**ABSTRACT**

Helminths like *Schistosoma mansoni* release excretory/secretory (E/S) products that modulate host immunity to enable infection. Extracellular vesicles (EVs) are among these E/S products, yet their molecular mechanisms and functionality of *S. mansoni* EV interaction with host immune cells is unknown. Here we demonstrate that EVs released by *S. mansoni* schistosomula are internalised by human monocyte-derived dendritic cells (moDCs). Importantly, we show that this uptake was mainly mediated via DC-SIGN (CD209). Blocking DC-SIGN almost completely abrogated EV uptake, while blocking mannose receptor (MR, CD206) or dendritic cell immunoreceptor (DCIR, CLEC4A) had no effect on EV uptake. Mass spectrometric analysis of EV glycans revealed the presence of surface N-glycans with terminal Gaβ1-4(Fucα1-3)GlcNAc (LewisX) motifs, and a wide array of fucosylated lipid-linked glycans, including LewisX, a known ligand for DC-SIGN. Stimulation of moDCs with schistosomula EVs led to increased expression of costimulatory molecules CD86 and CD80 and regulatory surface marker PD-L1. Furthermore, schistosomula EVs increased expression of IL-12 and IL-10 by moDCs, which was partly dependent on the interaction with DC-SIGN. These results provide the first evidence that glycosylation of *S. mansoni* EVs facilitates the interaction with host immune cells and reveals a role for DC-SIGN and EV-associated glycoconjugates in parasite-induced immune modulation.

**Introduction**

*S. mansoni* is one of the major helminth parasites of humans with over 200 million people infected [1,2]. In the initial phase of infection, schistosome larvae (cercariae) penetrate the skin of the host and transform into schistosomula. These schistosomula larvae will migrate to the circulatory system, develop into adult worms that mate and lay eggs [3]. *S. mansoni* can live up to 10 years in its host because it has developed successful mechanisms to evade immune responses. To establish this immune evasion, the parasite releases excretory/secretory (E/S) products that act on the host immune system [4]. Among these E/S products are extracellular vesicles (EVs) and evidence is accumulating that parasite-derived EVs contribute to parasite-host interaction [5]. The molecular composition of EVs from *S. mansoni* larvae and adult worms life stages has been partially characterized [6–8], but the interaction of their EVs with host cells remains unexplored.

For schistosomes to initiate and maintain infection, modulating host innate and adaptive immune responses is crucial [9]. Mouse models have shown that cercarial penetration triggers migration of innate antigen presenting cells (APCs), such as macrophages and dendritic cells (DCs), towards the skin draining lymph nodes [10]. These APCs have upregulated costimulatory molecules, like CD86 and major histocompatibility complex (MHC) class II, which are important for initiating adaptive immune responses. *In vitro* stimulation of mouse bone marrow-derived DCs (BMDCs) with E/S from schistosomula shows a similar pattern: increased costimulatory molecules and MHC class II expression and increased proinflammatory cytokine release (IL-6, IL-12, and TNF-α) [11]. Furthermore, cercarial secretions can upregulate the expression of IL-10 and programmed death ligand (PD-L)1 and 2 in human monocyte-derived (mo)DCs, suggesting that the parasite additionally utilizes regulatory pathways to dampen...
adaptive immune responses [12]. Relatively little is known about the exact molecular or structural components derived from schistosomula that induce immunomodulatory effects. Thus far, it has been shown that recombinant tetraspanins, transmembrane proteins associated to adult worm EVs [13], induce IL-10 and Th1 cytokine responses by peripheral blood mononuclear cells [14]. Whether schistosomula EVs have similar effects on human DCs has not been investigated.

Studies exploring the molecular content of schistosomula E/S have shown that part of the cytokine responses by mouse macrophages and whole blood of infected individuals is induced by glycosylated antigens in the E/S [15,16]. Schistosome glycoconjugates can be recognized by host pathogen recognition receptors (PRRs) on APCs, in particular the C-type lectin receptors (CLRs) such as the mannose receptor (MR, CD206) [17], dectin-1/2 [18], dendritic cell immunoreceptor (DCIR, CLEC4A, CD367) [19] and dendritic cell-specific ICAM-3-grabbing nonintegrin (DC-SIGN, CD209) [20]. A well-known immunogenic glycan-motif, Galβ1-4(Fucα1-3)GlcNAc or LewisX (LeX), is present on glycoproteins in schistosomula and egg E/S and can be recognized by DC-SIGN [21] as well as MR [22], leading to different effects depending on the structural context of the LeX motif. Pathogen-associated molecular patterns (PAMPs) containing high-mannose (oligomannose) structures, which form ligands for MR and DC-SIGN [23] have also been found on glycoproteins in E/S from schistosomes [24]. This raises the question whether schistosomula EVs expose glycans instrumental in targeting to host immune cells and whether these EV-associated glycans play a role in modifying subsequent immune responses. While it is known that mammalian EVs contain glycoconjugates, publications on the structure and function of EV glycans are very limited so far [25–29].

In order to gain insights into the interaction of schistosome EVs with the immune system we studied the interaction of schistosomula EVs with human moDCs and found that the EVs are internalised mainly via DC-SIGN. We show that N-glycans on the surface and glycolipids of the EVs contain DC-SIGN ligands, including LeX. Furthermore, we demonstrate that these EV preparations increase the activation status of moDCs, affecting both immunostimulatory and immunoregulatory pathways which were partly dependent on the interaction with DC-SIGN. Our study provides evidence for a specific CLR-mediated uptake of EVs that substantiates the importance of EV-associated glycoconjugates in pathogen-host interaction.

Materials and methods

Schistosomula culture

Infected Biomphalaria glabrata snails were incubated in water at 30°C for 2 h to shed cercariae of the Puerto Rican-strain of S. mansoni by exposure to light. The collected cercariae in water were stored on ice for 1.5 h to immobilize them and were subsequently pelleted by centrifugation at 440 × g. After removal of the supernatant, 12 mL of pre-warmed (37°C) DMEM (Dulbecco’s Modified Eagle Medium, high glucose with L-glutamine, Lonza, Basel, Switzerland) supplemented with 200 U/ml penicillin and 200 µg/ml streptomycin (Sigma-Aldrich, St. Louis, MO, USA), was added to transform the cercariae to schistosomula by providing mechanical force via pipetting and incubating for 20 min at 37°C [30]. Cercarial bodies were separated from their tails using an orbital shaker. The collected schistosomula were resuspended in DMEM at a concentration of 7,500 schistosomula/mL and cultured in 25 cm² polystyrene flasks (Greiner Bio-One, Alphen a/d Rijn, The Netherlands) at 37°C and 5% CO2 for 72 h.

EV isolation and staining

Schistosomula E/S products were enriched for EVs by differential centrifugation as previously described, with minor modifications [6]. Briefly, the collected culture supernatant (9–34 mL per culture, 7,500 schistosomula/mL) was centrifuged in 15 mL tubes (Greiner Bio-One) twice at 500 × g (k-factor 115,790.8) for 2 min (4°C) (SX4750A rotor and an Allegra X-15 R centrifuge) (Beckman Coulter, Brea, CA, USA) with low brake to remove remaining parasites. To remove any residual debris, the supernatant was subsequently centrifuged at 700 × g (k-factor 82,764.4) for 20 min (4°C, low brake). Next, an EV-enriched pellet was obtained by centrifugation of the supernatant at 31,000 rpm (average around 120,000 × g, k-factor 216.3) for 80 min at 4°C (max. acceleration and brake), followed by three wash steps with cold phosphate buffered saline (PBS) (B. Braun, Melsungen, Germany) in thin-wall polypropylene tubes using an SW41 Ti rotor and an Optima XE-90 ultracentrifuge (Beckman Coulter). For binding/uptake experiments, EVs were stained with PKH26 (Sigma-Aldrich) after the first ultracentrifugation step by addition of 80 µL Diluent C to the resuspended EV-enriched pellet and incubation of 93 µL diluted PKH26 (1.5 µL in 100 µL Diluent C) for 3 min at RT before addition of 11 mL PBS. Unconditioned culture medium incubated without
parasites was processed (and stained) following the same procedures and was used as (dye) control. EV-enriched pellets (from 66,300–253,200 schistosomula) for in vitro experiments were resuspended in 510 µL PBS, for transmission electron microscopy (TEM) and glycan analysis in 100 µL PBS, and for cryo EM in 40 µL PBS. All were stored at −80°C until further use except for 1 cryo EM sample, which was processed directly after EV isolation. We have submitted all relevant data of our experiments to the EV-TRACK knowledgebase (EV-TRACK ID: EV190032) [31].

Transmission electron microscopy

EV preparations were generated in Leiden and visualized by TEM at Aberystwyth University, as described previously [6]. Briefly, 10 µl of EV-enriched sample was fixed with an equal volume of 4% glutaraldehyde, adsorbed onto Formvar/carbon-coated copper grids (Agar scientific, Stansted, UK) for 40 min, and subsequently contrast stained with 2% uranyl acetate (pH 4) for 10 min. Processed samples were then visualized on a Jeol 1010 transmission electron microscope operated at 80 kV. Images were recorded with a Kodak MegaPlus camera Model 1.4i, other than the addition of scale bars, no further image processing was done. Sizes were measured by hand using Fiji/ImageJ software [32].

Nanoparticle tracking analysis (NTA)

EV-enriched suspensions were diluted 1:100 in PBS (to obtain 25–100 particles per frame at camera level 16) before analysing the concentration and size distribution by nanoparticle tracking analysis (NTA) using a NanoSight NS500 (Malvern Panalytical, Malvern, UK) equipped with an sCMOS camera. For each EV-enriched pellet, three videos of 30 seconds were recorded on three different camera levels: 12, 14 and 16. The analysis was done with NTA3.3 software and a detection threshold of 5. The average particle concentration of nine videos per EV-enriched sample, after subtraction of the NTA background data from PBS alone, was used for further experiments. We additionally measured the protein concentration of the EV-enriched pellets with microBCA according to the manufacturers protocol (Pierce, Thermo Fisher Scientific, Waltham, MA, USA).

Cryo electron microscopy

Previously frozen or freshly isolated EV preparations (EV from 86,00–112,00 schistosomula in 40 µL PBS) were visualized by cryo EM. Copper EM grids supporting a carbon film with regularly spaced 2 micron holes (R2/2, Quantifoil, Jena, Germany) were glow-discharged in air at 0.2 mbar for 2 min at 20 mA (EMITECH K950X with glow discharger unit). A 3 µL drop of sample was applied to the grid and transferred into the environmental chamber of a Leica EM grid plunger (Leica Microsystems, Wetzlar, Germany) operating at RT and between 92% and 94% humidity. Excess sample was blotted away for 1 second using filter paper (Whatman no.1) and without waiting plunged into a mixture of ethane/propane (63/37 v/v) cooled with liquid nitrogen to −193°C. After vitrification, grids were stored under liquid nitrogen and transferred into a cryo holder (type 626, Gatan, Germany). In total 238 cryo-EM projection images were recorded by 1 second exposures at spot 5 on a FEI Tecnai F12 at 120 kEV on a 4 k × 4 k CCD camera (Eagle, Thermo Fisher Scientific) at a magnification of 13,500 × (0.85 nm pixel size) and a defocus value ~−8 microns. Size measurement was performed by hand using Fiji/ImageJ software [32]. After removing double images and images with no EVs, 233 images were used to measure a total of 1056 EVs, which were subsequently quantified in segments of 20 nm (i.e. 21–40 nm, 41–60 nm, 61–80 nm ... 861–880 nm, 881–900 nm).

Human monocyte-derived dendritic cells (moDCs)

Venous blood of healthy volunteers who provided informed consent, approved by the Institutional Review Board of Leiden University Medical Centre, was used to isolate monocytes and differentiate to moDCs as previously described [33]. Immature DCs were harvested on day 5 or 6, counted, seeded at 5 × 10⁴ cells/well in a 96 well flat-bottom plate, and rested overnight at 37°C and 5% CO₂. Subsequently, cells were stimulated with or without a pre-incubation of 30 min with 10 mM EGTA (Sigma-Aldrich), 20 µg/mL αDC-SIGN/CD209 (clone AZN-D1, custom order without sodium-azide) (Beckman Coulter), 20 µg/mL αMR/CD206 (clone 15–2) (BioLegend, San Diego, CA, USA), 20 µg/mL αDCIR/CLEC4A (clone 111F8.04, Dendritics, Novus Biologicals, Centennial, CO, USA), and 20 µg/mL mouse IgG1 isotype control (clone P3.6.2.8.1) (Invitrogen, Thermo Fisher Scientific) in the presence of αFcgammaR-binding inhibitor (eBioscience, Invitrogen) and in the presence or absence of the maturation factors IL-1β (25 ng/mL) (BioLegend) and TNF-α (50 ng/mL) (Sino Biological, Beijing, P.R. China) or LPS (100 ng/mL) (InvivoGen, San Diego, CA, USA). As a positive control for the αMR, PF-647-labelled recombinant omega-1 was used [22,34]. EV-enriched pellets were thawed only once...
and several EV batches were pooled before incubation with the cells (6 × 10^6 EV/mL or mentioned otherwise). To investigate the effect of surface de-N-glycosylation, EVs were incubated with or without peptide N-glycosidase F (PNGase F) (4 U/100 µL, Roche Diagnostics, Almere, The Netherlands) at 37°C for 20 h before moDC incubation. Supernatants were collected from >85% CD1a^+ cell cultures after 24 h stimulation and IL-6 (Sanquin, Amsterdam, The Netherlands), IL-10 (BioLegend), and IL-12p70 (BD Biosciences, Franklin Lakes, NJ, USA) cytokine production was determined with ELISA according to the manufacturers protocols. Stimulated moDCs were washed, stained, and measured by flow cytometry on a FACSCanto II (BD Bioscience) and using the following antibodies: CD1a-BV421 (clone HI149) (BioLegend), HLA-DR-APC-eFl780 (clone LN3) (eBioscience), PD-L2/CD273-FITC (clone MIH18) (Miltenyi Biotec, Bergisch Gladbach, Germany), CD86-FITC (clone 2331 (FUN-1)), CD40-APC (clone 5C3), CD80-V450 (clone L307.4), PD-L1/CD274-PE-Cy7 (clone MIH1) (all BD Bioscience) with the addition of Fc receptor binding inhibitor (eBioscience) and Aqua live/dead staining (Invitrogen). Flow cytometric measurements were analysed with FlowJo (version 10, BD Bioscience).

**Confocal microscopy**

5 × 10^4 moDCs/chamber were seeded onto poly-l-lysine (Sigma-Aldrich) coated coverslips of a 4 chamber glass bottom dish (635 mm; Greiner Bio-One) for 24 h. Cells were pre-incubated with EGTA or aDC-SIGN+FcgammaR-binding inhibitor as described above, incubated with PKH-labelled schistosomula EV-enriched pellets for 5 h, subsequently washed, and treated with Hoechst (Sigma-Aldrich). Images were taken at 37°C and 5% CO_2_ on a Leica TCS (true confocal scanning SP8 WLL (white light laser) microscope (Leica Microsystems). The sequential scanning mode was applied to image Hoechst (excitation: 405 nm, emission: 420–470 nm) and PKH26 (excitation: 561 nm, emission: 570–630 nm). For imaging the uptake of EVs, a 63x objective (Leica HC PL APO 63x/1.40na OIL CS2) was used. The z-stacks were recorded and maximum projections of the recorded z-stacks were generated using the Leica software (LAS X version 1.1.0.12420; Leica Microsystems).

**N-glycan and glycolipid-glycan analysis**

For the N-glycan analysis, EV-enriched pellets (in PBS) from >100,000 cultured schistosomula were lyophilized, resuspended in 100 µL milliQ water, sonicated and subsequently reduced and denatured for 10 min at 95°C with the addition of SDS and β-mercaptoethanol which were neutralized by adding NP-40 (Sigma-Aldrich). Full details on N-glycan isolation has been described previously [35]. N-glycans were released by PNGase F (4 U/100 µL) incubation for 24 h at 37°C and cleaned up by collection in the flow through of reversed phase (RP) C18-cartridges (JT Baker, Phillipsburg, NJ, USA) followed by isolation on carbon cartridges (Supelclean ENVi-carb SPE, Sigma-Aldrich). In addition, directly after isolation, intact EV preparations in PBS were treated with PNGase F for 24 h at 37°C to release directly accessible N-glycans (surface glycans). The total suspension was subsequently transferred to a thin-wall polypropylene tube and topped up with PBS. The EVs were pelleted by ultracentrifugation at 42,000 rpm (average around 120,000 × g, k-factor 85.4) for 65 min at 4°C (max. acceleration and brake) using a TLS-55 rotor and an Optima TLX (Beckman Coulter). Next, the supernatant containing the PNGase F released N-glycans was collected and these N-glycans were isolated with C18- and carbon-cartridges. The EV-enriched pellet without PNGase F accessible surface N-glycans was resuspended in PBS, lyophilized, sonicated, reduced, denatured, and treated with PNGase F to isolate remaining N-glycans as above.

For glycolipid-glycans analysis, EV-enriched pellets were lyophilized, resuspended in milliQ water, sonicated, and subjected to extraction with chloroform and methanol (MeOH). The upper phase was collected after sonication and centrifugation. Similar volume as collected was replaced with 50% MeOH and the previous steps were repeated twice. All collected upper phases of the extraction were applied to an RP C18-cartridge and flow-through and wash fractions were combined and applied to another C18-cartridge. Glycolipids were eluted from the cartridges with chloroform/MeOH/water and dried under a flow of nitrogen. The glycolipids were subsequently dissolved in 200 µL 50 mM sodium acetate with 0.1% sodium taurodeoxycholate hydrate (Sigma-Aldrich), sonicated, and heated to 60°C for 10 min. 2 mU of recombinant endoglycoceramidase II (*Rhodococcus sp.*) (rEGCase II) (Takara-bio, Kusatsu, Shiga, Japan) was added to release the lipid-bound glycans. After 24 h at 37°C, another 2 mU was added and the sample was incubated at 37°C for another 24 h. The purification of released lipid-glycans was performed as described for the N-glycans using RP C18- and carbon cartridges.

To support glycan structure assignments, part of the isolated N-glycans and glycolipid-glycans were additionally treated with hydrofluoric acid (HF), which removes labile substitutions including α1-3 linked
fucoses. All isolated glycans were labelled with 2-aminobenzoic acid and purified by Biogel P10 (Bio-Rad, Hercules, CA, USA). The labelled glycans were measured by MALDI-TOF-MS with 2,5-dihydroxybenzoic acid (Bruker Daltonics, Bremen, Germany) as matrix using UltraflexXtreme mass spectrometers (Bruker Daltonics) in the negative-ion reflectron mode. When necessary, samples were cleaned up with ZipTip C18 (Merck Millipore, Burlington, MA, USA) before MALDI-TOF-MS. The obtained mass spectra were smoothed and the base-line was subtracted using FlexAnalysis (version 3.4, Bruker Daltonics). Glycan compositions were identified from the peak lists using GlycoWorkbench (Version 3) [36]. Peaks with a signal to noise ratio below 2 were excluded and masses are registered as deprotonated [M-H]-. 2-AA was taken into account as a fixed reducing-end modification.

For the interpretation of the relative most abundant signals for spectral assignments we used (when available) previously published structural data from *S. mansoni* glycans [35]. In the spectra, the structure of the most likely or most abundant isomer of the composition is indicated.

**RNA extraction and qPCR analysis**

Human moDCs were pre-incubated with or without αDC-SIGN, αMR, or IgG1 isotype and stimulated with IL-1β and TNF-α and *schistosomula* EV-enriched preparations as described above. After 6 h stimulation at 37°C and 5% CO₂, cells were stored on ice for 10 min, harvested, washed with cold PBS, snap frozen, and stored at −80°C till RNA extraction. RNA was extracted with the RNeasy Kit (Qiagen, Hilden, Germany) according to the manufacturer’s protocol. RNA was quantified using NanoDrop 1000 Spectrophotometer (Thermo Fisher Scientific) and cDNA synthesis was performed on 0.2 µg RNA according to standard procedures. Primer Express (Applied Biosystems, Waltham, MA, USA) was used to design primers that were synthesized by Biologio (Nijmegen, The Netherlands). Sequences of the primers were: β-actin_F: 5’-GCTACGAGCTGCTGACCG-3’; β-actin_R: 5’-CACGGAGGCGGATGGGACCC-3’; β-2-M_F: 5’-TGCCGTGTGAAACATGTA-3’; β-2-M_R: 5’-CCAAATGCGGCATCTTCAA-3’; RPLPO_F: 5’-GGCGACCTGGAAGTCCAACT-3’; RPLPO_R: 5’-CCATCAGCACACAGCCCTTC-3’; IL-10_F: 5’-ACCTGCCTAACATGCTCAGG-3’; IL-10_R: 5’-CCA GCTGATCCTTCATTTGGAAG-3’; TNF-α_F: 5’-TCTTCGAAACCCCGATGTA-3’; TNF-α_R: 5’-CCTCTGATGGCACCACAGG-3’; IL-12p35_F: 5’-CTCTGGAGCACCAGTTTG-3’; IL-12p35_R: 5’-TTGTCTGGCCTTCTGGAGC-3’. Quantitative real-time PCR (qPCR) was performed using CFX96 instruments (Bio-Rad Laboratories) and CFX Maestro (Bio-Rad) software. Technical duplicates with <1 Cq value difference were averaged and gene expression was calculated with the ΔΔCq method using the average Cq of the reference genes β-actin, β-2-M, and RPLPO to normalize [37].

**Statistical analyses**

All data were analysed using a paired student t-test or repeated measures One-way ANOVA (P values <0.05 were considered significant) with Tukey’s or Dunnett’s Multiple Comparison Test in GraphPad Prism 5.0 (GraphPad Software, Inc., La Jolla, CA, USA).

**Results**

**Cryo electron microscopy reveals ultrastructural characteristics of schistosomula EVs**

*Schistosomula* were cultured for 72 h and EV-enriched preparations were obtained from *schistosomula* E/S by sequential (ultra)centrifugation steps. TEM confirmed isolation of vesicles in the size-range between 35 and 190 nm (Figure 1(a–b)). NTA analysis showed a size-range of 30–650 nm with a minor peak around 40 nm and three major peaks around 110, 160, and 350 nm (Figure 1(c)). Culture medium without parasites that was processed similarly (medium control) and PBS alone only showed minor peaks between 60 and 150 nm as NTA background (Figure 1(c)). The average particle concentration measured with NTA was 2.33E10/100,000 schistosomula and the average protein concentration of EV-enriched preparations was 6 µg/100,000 schistosomula.

Although the TEM images corresponded with previous observations [6], there was a discrepancy in the sizes measured with the TEM and NTA. Therefore, we additionally analysed the EV preparations with cryo EM to visualise the near native state of the EVs and measure their size (Figure 1(d)). Interestingly, the cryo EM revealed thin filament-like structures covering the EV surface in 45.5% of all measured EV and in >70% of EVs when excluding EVs smaller than 100 nm. The filamentous structures ranged from 10 to 340 nm in length (average length of 128.5 nm) additional to the size of the EV diameter to the EV membrane (Figure 1(e)). These surface structures were most likely lost during the sample preparation for negative stained TEM (Figure 1(a)) and thus not observed, while NTA size measurement did include these structures as shown by similar size ranges between NTA and cryo EM.
Figure 1. TEM, NTA and cryo EM measurement of schistosomula EVs.

EV-enriched preparations analysed by transmission electron microscopy at 60,000 × (a) and 120,000 × magnification (b). Scale bars are 200 nm. EVs are pointed out with arrow heads, the small (<60 nm) structures (black arrows) in (b) are artefacts and were also observed in PBS only. In addition, EVs were analysed by nanoparticle tracking analysis (NTA) (c). The graph shows the average of 17 schistosomula EV-enriched preparations (black line), 10 medium controls (dark orange line), and 15 PBS (background) in which the preparations were resuspended (blue line), averages are shown with SEM (lighter areas). Cryo EM of EVs at 13,500 × magnification (d). Scale bars are 500 nm. The EV membrane and stretch of the filamentous structures are indicated in the middle pane with white dashed circles and black dashed circles, respectively. Right pane shows a close up of one EV with filaments. Quantification of the EV sizes excluding (EV diameter) or including (EV diameter + filaments) the thin filaments from a total of 1056 EVs (e). EM pictures are representative for four biological replicates.
Internalisation of schistosomula EV by human moDCs is calcium dependent

To investigate the interaction of schistosomula EVs with human moDCs, EVs were first labelled with the fluorescent dye PKH26. MoDCs (0.25E6 moDC/mL) were incubated with a maximum of 6E9 EV/mL, which is equivalent to approximately 10 moDCs receiving the number of EVs released by one schistosomulum during three days of culture. After 2 h of incubation, there was a dose-dependent increase in EV binding/uptake by the CD1a+ moDCs indicated by an increased geometric mean fluorescence intensity (geoMFI) relative to cells in medium only (Figure 2). Furthermore, the fluorescence of cells incubated with dye control was unchanged, indicating that the increase...
in MFI was due to binding/uptake of labelled EVs and not due to dye aggregates [38]. EV binding/uptake by moDCs did not change due to stimulation of the cells with LPS or in the presence of IL-1β+TNF-α. Incubation at 4°C instead of 37°C did not lead to binding/uptake of EVs by the moDCs (Figure 2), suggesting that the interaction with and uptake of the schistosomula EVs by target cells are active processes. We visually confirmed that EVs were internalised by moDCs with confocal microscopy (Figure 2(c)).

MoDCs are known to express C-type lectin receptors that bind glycan motifs present on schistosome E/S components in a calcium dependent manner [39]. To investigate whether moDCs also recognise schistosomula EVs via CLR-glycan motif interactions, moDCs were incubated with fluorescently labelled EVs in the presence of the calcium chelator EGTA and subsequently analysed by flow cytometry and confocal microscopy (Figure 2(d–e)). Pre-incubation of EGTA almost completely abrogated the fluorescence signal and EV internalisation of moDCs compared to EV-exposed moDCs without EGTA pre-incubation, indicating calcium dependent interaction with moDCs such as CLRs. Next, when EGTA was added after incubation with EVs in order to remove EVs bound to CLRs, this resulted in only a minor reduction of fluorescence signal (Figure 2(d)), confirming that most of the EVs were internalised by moDCs rather than bound to the surface.

**Schistosomula EVs contain CLR ligands on their surface**

We next analysed the glycosylation of schistosomula EVs to assess whether ligands for CLRs were present. First, the overall N-glycan content of the EV-enriched preparations was determined by mass spectrometry (Figure 3(a)). MALDI-TOF mass spectra of PNGase F released N-glycans were assigned based on the detailed glycan structure descriptions available for overall N-glycan preparations of S. mansoni schistosomula [35]. Interestingly, the spectrum of N-glycans from the EV preparation was highly similar to previously published spectra of total extracts of three-day cultured schistosomula [24,35]. The major signals are from oligomannosidic structures of complex glycans with a core(a6)-fucose and one or two antennae consisting of Galβ1–4GlcNAc (LacNAc, LN) and/or LeX. Structures with a core-xylene modification or GalNAcβ1–4GlcNAc (LacDiNAc or LDN) antennae with five to nine fucose residues were observed at relatively low levels (Figure 3(b)). These results show that schistosomula EVs contain similar N-glycans as previously found in total schistosomula extracts, including LeX and oligomannose motifs, which are both ligands for DC-SIGN and MR [21,23].

To identify specific N-glycans that might be available for interaction with CLRs on the surface of EVs, we treated intact EVs with PNGase F to release all ("surface") N-glycans that were accessible to the enzyme. MALDI-TOF-MS analysis was performed on both the released "surface" N-glycans as well as on the remaining N-glycans of the PNGase F-treated ("shaved") EVs (Figure 3(c–d)), which likely represent glycans on the inside of the EVs. Interestingly, the most abundantly detected N-glycans on the EV surface contained one or two LeX antennae (Figure 3(c)) while the major glycans of the "shaved" EVs were the oligomannosidic structures (Figure 3(d)). These results show that the glycoproteins on the surface of schistosomula EVs carry a specific subset of N-glycans with LeX motifs, which are potential ligands for DC-SIGN or MR in the context of pathogen-host interactions [19,40,41].

**EVs are internalised via DC-SIGN but not MR or DCIR**

To investigate whether DC-SIGN or MR on moDCs were involved in uptake of schistosomula EVs via glycan motifs on these EVs, moDCs were pre-incubated with antibodies blocking DC-SIGN or MR and subsequently incubated with labelled schistosomula EVs for 2 h (Figure 4(a–b)). In addition we investigated EV binding after blocking DCIR, a receptor expressed on moDCs that does not bind LeX containing glycans but has been shown to bind S. mansoni cercarial extract [19]. Blocking DC-SIGN led to almost complete inhibition of EV uptake while blocking the MR, DCIR, or pre-incubation with the isotype control did not reduce EV internalisation. Inhibition of EV internalisation after DC-SIGN block was confirmed by confocal microscopy (Figure 4(c)). This reveals that schistosomula EVs were internalised by moDCs via DC-SIGN.

**EV-associated glycolipid-glycans include DC-SIGN ligands**

Given that the EV surface contained DC-SIGN ligands, including the LeX motif, we investigated whether EV deprived from surface N-glycans by PNGase F treatment would still be internalised by moDCs. Interestingly, the PNGase F-treated EVs showed a minor but not significant reduction in internalisation, and the EV uptake could still be inhibited by blocking DC-SIGN (Figure 5(a)). Since it has been shown that cercariae produce glycolipid-glycans that contain LeX...
Schistosomula EV-surface N-glycans include DC-SIGN ligands. 

and other potential DC-SIGN ligands [35,42], we additionally determined the total lipid-derived glycan profile of the schistosomula EVs (Figure 5(b)). Here we detected a heterogeneous set of highly α3-fucosylated glycolipid structures, mostly similar to the glycolipid-glycans found in total schistosomula extract [35]. Interestingly, these lipid-glycans had terminal motifs that contained LeX, Fuca1-3Galβ1-4(Fuca1-3)GlcNAc (pseudo-LewisY
(LeY) and GalNAcβ1–4(Fucα1–3)GlcNAc (LDN-F) (Figure 5(b) insert), which are all previously described ligands of DC-SIGN [42, 43]. These data suggest that additional to surface N-glycans, EV-associated lipid-glycans play a role in interaction with DC-SIGN.

**Schistosomula EVs induced increased cytokine release and costimulatory molecule expression on moDCs**

Next, we investigated whether incubation of moDCs with schistosomula EVs affect their activation status and cytokine production. We first incubated human moDCs for 24 h with EV-enriched preparations but observed no or very low release of cytokines. Therefore, in addition to EVs, moDCs were co-cultured with IL-1β and TNF-α, cytokines that are released by immune cells upon schistosome infection [3], allowing us to further investigate whether EVs could change the activation status of moDCs. Under these conditions, schistosomula EVs significantly increased IL-6, IL-10 and IL-12 secretion by moDCs (Figure 6(a)).

Furthermore, a significant upregulation of the costimulatory molecules CD80, CD86 and regulatory surface molecule PD-L1 was observed, while there was no significant effect on CD40, HLA-DR and PD-L2 expression (Figure 6(b)). These data suggest that the EV-enriched preparations were mostly synergizing or augmenting other inflammatory signals.

**Role for DC-SIGN in EV-augmented immune responses**

Since schistosomula-derived EVs were mainly internalised via DC-SIGN, we hypothesized that inhibiting this receptor would alter the observed augmented immune responses of moDCs by the EVs. Blocking DC-SIGN during a 24 h EV stimulation, however, did not show a significant effect on IL-6, IL-10 and IL-12 release (Figure 7(a)) nor on the expression of the co-stimulatory surface markers CD80, CD86, HLA-DR, PD-L1 and PD-L2 after 48 h stimulation (Figure 7(b)). To understand why the blocking of DC-SIGN did not consistently influence the enhanced immune responses of moDCs by the EVs whereas it almost completely blocked EV internalisation after 2 h, we studied EV uptake in the presence of DC-SIGN blocking antibodies after 48 h of
incubation. Blocking of DC-SIGN significantly reduced EV internalisation after 48 h (Figure 7(c)). However, the vast majority of the moDCs were still positive for the fluorescent dye, although the fluorescent intensity was lower when DC-SIGN was blocked (Figure 7(d)). Possibly other mechanisms than those mediated by DC-SIGN play a role in EV internalisation by moDCs after prolonged exposure. To investigate whether blocking of DC-SIGN does interfere with moDC function during shorter incubations with schistosomula-derived EVs, moDCs were stimulated with EVs in the absence or presence of blocking antibodies for 6 h, after which cytokine mRNA expression was determined. Indeed, at this time point, the effect of blocking DC-SIGN was more prominent with a significant reduction in TNF-α mRNA and a trend for lower IL-10 mRNA expression (Figure 7(e)). IL-12p35 mRNA levels were also reduced in most donors, though not significant. These results show that, within the first hours of exposure but not at a longer timescale, the moDC immune profile is influenced by schistosomula EVs through internalisation via DC-SIGN.

Discussion

Schistosome parasites are master regulators of host immune responses and they release various molecules
and products to achieve this. Here we demonstrate that *S. mansoni* schistosomula release EVs that contain LeX antigens on their surface. Interaction of the glycosylated EVs with DC-SIGN on moDCs lead to internalisation and enhanced expression of both immunostimulatory and regulatory effector molecules. Thus, schistosome EVs with DC-SIGN on moDCs lead to internalisation and enhanced expression of both immunostimulatory and regulatory effector molecules. Thus, schistosome

**Figure 6.** Schistosomula EVs augment moDC immune responses.

Released cytokines (a) and surface marker expression (b) of moDC incubated with EVs. The control is medium without parasites that has been cultured and processed exactly like the schistosomula EVs. Data points are shown as fold change relative to moDC with IL-1β and TNF-α for that specific donor. MoDC from 7 to 11 donors. Mean±SEM ***p < 0.001, using repeated measures ANOVA with Dunnett’s Multiple Comparison Test.
Figure 7. Role for DC-SIGN in augmented immune responses by schistosomula EVs.

Released cytokines (a) and surface marker expression (b) of moDC with indicated pre-incubations (x-axes) before EV incubation. Linked points represent data from one donor. Data is shown as fold change relative to moDC with IL-1β and TNF-α from that specific donor (n = 7). Summary of geometric mean fluorescent intensity (geoMFI) of moDC incubated with PKH26-labelled EVs for 48 h, relative to medium control (c) (n = 6). Histograms of moDC from one representative donor from data plotted in c (d). Relative expression of mRNA from moDC after 6 h EV incubation and indicated pre-incubations (x-axes) (e) (n = 5). *p < 0.05, ***p < 0.001, using repeated measures ANOVA with Tukey’s Multiple Comparison Test (c) or paired student t-test (a, b, e)
EVs appear to contribute to immune modulation by the parasite.

Schistosomula are multicellular organisms, having various organs and cells as potential sources of their released EVs. It has been shown that micron-sized vesicles are released from their acetabular glands [11, 44] and it is suggested that EVs can also derive from the tegument [6]. Since multicellular organisms release heterogeneous populations of EVs, it is likely that the moDCs can internalise these EVs via various receptors and/or routes [13]. The schistosomula-derived EV populations described here were heterogeneous in size, which we analysed with NTA (Figure 1(c)) and cryo EM (Figure 1(e)). The size of EVs may influence uptake routes, as has been shown for Helicobacter pylori outer membrane vesicles (OMVs): small OMVs were taken up via caveolae-mediated endocytosis and bigger OMVs via micropinocytosis and endocytosis [45]. The major population of EVs >300 nm we observed have not been described before for S. mansoni schistosomula [6]. Although we used a similar schistosomula EV isolation protocol as Nowacki et al [6], the isolated EVs in that paper were only visualised by TEM of negatively stained EVs and not by NTA. With this same technique, our isolated schistosomula EVs also showed similar sizes <200 nm (Figure 1(a–b)) but only with the NTA and cryo EM the EV population >300 nm was detectable. EVs >300 nm have not been found among adult worms EVs [7, 8, 46]. Schistosomes are organisms with a complex life cycle and the schistosomula and adult worm life stages differ in size, shape, molecular, and cellular make up, and interaction with the host and its immune system [2, 47]. Therefore, it is likely that different life stages of the parasite produce different EV populations regarding molecular content [6, 48] as well as size. Furthermore, the adult worm EVs published have been isolated by different protocols which may further explain differences between the observed sizes of EVs derived from various schistosome different life stages.

Interestingly, our cryo EM data showed that around half of the schistosomula-derived EVs were covered with thin filament-like structures (Figure 1(d)). These structures have not been described for eukaryotic EVs so far. It remains to be explored what the molecular composition of these EV-associated structures is and whether these structures are specific for S. mansoni and/or specifically related to the schistosomula life stage. These thin filaments resemble the electron-dense surface layer of extracellular proteins and LPS or glycoalyx on some bacteria [49, 50] and their released OMVs [51]. We hypothesize that the structures on schistosomula EVs are composed of large, complex glycoconjugates and/or include proteins with attached glycan polymers, similar to the mammalian glycocalyx consisting of proteoglycans with attached glycosaminoglycans [25]. Altogether, the EM analyses emphasize that cryo EM provides improved visualisation of the near native state of isolated EVs compared to TEM of negatively stained EVs.

EVs released by S. mansoni schistosomula were internalised by human moDCs (Figure 2). Previous studies have demonstrated that S. japonicum EVs derived from adult worms and eggs are also internalised by host cells [46, 52–54]. This indicates, together with our data, that at least three lifecycle stages of Schistosoma in humans release EVs that interact with the host. By chelating extracellular calcium with EGTA, schistosomula EV could not bind to or be taken up by the moDC (Figure 2(d–e)). This suggests that the interaction of the EVs with the moDC was via the ligation of glycoconjugates to CLRs, which is calcium dependent, and not via protein-protein interaction. By inhibiting specific CLRs we show that internalisation of schistosomula EVs by moDCs is primarily facilitated by DC-SIGN (Figure 4). This CLR is known to bind schistosome egg [21], cercarial [43], and worm antigens [55], via glycan motifs including mannnotriose, LeX, LDN-1, and pseudo-LeA [19, 42, 43]. In contrast, CLR DCIR mainly binds LeB (LewisB) and sulpho-LeA (LewisA) motifs [19], and indeed, blocking this receptor did not reduce EV uptake by the moDC. Previously, it was shown that the egg-derived glycoprotein omega-1, which carries LeX containing N-glycans highly similar to those found here on the EVs (Figure 3), is mainly internalised by moDCs via the MR [22]. However, when blocking the MR, EV internalisation was not inhibited. This difference of internalisation routes between omega-1 and EVs could be explained by the characteristics of the two CLRs: DC-SIGN has one carbohydrate recognition domain (CRD), is present as tetramers, and it clusters in (lipid raft) nanodomains that are distributed on the cell membrane. Clustering increases the avidity of low affinity glycan-lectin interactions and allows this receptor to interact at multiple sites with pathogens that differ greatly in size [56]. On the other hand, the MR consists of multiple CRDs, which results in binding of multivalent or repetitive ligands to a single MR monomer [57]. Considering these aspects, larger particles such as EVs would favour DC-SIGN mediated uptake while smaller glycosylated proteins are more likely to be internalised via other receptors, such as the MR. A very recent publication substantiates this theory by showing that tumour-derived apoptotic EVs with high-mannose glycans on their surface were mainly internalised via interaction with DC-SIGN and not the MR [58]. Furthermore, it
suggests that, together with our findings, DC-SIGN mediated EV uptake may be a widely occurring mechanism, across a broad spectrum of species and may not be exclusive to schistosomula-derived EVs. Of note is that helminths, including S. mansoni, lack sialic acid in their glycan repertoire, which is a fundamental difference with mammalian glycans. It was recently found that EVs released by human glioblastoma cells had complex sialic acid-capped N-glycans on their surface that mainly bound to Siglec-9 on moDCs [59]. When sialic acids on the EV surface were enzymatically removed and LeY was inserted, EV uptake by moDC and binding of the EVs to DC-SIGN were increased. Furthermore, EVs from murine hepatic cell lines expose sialyl-LeX on the EV surface and removing the sialic acids with neuraminidase significantly increased EV uptake by M1 cell lines [28]. However, no DC-SIGN blocking experiments were performed in these studies. Thus, the question remains whether EV internalisation via DC-SIGN is specific for schistosomula EVs, or whether it is a more general mechanism via which EVs from both pathogens and mammalian cells are internalised. However, we cannot exclude the possibility that internalisation of schistosomula EVs may be facilitated via different CLRs in a different host. For example, the murine macrophage galactose-type lectin (MGL-1) can bind to LeX [60] while the human MGL cannot [61]. Human MGL binds to LDN motifs, which were absent in our schistosomula EV glycan analyses.

So far, most research on glycosylation of pathogen-derived EVs focussed on lipopolysaccharides of bacterial OMVs [62–65]. Only very recently, the presence of glycoconjugates on the surface of EVs released by the helminth Fasciola hepatica was studied using lectin microarrays [66]. Here we performed a more detailed structural analysis of glycans associated with EVs released by a helminth parasite. Using mass spectrometry we showed the presence of mainly complex type N-glycans with LeX motifs as well as oligomannose and high mannosylated glycans (Figure 3(b)) and the presence of lipid-linked glycans with LeX, pseudo-LeY, and other α3-fucosylated glycan motifs (Figure 5(b)). The EV N-glycan profile was qualitatively very similar to that of whole schistosomula [35]. The overall glycosylation pattern associated with this particular schistosome life stage was reflected in the EVs, however, with differences in relative abundances of the glycans. Biological replicates of the parasite culture and EV isolation generated at two different laboratories (Leiden and Aberystwyth) showed similar N-glycan profiles, confirming the reproducibility of the EV isolation protocol and glycan patterns [6]. Mostly complex glycans with LeX motifs were cleaved from intact EVs by incubation with PNGase-F (Figure 3(c)), implicating that these structures were on the EV surface. Glycoconjugates on the EV surface can influence their cellular internalisation, as was shown for bacterial OMV [65], murine hepatic cell line EVs [28], tumour-derived EVs [58,59] and F. hepatica adult worm EVs [66]. Interestingly, PNGase F treated EVs were still internalised by moDC via DC-SIGN (Figure 5(a)). This indicates that other glycans with DC-SIGN ligands such as the fucosylated lipid-linked glycans play a role in this process, either specifically or in addition to the N-glycans. Mass spectrometry analysis showed the presence of EV-associated glycolipid-glycans (Figure 5(b)) containing several structures that were reported previously for schistosomula [35], but with more extended higher molecular weight structures. Our data indicate that many of the EV lipid-derived glycans contain motifs such as LeX, pseudo-LeY and other α3-linked fucose containing motifs that can bind to DC-SIGN [42,43], which may explain why PNGase-F treated EVs could still be internalised via DC-SIGN. Furthermore, S. mansoni produce various O-glycans [35], which are possibly also present on EVs and contribute to EV-CLR interaction. We therefore suggest that both N-glycans and glycolipid-glycans, and possibly O-glycans, contribute to the interaction of the EVs with DC-SIGN. This interaction is most likely via LeX motifs, which are abundant in these glycan types, with the possible contribution of other α3-fucosylated lipid-glycans.

Interestingly, the complex type N-glycans on the EV surface and many of the glycolipids of the schistosomula EVs contain antigenic glycan motifs that were previously shown to be the target of antibodies of various isotypes during schistosome infection. These motifs include N-glycan core-xylose, LeX, and the various fucosylated glycolipid motifs [67–69]. It is therefore tempting to speculate that EVs can either elicit these antibodies and/or that EVs are targeted by antibodies generated against other, similarly glycosylated, antigens produced by schistosomes during an infection. Antibodies that recognize and bind molecules on the EV surface can facilitate internalisation by APCs, for example via Fc receptors. Antibody-bound EVs can be targeted to different intracellular compartments compared to EVs without antibodies. This has been observed for EVs from the helminth Heligmosomoides polygyrus [70]. H. polygyrus EVs pre-incubated with antisera were targeted to lysosomes. However, lysosome targeting has also been observed for antigens internalised by DC-SIGN [71]. Another possibility is that EV uptake is enhanced after incubation with antisera, which was observed for F. hepatica EVs and RAW264.7 macrophages [66]. Differences in route of
uptake can possibly alter the fate of the EVs and thus possibly influence EV-induced immunomodulation.

Cross-species communication via EVs that contributes to modulation of host immune responses has been described previously for helminths, including *H. polygyrus* [70] and *Nippostrongylus brasiliensis* [72]. In this study, we observed that *S. mansoni* schistosomula EVs are capable of augmenting activation-induced cytokine secretion and surface molecule expression by human moDCs, including both immunostimulatory and regulatory factors (Figure 6). It has been suggested that a delicate balance between benefit for the host and benefit for the parasite contributes to overall survival of the parasite within the host with limited pathology [3]. The induction of pro-inflammatory cytokines is a natural response of the host to the skin-invading pathogen, however, this response is transient. Priming of a protective adaptive immune response is hampered, probably via the induction of regulatory responses by the parasite, such as increased IL-10 release and PD-L1 expression [12], allowing the parasite to develop into mature worms and start egg laying. Different molecules present in the heterogenous EV population may have contributed to the observed augmented immune responses. It is known that DC-SIGN signalling via fucose ligands, which are motifs found on the N-glycan and lipid-glycan structures of the EVs, mainly increases IL-10 and decreases pro-inflammatory responses [73]. In contrast, we observed that blocking DC-SIGN actually decreased pro-inflammatory TNF-α and IL-12 mRNA and did not fully reduce the responses to baseline (Figure 7(e)). Thus, it is likely to assume that schistosomula EVs contain a mix of various (glycosylated) proteins and RNAs (amongst other biological molecules, such as lipids [74]) that all may contribute to a combined effect on host immunity [6]. Indeed, the NTA data as well as the cryo EM show at least a variation in EV size and with or without the thin filaments, and it is tempting to speculate that these different EVs may show variation in their surface glycan profile and may have specific activities on host immunity. Interestingly, since total E/S from schistosomula increases IL-12, IL-6 and IL-10 release as well as CD86 expression by mouse BMDCs [11], it is tempting to suggest that part of the effects of the E/S are mediated by the EVs in that secretion.

In contrast to the strong blocking effect of anti-DC-SIGN antibodies on EV internalisation (Figure 4 and Figure 7(c)), however, blocking DC-SIGN did not significantly alter EV-augmented immune responses during prolonged stimulation (Figure 7(a–b)). The variation we detected among donors in reduction or increase of IL-10 release in the presence of blocking DC-SIGN (Figure 7(a)) was also observed in another study that examined glycan-mediated effects by larval E/S [75]. These donor-specific variances could be associated with intrinsic DC-SIGN levels, which vary highly between and within donors [76]. Furthermore, residual uptake of EVs via other processes such as (macro-) pinocytosis or protein–protein interaction could still have affected the immune activation of moDCs upon prolonged culture (Figure 7(d)). Shorter incubation, however, showed that part of the augmented immune profile of moDCs by schistosomula EVs was indeed dependent on interaction with DC-SIGN (Figure 7(e)) and provides evidence that interaction of glycans on the EV surface with DC-SIGN does play a role in immune modulation of host responses.

In conclusion, our study demonstrates that *S. mansoni* schistosomula release glycosylated EVs that carry LeX, pseudo-LeY, and other fucosylated motifs, and we reveal a distinct role for DC-SIGN in glycan-mediated internalisation of EVs by host immune cells. This interaction contributes to increased pro- and anti-inflammatory responses, substantiating that EVs play a role in host immune regulation by helminths to establish and control infection. Future studies on how EV-associated molecules contribute to immune modulation will further our understanding of parasite-host interactions and may provide insights for vaccine development.

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Authors’ contributions

M.E.K. contributed to conceiving the study, performed the schistosomula cultures, EV isolations, NTAs, cryo EM measurements, moDC isolations/stimulations, ELISAs, flow cytometry, RNA extraction, qPCR, data analysis and drafting of the manuscript. A.J.H. assisted with moDC isolations/differentiation and constructing flow cytometry panels. A.O.F. performed shedding of snails to obtain cercariae and assisted with moDC isolations. D.L.N. performed the N-glycan characterization and assisted with the analysis of the spectra. C.M. K. performed the confocal microscopy and generated the confocal images. R.I.K. performed the cryo EM imaging and assisted with the subsequent analysis. J.J.T. performed the TEM imaging and contributed to the TEM analysis. K.F. H. contributed to the design of experiments, interpretation of the results and correcting the manuscript. E.N.-tH. contributed to conceiving the study, the design of the experiments, overseeing the EV isolation, interpretation of the results and drafting of the manuscript. H.H.S. and C.H.H. both participated in conceiving the study, the design of the experiments, interpretation of the results and drafting of the manuscript. All authors read and approved the final manuscript.

Disclosure of interest

The authors report no conflict of interest.

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