Biotinylation of the Neospora caninum parasitophorous vacuole reveals novel dense granule proteins

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Abstract

Background: Neospora caninum is an obligate intracellular parasite that invades host cells and replicates within the parasitophorous vacuole (PV), which resists fusion with host cell lysosomal compartments. To modify the PV, the parasite secretes an array of proteins, including dense granule proteins (GRAs). The vital role of GRAs in the Neospora life cycle cannot be overestimated. Despite this important role, only a subset of these proteins have been identified, and most of their functions have not been elucidated. Our previous study demonstrated that NcGRA17 is specifically targeted to the delimiting membrane of the parasitophorous vacuole membrane (PVM). In this study, we utilize proximity-dependent biotin identification (BioID) to identify novel components of the dense granules.

Methods: NcGRA17 was BirA* epitope-tagged in the Nc1 strain utilizing the CRISPR/Cas9 system to create a fusion of NcGRA17 with the biotin ligase BirA*. The biotinylated proteins were affinity-purified for mass spectrometric analysis, and the candidate GRA proteins from BioID data set were identified by gene tagging. To verify the biological role of novel identified GRA proteins, we constructed the NcGRA23 and NcGRA11 (a–e) knockout strains using the CRISPR/Cas9 system and analyzed the phenotypes of these mutants.

Results: Using NcGRA17-BirA* fusion protein as bait, we have identified some known GRAs and verified localization of 11 novel GRA proteins by gene endogenous tagging or overexpression in the Nc1 strain. We proceeded to functionally characterize NcGRA23 and NcGRA11 (a–e) by gene knockout. The lack of NcGRA23 or NcGRA11 (a–e) did not affect the parasite propagation in vitro and virulence in vivo.

Conclusions: In summary, our findings reveal that BioID is effective in discovering novel constituents of N. caninum dense granules. The exact biological functions of the novel GRA proteins are yet unknown, but this could be explored in future studies.

Keywords: Neospora caninum, Dense granule protein, BioID, Parasitophorous vacuole, CRISPR/Cas9

Background

Neospora caninum, a member of the phylum Apicomplexa, is an obligate intracellular parasite that causes neosporosis, with worldwide distribution [1]. The parasite has been recognized as one of the most important infectious causes of reproductive issues and abortion in cattle, and 12–42% of aborted bovine fetuses worldwide are infected [2, 3]. Given the high prevalence of the parasite, neosporosis is a major threat facing dairy and livestock industries and results in significant economic losses [3, 4].

Neospora caninum has three specific secretory organelles, namely micronemes, rhoptries, and dense granules, which sequentially discharge effectors during invasion. The dense granule (GRA) proteins are essential
in the formation of parasitophorous vacuoles (PVs) within which the parasites multiply and grow [5, 6]. There are three localizations in PVs within the infected cell where *Neospora* dense granule proteins (NcGRAs) are exhibited: the vacuolar space, intravacuolar membranous nanotubular network (MNN), and parasitophorous vacuole membrane (PVM). Although the dense granule proteome is hypothesized to be composed of hundreds of proteins, only nine GRA proteins (NcGRA1, NcGRA2, NcGRA6, NcGRA7, NcGRA9, NcGRA12, NcGRA14, NcGRA17, and NcGRA23) [7–15] have been previously identified in *N. caninum*; however, the precise roles of these GRA proteins in the intracellular survival and growth of the parasites are not well elucidated.

To date, most NcGRAs have been discovered individually by bioinformatics searches for proteins exhibiting homology to known *Toxoplasma* dense granule proteins (TgGRAs). Although the bioinformatics approach has been successful, we sought a method to identify a large number of dense granule components more rapidly. In this study, we adapted the proximity-dependent biotin identification (BioID) system for *Neospora* and used the dense granule protein NcGRA17 as bait to identify novel NcGRA proteins. We demonstrate that the NcGRA17-BirA* fusion localizes accurately to the dense granules, is catalytically active, and labels previously described NcGRA proteins as well as novel targets. A number of these targets were confirmed by gene overexpression or endogenous tagging in the Nc1 strain, thereby validating our approach. Subsequently, six of them were characterized by gene disruption. The discovery of novel NcGRA proteins provides a more comprehensive basis for the understanding of host–parasite interactions and *N. caninum* parasitism.

**Methods**

**Parasites and cell cultures**

All *Neospora* tachyzoites used in this study were propagated in a human foreskin fibroblast (HFF) line cultured in complete Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS, Gibco, USA), as described previously [14, 16]. Both cells and parasites were incubated at 37°C with 5% CO₂ in a humidified incubator.

**Plasmid construction**

All primer sequences are shown in Additional file 3: Table S1. The NcGRA17-targeting CRISPR plasmid (pNC_Cas9CRISPR::sgNcGRA17) was constructed as described previously [14]. All other CRISPR/Cas9 plasmids were obtained through FastPfu (TansGen Biotech Co., Ltd., China) mutagenesis of the above plasmid to change the NcGRA17 targeting gRNA to other specific gRNAs. To generate clean knockouts of NcGRA11 (a-e) genes (62240 bp), a double-gRNA CRISPR/Cas9 system was designed in which the first gRNA sequence (gRNA1) was placed close to the start codon of the first gene (NcGRA11a), with the second gRNA (gRNA2) near the stop codon of the last gene (NcGRA11e), as described previously [17]. To express two gRNAs from a single plasmid pNC_Cas9CRISPR::sgNcGRA11(a–e), the gRNA2 expression cassette (NcU6 promoter-gRNA2-RNA scaffold TTTTT) was amplified using a common set of primers, 2x gRNA NcU6 F and 2x gRNA NcU6 R, and cloned into the pNC_Cas9CRISPR::sgNcGRA11(a–e) plasmid using a one-step cloning kit (Vazyme Biotech Co., Ltd., China). For disruption of NcGRA11 (a–e) genes, CRISPR/Cas9 double-gRNA plasmid pNC_Cas9CRISPR::sgNcGRA11(a–e) was co-transfected with its corresponding DHFR-TS ampiclon containing 60-bp homology regions matching the NcGRA11 (a–e) genes. To generate a construct (pLIC-BirA*-HA-DHFR-NcGRA17) for inserting a BirA*-HA tag into the NcGRA17 gene 3’ end, upstream (830-bp) and downstream (950-bp) regions directly adjacent to the gRNA target sequence and BirA*(963 bp) sequence were cloned into the pLIC-HA vector. pDMG plasmid [16, 18, 19] was used to express NcGRA11 (a-e) genes in the Nc1 strain. The coding sequences of NcGRA11 (a–e) were cloned into the backbone plasmid pDMG with GFP replaced by HA [18] using a one-step cloning kit (Vazyme Biotech Co., Ltd., China). The vector p6 × HA-HXGPRT used for in situ C-terminal tagging, as described previously [20], was generously provided by Prof. Shaojun Long of China Agricultural University. This plasmid was modified by replacing the 6HA tag with the spaghetti monster tag sHxA [21] and replacing the selectable marker HXGPRT with DHFR. The construct (pTCR-NcGRA23-CD KO) for disrupting the NcGRA23 locus was designed by inserting the 5’ (990-bp) and 3’ (930-bp) regions flanking the NcGRA23 gene region into the backbone plasmid pTCR-CD, which has been described previously [18].

**Generation of NcGRA17-BirA* fusions**

C-terminal BirA*-HA-tagging of NcGRA17 was performed via homologous recombination. Nc1, as the parent strain, was co-transfected with the pNC_Cas9CRISPR::sgNcGRA17 vector and homologous repair ampiclon. The ampiclon was amplified using primers listed in Additional file 3: Table S1 and pLIC-BirA*-HA-DHFR-NcGRA17 plasmid as a template. The parasites were cultured to the third generation in the presence of pyrimethamine (1 µM) and then screened to confirm the purity of the selected strains. The selected NcGRA17-tagged strain, named NcGRA17-BirA*-HA,
was identified by polymerase chain reaction (PCR), western blotting, and immunofluorescence assay.

Affinity capture of biotinylated proteins
BirA*-tagged and parent lines were used to infect HFF monolayers and grown in medium containing 150 µM biotin for 48 h prior to parasite egress [22]. Infected cells were collected, washed in phosphate-buffered saline (PBS), and lysed in radioimmunoprecipitation assay (RIPA) buffer (Beyotime, China) supplemented with protease inhibitor cocktail (Sigma-Aldrich, USA) for 30 min on ice. Lysates were centrifuged at 15 min at 14,000×g to pellet insoluble debris and then incubated with streptavidin magnetic beads (Beaver, China) at room temperature for 4 h under gentle agitation. Beads were collected using magnets and washed five times in RIPA buffer, followed by three washes in 8 M urea buffer (50 mM Tris–HCl [pH 7.4], 150 mM NaCl). Ten percent of each sample was boiled in Laemmli sample buffer, and eluted proteins were analyzed by western blotting with streptavidin-horseradish peroxidase (HRP) before mass spectrometry (MS) analysis.

Mass spectrometry
Purified proteins bound to streptavidin beads were reduced, alkylated, and digested by the sequential addition of Lys-C and trypsin proteases [22]. The peptide mixture was desalted using C18 tips and fractionated by a 75 µm inner diameter filter fused to the silica capillary column with a 5 µm pulled electrospray tip, and packed in-house with 15 cm of a Luna C18 column with 3 µm reversed-phase particles. Delivery of the gradient was performed by the EASY-nLC 1000 ultrahigh-pressure liquid chromatography (UHPLC) system (Thermo Scientific). Tandem mass spectrometry (MS/MS) spectra were collected on a Q Exactive mass spectrometer (Thermo Scientific). Data analysis was performed using ProLuCID and DTASelect2 implemented in the Integrated Proteomics Pipeline (IP2) platform (Integrated Proteomics Applications, Inc., San Diego, CA, USA). Protein and peptide identification was filtered using DTASelect and required a minimum of two unique peptides per protein and a peptide-level false-positive rate of less than 5%, as estimated by a decoy database strategy. Normalized spectral abundance factor (NSAF) values were calculated as described previously [22].

Generation of NcGRA-HA parasites
As few data are available from Neospora, we utilized homologous gene expression data of Toxoplasma to filter hits from our BioID data set for further investigation. To epitope-tag the candidate NcGRA proteins, CRISPR/Cas9-sgNcGRA plasmid was co-transfected with their corresponding smHA tag amplicon containing 42-bp homology regions matching the gene of interest. The amplicon was amplified using primers listed in Additional file 3: Table S1 and psmHA-DHFR plasmid as a template. As we failed to endogenous HA-tag the NcGRA11b gene, the vector pDMG-NcGRA11b-HA was electroporated into Nc1. The transfected parasites were grown in medium containing 1 µM pyrimethamine, and identified by immunofluorescence assay (IFA). All confirmed NcGRAs were examined in silico using the Basic Local Alignment Search Tool (BLAST) to search for other Neospora proteins with sequence similarity.

Deletion of the genes encoding NcGRA23 and NcGRA11 (a–e)
NcGRA23-KO parasites were generated via homologous recombination. The parent Nc1 strain was co-transfected with the pNc_Cas9CRISPR::sgNcGRA23 and linearized complete knockout plasmid pTCR-NcGRA23-CD KO (primers listed in Additional file 3: Table S1). The transgenic parasites were grown under chloramphenicol (20 µM) and 5-fluorine cytosine (40 µM) selection pressure. The selected NcGRA23-deficient strain, named Δncgra23, was identified by PCR and western blotting. For disruption of NcGRA11 (a–e) genes, CRISPR/Cas9 double-gRNA plasmid pNc_Cas9CRISPR::sgNcGRA11(a–e)1–2 was co-transfected with its corresponding DHFR-TS amplicon containing 60-bp homology regions matching the NcGRA11 (a–e) genes. The parasites were cultured in the presence of pyrimethamine (1 µM) to the third generation. Resistant clones were isolated and confirmed by PCR and western blotting.

Western blotting
Western blotting was performed as described previously [14]. The parent and modified strains of whole-parasite lysates were resolved using sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) on a 12% (w/v) gel. Samples were transferred onto polyvinylidene fluoride membranes and probed with the mouse anti- HA antibody (Sigma-Aldrich, 1:500), the mouse anti-NcGRA23 antibody (prepared in our laboratory, 1:200), or the mouse anti-NcGRA11c antibody (prepared in our laboratory, 1:200). The N. caninum F-actin subunit beta (NActin) was used as a loading control and was incubated with the rabbit anti-NcActin antibody (prepared in our laboratory, 1:2000). For all secondary antibody incubations, HRP-labeled goat anti-mouse IgG (H+L) antibody (Sigma, USA) was diluted 1:5000, an HRP-labeled tablet was used with a 3-mm diameter circle filter.
goat anti-rabbit IgG secondary antibody (Sigma, USA) was diluted 1:10,000, or streptavidin-HRP (4A Biotech Co., Ltd., China) was used at a 1:1000 dilution. Following secondary incubation, enhanced chemiluminescence reagents (CoWin, China) were used for the detection of HRP activity.

**Immunofluorescence assay**

Immunofluorescence assay (IFA) to detect the biotinylated proteins of BirA*-tagged parasites was performed as described previously [22]. Parasites were used to infect HFF monolayers on coverslips in a 12-well plate and grown for 30 h with the addition of 150 μM biotin. Infected cells were fixed for 15 min in 4% formaldehyde, and then permeabilized with 0.25% Triton X-100 for 15 min and blocked with 3% bovine serum albumin (BSA) for 30 min. Subsequently, the cells were incubated with a rabbit anti-HA polyclonal antibody (CWBIOTECH, China, 1:50); then incubation was performed with Cy3-conjugated goat anti-rabbit IgG (H+L) (Sigma, USA) and streptavidin/fluorescein isothiocyanate (FITC) (Solarbio, 1:100). The images were obtained using a Leica confocal microscope system (Leica TCS SP52, Germany).

To analyze NcGRA localization, parasite-infected HFFs or freshly released parasites were processed for IFA and stained with a rabbit anti-HA polyclonal antibody, mouse anti-NcGRA6 polyclonal antibody (prepared in our laboratory, 1:50), or mouse anti-NcGRA23 polyclonal antibody (1:50). To determine the trafficking route of the protein following secretion into the PV, freshly harvested parasites (NcGRA17-HA or NcGRA11c-HA) were used to pulse-infect monolayers of HFF cells for 3 min at 37 °C. Monolayers were washed with PBS and further incubated at 37 °C for 5 min, 10 min, 15 min, or 1 h, and then fixed and permeabilized in 0.002% saponin [23]. Localization of the indicated NcGRAs was performed by double immunofluorescence labeling, using both the rabbit serum anti-HA and the mouse serum anti-NcGRA6 or anti-NcGRA23. Primary antibodies were revealed using FITC-conjugated goat anti-mouse IgG (H+L) and Cy3-conjugated goat anti-rabbit IgG (H+L). The IFA process and image collection were performed as described above.

**Plaque assay**

Plaque assays were performed on HFF cells cultured in 12-well plates (Corning Costar, USA) as described previously [14, 16]. Briefly, 200 freshly isolated parasites were seeded into HFF monolayers and incubated at 37 °C with 5% CO₂ for 9 days undisturbed. The plaques were scanned using a Canon digital scanner (model F917500, Japan), and the plaque area was measured as described previously [18].

**Parasite virulence assay**

The virulence of the parasites was investigated as described previously [14]. Eight-week-old female BALB/c mice (Laboratory Animal Center of Academy of Military Medical Sciences, Beijing, China) were injected intraperitoneally with Δncgra23, Δncgra11(a–e), or parent Nc1 parasites at doses of 5 × 10⁶ and 8 × 10⁶ parasites (n = 5 mice/dose). Mice were carefully monitored every 8 h based on their clinical signs and mortality, and humanely euthanized via cervical dislocation when they were unable to reach food or water for more than 24 h or lost 20% of normal body weight. Survival was monitored for 60 days.

**Statistical analysis**

All statistical data were analyzed using GraphPad Prism 5 (version 5.01; GraphPad Software, San Diego, CA, USA) except as otherwise indicated above. The results were expressed as mean ± SD and evaluated by non-parametric tests. Values of P < 0.05 and P < 0.01 were considered statistically significant.

**Results**

**Biotinylation of the PV using NcGRA17 as bait**

To identify novel NcGRA proteins, we adapted the BioID approach using NcGRA17 as bait. NcGRA17 was BirA* epitope-tagged in the Nc1 strain utilizing the CRISPR/Cas9-directed genome editing system to create a fusion of NcGRA17 with the biotin ligase BirA* (Fig. 1a; Additional file 1: Figure S1a–c). The localization of the NcGRA17-BirA* fusion was assessed by IFA. As expected, we observed staining in the PV (Fig. 1b), a pattern consistent with NcGRA7, indicating that the fusion did not alter the trafficking of NcGRA17. To determine which proximal proteins in the PV were labeled by the NcGRA17-BirA* tachyzoites, we grew parasites in medium supplemented with biotin and stained for biotinylated proteins using fluorophore-conjugated streptavidin. NcGRA17-BirA* tachyzoites showed a robust streptavidin staining in the PV, indicating that the biotin ligase fusion was active and that it labeled proteins in the PV (Fig. 1c, bottom). We then assessed the labeling of N. caninum proteins by NcGRA17-BirA* in whole-cell lysates by western blotting with a streptavidin-HRP probe. There was a significant increase in biotinylated proteins in lysates of NcGRA17-BirA* parasites, and only when biotin was present in the growth medium (Fig. 1d). Collectively, these data demonstrate that NcGRA17-BirA* is active and biotinylates proteins in the PV.

**Identification of novel dense granule proteins**

The target labeled was identified by NcGRA17-BirA* as we affinity-purified biotinylated proteins from
parasite lysates for MuDPIT mass spectrometric analysis. Biotinylated proteins from both parent and NcGRA17-BirA* lysates were significantly enriched using streptavidin-conjugated magnetic beads. Proteins identified by MS that were unique to NcGRA17-BirA* samples were scored as hits based on the number of identified spectra and unique peptides (see Additional file 4: Table S2 and Additional file 5: Table S3). Compared to a control lysate derived from Nc1 supplemented with biotin, the NcGRA17-BirA* purified fraction revealed two known dense granule proteins: NcGRA2 and NcGRA23 [7, 8]. As few data are available from Neospora, we utilized homologous gene expression data of Toxoplasma to filter hits from our BioID data set for further investigation.

We selected a group of hypothetical proteins whose homologous protein in Toxoplasma had no signature cyclical expression pattern, and were constitutively expressed and highly expressed at the tachyzoite stage. To localize these candidates, we used CRISPR/Cas9 to add smHA epitope tags to the C termini (Fig. 2a; Additional file 2: Figure S2), as described previously [20]. The overexpression plasmid pDMG-HA was also used to add 3×HA epitope tags to the C termini. Among our NcGRA17-BirA* hits, we epitope-tagged seven proteins that shared the same localization in extracellular parasites as NcGRA6: NCLIV_045870 (NcGRA3), NCLIV_012020 (NcGRA11b), NCLIV_042680 (NcGRA25), NCLIV_043760 (NcGRA27), NCLIV_055980 (NcGRA38), NCLIV_058760 (NcGRA45), and NCLIV_057330 (NcGRA61) (Fig. 2b). NcGRA3, NcGRA25, and NcGRA61 were located at the PVM, and the other proteins were located within the PV in intracellular parasites (Fig. 2b). Among the seven novel dense granule proteins, NcGRA3, NcGRA11b, NcGRA25, NcGRA38, and NcGRA45 have been previously described as orthologs in Toxoplasma.

Identification and infectivity potential of NcGRA23
NcGRA23 was chosen for analysis because it was the closest in sequence similarity (47.8% amino acid sequence similarity) to NcGRA17 and scored highly in the mass spectrometry data set in our data (Table 1; Additional file 4: Table S2). To examine the subcellular localization, the full-length NcGRA23 protein fused with a histidine-tag was successfully expressed and used to generate anti-rNcGRA23 serum in mice. IFA analysis showed that the NcGRA23 protein of the parasites overlapped with NcGRA17 and the protein of intracellular
tachyzoites was localized on the PVM (Fig. 3a). To determine the secretion characteristic of the protein, we also analyzed the secretion kinetics of the NcGRA23 at different time points (5 min to 1 h) after invasion. NcGRA23 was secreted into the PV at the apical end of the parasite following host cell invasion, consistent with the release of NcGRA17. At 10 min post-invasion, NcGRA23 was localized on both sides of the parasite, and then it was
gradually dispersed throughout the vacuole by 60 min (Fig. 3b).

In *Toxoplasma*, the TgGRA17 and TgGRA23 proteins, which share amino acid sequence similarity with NcGRA17 and NcGRA23, function to facilitate the movement of small molecules or nutrients across the PVM [24]. NcGRA23 lacks a signal peptide, a transmembrane domain, and sequence motifs that suggest a functional role. To verify the biological role of NcGRA23, we successfully constructed the NcGRA23 knockout strain (Fig. 4a). We isolated the NcGRA23 knockout clones. Several strains with a complete knockout of the NcGRA23 gene (∆ncgra23) were generated. PCR and western blotting confirmed the deletion of the NcGRA23 gene (Fig. 4b, c). The growth of the NcGRA23 gene knockout in the Nc1 strain was assayed by plaque formation on HFF monolayers as described above. The knockout mutant had no significant reductions in plaque formation (Fig. 4d, e). The knockout mutant was also examined for virulence in BALB/c mice. All of the mice inoculated with the wild-type or NcGRA23 gene knockouts succumbed to infection with $8 \times 10^6$ tachyzoites, with approximately equal time to death (Fig. 4f), and there was no significant difference in survival time between mice in the group infected with $5 \times 10^6$ tachyzoites, indicating that this protein does not affect parasite virulence in mice.

**Identification and infectivity potential of NcGRA11**

Examination of novel identified dense granule proteins by BLAST analysis revealed four other proteins

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**Table 1** Summary of known and novel NcGRAs found by NcGRA17-BioID

| Gene ID (NcLIV no. [ToxoDB release 37]) | Protein name |
|---------------------------------------|--------------|
| 006780                                | NcGRA23      |
| 057330                                | NcGRA61      |
| 045650                                | NcGRA32      |
| 005560                                | NcGRA17      |
| 043760                                | NcGRA27      |
| 045870                                | NcGRA3       |
| 012020                                | NcGRA11b     |
| 042680                                | NcGRA25      |
| 055980                                | NcGRA38      |
| 058760                                | NcGRA45      |

Novel NcGRAs are highlighted in boldface.

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**Fig. 3** The localization of the NcGRA23. **a** IFA results showing that HA-tagged NcGRA17 co-localizes with NcGRA23 in dense granules and PVM (intracellular, scale bar = 5 μm; extracellular, scale bar = 2 μm). **b** Localization analysis of the distribution of NcGRA17-HA and NcGRA23 following secretion into the PV at 5 min, 10 min, and 1 h post-invasion (scale bar = 2 μm).
not found in the NcGRA17-BioID data set with similarity to NcGRA11b (NCLIV_012010, NCLIV_069640, NCLIV_012210, and NCLIV_012220). To verify the biological role of the four genes, we constructed the overexpression strains (Fig. 5b). The IFA results showed that proteins appeared to overlap with NcGRA6. Therefore, NCLIV_012010, NCLIV_069640, NCLIV_012210, and NCLIV_012220 were named NcGRA11a, NcGRA11c, NcGRA11d, and NcGRA11e, respectively (Fig. 5c). To determine the secretion characteristic of NcGRA11, we also analyzed the secretion kinetics of the NcGRA11c at different time points (5 min to 1 h) after invasion (Fig. 5d). At the earliest time when secretion was detected (5 min post-invasion), NcGRA11c-HA and NcGRA6 were secreted into the vacuole as concentrated foci on either side of the anterior part of the parasite. Following secretion into the vacuole, NcGRA11c and NcGRA6 were rapidly localized to the posterior end of the parasite cell (15 min post-invasion). By 60 min after invasion, NcGRA11c and NcGRA6 were found throughout the vacuole, forming an accumulation surrounding the parasite cell and with a distinct concentration at one pole.
The patterns of secretion suggest that the exocytosed NcGRA11c and NcGRA6 may be rapidly translocated from the anterior to the posterior end to participate in network formation.

The five genes were then verified to be located together on chromosome 4 for 62,240 bp (Fig. 5a). The encoded proteins contain a predicted signal peptide and three transmembrane domains, and lack known functional domains. To generate clean knockouts of NcGRA11 (a-e) genes, a double-gRNA was used in targeting the 5′ and 3′ regions of the sequence to delete the entire sequence, followed by selection for insertion of the DHFR-TS selectable marker flanked by short, gene-specific homology regions (Fig. 6a), as described previously [17]. We were able to easily generate NcGRA11 (a–e) gene knockout strain (Δncgra11 (a–e)), as shown using diagnostic PCR and western blotting (Fig. 6b, c). We investigated the replication and virulence of Δncgra11 (a–e). The lack of NcGRA11 (a–e) did not affect parasite growth in vitro, as the plaque areas of Δncgra11 (a–e) were equivalent to Nc1 (Fig. 6d, e). We observed no significant difference in survival time between mice in the group infected with 5 × 10^6 or 8 × 10^6 tachyzoites of the different strains (Fig. 6f). The results above suggest that NcGRA11 (a–e) is not involved in parasite growth and replication.

**Discussion**

*Neospora* dense granules are necessary storage secretory organelles. Today, the full proteome of dense granules remains unknown, and only a small number of NcGRA proteins (~9) have been identified. The analysis of protein content of dense granules is an important aspect of biological function study of *Neospora*. Biochemical fractionation approaches were used to isolate *Toxoplasma* dense granules, but the excessive parasite and/or host contamination limited its application [25]. Recently, the BioID technique has been used to screen proteins interacting with or in proximity to bait proteins, with high screening efficiency [20, 22, 26, 27]. Therefore, in this study, we adapted the BioID technique for *N. caninum* and demonstrated that BirA* fusions can biotinylate a robust number of proteins in the NcGRA.

We chose the dense granule protein NcGRA17 as bait for the initial BirA* fusion, because NcGRA17 is highly expressed and can be fused to an endogenous HA tag and traffic correctly to the dense granules [14]. Because NcGRA17 localized at the PV, we hypothesized that
NcGRA17-BirA* should label proteins in the dense granules and PV. Indeed, our NcGRA17 BioID data set contains known and novel proteins from both sub-organelle compartments (see Table 1). The high success rate of our verifications by gene tagging supports specific labeling of NcGRAs secreted into the dense granules. We did identify eight NcGRA proteins within the PV and three NcGRA proteins associated with the PVM. The extent to which these are truly proteins that interact with NcGRA17 or merely proximal at the vacuolar remains to be determined.

Our study on the endogenous tagging and deletion of the NcGRAs coding sequences takes advantage of the improved efficiency of CRISPR/Cas9 for targeted gene editing in *Neospora* [14]. The target genes were rapidly tagged or deleted using a cassette approach where amplicons are generated from separate plasmids bearing distinct tags or resistance markers using a common set of primers containing short homology regions (42 or 60 bp).
for gene targeting in the Nc1 background, as described in Toxoplasma previously [17]. The smHA tagging plasmids and resistance markers encoding DHFR plasmids described here are easily adapted to other genes of interest by designing new primer pairs and modifying the CRISPR/Cas9 plasmid or CRISPR/Cas9 double-gRNA plasmid to contain different gRNAs, as described previously [28].

As a known Neospora dense granule protein, NcGRA23 was previously found to be localized at the PVM and shared sequence similarity with NcGRA17 [7, 14]. In agreement with this, NcGRA23 scored highly in the mass spectrometry data set (Table 1; Additional file 4: Table S2). In this study, NcGRA23 co-localized with NcGRA17 within the dense granules and at the PVM, and had same patterns of secretion with NcGRA17. To verify the biological role of NcGRA23, our attempts to disrupt the gene using CRISPR/Cas9 confirmed that NcGRA23 is nonessential for in vitro growth or in vivo virulence. Unfortunately, we were unable to generate a Δncgra23/Δncgra23 strain, which suggests that they may be essential for parasite growth. A tetracycline repressor-based system and plant-like auxin-induced degron (AID) system have been used in Toxoplasma essential gene study [20, 29]; these conditional gene knockout approaches may be used to explore the function of Neospora essential genes in the future.

While most of the dense granule proteins identified to date lack paralogs in Neospora, we identified a group of new NcGRA proteins (NcGRA11 a–e) that have shared sequence similarity. We then verified that the five genes were located together on chromosome 4 for 62,240 bp and had same patterns of secretion with NcGRA17. To verify the biological role of NcGRA23, our attempts to disrupt the gene using CRISPR/Cas9 confirmed that NcGRA23 is nonessential for in vitro growth or in vivo virulence. Unfortunately, we were unable to generate a Δncgra17/Δncgra23 strain, which suggests that they may be essential for parasite growth. A tetracycline repressor-based system and plant-like auxin-induced degron (AID) system have been used in Toxoplasma essential gene study [20, 29]; these conditional gene knockout approaches may be used to explore the function of Neospora essential genes in the future.

Conclusion
Our results highlight that biotinylation of proximal proteins using the BioID system is a powerful approach to rapidly identify novel NcGRA proteins. Using NcGRA17-BirA* fusion protein as bait, we have identified 11 novel GRA proteins. In addition, our findings indicate that NcGRA23 and NcGRA11 (a–e) are not involved in parasite propagation in vitro and virulence in vivo. The exact biological functions of novel identified GRA proteins need to be examined further in future studies.

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Authors’ contributions
Conceptualization, QL and CSY; methodology, JL, CRW, and CSY; software, CSY and CRW; validation, CSY and CRW; formal analysis, CSY and CRW; investigation, CSY; resources, CSY; data curation, CSY; writing—original draft preparation, CSY; writing—review and editing, QL, JL, CRW, and CSY; visualization, QL and JL; supervision, QL and JL; project administration, QL and JL; funding acquisition, QL, JL, and CSY. All authors read and approved the final manuscript.
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Availability of data and materials
All data analyzed or generated during this study are included in this published article.

Declarations

Ethics approval and consent to participate
This research was approved by the Beijing Laboratory Administration Committee based on the recommendations and the Guide for the Care and Use of Laboratory Animals of the Ministry of Science and Technology of China (Approval No.: 18049).

Consent for publication
Not applicable.

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

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