H11-induced immunoprotection is predominantly linked to N-glycan moieties during Haemonchus contortus infection

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Nematodes are one of the largest groups of animals on the planet. Many of them are major pathogens of humans, animals and plants, and cause destructive diseases and socioeconomic losses worldwide. Despite their adverse impacts on human health and agriculture, nematodes can be challenging to control, because anthelmintic treatments do not prevent re-infection, and excessive treatment has led to widespread drug resistance in nematode populations. Indeed, many nematode species of livestock animals have become resistant to almost all classes of anthelmintics used. Most efforts to develop commercial anti-nematode vaccines (native or recombinant) for use in animals and humans have not succeeded, although one effective (dead) vaccine (Barbervax) has been developed to protect animals against one of the most pathogenic parasites of livestock animals – Haemonchus contortus (the barber’s pole worm). This vaccine contains native molecules, called H11 and H-Gal-GP, derived from the intestine of this blood-feeding worm. In its native form, H11 alone consistently induces high levels (75-95%) of immunoprotection during H. contortus infection. Here, to test the hypothesis that post-translational modification (glycosylation) of H11 plays a crucial role in achieving such high immunoprotection, we explored the N-glycoproteome and N-glycome of H11 using the high-resolution mass spectrometry and assessed the roles of N-glycosylation in protective immunity against H. contortus. Our results showed conclusively that N-glycan moieties on H11 are the dominant immunogens,
which induce high IgG serum antibody levels in immunised animals, and that anti-H11 IgG antibodies can confer specific, passive immunity in naïve animals. This work provides the first detailed account of the relevance and role of protein glycosylation in protective immunity against a parasitic nematode, with important implications for the design of vaccines against metazoan parasites.

**KEYWORDS**
parasitic nematode, vaccine, H11 antigen, N-glycosylation, N-glycan, IgG antibodies, immunoprotection

**Introduction**

Roundworms (nematodes) represent one of the largest groups of animals (phylum Nematoda). While most nematodes are free-living, many of them are major pathogens of plants and animals worldwide (1). Nematode diseases (nematodiases) of plants cause major production losses, equating to ~ US$ 80 billion per annum, and the adverse economic impact of nematodiases on agricultural animals is estimated at ~ US$ 30 billion worldwide. Humans are also affected by a range of nematodes, such as soil-transmitted helminths (STHs), filarioids and food-borne worms, which cause neglected diseases, imposing a global burden of ~10 million disability-adjusted life years (DALYs) per annum (2, 3). In spite of their very substantial, often chronic impact, nematodes can be very challenging to control, because anthelmintic treatments do not prevent re-infection, and repeated treatments lead to genetic resistance in nematode populations (4, 5). Indeed, many nematode species of livestock animals have become resistant to almost all of the main classes of anthelmintics (or pesticides) sold commercially.

To circumvent the anthelmintic resistance problem, there have been major efforts to develop anti-nematode vaccines (native or recombinant) for use in animals and humans (6, 7). However, the majority of these efforts have not been successful due to limited insight into the protective epitopes and underlying immune response. Presently, only one effective (dead) anti-nematode vaccine (called Barbervax) has been developed and commercialised that protects livestock animals against the disease (= haemonchosis) caused by *Haemonchus contortus* (the barber’s pole worm). This vaccine comprises native molecules and molecular complexes, including H11 (8) and H-Gal-GP (9), derived from the intestinal tract of this worm. Although native H11 consistently achieves high levels (75-95%) of protection (8, 10, 11), the substantial efforts attempting to produce recombinant forms of some of these molecules, particularly H11, in an academic or commercial context were consistently unsuccessful (12, 13). Also attempts to develop a well-defined, recombinant vaccine molecules, employing a range of bacterial (14), yeast (15), insect cell (16) and *Caenorhabditis elegans* expression systems (17, 18), have not been successful.

Although H11 has remained a prime vaccine molecule candidate for more than three decades, there has been no detailed investigation of the reason(s) why recombinant proteins expressed in prokaryotic and eukaryotic organisms have failed to protect at levels achieved using native H11. Here, we hypothesise that post-translational modification(s) play a crucial role in immunoprotection against *H. contortus* – which represents a powerful model system for parasitic nematodes more generally (19). A critical appraisal of current literature reveals evidence that glycosylation can associate with immunogenicity and/or antigenicity in hosts infected with particular nematodes (e.g., *Dictyocaulus*) (20), but detailed information is very scant. In other host-pathogen systems, N-glycosylation has been associated with the immunogenicity of proteins and immune modulation (21–23), thereby elevating the relevance of using glycomic and glycoproteomic tools to discover the role of carbohydrate antigens in inducing immune responses (24–27). Despite these developments, no detailed information is available on parasitic nematodes.

Here, we explore the N-glycoproteome and N-glycome of H11 using high-resolution mass spectrometry and assess the roles of N-glycosylation in protective immunity against *H. contortus*. This work provides the first detailed insight into the relevance and role of protein glycosylation in protective immunity against a parasitic nematode. It should have important implications for developing vaccines against metazoan parasites and understanding host immune responses to these parasites.

**Materials and methods**

**Parasite materials**

Different developmental stages of *H. contortus* were maintained, collected, processed and stored using well-established protocols (28, 29). Specifically, infective third-stage
lacte (iL3s, Haecon-5 strain) were obtained from coproculture (28), isolated and maintained at 15°C. Exsheathed L3s (xL3s) were produced by incubation with 0.15% (v/v) sodium hypochlorite at 37°C for 10 min (29). Adult H. contortus were obtained from the abomasum of infected goats.

**H11 antigen**

Native H11 was isolated from adult *H. contortus* using concanavalin A–sepharose (GE Healthcare), as previously described (8). Briefly, 15 g of the worms were homogenized in ice-cold phosphate-buffered saline (PBS, pH 7.4) for 25 min using a glass homogenizer. The homogenate was centrifuged (12,000 g for 25 min) and the pellet was extracted four times with 1% (v/v) Triton X-100. The resultant solution was filtered (0.22 µM). H11 was isolated using concanavalin A–sepharose columns. The column-bound H11 was washed 3 times (20 mM Tris-HCl), followed by elution using a buffer containing 200 mM of methyl-D-mannopyranoside and methyl-D-glucopyranoside. The resultant solution was enriched and filtered (0.22 µM). This final filtrate was designated the native H11 antigen. Protein concentration was estimated using a bicinchoninic acid assay (BCA) kit (Beyotime). Protein ratio of 1:50. After tryptic digestion, the hydrophilic protein sample was digested with trypsin at an enzyme-to-protein ratio of 1:50. Total protein (400 µg) was reduced with 5 mM dithiothreitol and then alkylated with 11 mM iodoacetamide. The reduced protein sample was digested with trypsin in an enzyme-to-protein ratio of 1:50. After tryptic digestion, the hydrophilic interaction chromatography (HILIC) enrichment of N-glycopeptides was carried out as reported previously (30). Glycopeptides were subjected to de-glycosylation by PNGase F and PNGase A (31). Brieﬂy, 2 µL of PNGase F in 50 µL of 50 mM NH₄HCO₃ buffer was incubated at 37°C overnight. The remaining peptides were subjected to HILIC enrichment and reconstituted in 2 µL of PNGase A in 50 µL of 50 mM sodium acetate solution overnight at 37°C. Then PNGase F- and A-deglycosylated peptides were acidified with formic acid, puriﬁed using an EASY-nLC 1000 UPLC system and subjected to liquid chromatography-tandem mass spectrometry (LC-MS/MS) analysis using an Orbitrap Fusion™-TrapID™ instrument (Thermo Fisher Scientific, Bremen, Germany).

MS data were analysed using the software package MaxQuant to identify peptides, followed by searches against the UniProtKB *H. contortus* database (24,277 proteins), concatenated using the reverse decay database. Glycosylation sites were inferred using the consensus of two biological replicates. Amino acid sequence motifs comprising at least 20 residues (+/-10) of the N-glycosylation sites were analysed using the MoMo program (https://meme-suite.org/meme/tools/momo). Functional annotation relied on the InterPro domain database (http://www.ebi.ac.uk/interpro/).

**N-glycoproteome analysis**

Total protein (400 µg) was reduced with 5 mM dithiothreitol and then alkylated with 11 mM iodoacetamide. The reduced protein sample was digested with trypsin in an enzyme-to-protein ratio of 1:50. After tryptic digestion, the hydrophilic interaction chromatography (HILIC) enrichment of N-glycopeptides was carried out as reported previously (30). Glycopeptides were subjected to de-glycosylation by PNGase F and PNGase A (31). Brieﬂy, 2 µL of PNGase F in 50 µL of 50 mM NH₄HCO₃ buffer was incubated at 37°C overnight. The remaining peptides were subjected to HILIC enrichment and reconstituted in 2 µL of PNGase A in 50 µL of 50 mM sodium acetate solution overnight at 37°C. Then PNGase F- and A-deglycosylated peptides were acidified with formic acid, puriﬁed using an EASY-nLC 1000 UPLC system and subjected to liquid chromatography-tandem mass spectrometry (LC-MS/MS) analysis using an Orbitrap Fusion™-TrapID™ instrument (Thermo Fisher Scientific, Bremen, Germany).

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**N-glycome analysis**

This analysis of H11 was conducted using an established protocol (31). N-glycans were released from glycopeptides by PNGase F. PNGase F-resistant N-glycans were subjected to PNGase A digestion. Following porous graphitic carbon (PGC) puriﬁcation (32), native glycans were permethylated by adding 100 µL of DMSO-NaOH slurry. Subsequently, the glycan sample was mixed with freshly-prepared 2,5-dihydroxybenzoic acid (DHB) before crystallisation at room temperature. N-glycome spectra were obtained in the positive ionisation mode using an AB 5800 MALDI-TOF-TOF instrument (SCIEX, Concord, Canada). The data were processed using Data Explorer 4.0 (SCIEX) and GlycoWorkbench (v.2.1).

**Treatment of H11 antigen**

For experimentation, H11 was used in a native (NA) form; denatured (DN) at 85°C for 25 min (33); periodate (PI)-treated (34), or digested with PNGase F and PNGase A (F+A) (see N-glycoproteome analysis) to remove N-glycans (31). Glycan moieties of the H11 antigen were destroyed using an established periodate oxidation procedure (34). Native H11 (1 mg/mL) was incubated in 10 mM sodium periodate (pH 5.0, 45 min at 24°C in the dark) to disrupt glycans without affecting protein (35), and then reduced by 50 mM sodium borohydride for 25 min. Following protein quantitation using the BCA method, all samples were stored at -80°C until use. The H11 samples (NA, DN, PI and F+A) were quantiﬁed and concentrations adjusted to 4 µg/mL; the effectiveness of periodate treatment (oxidation) was assessed by direct ELISA (36) and immunoblot (34) using horse radish peroxidase (HRP)- conjugated concanavalin A (EY Laboratories Inc., San Mateo, USA).

**Immunisation trial**

To test whether native, denatured- or periodate-treated H11 proteins could induce protective immunity against *H. contortus*, immunisation and infection experiments were conducted in goats (raised helminth-free). All animal experimentation was approved by the Animals Ethics Committee of Huazhong Agricultural University (permit HZAUGO-2019-002). Four-month-old Boer goats, procured from the Hubei Academy of Agricultural Sciences (Wuhan, China), were housed indoors under parasite-free conditions, and provided with fodder twice.
a day with water ad libitum. Four groups were formed with five goats in each group (age- and weight-matched). Goats in group AJ were injected with a saponin-based adjuvant (Quil-A®, In vivoGen, San Diego, California, USA; negative control); those in groups NA, DN and PI with native H11 (positive control), denatured H11 or periodate-treated H11 (Figure 4A); one animal in group PI died of an unrelated cause on day 56 of the immunisation/challenge experiment. Each goat was immunised subcutaneously thrice at three-week intervals with 150 µg/mL protein equivalent in Quil-A® (37). On the same day after the third immunisation (day 42), all goats were challenged with 7000 infective, third-stage larvae (L3s); on this day, none of the goats excreted nematode eggs in faeces. During this experiment, blood was taken at eight time points (days 0, 10, 21, 31, 42, 49, 63 and 77). Faecal samples were collected rectally from individual goats three times each week, commencing two weeks after the challenge infection (day 56) until the end of the experiment (day 80) (Figure 4A). On day 80, goats were euthanised, and adult worms in individual abomasas were enumerated (Figure 4A). Tissues were prepared for haematoxylin and eosin (HE) staining and histopathological examination.

Serum antibody ELISA

Specific anti-H11 IgG, IgM and IgA serum antibody levels (titers) were measured by ELISA (38). In brief, microtitration plates were coated with native H11 (4 µg/mL), blocked with phosphate-buffered saline (PBS; pH 7.4) containing 1% (w/v) bovine serum albumin (PBS-BSA) and washed with the same solution. Individual goat sera were tested at 1/200 dilution, incubated with HRP-conjugated donkey anti-goat IgG (1/500), rabbit anti-goat IgM (1/10,000) or rabbit anti-goat IgA (1/10,000) in PBS-BSA. Plates were washed and incubated with tetramethylbenzidine (for 15 min), and plates were read at 630 nm.

IgG purification and immunoblot

Serum samples from the immunisation trial were examined for IgG antibody responses in the goat groups AJ, NA, DN and PI. First, the serum samples collected from each group on day 42 were pooled (10 mL each). Second, IgG antibodies were purified from each of the four pooled serum samples using Protein A+G agarose (Beyotime Biotechnology, Shanghai, China), according to the manufacturer’s protocol (eluting into 5 mL). Third, these antibody samples were each diluted 1/400 and used to probe immunoblot strips (polyvinylidene fluoride, PVDF) each containing 10 µg of sodium dodecyl sulfate (SDS)-denatured H11, separated in a gradient (4-12%) SDS-PAGE gel (blocked with PBS-BSA) (17); each strip was washed in PBS-BSA and then incubated in HRP-conjugated donkey anti-goat IgG (1/1000), developed for 90 min and scanned on a chemiluminescence imaging system (Tanon 4600, Shanghai, China) and processed using ImageJ software (NIH, USA).

Passive transfer of IgG antibodies

To evaluate whether IgG antibodies (from goats actively immunised with H11; i.e. group NA) passively transferred to naive goats (maintained under parasite-free conditions) could achieve protection against H. contortus infection, we used the same experimental design as for the immunisation experiment (i.e. assigned four goats to group AJ, and five to each of the groups NA, DN and PI). The amount of IgG antibodies transferred (dose) was as reported in the literature (39). Each goat received IgG antibodies (5 mg) in 50 mL physiological saline intravenously on days 0 and 4, and was infected with 7000 L3s on day 1 of the experiment. Faecal samples were collected six times from day 25 after the challenge to day 35 so that H. contortus eggs per gram of faeces could be enumerated. On day 35, all goats were euthanised, and the numbers of worms in individual abomasas were counted (Figure 6A).

Lectin histochemistry and immunohistochemistry

To investigate whether purified IgG antibodies specifically bound native H11 in the gut of H. contortus, lectin histochemistry and immunohistochemistry were performed by fluorescent localisation analysis (40). The adult female worms were sliced (4 µm thick), mounted on the slides, then incubated with fluorescein-labelled concanavalin A (Vector Laboratories) and stained with DAPI to detect cell nuclei. For co-localisation, slides were incubated first with fluorescein-labelled concanavalin A and then with IgG antibodies representing the goat-groups AJ, NA, DN or PI, respectively. After extensive washing, the samples were incubated with Cy3-conjugated donkey anti-goat IgG (Beyotime Biotechnology) and then 4′,6-diamidino-2-phenylindole (DAPI). The slides were mounted with an anti-fading solution for fluorescence microscopy (Olympus, Tokyo, Japan).

IgG antibody inhibition assays

To assess whether IgG antibodies from goats immunised with native or treated H11 could inhibit aminopeptidase activity, the intestinal tracts from 20 fresh, live adult H. contortus were collected (41), homogenized in 200 µL of PBS. The homogenate (10 µg protein per well) (42) was mixed with 0.2 mM L-leucine-p-nitroanilide (L-Leu-pNA) at pH 3.0 to 8.0 to assess the optimal
pH of aminopeptidase. To investigate the inhibition effect of respective IgG antibodies at an optimal pH, protein samples were pre-incubated with 5 μL of IgG antibodies (1 mg/mL) purified from the normal pre-immunisation sera or sera from goats in groups A1, NA, DN or PI (see IgG purification and immunoblot) for 15 min, and then incubated with L-Leu-pNA substrate for 2.5 h. The inhibitor sensitivity assay (positive control) was carried out by pre-incubation with 10 μM bestatin before adding substrate (17). The absorbance was determined at 405 nm using a multi-mode plate reader (BioTek Cytation 5, Winooski, Vermont, USA).

To assess whether IgG antibodies could inhibit larval development, xL3s were cultured in 24-well plates (100 xL3s/well) in 200 μL of sterile Luria Bertani (LB) medium supplemented with 100 IU/mL of penicillin, 100 μg/mL of streptomycin and 0.25 μg/mL of amphotericin (Antibiotic-Antimycotic, Sigma-Aldrich, St. Louis, Missouri, USA) (43) and exposed to IgG antibodies (1 μg/mL; 50 μL aliquots) from each group (normal, A1, NA, DN or PI; in triplicate), and incubated at 39°C and 20% CO₂. The developmental rate was assessed on days 3 and 4 by examining the presence of a buccal capsule, characteristic of the L4 stage (44). The length and width of individual L4s were measured on day 7.

**Statistical analyses**

All statistical analyses were conducted using Premier 8.0 software (GraphPad, La Jolla, USA), and standard deviation (SD) or standard error of mean (SEM) calculated. The statistical differences of worm burdens and cumulative FEC were compared by one-way analysis of variance (ANOVA) followed by Dunnett test. The *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001 and ns (nonsignificant) indicate the degree of significance.

**Results**

**N-glycoproteome, and glycan profiles and moieties**

A glycoproteomic assessment of native H11, corresponding to bands of 100-130, 40-55 and 30-35 kDa in SDS-PAGE analysis (Figure 1A), identified 85 distinct proteins with 125 N-glycosylated sites (Figures 1B, C; Supplementary Table 1). Two conserved motifs N-x-T and N-x-S were associated with glycosylated asparagine residues (−10 to +10), where x represents all amino acids, except proline (Figure 1D). Domain enrichment analysis revealed cysteine protease and peptidase M1 aminopeptidase as the two most abundant glycoproteins (Figure 1E; Supplementary Table 2).

MalDI-TOF-MS analysis of H11 identified 38 major signal peaks for permethylated, native glycanics. PNGase F-released glycans (n = 38) corresponded to the pauci-mannosidic structures with or without fucose (Hex₃,HexNAc₂,Fuc₀₋₂), high-mannose moieties (Hex₅₋₁,HexNAc₂) and some complex glycans (Hex₃₋₂,HexNAc₂,Fuc₀₋₃) harbouring the major core α1,6 fucose and antenna motifs, including the putative GalNAc-GlcNAc (LDN), fucosylated LDN (LDNF) and/or galactosylated LDNF (Figure 2A; Supplementary Table 3). PNGase A-released glycans (n = 9) represented structures containing core α1,3 fucose residues (Figure 2B; Supplementary Table 4). The remainder of N-glycans (n = 9) had incompletely removed moieties from glycopeptides following PNGase F treatment. Most N-glycan peaks were consistent with those published previously (31), except for novel signals at m/z 2081.5, 2125.6, 2224.6, 2388.7, 2418.7, 2459.8, 2500.8, 2633.9, 2663.9, 2674.9, 2705.0, 2879.1 and 3083.2, following PNGase F-treatment (Supplementary Table 3). Extending this work, we showed that native and denatured H11 bound to concanavalin A (a lectin that binds to mannose on glycoproteins). In contrast, sodium periodate-treated or PNGase (F+A)-treated H11 did not, demonstrating a loss of N-glycan moieties in H11 following each of these treatments (Figures 3A, B).

**N-glycan moieties in H11 induce substantial protective immunity and a specific IgG antibody response**

Given that vaccination with native H11 achieves high levels of protection in sheep (45), we assessed whether this protection is linked to protein and/or glycan moieties. To do this, we injected goats subcutaneously three times at three-weekly intervals with either adjuvant, native, denatured or periodate-treated H11 and challenged them (on the same day after the last injection) with 7000 infective third-stage larvae (L3s) of H. contortus (Figure 4A). Goats immunised separately with native and denatured H11 exhibited a significant decrease (> 93% and 87%, respectively) in the numbers of H. contortus eggs in faeces (Figure 4B; Table 1), and harboured 86% and 80% fewer worms at necropsy on day 38 (following challenge infection) compared with goats that received periodate-treated H11 (37% and 31%, respectively) (Figure 4C; Table 1); in the latter group of goats, pathological alterations in the abomasum were more pronounced (Figure 4D).

We also measured serum antibody levels, and showed that goats immunised with either native or denatured H11 had significantly (p < 0.0001) high-specific anti-H11 IgG levels (days 42, 49, 63 and 77) compared with control goats administered either periodate-treated H11 or adjuvant alone (Figure 5A). Specific anti-H11 IgM levels were detected on day 10 and decreased gradually from day 31 onward, whereas anti-
H11 IgA was not detectable in the sera from any of the four animal groups (Figures 5B, C). Immunoblot analysis revealed that bands of 110-130 kDa of high intensity were recognised by IgG serum antibodies from goats immunised with native or denatured H11 compared with controls (Figures 5D, E). Overall, these findings provided evidence that the specific IgG antibody response recorded in immunised animals relates to glycan epitopes of H11.

**FIGURE 1**
N-glycoproteome analysis of *Haemonchus contortus* H11. (A) SDS-PAGE analysis of concanavalin A lectin-purified *H. contortus* native H11 (10 µg). (B) N-glycoproteome identified in native H11. H11 sample was sequentially digested with PNGase F and PNGase A, and their N-glycoproteins and N-glycosylation sites were identified by liquid chromatography-tandem mass spectrometry (LC-MS/MS). Data are derived from two independent experiments (cf. Supplementary Table 1). (C) Comparison between the PNGase F- and PNGase A-released N-glycosylation sites and total numbers of N-glycoproteins and N-glycosylation sites are shown. (D) Conserved sequence motifs analyses around glycosylated asparagine residues (−10 to +10; N, asparagine; T, threonine; S, serine). (E) Domain enrichment analysis of the identified N-glycoproteins (p < 0.05). Each domain is bubble-coded; red and blue represent the enrichment and depletion (Fisher’s exact test p value); size reflects the number of glycoproteins (cf. Supplementary Table 2).
Passive transfer of protective immunity

Having shown that immunoprotection relates predominantly to glycan moieties, we explored whether a passive transfer of immunity could be achieved. To do this, we transferred IgG antibodies purified from sera from goats with known levels of protection (31-93%) against *H. contortus* (Figure 4; Table 1) to naïve goats (raised under parasite-free conditions) (Figure 6A). We showed that naïve goats that received IgG antibodies from goats immunised with either native or denatured H11 exhibited significantly \( p \leq 0.0001 \) lower *H. contortus* egg numbers in their faeces (Figure 6B) and a very \( p \leq 0.0006 \) lower infection intensity (Figure 6C) compared with controls receiving IgG antibodies from goats injected with either periodate-treated H11 or adjuvant alone. These results indicated that IgG serum antibodies induced by native H11 antigen could confer passive immunity to naïve goats.
IgG serum antibodies from immunoprotected goats bind glycoconjugates in the microvilli of the *H. contortus* gut, and inhibit aminopeptidase activity and worm development

Extending previous findings (10, 11), we specifically identified glycoconjugates in the intestine of *H. contortus* adults with fluorescein-labelled concanavalin A. Fluorescence was pronounced in the cytoplasm and microvilli throughout the entire intestine (Figures 7A, B). We also showed that IgG antibodies from the serum from goats immunised with native or denatured H11 colocalized with these glycoconjugates, with significantly higher binding intensity compared with antibodies from the periodate-treated and adjuvant controls (Figure 7C). For adult worms, we also demonstrated that intestinal aminopeptidase activity could be markedly inhibited by the same IgG antibodies from immunoprotected goats compared with respective controls (Figures 8A, B), and that inhibition levels were correlated with the reductions in *H. contortus* egg numbers in faeces and infection intensity (R² of 0.98 and 0.96 respectively; Figures 8C, D).

Subsequently, we assessed whether IgG serum antibodies from immunoprotected goats inhibited the development of *H. contortus in vitro*. We showed that IgG antibodies from goats immunised with native or denatured H11 inhibited the development of exsheathed third-stage larvae (xL3s) to fourth-stage larvae (L4s) (Figure 8E), compared with IgG antibodies from control goats either immunised with periodate-treated H11 or injected with adjuvant alone (Figure 8F). L4s incubated with IgG antibodies from immunoprotected goats were significantly smaller than those incubated with antibodies from these respective control animals (Figures 8G, H).

**Discussion**

H11 is the best-studied immunogen (vaccine molecule) from a parasitic nematode (12, 46). In its native form, H11 achieves high levels (usually 75-95%) of protection in animals (8). In the late 1980s, the advent of recombinant DNA technology stimulated researchers in the UK and Australia to focus on developing a well-defined recombinant, subunit vaccine representing H11, in order to be prepared to meet the supply, quality and demand for a commercial vaccine. Despite major efforts in academic-industry collaborations, the select H11 proteins expressed in prokaryotic and eukaryotic expression systems (14, 16, 17) did not achieve significant immuno-protection in animals (13, 47, 48). After more than three decades of research on H11, this result raised pertinent questions about precisely which component(s) of native H11 confer(s) reproducible immunoprotection.

Here, we explored whether particular post-translation modifications are responsible for the immuno-protection induced by H11. Using advanced experimental and molecular tools, we elucidated the N-glycoproteome and N-glycome of H11, and demonstrated that N-glycans are a predominant component that induces protection and an associated IgG serum antibody
response in immunised animals (i.e. goats), and that this antibody response leads to a marked inhibition of aminopeptidase activity in the nematode’s intestine as well as worm development. We also showed that the H11 glycoproteome contains at least 85 proteins with 125 N-glycosylated sites, contrasting prior work identifying 23 proteins (49). The high analytical sensitivity of the methodology (including HILIC enrichment) employed here allowed us to discover at least six high-confidence aminopeptidases with 13 N-glycosylation sites relating proteins of 100-130 kDa, some of which are consistent with isoforms found in H. contortus adults (upper panel) and hematoxylin and eosin(HE) staining (lower panel) in individual abomasas of goats from four groups (AJ, NA, DN and PI). Evident pathological alterations in the abomasum show atrophy and cell necrosis of glands, enhanced eosinophilia and lymphocyte infiltration. Scale bars: 100 µm.

FIGURE 4
N-glycan moieties on H11 induce prominent protective immunity in goats against Haemonchus contortus infection. (A) Design of the animal trial. Four groups of five goats each were injected with Quil A® adjuvant alone (AJ), native H11 (NA), denatured H11 (DN) or periodate-treated H11 (PI), respectively, each with 150 µg proteins equivalent in Quil-A®. Each goat was immunised subcutaneously three times at three-week intervals (days 0, 21 and 42; red arrow) and then challenged with 7000 infective third-stage larvae (L3s) on day 42 (blue arrow). One goat in group PI died of an unrelated cause. The time points for serum collection (eight times), IgG isolation (twice), faecal sample collection (eight times), worm sample collection (once) and abomasum collection (once) are indicated with black diamond. The numbers of eggs per gram faecal sample (faecal egg count, FEC) were counted under microscopy and normalised to the numbers of adult worms. (B) Mean group FEC (mean ± SD; AJ, n = 5; NA, n = 5; DN, n = 5 and PI, n = 4) was monitored at eight time points (days 14, 18, 21, 25, 28, 31, 35 and 38 after challenge infection). (C) Mean group worm burden (mean ± SD; AJ, n = 5; NA, n = 5; DN, n = 5 and PI n = 4) was counted at the end of the experiment (day 80). Statistical significance was determined using one-way ANOVA, ***p < 0.001. (D) Representative images showing the H. contortus adults (upper panel) and hematoxylin and eosin(HE) staining (lower panel) in individual abomasas of goats from four groups (AJ, NA, DN and PI). Evident pathological alterations in the abomasum show atrophy and cell necrosis of glands, enhanced eosinophilia and lymphocyte infiltration. Scale bars: 100 µm.
N-glycan moieties on H11 induce a highly-specific anti-H11 IgG antibody response in goats immunised with H11 antigens of Haemonchus contortus. Specific anti-H11 subtype antibody (IgG, IgM and IgA) responses in four groups of goats immunised with adjuvant (AJ), native H11 (NA), denatured H11 (DN) and periodate-treated H11 (PI), were determined by enzyme-linked immunosorbent assay (ELISA) and immunoblot analysis (for IgG only). (A–C) Dynamics of anti-H11 IgG (A), IgM (B) and IgA (C) serum antibody responses in four groups of goats (AJ, NA, DN and PI) at eight time points (days 0, 10, 21, 42, 49, 63 and 77, cf. Figure 4A). Each data point (absorbance at 630 nm) denoted the mean group antibody titers (mean ± SD; AJ, n=5; NA, n=5; DN, n=5 and PI, n=4). The black triangles indicated three time points of goat immunisation (days 0, 21, and 42, cf. Figure 4A) and asterisks represented statistically significant differences compared to Group PI (***p < 0.01, ****p < 0.0001). (D) Immunoblot of IgG antibodies binding to native H11. Four IgG antibodies were purified from pooled sera samples (collected on day 42 of the immunisation trial, cf. Figure 4A) from individual animal groups (AJ, NA, DN or PI). (E) Densities of four immunoblot bands were quantitatively analysed using the ImageJ program (NIH, USA). Data represent the mean ± SEM from three independent analyses. Statistical significance was determined using one-way ANOVA, ***p < 0.001, ****p < 0.0001.

**TABLE 1** Results from the immunisation trial in goats. Four groups of five goats each were injected with adjuvant alone (AJ), native H11 (NA), denatured H11 (DA) or periodate-treated H11 (PI), respectively and then challenged with 7000 third-stage larvae (L3s) of *H. contortus* (day 42, cf. Figure 4A). Faecal samples were taken from individual goats at eight time points (cf. Figure 4A), and the number of *H. contortus* eggs per gram was counted (faecal egg count, FEC). Cumulative FEC of individual goats and mean cumulative FEC (with standard deviations, SD) were calculated for each group. At the end of the experiment (day 80), the numbers of worms in the stomachs (abomasum) were counted, and the reduction in the intensity of infection was calculated for each group.

| Group | Faecal egg count (FEC) | Intensity of infection |
|-------|------------------------|------------------------|
|       | Cumulative FEC | Mean FEC | SD | Reductiona (%) | Number of worms | Mean | SD | Reductionb (%) |
| AJ    | 22700 | 15720.0 | 5900.7 | Not | applicable | 1355 | 1260.2 | 295.3 | Not |
|       | 9000 | | | applicable | 939 | applicable |
|       | 13150 | | | | 1550 | |
|       | 13350 | | | | 954 | |
|       | 20400 | | | | 1503 | |
| NA    | 700 | 1080.0 | 555.5 | 93.1*** | 178 | 177.0 | 54.5 | 86.0*** |
|       | 1350 | | | | 201 | |
|       | 400 | | | | 94 | |
|       | 950 | | | | 169 | |
|       | 2000 | | | | 243 | |

(Continued)
Specific anti-H11 IgG antibodies confer the protection in passive immunisation trials against *Haemonchus contortus* infection. (A) Experiment design of passive animal immunisation. Each of four groups of goats (AJ, n = 4; NA, n = 5; DN, n = 5 and PI, n = 5) received intravenously twice (days 0 and 4, red arrow) IgG antibodies (5 mg) purified from pooled sera samples (collected on day 42 of the immunisation trial, cf. Figure 4A) from individual animal groups, and infected with 7000 infective third-stage larvae (L3s) on day 1 of the experiment (blue arrow). The numbers of eggs per gram faecal sample (faecal egg count, FEC) were microscopically counted at six time points (days 25, 27, 29, 31, 33 and 35, black diamond) and worm burden was counted at the end of the experiment (day 35, black diamond). (B) Mean group FEC (mean ± SD; AJ, n = 4; NA, n = 5; DN, n = 5 and PI, n = 5) was calculated by the cumulative FEC of each goat. (C) Mean group worm burden (mean ± SD; AJ, n = 4; NA, n = 5; DN, n = 5 and PI, n = 5) was counted by the numbers of adult worms of each goat. Statistical significance was determined using one-way ANOVA and indicated by asterisks, ***p < 0.001, ****p < 0.0001 and ns (not significant).
previously (14, 50) (cf. Supplementary Table 5). The aminopeptidases represent type II integral membrane proteins with aminopeptidase A and M activities, which are located in the intestinal microvilli of *H. contortus* (10). Aminopeptidase activities have been reported in the blood-feeding parasites, indicating an association with the degradation of haemoglobin peptides into free amino acids (51). Notably, our findings revealed that multiple N-glycosylation sites in aminopeptidases have α1,3-fucose in asparagine-linked GlcNAc residues, known to be highly antigenic in other helminth-host systems (52, 53).

![FIGURE 7](image)

Anti-H11 IgG antibodies from immunoprotected goats bind glycoconjugates in the microvilli of *Haemonchus contortus* gut. (A, B) Lectin histochemistry analysis to determine the localisation of native H11 in adult *H. contortus*. Paraffin-embedded slides of adult worm (A) and isolated intestinal tissue (B) were probed with fluorescein-labelled concanavalin A (Con A, green), and then stained with 4,6-diamidino-2-phenylindole (DAPI, blue) to detect cell nuclei. Panel 1, DAPI; Panel 2, Con A; Panel 3, merged signals from DAPI and Con A; Panel 4, merged image magnified from panel 3. (C) Lectin histochemistry and immunohistochemistry to determine the co-localisation of Con A and IgG antibodies bound native H11 in adult *H. contortus*. Slides of adult worms were incubated with fluorescein-labelled Con A (green) and then with four IgG antibodies purified from pooled sera samples (collected on day 42 of the immunisation trial, cf. Figure 4A) from individual animal groups (AJ, NA, DN or PI). Washing samples were incubated with Cy3-conjugated donkey anti-goat IgG (red) and then stained with DAPI (blue). Panel 1, DAPI; Panel 2, Con A; Panel 3, individual IgG antibodies; Panel 4, merged signals from DAPI, Con A and individual IgG antibodies. Scale bars: A = 50 µm or 200 µm; B = 50 µm; C = 200 µm. (ic = intestinal cytoplasm; mv = microvilli). The images shown here were derived from two independent experiments.
Anti-H11 IgG antibodies from immunprotected goats inhibit intestinal aminopeptidase activity and Haemonchus contortus development. IgG antibody inhibition assays were assessed by inhibiting intestinal aminopeptidase activity and H. contortus development in vitro. Five IgG antibodies: one was purified from the normal pre-immunisation sera; four were purified from pooled sera samples (collected on day 42 of the immunisation trial, cf. Figure 4A) from individual animal groups (AJ, NA, DN or PI). (A) Determination of intestinal aminopeptidase activity at pH 3.0 to 8.0 to assess the optimal pH of aminopeptidase. Intestinal homogenate from adult H. contortus was incubated with 0.2 mM L-leucine-p-nitroanilide (L-Leu-pNA) substrate. Enzyme activity was detected at 405 nm and expressed as nanomole of pNA/min/µg. (B) Inhibition assays of intestinal aminopeptidase activity were performed in the optimal pH by pre-incubation with each group of IgG antibodies (1 µg/µL; 5 mL aliquots) and followed by incubation with L-Leu-pNA substrate. The inhibitor sensitivity assay was carried out by incubation with 10 µM bestatin (Bes). (C, D) Correlations of aminopeptidase activity inhibition by each groups of IgG antibodies (AJ, NA, DN or PI) with mean faecal egg count (FEC) and mean intensity of infection (D) in passive immunisation trials. (E, F) Exsheathed L3s were cultured in 200 mL of sterile Luria Bertani (LB) medium and then five groups of IgG antibodies (1 µg/µL; 50 mL aliquots) were added to the culture media, respectively. Developmental rate (%) of L4 (n = 50) was assessed on day 3 and day 4 (F) by examining the presence of a buccal capsule (E; arrowhead; Scale bars = 20 µm). (G, H) On day 7 of the co-culture with indicated five groups of IgG antibodies, body length (G) and width (H) of L4 (n = 10) were measured (µm). Experiments were independently repeated three times in (A, B, F, G, H). Data show as mean ± SEM in (A, B, F) or mean ± SD in (G, H), and analyse by one-way ANOVA, *p < 0.05, **p < 0.01, ***p < 0.001 and ns (not significant).
Advanced MALDI-TOF-MS analysis, combined with PNGase F/A digestion of N-glycans, allowed us to infer at least 38 N-glycan structures within H11, which correlates with N-glycans in previous studies. The detection of 13 additional high-mass glycans relates to the use of concanavalin A for enrichment before analysis (31, 54, 55). The MS/MS spectra of m/z 2674.9, 2879.1 and 3083.2 inferred the presence of two LDNF antennae and two galactosylated LDNF antennae (56) (Supplementary Figures 1A–C), which have been reported previously to be associated with anti-parasite responses (57). Interestingly, the proportion (67%) of fucose-containing glycans in the H11 N-glycome is high, harbouring α1,3- and/or α1,6-linked monofucosylated, difucosylated, and uncommon trifucosylated glycan structures. The anti-glycan antibody response induced by recombinant *C. elegans* H11, but the α1,3- and/or α1,6-linked difucosylated and trifucosylated N-glycans did not achieve a protective effect (17). This may be explained by the distinctiveness in the N-glycans compared with native H11. Alternatively, additional glycan moieties present in native H11-enriched extract (compared with recombinant *C. elegans* H11) may induce a protective antibody response. In immunobiological studies of the blood fluke *Schistosoma mansoni* (class Trematoda), core α1,3- and α1,6-fucose di-fucosylation and LDNFs have been detected on the egg-derived antigen omega-1 (58). This fucose-linked epitope is reported to play a critical role in stimulating a Th2 phenotype by mediating the internalisation of omega-1 via a mannose receptor on dendritic cells (59). Similar, immunologically relevant functions of glycan structures have been inferred for other parasitic helminths, such as *Fasciola hepatica* (liver fluke – a trematode) and *Dictyocaulus viviparus* (lungworm – a nematode) (60, 61).

The present immunisation experiments in goats demonstrated that the de-glycosylation of H11 by periodate treatment significantly impaired immunoprotection and reduced specific anti-H11 IgG serum antibody levels. We infer that glycan moieties present in the glycoproteins of intestinal microvilli induce a particular IgG response that, in turn, targets and destroys the integrity of the nematode’s intestine in the immunised animal, and we demonstrate that IgG antibodies target glycans in the intestine and inhibit the aminopeptidase activity, which correlates with immunoprotection. Importantly, we provide the first experimental evidence that N-glycosylation of the H11 glycoprotein complex in the intestinal microvilli of *H. contortus* is crucial for immunoprotection. Taken together, these findings provide valuable insights and a solid foundation for the future development of vaccines against parasitic nematodes more broadly.

Data availability statement

The datasets presented in this study can be found in online repositories. The name of the repository and accession number can be found below: EMBL-EBI PRIDE Archive; PXD037140.

Ethics statement

This study was reviewed and approved by the Animals Ethics Committee of Huazhong Agricultural University (permit HZAUOGO-2019-002).

Author contributions

MH conceived and designed the project with the input from CW, NG, and RBG. CW performed the experiments and analyzed data. LL and TW assisted with animal experiments. XL contributed to N-glycan analysis. WP, RKS, and X-QZ contributed materials. CW, MH, and RBG wrote the manuscript, with inputs from X-QZ and NG. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2022.1034820/full#supplementary-material

SUPPLEMENTARY FIGURE 1 MALDI-TOF-MS/MS spectra of the permethylated N-glycans at m/z 2675, 2879 and 3083. (A–C) Permethylated N-glycans at m/z 2675 (A), 2879 (B) and 3083 (C) MALDI-TOF-MS/MS analyses: the predicted N-glycan configurations are symbol nomenclature: green circle = mannose;
yellow circle = galactose; blue square = GlcNAc; yellow square = GalNAc; red triangle = fucose. Peaks indicated by red arrows with dotted lines represent the loss of indicated fragments from the N-glycan ion.

SUPPLEMENTARY TABLE 1
List of N-glycoproteins and N-glycosylation sites identified in native H11 from Haemonchus contortus.

SUPPLEMENTARY TABLE 2
Results from domain enrichment analysis of N-glycoproteins of native H11 from Haemonchus contortus.

SUPPLEMENTARY TABLE 3
N-glycan configurations predicted for native H11 from Haemonchus contortus released by PNGase F.

SUPPLEMENTARY TABLE 4
N-glycan configurations predicted for native H11 from Haemonchus contortus released by PNGase A.

SUPPLEMENTARY TABLE 5
List of aminopeptidases and their N-glycosylation sites identified in native H11 from Haemonchus contortus.

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