μL RPMI 1640 medium containing 1:100 dilutions of the following rabbit polyclonal Abs: α-P1, α-P2, α-recNcROP2Fam-1 (kinase-like domain) or rabbit α-NC1 (whole N. caninum strain NC1 lysate). As controls, tachyzoites were pre-incubated in rabbit pre-immune serum (1:100) or in RPMI 1640 medium.

Parasite suspensions were added to HFF monolayers in 6-well plates for 30 min at 37°C/5% CO₂. Monolayers were washed 3 times with cold RPMI 1640 and incubated with RPMI 1640 medium containing 100 μM pyrrolidine dithiocarbamate (PDTC), 0.2 μM CuSO⁴ and α-N. caninum hyper-immune serum (1:200) for 2 h at 37°C/5% CO₂ to permeabilize adherent, non-invaded (extracellular) tachyzoites. Wells were then washed 3 times with cold RPMI 1640 and incubated for 1 h at 37°C/5% CO₂ with RPMI 1640 containing DNaseI (Roche; 1 mg mL⁻¹). Wells were then washed twice with cold RPMI 1640 and a third time with RPMI 1640 containing 1 mM EDTA. Finally, monolayers were trypsinized and the DNA was purified (High Pure PCR Purification kit; Roche Diagnostics). DNA concentrations were measured by Hoechst 33258 (Sigma) fluorimetry (Auszubel et al. 1997) on a Synergy HT plate reader (Biotek Instruments, Winooski, VT, USA). The number of parasite equivalents in each DNA sample (200 ng per assay, each in triplicate) was assessed by real-time PCR on a Light Cycler instrument (Roche Diagnosis, Basel, Switzerland) using an established protocol (Müller et al. 2002).

**Cloning and sequence analysis of NcROP2Fam-1**

NcROP2Fam-1 cDNA was submitted to the DDBJ/EMBL/GenBank databases under accession number HM587954.

**RESULTS**

**Cloning and sequence analysis of NcROP2Fam-1**

A sequence homology search against N. caninum expressed sequence tags (ESTs) in the ApiDots database (Li et al. 2003) using T. gondii ROP2 (TgROP2) cDNA (Beckers et al. 1994) as query sequence identified DT.92484732 as best hit. This contig spans ~2.8 kb and potentially encodes a 594 amino acid (aa) protein with an overall 47% identity to TgROP2.

To confirm the DT.92484732 data, we initially planned to clone and sequence the corresponding cDNA as 2 overlapping 5’- and 3’-RACE products. First-strand cDNA synthesized from total RNA isolated from N. caninum Liverpool tachyzoites was prepared for this purpose. Several attempts to amplify the 5’-end of the transcript by 5’-RACE after tailing the first-strand cDNA with either (dG)₅₉ or (dA)₅₉ homopolymers were not successful and resulted in 5’-truncated products (data not shown). Finally, we amplified a nearly complete 5’-end cDNA
by conventional RT-PCR. The 3′-end of the DT.92484732 cDNA was obtained by 3′-RACE. We cloned the 5′-end RT-PCR and 3′-RACE products and sequenced 5 clones of each. A composite, consensus 2278 bp cDNA sequence made out of the 5′-end RT-PCR and 3′-RACE overlapping products was submitted to the DDBJ/EMBL/GenBank databases under accession number HM587954.

The cDNA possesses a 1758 nucleotides open reading frame (ORF) starting with an ATG initiation codon that is part of a sequence (TTTACCATGG) matching 7 positions of the previously described T. gondii translation initiation consensus sequence (gNCAAAATGG) (Seeber, 1997). The ORF potentially encodes a 594 aa protein matching perfectly the predicted translated product of ApiDots DT.92484732/ToxoDB NCLIV_001970 transcripts. In addition to the ~47% identity shared with T. gondii ROP2 and ROP8, the most closely related rhoptry proteins in Toxoplasma (Fig. S3 – in Online version only), the predicted protein also shares 52% identity with the predicted protein encoded by NCLIV_001950, a tandem, expressed gene (Figs S3 and S4B – in Online version only) located on N. caninum Liverpool chromosome Ia. Both genes are located in a region of synteny shared with the region around the ROP4/7 locus on Toxoplasma ME49 chromosome Ia (Fig. SSA – in Online version only, left panel). Such a syntenic region does not exist on chromosome X between Neospora NCLIV_001970/NCLIV_001950 and the Toxoplasma ROP2/8 locus (Fig. S5A, right panel). Therefore, in spite of a higher homology of Neospora NCLIV_001970 and NCLIV_001950 predicted proteins with TgROP2 and TgROP8 (Fig. S3), we do not believe these gene pairs are orthologous. A recent report (Reid et al. 2012) established NCLIV_001950 and NCLIV_001970 as the syntenic orthologues of Toxoplasma ROP4 and ROP7, respectively. While this might be correct for the pair NCLIV_001950/ROP4, our genomic analysis suggests that it is unlikely that ROP7 and NCLIV_001970 are orthologues (Fig. SSB). For this reason, in the present work, NCLIV_001970 was named NcROP2Fam-1 (N. caninum ROP2 family member 1) and NCLIV_001950 was named NcROP2Fam-2.

NcROP2Fam-1 possesses all features of ROP2 family proteins

The predicted NcROP2Fam-1 precursor is 67 kDa and its deduced primary sequence exhibits all characteristics of ROP2 family members (Fig. S3A). The protein contains an N-terminal predicted signal peptide with a computed cleavage site between residues 28 and 29. In its N-terminal domain, a sequence is found (SWLQ, aa 82–85), that is reminiscent of the SΦxΦ consensus (where Φ is a hydrophobic residue and x is any aa) proposed in T. gondii as a likely processing site for the TgSUB2 protease by Miller et al. (2003) and that was recently re-defined as SΦxx (E/D) by Hajagos et al. (2011). Whether this site in NcROP2Fam-1 actually represents a cleavage site for a TgSUB2-like Neospora maturase is uncertain, since there is a negatively charged (E) to neutral (Q) amino acid substitution at position P1. A second, although perfect match to the TgSUB2 consensus (SWDD, aa 39–42) was identified upstream in the NcROP2Fam-1 precursor. This site is not conserved in other ROP2 family members (Fig. S3A). Three arginine-rich sequences (R-rich 1–3), collectively referred to as the RAH (R-rich Amphipathic Helix) domain, were also found in the N-terminal domain of NcROP2Fam-1 (aa 111–126, 138–159 and 168–179; precursor numbering; Fig. S3A and Fig. 1). This domain was recently shown to mediate the attachment of ROP2 family proteins to membranes and especially to the PVM (Reese and Boothroyd, 2009; Fentress et al. 2012). Combining secondary structure predictions and α-helix properties analysis, we inferred the presence of 2 amphipathic α-helices within the portion of the RAH domain containing R-rich 2 and R-rich 3 (Fig. 1). Whereas R-rich 1 harbour a predicted α-helix (aa 110–122) it did not have a marked hydrophobic face (Fig. 1).

Additionally, a serine/threonine protein kinase-like domain was detected in the C-terminal moiety of NcROP2Fam-1 (aa 257–546; Fig. 1 and Fig. S3A). However, this domain lacks 3 of the most invariant key residues (K, D and D residues of Hanks motifs II, VIb and VII, respectively) commonly found in ATP-binding and catalytic sites of active kinases (Hanks and Hunter, 1995). It is therefore unlikely that NcROP2Fam-1 is an active kinase. NcROP2Fam-1 also possesses a hydrophobic stretch (aa 473–489; Fig. S3A) encompassing Hanks domain IX that is buried within the core of other rhoptry proteins adopting a kinase fold (El Hajj et al. 2006, 2007a, 2007b; Labesse et al. 2009; Qiu et al. 2009).

Lastly, a non-canonical motif (FEN1; aa 546–549, Fig. S3A), reminiscent of the YxxΦ motif (where x is any aa and Φ is hydrophobic), as well as 7 dileucine motifs (LL; not shown) that may function as sorting signals mediating trafficking to the rhoptries (Hoppe et al. 2000; Ngo et al. 2003) were found in NcROP2Fam-1.

NcROP2Fam-1 undergoes a proteolytic maturation

On Western blots of lysates of free N. caninum tachyzoites, α-P2 antibodies, raised against the C-terminal tail of NcROP2Fam-1, detected 2 major bands of 55 and 49 kDa, as well as a fainter band of 64 kDa, the latter being compatible with the predicted size of the pro-NcROP2Fam-1 precursor (Fig. 2). The 55 and 49 kDa bands were repeatedly observed in all lysates of tachyzoites, regardless of
which of the 3 antibodies was used (Fig. S2C—in Online version only) or if parasites were directly boiled in sample buffer or not. We therefore exclude that the major 55 kDa or the fainter 49 kDa bands may be artifactual (either resulting from protein degradation in a sample or from the cross-reactivity of one antibody with a ROP protein or any unrelated *Neospora* protein). The same protein pattern was also observed in lysates of parasite-infected cells, although the 64 kDa pre-pro-protein was not visible in all samples (data not shown). Additional faint bands of 30–40 kDa, not present in lysates of free tachyzoites, were detected in infected cells lysates and are likely to be due to protein degradation. There was no visible cross-reactivity of α-P2 with uninfected Vero cell lysate (Fig. 2).

The positions of the processing sites inferred from the sizes of the observed mature NcROP2Fam-1 are not compatible with a cleavage at any of the putative TgSUB2 sites mentioned above. Indeed, a cleavage of pro-NcROP2Fam-1 would give rise to a 62·5 kDa or a 58 kDa mature protein, depending on whether the upstream or the downstream site is used, respectively. Instead, our results indicate that pro-NcROP2Fam-1 is cleaved around F 112 (pre-pro-protein numbering; major 55 kDa band), leaving the whole RAH domain intact, or around F 157 (minor 49 kDa band), leaving only R-rich 3 on mature NcROP2Fam-1 (Fig. 2).

*NcROP2Fam-1 is associated with rhoptry bulbs in parasites*

Affinity-purified antibodies were used to localize NcROP2Fam-1 by IF and TEM. In routine cultures of infected cells, after membrane permeabilization, NcROP2Fam-1 was detected within the apical region of intracellular parasites, at locations compatible with a rhoptry staining (Fig. 3A). In some infected cells, and only with a standard but not with a confocal microscope, we observed a punctate staining within the PV and in the host cell cytosol (not shown).
Immunogold-TEM was performed on *N. caninum* -infected keratinocytes using α-P1 or α-P2 antibodies. In tachyzoites, labelling was clearly associated with rhoptry bulbs (Fig. 3B; Fig. S2D and Table S1 in Online version only).

*NcROP2Fam-1* is secreted and associates with the PVM and the surface of invaded and adherent parasites

*N. caninum* tachyzoites were allowed to interact with host cells for 3 min and non-invaded/non-adherent parasites were washed away. This time-point was designated as time-point zero of infection. At 5 min post-infection (p.i.) and without permeabilization, anti-NcROP2Fam-1 antibodies revealed a punctate staining pattern on the surface of a small amount of parasites per coverslip, while a monoclonal Ab directed against cellular alpha-tubulin applied simultaneously failed to stain the host cell cytoskeleton, confirming that under our experimental conditions the cellular membrane was not compromised and therefore that only adherent, but not already invaded, parasites were accessible to antibodies (Fig. 4A). The labelling on the parasite pictured presents a truncation at the anterior end, where it already has penetrated the host cell; this indicates that the parasite membrane was also impermeant to antibodies and that NcROP2Fam-1 is indeed on its surface. The small number of parasites we detected per slide is most certainly due to the fact that invasion processing rapidly and therefore most parasites were already within the host cells at 5 min p.i. After 5 min p.i., tachyzoites were not detected anymore without permeabilization, indicating that they all had completely invaded the host cells (data not shown), suggesting that the parasites that were labelled with NcROP2Fam-1 antibodies at 5 min p.i. were viable and not arrested in the adhesion phase.
In permeabilized preparations, antibodies strongly labelled the newly formed PV (Fig. 4B). In addition, NcROP2Fam-1 was also observed as punctuations associated with cytoplasmic filaments that we identified as evacuoles (Hakansson et al. 2001; Saeij et al. 2006; Taylor et al. 2006; Turetzky et al. 2010) (Fig. 4B, filled arrowheads). At 30–60 min p.i., most parasites appeared closely associated with the host cell perinuclear region. In addition, NcROP2Fam-1-immunoreactive evacuoles were longer, sometimes branched, and most of them surrounded the host cell nucleus (not shown). On some occasions we clearly observed that NcROP2Fam-1 was not only detected on the nascent PVM, but also on the surface of intracellular parasites (Fig. 4, empty arrowheads). NcROP2Fam-1 staining intensity decreased progressively starting from 60–90 min p.i. and at 24 h p.i., the protein was exclusively distributed at the parasite apex, consistent with its localization in rhoptries (not shown).

Tachyzoites and bradyzoites express similar levels of NcROP2Fam-1

_Neospora caninum_ bradyzoites were generated in vitro by growing parasites for 6 days in the presence of SNP added to the culture medium. Preparations of were left in contact with cells. (A) NcROP2Fam-1 is present on the parasite surface of some adherent _Neospora caninum_ tachyzoites. The following primary antibodies and appropriate Alexa Fluor-labelled secondary Abs were used: 12G10, a monoclonal Ab reacting against a broad range of alpha-tubulins (α-tub., red) was used to check the integrity of the host cell membrane in the absence of permeabilization and α-P2 Ab (green) was used to stain NcROP2Fam-1. Nuclei were counterstained with DAPI. Only the merged (3-colours) pictures are presented. Top picture: no permeabilization prior to staining. The filled arrowhead points towards an adherent parasite coated with NcROP2Fam-1. A higher magnification view of the parasite is shown in the box. Note the truncated aspect of the extracellular NcROP2Fam-1 staining in the apical region (empty arrowheads), indicating that (i) this adherent parasite was actively invading the host cell when invasion was arrested by fixation and (ii) the parasite body was itself impermeant to antibodies. Bottom: a permeabilized control slide was processed in parallel to check the efficiency of alpha-tubulin staining by 12G10 mAb (red). Scale bar = 5 μm; (B) NcROP2Fam-1 is present on the PVM, in evacuoles and on the surface of intracellular parasites after invasion. Images were taken using a laser scanning confocal microscope showing a unique plane. NcROP2Fam-1 was detected with α-P2 Ab (green) and α-SAG1 was used to stain the parasites (red in Merge panel) after permeabilization. DIC images are presented on the left. Filled arrowheads: staining of evacuoles; empty arrowheads: staining of both the PVM and the parasite surface. Scale bars = 5 μm.
tachyzoites and bradyzoites were split into 2 aliquots each. The first aliquots of infected cells were used for IF experiments. Free parasites were purified from the second aliquots and were used for controlling the tachyzoite to bradyzoite stage conversion by IF, for the assessment of \textit{NcROP2Fam-1} levels by real-time RT-PCR and for Western blot experiments. The efficiency of the stage conversion was controlled at day (d) 6 by checking the expression of \textit{NcBag1} (a classical bradyzoite marker) in parasites isolated from the culture. Nearly 70\% of individual d6 parasites displayed a stronger BAG1 staining than d0 parasites (tachyzoites), while \textit{BAG1} transcripts increased by 15-fold (Fig. S1).

\textit{NcROP2Fam-1} transcripts in tachyzoites and bradyzoites were quantified by real-time RT-PCR. \textit{NcROP2Fam-1} steady-state levels were very similar in tachyzoites (d0), at d3 of stage conversion and in cultures containing mainly bradyzoites (d6) (Fig. 5A), indicating that \textit{NcROP2Fam-1} is not a gene with a stage-specific pattern of transcription. Using the same approach we could show that \textit{NcROP2Fam-2} is also transcribed in both tachyzoites and bradyzoites, but appears to be 2-to3-fold more expressed in d6 bradyzoites than in d0 tachyzoites (Fig. S4B).

Western blots of lysates prepared from purified tachyzoites (d0) and from d6 bradyzoites indicated...
that both stages expressed the two major forms (55 and 49 kDa) of mature NcROP2Fam-1 (Fig. 5B).

Next, we compared by IF the localization of NcROP2Fam-1 in infected cells cultured in the presence of SNP for 6 days (bradyzoite conditions) to its localization in infected cells grown in tachyzoite conditions (no SNP, 48 h p.i.). NcROP2Fam-1 staining was observed in the rhoptries of all intracellular parasites, irrespective of the culture conditions. We also noticed that NcROP2Fam-1 staining was much more prominent in d6 cysts (bradyzoite culture) than in PVs (tachyzoite culture; staining not visible with exposure settings in Fig. 5C). NcROP2Fam-1 and the bradyzoite-specific CC2-reactive cyst wall antigen are both found at the periphery of cysts (Fig. 5C).

Selected specific anti-peptide antibodies directed against NcROP2Fam-1 partially neutralize host cell invasion in vitro

In a previous study from our group (Debache et al. 2008), incubation of Neospora tachyzoites in the presence of antibodies against rec-NcROP2Fam-1 significantly reduced invasion in vitro, thus suggesting that NcROP2Fam-1 may have a previously unsuspected function in adhesion and/or invasion and that it may also be accessible to antibodies. Since rec-NcROP2Fam-1 likely shares common epitopes with other ROP2 family members in Neospora (Fig. S3A), the neutralization effect observed could possibly have been due to recognition of a cross-reacting antigen on the parasite. We therefore tested whether anti-antigen are both found at the periphery of cysts (El Hajj et al. 2006), yet no protein of this family has so far been characterized in the related genus Neospora. Using a reverse genetics approach, we identified and cloned a cDNA encoding a ROP2 family member in N. caninum. The cDNA sequence was in perfect agreement with the original NCLIV_001970 in ToxoDB v. 7.1 (Kissinger et al. 2003; Gajria et al. 2008). The NCLIV_001970 deduced primary amino-acid sequence showed most significant homology to NCLIV_001950, an expressed, tandemly arranged gene on Neospora chromosome Ia, and to a lesser extent to Toxoplasma rhoptry proteins (ROP2, 4, 7, 8 and 18). It was also surprising to notice that apparently Neospora does not possess a ROP18 gene, hence NcROP18 is pseudogenized, as recently demonstrated by Reid et al. (2012). In addition, within the same clade only two genes (NcROP2Fam-1 and NcROP2Fam-2) were found in Neospora, whereas Toxoplasma has a set of four to five (Tg ROP2 (a/b), 4, 7 and 8). These differences may reflect a degree of redundancy in the functions of Toxoplasma proteins or could contribute to different strategies of adaptation of the two species in their respective hosts (Reid et al. 2012). The predicted NcROP2Fam-1 protein possesses the hallmarks of ROP2 family proteins (El Hajj et al. 2006), each of which will be further discussed. In the N-terminal moiety of the predicted NcROP2Fam-1 we found a potential signal peptide, as well as an approximate match (SWLQ) and a perfect match (SWDD) to the...
SΦx(E/D) consensus for the TgSUB2 maturase (Bradley and Boothroyd, 1999; Miller et al. 2003; Hajagos et al. 2011), suggesting that NcROP2Fam-1 might be synthesized as a pre-pro-protein. The N-terminal moiety of NcROP2Fam-1 also contained an RAH domain made out of three R-rich regions and a serine/threonin protein kinase-like (PK-L) domain.

Due to the high overall homology of the NcROP2Fam-1 PK-L domain with those of TgROP2 and TgROP8, for which the structure has been established (Labesse et al. 2009; Qiu et al. 2009), we anticipate that the PK-L domain of NcROP2Fam-1 certainly also adopts the kinase fold. Unlike TgROP18, the only ROP2 family protein with a demonstrated kinase activity (El Hajj et al. 2007b) and similar to most other members of the ROP2 family in *Toxoplasma* (El Hajj et al. 2006), NcROP2Fam-1 lacks essential, conserved residues present within the catalytic core of classical kinases (Hanks and Hunter, 1995). NcROP2Fam-1 can therefore be qualified as a pseudokinase (Boudeau et al. 2006). However, it would be difficult to make any assumption as to whether NcROP2Fam-1 might nevertheless have a kinase activity since a number of pseudokinases, such as WNKs (with-no-lysine (K)) or CASK (Ca²⁺/calmodulin-dependent serine kinase) have evolved alternative ways of binding ATP and performing the phosphoryl transfer reaction (Xu et al. 2000; Mukherjee et al. 2008).

In order to proceed with the characterization of NcROP2Fam-1, it was necessary to obtain specific antibodies that would not cross-react with e.g. other ROP2 family members. The selection process we used in choosing unique regions of NcROP2Fam-1 for the generation of specific anti-peptide antibodies is fully detailed in the Materials and Methods section. The fact that both anti-NcROP2Fam-1 anti-peptide antibodies (α-P1 and α-P2) recognized the same patterns in Western blot and immunolocalization (IF and TEM) experiments provides strong evidence that our selection strategy was successful.

Most rhoptry proteins studied are synthesized as pre-pro-proteins (Sadak et al. 1988; Bradley and Boothroyd, 1999; Carey et al. 2004; El Hajj et al. 2006, 2007b; Turetzky et al. 2010; Hajagos et al. 2011). The signal peptide is cleaved as soon as the precursor enters the secretory pathway; however, the removal of the pro-domain occurs much later, within the nascent rhoptries (Soldati et al. 1998).

Western blot analysis of purified tachyzoite lysates with three different antibodies raised against the C-terminal moiety of the protein revealed that at least two N-terminal processing events must occur following the synthesis of the NcROP2Fam-1 precursor. A faint band of 64 kDa was observed, the size of which was compatible with the predicted pro-NcROP2Fam-1. In addition, we observed two additional bands of 55 and 49 kDa (the 55 kDa band being prominent), indicating a further maturation of pro-NcROP2Fam-1. TgSUB2, a subtilisin-like protease present in rhoptry bulbs, is viewed as the most likely maturase candidate for N-terminal maturation of rhoptry proteins (Miller et al. 2003). In earlier studies, it was shown that the cleavage occurs at SΦxE sites on rhoptry proteins (Bradley and Boothroyd, 1999; Miller et al. 2003) and that mutations at the P1 position of the cleavage site (E in SΦxE) had a deleterious effect on the processing (Bradley et al. 2002; Miller et al. 2003; Turetzky et al. 2010). However, a recent study by Hajagos et al. (2011) showed that the P1 site could accommodate a conservative E to D substitution, thus re-defining the TgSUB2 consensus as SΦx(E/D). In NcROP2Fam-1, we found a perfect match to the cleavage consensus (SWDD (aa 39–42)) at a position not conserved in ROP2 family proteins and a poorer match (SWLQ (aa 82–85)) at a conserved position, but with a negatively charged (E) to neutral (Q) substitution at the P1 site that would likely impair cleavage. Remarkably, neither the 55 nor the 49 kDa bands that we observed in Western blots are compatible with a maturation of NcROP2Fam-1 at any of the above-mentioned sites (the expected sizes would be 62.5 and 58 kDa). We therefore conclude that the processing of pro-NcROP2Fam-1 in *Neospora* involves a protease with a different substrate specificity than that of TgSUB2. The bands we observed also do not fit with the yet unidentified rhoptry protein maturase acting at RAMA1-like sites (SxL) in *T. gondii* (Hajagos et al. 2011). We therefore cannot determine the exact locations of the two maturation events that take place in pro-NcROP2Fam-1. In lysates of infected cells, additional bands of 30–40 kDa, which were not present in lysates of cell-free, purified parasites, were detected. This indicates that NcROP2Fam-1 may be subjected to a variety of additional proteolytic activities once it has been secreted by the parasites.

In eukaryotic cells, sorting of integral proteins to subcellular organelles involves conserved mechanisms mediated by cytoplasmic adaptor complexes (APs), which recognize short motifs, usually either tyrosine-based (YxxΦ) or dileucine-based (LL), exposed on the cytoplasmic tail of the cargo (Bonifacino and Lippincott-Schwartz, 2003; Robinson, 2004). *Toxoplasma* rhoptry proteins are also sorted through the secretory pathway (Joiner and Roos, 2002; Sheiner and Soldati-Favre, 2008) using similar mechanisms. Indeed, it was shown that targeting of TgROP2 to the rhoptries is mediated by the clathrin/adaptor protein 1 (AP1) complex (Ngo et al. 2003). Likewise, a number of rhoptry-addressing signals have been identified either in the pro-region or in mature rhoptry proteins (Hoppe et al. 2000; Bradley and Boothroyd, 2001; Striepen et al. 2001). In the case of TgROP2, there is strong evidence that these rhoptry-addressing signals are tyrosine-based (YxxΦ) or dileucine-based (LL) (Hoppe et al. 2000; Ngo et al. 2003). Although NcROP2Fam-1 harbours...
a non-canonical, most likely non-functional, tyrosine-based-like motif, it also possesses 7 LL motifs, one of which (aa 535–536) is conserved across most ROP2 family members and was shown to be involved in the targeting of TgROP2 to the rhoptries (Ngo et al. 2003). However, structural determination of PK-L domains of TgROP2 and TgROP8 has established that these domains do not contain any transmembrane helix (Labesse et al. 2009; Qiu et al. 2009), thus questioning the possibility of a direct interaction of the previously identified C-terminal sorting motifs (Hoppe et al. 2000; Ngo et al. 2003) with cytosolic APs. Interestingly, in P. falciparum most rhoptry proteins are not integral proteins with a cytoplasmic tail but they are nonetheless targeted to the rhoptries (Kats et al. 2006). In Plasmodium, RAP1 (Rhoptry Associated Protein 1) is first directed to the rhoptries via its interaction with the glycosphatidyl inositol-anchored RAMA1 (Rhoptry Associated Membrane Antigen 1), which in turn interacts with components of the trafficking machinery, possibly via a transmembrane escorter (Richard et al. 2009). Such a complex scaffolding mechanism may also exist for members of the ROP2 family in Toxoplasma and Neospora.

We confirmed by IF and TEM that NcROP2Fam-1 indeed belongs to the set of rhoptry proteins. NcROP2Fam-1 was found in the subapical region of free or intracellular parasites and was associated with rhoptry bulbs, consistent with the findings of other ultrastructural localization studies of ROP2 family proteins in T. gondii (Sadak et al. 1988; Saffer et al. 1992; Soldati et al. 1998; Lee et al. 2001; Carey et al. 2004; Taylor et al. 2006; Slove et al. 2012).

The invasion of a cell by T. gondii is accompanied by a rapid discharge of the contents of rhoptries that is concomitant with the formation of the PV. The trigger for the secretion is presently unknown but it is widely accepted that it involves an interaction between the parasite apex and one or more cellular receptor(s). During the secretion process, RONs serve as ducts that direct the contents of the rhoptries towards an opening at the apex of parasites (Nichols et al. 1983; Porchet-Hennere and Nicolas, 1983; Dubremetz et al. 1993; Carruthers and Sibley, 1997; Alexander et al. 2005; Boothroyd and Dubremetz, 2008). Upon secretion, ROP proteins either remain in the lumen of the PV, associate with the PVM, or are injected into the host cell cytoplasm and may then reach further locations e.g. the host cell nucleus (Hakansson et al. 2001; Carey et al. 2004; Hajj et al. 2006; Saeij et al. 2006; Taylor et al. 2006; El Hajj et al. 2007a; Turetzky et al. 2010). We detected NcROP2Fam-1 on the surface of intracellular parasites, within the nascent PVM and cysts, and antibodies also stained weakly the PVM of mature vacuoles. This indicates that the protein is secreted and is able to bind a variety of biological membranes. It was recently shown that the RAH domain of ROP2 family members is composed of R-rich, amphipathic helices that individually confer rhoptry proteins the property to associate with membranes and act in concert to bind preferentially to the PVM (Labesse et al. 2009; Reese and Boothroyd, 2009). An RAH domain composed of three R-rich regions was found in NcROP2Fam-1. In NcROP2Fam-1, the amphipathic characteristics of the α-helices predicted within R-rich regions 1–3 were not as obvious as with Toxoplasma ROP2 family proteins. In fact, only the predicted α-helices within R-rich 2-R-rich 3 clearly appeared amphipathic. Interestingly, whereas each region can individually bind cellular membranes, R-rich 2 is the main determinant of the preferential association with the PVM (Reese and Boothroyd, 2009).

By Western blot analysis, we observed that the processing of pro-NcROP2Fam-1 led to 2 mature proteins of 55 and 49 kDa, predicted to have retained either the whole RAH domain or only R-rich 3, respectively. It would be of interest to test which form(s) of mature NcROP2Fam-1 can actually associate with membranes and if the short, 49 kDa mature NcROP2Fam-1, having lost R-rich 2, possesses a skewed membrane selectivity compared with the 55 kDa protein.

NcROP2Fam-1 was also found on intracytoplasmic ribbon-like structures in the host cell. We identified these structures as being evacuoles since they were similar to the evacuoles observed with other rhoptry proteins in the absence of treatment by cytochalasin D (Hakansson et al. 2001; Saeij et al. 2006; Taylor et al. 2006; Turetzky et al. 2010). The parasite migration towards the host cell nucleus followed the growth of evacuoles in the direction of the perinuclear region, suggesting that NcROP2Fam-1 may play a role in parasite migration or in the establishment of the PV in the vicinity of the host cell nucleus. Finally, and certainly most intriguing, was the fact that NcROP2Fam-1 was also detected on the surface of extracellular, adherent tachyzoites that had partially invaded the host cell, as well as on the surface of intracellular parasites. To the best of our knowledge, this is the first time a rhoptry protein is observed on the surface of tachyzoites at the time of entry and immediately after invasion of the host cell. Rhopty protein secretion takes place exclusively in the apical region of the parasite and is triggered by contact with a host cell (Joiner and Roos, 2002). In this context, it is likely that some of the NcROP2Fam-1 proteins exocytosed during the secretion burst that follows adhesion to the host cells bind to the parasite membrane in a fashion reminiscent of T. gondii micronemal proteins MIC2 (Carruthers et al. 1999, 2000) and MIC3 (Garcia-Reguet et al. 2000), both of which are exocytosed at the parasite apex but redistribute to the whole surface of the parasite body.
Since similar levels of NcROP2Fam-1 transcripts were found in tachyzoites and bradyzoites, transcription of the NcROP2Fam-1 gene is most certainly not developmentally regulated in these stages. We also showed that tachyzoites and bradyzoites express the same forms of NcROP2Fam-1 proteins. NcROP2Fam-1 IF labelling of the nascent PVM was always strong shortly after invasion but rapidly fell at levels close to the detection threshold past 90 min p.i., thus requiring much longer exposure times. The protein could not usually be detected on the PVM of older vacuoles. The decrease of NcROP2Fam-1 staining intensity associated with the PVM that we observed over time has also been shown for T. gondii rhoptry proteins in the hours following invasion. It was usually attributed to protein turnover and to the dilution of PVM-bound ROP proteins consequent to the expansion of the PV surface as the vacuole grows. Although staining of the PVM fell below detectable levels past 90 min p.i., NcROP2Fam-1 labelling reappeared as punctuations on the PVM of a few mature vacuoles containing a few parasites (tachyzoite growth conditions), but was readily visible on cyst walls (bradyzoites). Taken together, these observations strongly suggest that a continuous secretion of NcROP2Fam-1 by intracellular parasites still takes place within the PV and the bradyzoite-containing cysts. The stronger staining intensity observed in cysts may be due to a higher affinity of NcROP2Fam-1 for cyst wall components and/or to a difference in protein secretion rate or turnover between these stages. We propose that the few mature PV that showed some NcROP2Fam-1 staining in tachyzoite culture conditions belonged to vacuoles containing parasites having acquired some bradyzoite features.

The region of NcROP2Fam-1 including the PK-like domain and the C-terminal tail that was bacterially expressed as the recNcROP2Fam-1 recombinant protein appears to play an unsuspected role in the invasion process. Indeed, the recombinant protein itself ((Debache et al., 2008) and this work) and antibodies directed against recNcROP2Fam-1-derived peptides (this work) partially neutralized N. caninum tachyzoite host cell invasion in vitro. This observation provides further support that the protein may be accessible on the surface of parasites during the invasion process. RecNcROP2Fam-1 is an efficient antigen against cerebral infection by N. caninum in adult mice and against a vertical transmission of the infection from the mother to the offspring (Debache et al., 2008, 2009, 2010). Classically, Th1 responses are elicited against intracellular antigens while Th2 responses are elicited against extracellular antigens. In all vaccination studies in mouse, recNcROP2Fam-1 induced predominantly either IgG2a (Th1-biased) or IgG1 (Th2-biased) Abs, depending on the adjuvant used but, nevertheless, the type of response elicited did not affect the protectiveity of the antigen against lethal challenge. This indicates that the protective effects of recNcROP2Fam-1 can be attributed to a combination of cellular (Th1) and humoral (Th2) responses.

We have demonstrated that highly immunogenic NcROP2Fam-1-specific peptides (P1 and P2), displaying no cross-reactivity with mammalian cells, could be used to produce antibodies specific to NcROP2Fam-1 that inhibit host cell invasion in vitro. This suggests that peptides P1 and P2, as well as yet-to-be-discovered similar peptides, could be included in future experimental vaccine formulations. In addition, NcROP2Fam-2, which is also transcribed in tachyzoites and bradyzoites, should be investigated for its potential role as a vaccine candidate, either alone or synergistically with NcROP2Fam-1.

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Molecular characterization of NcROP2Fam-1


