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The Extracellular Vesicles of the Helminth Pathogen, *Fasciola hepatica*: Biogenesis Pathways and Cargo Molecules Involved in Parasite Pathogenesis*

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Extracellular vesicles (EVs) released by parasites have important roles in establishing and maintaining infection. Analysis of the soluble and vesicular secretions of adult *Fasciola hepatica* has established a definitive characterization of the total secretome of this zoonotic parasite. *Fasciola* secretes at least two subpopulations of EVs that differ according to size, cargo molecules and site of release from the parasite. The larger EVs are released from the specialized cells that line the parasite gastrodermus and contain the zymogen of the 37 kDa cathepsin L peptidase that performs a digestive function. The smaller exosome-like vesicle population originate from multivesicular bodies within the tegumental syncytium and carry many previously described immunomodulatory molecules that could be delivered into host cells. By integrating our proteomics data with recently available transcriptomic data sets we have detailed the pathways involved with EV biogenesis in *F. hepatica* and propose that the small exosome biogenesis occurs via ESCRT-dependent MVB formation in the tegumental syncytium before being shed from the apical plasma membrane. Furthermore, we found that the molecular “machinery” required for EV biogenesis is constitutively expressed across the intramammalian development stages of the parasite. By contrast, the cargo molecules packaged within the EVs are developmentally regulated, most likely to facilitate the parasites migration through host tissue and to counteract host immune attack.

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The trematode parasite, *Fasciola hepatica* is the causative agent of liver fluke disease (fasciolosis) in domestic animals. *F. hepatica* infects more than 300 million cattle and 250 million sheep worldwide resulting in losses of over $3 billion to global agriculture through lost productivity (1). Fasciolosis is also an important zoonotic disease with an estimated 2.6 million people infected worldwide; human infection is particularly highly prevalent in Egypt and the Andes of South America, while outbreaks regularly occur in Northern Iran (2).

Infection in animals and humans can last many years while adult parasites reside in the bile ducts. *F. hepatica* are obligate blood feeders and obtain blood by puncturing the bile duct wall. The regular regurgitation of parasite gut contents is thought to inject molecules into the bloodstream where they can exert an immunosuppressive activity on the host immune system (3). Other molecules are secreted from the gut, excretory pores and surface tegument into the bile where they may be carried to the exterior via the host intestine (4–6). We, and others, have previously used proteomic techniques to profile the secretome of adult *F. hepatica* (7–10) but these studies had a number of limitations: (1) they relied on very limited transcriptome data sets with considerable redundancy; (2) because genome sequence data was lacking it was not pos-
sible to determine precisely the identity and/or the number of the genes contributing to various secreted protein families; (3) the contribution of extracellular vesicles to fluke secretions was not investigated. However, our recent report of the Fasciola genome and associated transcriptome data sets (11) has now allowed us to address these limitations and perform a definitive characterization of the total secretome of adult F. hepatica. This is timely, given the report of exosome-like EVs released by adult F. hepatica (12).

Extracellular vesicles (EVs) are small membrane bound organelles that are shed by most cell types. Although once considered to be “cellular garbage cans” with the sole purpose of discarding unwanted cellular material (13), EVs are now recognized as important mediators of intercellular communication by transferring molecular signals, including proteins, lipids, mRNA, microRNA and other non-coding RNA species (14, 15). Variousy described as exosomes or microvesicles depending on their cellular origin and mode of biogenesis, EVs perform a variety of roles in the maintenance of normal physiology such as blood coagulation, immune regulation and tissue repair (16–18), but also participate in pathological settings, notably in tumor progression (19). A number of recent studies have shown that parasite-derived EVs play an important role during infection. For instance, EVs released by the helminth parasites Heligmosomoides polygyrus and Schistosoma japonicum are capable of modulating the response of host innate immune cells (20, 21).

Here, we report the deep analysis of the soluble and vesicular components of the adult F. hepatica secretome. We found that F. hepatica secretes at least two subpopulations of EVs that differ according to size, cargo molecules and potential site of release from the parasite. Using a proteomic approach, integrated with newly available F. hepatica genome and transcriptome resources (11), we have defined the protein composition of these two classes of vesicle and non-vesicle associated proteins. The larger 15K EVs are released from the specialized cells that line the parasite gastrodermus and contain, as a marker, the 37 kDa inactive zymogen of the cathepsin L1 digestive peptidase. In contrast, our analysis of the smaller 120K exosome-like vesicle population suggest that they are shed from the tegument of adult fluke. Although the 120K EV biogenesis “machinery” is constitutively expressed across the intramammalian developmental stages of the parasite, the cargo molecules packaged within the EVs are highly regulated, most likely to facilitate the parasites migration through host tissue and to counteract host immune attack.

**EXPERIMENTAL PROCEDURES**

**Experimental Design and Statistical Rationale**—Western blots and fluorogenic assays were performed with at least two biological replicates and three technical triplicates. For the proteomics study, whole Fasciola EVs were initially analyzed by LC-MS/MS (two technical replicates). These identifications were subsequently mapped to the surface/membrane/lumen of the EVs following analysis of the EV fractions, which were run without replicates because of the paucity of parasite material. The aim of this experiment was to profile the molecules associated with fluke EVs and no quantitation was performed.

**Isolation of Adult F. hepatica Extracellular Vesicles**—Adult F. hepatica parasites were obtained from bovine livers from local abattoirs. To prepare secretions, adult flukes were thoroughly washed with PBS to void their gut contents and then maintained in RPMI 1640 culture medium containing 0.1% glucose, 100 U penicillin and 100 mg/ml streptomycin (Sigma), at 2 worms/ml for 5 h at 37 °C. EVs were purified according to the differential centrifugation protocol described by Marcilla et al. (12). Briefly, after the incubation period, the parasite culture media was collected and centrifuged at low speed (first at 300 × g/10 min, and then at 700 × g/30 min) to remove large debris. The resulting supernatant was centrifuged at 15,000 × g for 45 min at 4 °C to obtain large vesicles. Culture supernatants were then filtered using a 0.2 μm ultrafiltration membrane, and centrifuged at 120,000 × g/1 h at 4 °C to recover smaller vesicles that were subsequently washed with PBS.

**Immunoblot Analysis of the EVs and Secretome**—The protein content of adult fluke soluble secretions and EVs was determined using the BCA protein assay kit (Pierce, Loughborough, UK). Equal amounts (5 μg) of each sample were run on reducing NuPage Novex 4–12% Bis-Tris gels (Life Technologies), and transferred to nitrocellulose membranes (GE Healthcare, Paisley, UK) at 120 mA for 45 min. Following transfer, the membranes were incubated in blocking solution (TBST: 20 mM Tris-HCl, 150 mM NaCl, 1% Tween-20, pH 7.6) containing 5% skimmed milk for 2 h at room temperature (18–21 °C). Blots were probed with a 1:1,000 dilution of sheep antiserum raised against recombinant F. hepatica peroxiredoxin (FhPPIX; 22) or recombinant cathepsin L1 (FhCL1; Dalton, unpublished); a 1:1,000 dilution of rabbit antiserum raised against recombinant Schistosoma mansoni leucine aminopeptidase which is reactive to F. hepatica leucine aminopeptidase (SmLAP; 24, Dalton unpublished) for 2 h at room temperature (18–21 °C). After washing in TBST (3 × 10 min), an appropriate alkaline phosphatase-conjugated IgG secondary antibody was applied to the membranes for 1 h at room temperature (18–21 °C) before detection using the BCIP/NBT substrate (Sigma).

**Trypsin Shaving and Sequential Extraction of Purified F. hepatica EVs**—Peptides from surface-accessible proteins were released by trypsin hydrolysis of EVs isolated from adult F. hepatica using our previously described protocol (25). Briefly, sequencing grade trypsin (Promega, Southampton, UK) was added to purified F. hepatica EVs at a final concentration of 50 μg/ml for 5 min at 37 °C. The treated EVs were then centrifuged at 100,000 × g for 1 h at 4 °C, and the released peptides recovered from the supernatant.

To investigate subvesicular protein localization, purified F. hepatica EVs were sequentially extracted to identify proteins associated with the EV membrane or those contained within the EV lumen. First, the EVs were resuspended in water; the resulting osmotic shock causes

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1 The abbreviations used are: EV, extracellular vesicle; NEJ, newly excysted juvenile; HDM, helminth defence molecule; CB, cathepsin B; CL, cathepsin L; LAP, leucine aminopeptidase; Ppx, peroxiredoxin; FaBP, fatty acid-binding protein; GST, glutathione transferase; MVB, multivesicular body; ESCRT, endosomal sorting complex required for transport; DAMP, damage-associated molecular pattern.

Data access statement: Genomic and transcriptomic data are available from the European Nucleotide Archive under the accession numbers PRJEB6687 and PRJEB6904 respectively. The mass spectrometry proteomics data have been deposited in the ProteomeXchange Consortium via the PRIDE partner repository with the data set identifier PXD002570.
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...the EVs to burst and release the soluble contents held within the EV lumen. The EV membranes were collected by centrifugation at 100,000 x g for 1 h at 4 °C, while the soluble contents of the EV lumen were recovered in the supernatant. The membrane pellet was sequentially extracted with 0.1 M Na2CO3 (pH 11) on ice for 30 min to extract peripheral proteins and the sample centrifuged at 100,000 x g for 1 h at 4 °C. The final pellet was then solubilized in 1% Triton X-100/2% SDS for 15 min at 37 °C to produce integral membrane protein fractions (26).

Mass Spectrometry Analysis of EV and Secretome Fractions—Proteins in the various EV fractions were reduced with 2 mM DTT in 50 mM NH4HCO3 (60 °C/20 min) and alkylated with 5 mM iodoacetamide (room temperature (18–21 °C) in the dark/30 min). Samples were then incubated with 100 ng/μl sequencing grade trypsin (Promega) (37 °C/overnight). The digestions were stopped by the addition of trifluoroacetic acid (TFA) to a final concentration of 0.1% and dried in a vacuum centrifuge. The final mixtures were reconstituted with 10 μl of 0.1% TFA before analysis by LC-MS/MS. Five microliters of the resulting suspension were delivered to an analytical column (Eksigent C18-CL NanoLC Column, 3 μm; 75 μm x 15 cm) equilibrated in 5% acetonitrile/0.1% formic acid (FA). Elution was carried out with a linear gradient of 5–35% buffer B in buffer A for 30 min (buffer A: 0.1% FA; buffer B: acetonitrile, 0.1% FA) at a flow rate of 300 nL/min. Peptides were analyzed in a nanoESI qTOF mass spectrometer (5600 TripleTOF, ABSCIEX) operating in information-dependent acquisition mode, in which a 0.25-s TOF MS scan from 350–1250 m/z, was performed, followed by 0.05-s product ion scans from 100–1500 m/z on the 50 most intense 2–5 charged ions. Peak list files were generated by Protein Pilot v4.5 (Applied Biosystems) using default parameters and exported to Mascot v2.4.1 (Matrix Science) for database searching.

Database Searching—All MS/MS samples were analyzed using Mascot v2.4.1 (Matrix Science). Mascot was set up to search a database comprised of the gene models identified within the F. hepatica genome (version 1.0, 101,780 entries; 11) assuming trypsin digestion with 1 missed cleavage permitted. The F. hepatica gene model sequences can be accessed through WormBase ParaSite (http://parasite.wormbase.org/) under accession PRJEB6687 (genomic read data and gene model transcripts). Mascot was searched with a fragment ion mass tolerance of 0.1 Da and a parent ion tolerance of 0.1 Da. Iodoacetamide derivative of cysteine was specified in Mascot as a fixed modification. Glu->pyro-Glu of the N terminus, Gin->pyro-Glu of the N terminus, deamidation of Asn and Gin and oxidation of Met were specified in Mascot as variable modifications.

Criteria for Protein Identification—Scaffold (version Scaffold 4.3.0, Proteome Software Inc.) was used to validate MS/MS based peptide and protein identifications. Peptide identifications were accepted as they could be established at greater than 95.0% probability by the Peptide Prophet algorithm (27) with Scaffold delta-mass correction. Protein identifications were accepted if they could be established at greater than 95.0% probability and contained at least two identified peptides. Protein probabilities were assigned by the Protein Prophet algorithm (28). Proteins that contained similar peptides and could not be differentiated based on MS/MS analysis alone were grouped to satisfy the principles of parsimony.

Analysis of the F. hepatica Transcriptome and Gene Models—RNA sequencing and gene expression analysis was carried out for multiple lifecycle stages, including metacercariae, NEJ 1, 3 and 24 h postexcretory, juvenile 21-day-old and adult liver flukes, as described by Cwiklinski et al. (11). Briefly, illumina TruSeq RNA libraries were prepared and sequenced, with the resulting RNAseq data being mapped to F. hepatica genome scaffolds and gene models. Differential expression analysis in edgeR was carried out for all genes to which at least five Illumina reads across all libraries could be mapped. Discovery of F. hepatica genes associated with EV machinery was carried out using BLAST analysis against the F. hepatica genome. Briefly, human proteins, inferred from the literature to have roles in exosome biogenesis, cargo sorting or uptake, were used as BLAST queries to identify homologs in closely-related trematode species (NCBI; trematoda). Both the human proteins and trematode homologs were used as BLAST queries to interrogate the F. hepatica genome (NCBI v2.2.29). Each match was manually validated and confirmed using InterPro (http://www.ebi.ac.uk/interpro/) which detects the presence of conserved protein domains. Expression of these identified genes across the F. hepatica lifecycle was investigated using the differential expression analysis of all the gene models identified within the genome. Hierarchical clustering using Gene Cluster 3.0 (29) was used to group differentially expressed genes by similarity of expression, graphically represented using heat-maps.

Analysis of Peptidase Activity in EVs—The soluble contents of F. hepatica EVs recovered following the 15,000 x g and 120,000 x g centrifugation of fluke culture media were isolated by osmotic shock. Briefly, the vesicle pellets were resuspended in hypotonic buffer (20 mM HEPES-KOH; pH 7.3) and placed on ice for 30 min with intermittent vortexing followed by sonication (5 x 20s at 20% power). The vesicle membranes were then removed by centrifugation at 20,000 x g for 1 h at 4 °C. Auto-activation of the native procathepsin L was carried out by incubating the soluble extract from the 15,000 x g vesicle pellet (50 μg total protein) at 37 °C in 100 mM citrate phosphate buffer, pH 4.5, containing 1 mM DTT. Aliquots (15 μl) were removed at time intervals and added to tubes containing 1 μl of 1 mM E-64 to stop the reaction. Proteolytic cleavage of the procsegment was visualized by immunoblotting as described above. Cathepsin L activity was also monitored in the presence of the fluorogenic peptide substrate Z-Leu-Arg-NHMec (20 μM) by measuring the release of fluorescence over time using a Polarstar Omega micro-plate reader (BMG Labtech) in 96-well fluorescent plates as previously described (30). Cathepsin B activity was measured by incubating the soluble vesicular extracts with Z-Arg-Arg-NHMec (10 μM) in 100 mM citrate phosphate buffer (pH 6.5) containing 1 mM DTT and 0.01% Triton X-100. E-64 (10 μM) was used to inhibit cathepsin L and B activity. Leucine aminopeptidase activity was measured by incubating the vesicular extracts with the substrate H-Leu-NHMec (10 μM) in 100 mM Tris-HCI buffer (pH 7.5) containing 1 mM MnCl2 and 0.01% Triton X-100. Bestatin (50 μM), a broad-spectrum aminopeptidase inhibitor (31) was used to demonstrate inhibition of the reaction.

Transmission Electron Microscopy—Transmission electron microscopy (TEM) was performed on adult flukes as previously described (32). Briefly, adult flukes were fixed for 1 h in 2% double-distilled glutaraldehyde (DDG; Agar Scientific) in 0.1 M sodium cacodylate buffer (pH 7.2) containing 3% sucrose at 4 °C. After washing in buffer, specimens were dehydrated in ethanol and embedded in Agar 100 resin (Agar Scientific, Stansted, UK). Ultrathin sections (60–70 nm) were cut on a Reichert Ultracut E ultramicrotome and collected on specimen were dehydrated in ethanol and embedded in Agar 100 resin (Agar Scientific, Stansted, UK). Ultrathin sections (60–70 nm) were cut on a Reichert Ultracut E ultramicrotome and collected on...
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RESULTS

Profiling the Total Secretome of Adult F. hepatica—Taking advantage of our recently reported F. hepatica genome and transcriptome data sets for the major intramammalian lifecycle stages (newly excysted juveniles, NEJs; 21-day old immature liver stage juveniles, and adult flukes) (11), we were able to perform a definitive characterization of the secretions of adult F. hepatica. Sixty-nine proteins secreted by adult F. hepatica were identified in this analysis (supplemental Tables S1 and S2). In accordance with previous transcriptomic analyses (10, 11, 33) peptidases were highly represented (12 matches), including Pro-Xaa carboxypeptidases, cathepsin L1 and L2 clade members, dipeptidylpeptidase III, cathepsin A, legumain 2, metalloendopeptidases, and a cathepsin B. Inhibitors of serine peptidases, serpin, and cysteine peptidases, cystatin 1, were also identified. Another well represented functional group included molecules involved in defense against immune effector cells (10 matches), comprising helminth defense molecule (HDM), GST-mu and -sigma classes, fatty acid binding protein (FABP Fh 2), thioredoxin and thioredoxin-glutathione reductase. Other functional groups included membrane structure (12 matches), receptors/transporters (five matches), enzymes (seven matches), metabolism (six matches), and 14 hits corresponded to proteins with various/unknown functions. A large proportion of the identified proteins (30%) possessed a predicted N-terminal signal peptide, including the majority of the peptidases. Of the remaining proteins, 57% did not contain a predicted signal peptide, whereas 13% could not be predicted because of 5’ truncation of the transcripts.

Adult Fluke Culture Supernatants Contain Distinct Subpopulations of EVs—Following ultracentrifugation, a homogeneous group of exosome-like EVs (post-120,000 × g spin) 30–100 nm in diameter could be isolated from F. hepatica culture media as confirmed by transmission electron microscopy (supplemental Fig. S1A, and 12). However, in the present study we also recovered a pellet following the 15,000 × g centrifugation step that had been previously overlooked. Transmission electron microscopy also revealed that this pellet contained membrane-bound vesicles 50–200 nm in diameter (supplemental Fig. S1B–S1E). We performed a protein assay to determine the relative contributions of the 15 K pellet, the 120 K pellet, and the final post-120 K supernatant to the total protein secreted by adult flukes. We determined that adult F. hepatica secretes total protein at a rate of ~17.3 μg/fluke/hr (100%) of which 0.6 μg/fluke/h (3.6%) is associated with the 120 K pellet, 1.5 μg/fluke/hr (8.6%) is associated with the 15 K pellet whereas the majority 15.2 μg/fluke/hr (87.8%) remains in solution following the 120 K spin. The complexity of each fraction, as determined by 1D gel electrophoresis, can be observed in Fig. 1A. Each sample is defined by a distinct protein profile indicating a compartmentalization of the total secretome that can be separated by centrifugation.

To investigate these fractions further, each fraction (5 μg total protein) was probed with specific antisera available in our laboratory against a range of secreted Fasciola secreted molecules (Fig. 1B). Immunoblotting showed that bands corresponding to helminth defense molecule (FhHDM; 6 kDa) were detected across all samples with the most intense HDM staining observed in the 15 K pellet. We found the F. hepatica leucine aminopeptidase (FhLAP; 55 kDa) and peroxiredoxin (FhPrx; 24 kDa) in both the 15 K and 120 K vesicle pellets but not in the total secretome or the 120 K supernatant. F. hepatica cathepsin L1 (FhCL1), which is synthesized as an inactive 37 kDa zymogen (9), was revealed as an intense reactive band in the 15 K vesicle pellet, but was barely visible by comparison in the 120 K pellet. Although the 37 kDa zymogen was not observed in the total secretome and final supernatant, a processed mature 24 kDa cathepsin L containing the catalytic domain was.

To confirm the above data, we used fluorogenic peptide substrates to determine the relative peptidase activities associated with the vesicle subpopulations (i.e. 15 K versus 120 K pellets). Although leucine aminopeptidase activity, measured using the substrate H-Leu-NHMec, was not detected in soluble extracts from either pellet, strong H-Leu-NHMec hydrolysis was observed when the membranes from both the 15 K and 120 K pellets were incubated with the substrate. The activity associated with membranes from the 120K pellet was fourfold higher than that associated with the 15 K pellet membranes in accordance with Acosta et al. (34). In all assays, this leucine aminopeptidase activity was inhibited by bestatin, a broad-spectrum aminopeptidase inhibitor (data not shown).

Equal amounts of total protein from each vesicle sample were incubated with diagnostic substrates for cathepsin L (Z-Leu-Arg-NHMec) and cathepsin B (Z-Arg-Arg-NHMec) activities. Comparison of the hydrolysis of each fluorogenic substrate within the linear range of the reactions (15 min) by each extract is shown in Fig. 1C. Strong cathepsin activity was detected in both 15 K and 120 K vesicle pellets. However, both cathepsin B- and L-specific activity in extracts of the 120 K pellet were on average 60% less than that observed in extracts of the 15 K pellet. In all assays, cathepsin B and L activity were completely inhibited by 10 μM E-64, a cysteine peptidase inhibitor (data not shown).

The 15 K EVs are Derived from Gastrodermal Cells and Contain Inactive Cathepsin L Zymogens that Can Be Activated at low pH—Our immunoblot analysis of the centrifugation fractions indicated that adult F. hepatica secretes a population of relatively large vesicles (15 K pellet) enriched
for the inactive 37 kDa cathepsin L zymogen (Fig. 1B). To determine whether the native zymogen packaged within EVs could undergo pH-dependent autocatalytic activation similar to our purified recombinant cathepsin Ls (35–37), soluble extracts of the 15 K vesicle pellet were incubated at low pH (pH 4.5) for up to 3 h at 37 °C. Western blot analysis of the in vitro auto-activation process shows that the 37 kDa zymogen diminishes over time, concomitant with the appear-
Fig. 2. Gastrodermal cell-derived EVs contain inactive cathepsin Lzymogens. A, Soluble extracts from the 15K vesicle pellet were incubated in 100 mM citrate phosphate buffer (pH 4.5) for 3 h at 37 °C. Aliquots of the reaction mixtures were removed after 0, 30, 60, 120, and 180 min and halted with 10 μM E-64 on ice. Samples were then transferred to nitrocellulose membranes and probed with an anti-Fasciola cathepsin L1 antibody. The inactive zymogen (37 kDa) showed a progressive decrease in intensity concurrent with the appearance of a band representing the 24 kDa mature enzyme (arrowhead). The rate of formation of an active mature enzyme from the inactive zymogen at pH 4.5 was rapid indicating that autocatalytic activation of the native enzyme occurs readily in an acidic environment (Fig. 2B).

Transmission electron microscopy showed the presence of secretory vesicles densely packed within the specialized gastrodermal cells that line the fluke gut (Fig. 2C). Immunogold labeling using anti-cathepsin L antibodies revealed that these vesicles were intensely, and specifically, stained for Fasciola cathepsin L1 (Fig. 2D). No labeling was observed in any other structures which indicates that the large vesicles of the 15 K pellet that are enriched for the 37 kDa cathepsin L1 zymogen originate from the gastrodermal cells of the fluke gut.

The Proteome of F. hepatica 120 K Exosome-like EVs—A proteomics analysis of the small exosome-like EVs (120 K vesicle pellet) was performed taking advantage of the newly available genome and transcriptome resources for Fasciola hepatica (11), to gain a deeper coverage of the EV proteome than previously reported (12). Here, we performed a sequential treatment to identify differentially located proteins in the EVs. We first employed trypsin shaving (25) to release proteins from the surface of the EVs, before sequentially extracting them to identify proteins associated with the EV membranes as well as soluble proteins packaged within the EV lumen. A total of 180 different EV proteins were identified using this approach and their functional annotation is shown in Tables I and II.

A number of well-known exosomal markers, including Hsp70, ALIX and members of the tetranspin family, were identified during the proteomics analysis. This supports the classification of the vesicles derived from the 120K pellet as “exosome-like.” Peptidases were highly represented in the EV proteome. Peptides corresponding to legumain 4/5, cathepsin A, cathepsin B6/8, and a Pro-Xaa carboxypeptidase were released following trypsin shaving of the EV surface. Other proteases, such as leucine aminopeptidase, various subunits of the Fasciola proteasome and several cathepsin Bs were either found associated with the EV membrane (present in carbonate washes), or within the EV lumen (present in the soluble extract).

Several membrane channels, transporters and other molecules associated with membrane structure were identified. These included the ATPase pumps ABCB1/p-glycoprotein, Na+/K+ ATPase, and a phospholipid-transporting ATPase (flippase). Membrane transporters such as anoctamin (a calcium-activated chloride channel), Niemann-Pick C1, and C2 proteins (cholesterol transporters), an amino acid permease, an inositol transporter and a glucose transporter were also found. Other proteins with potential roles in membrane structure included various annexins and the Fasciola T1 tegumental antigen (38). In addition to components of the cytoskeleton, peptides corresponding to several metabolic enzymes were released following trypsin shaving of the EV.
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| Proteins associated with the membrane of EVs from adult *F. hepatica*. Figures represent number of unique peptides |
|---------------------------------------------------------------|
| **Protein Identifier** | **Extract** | **Trypsin** | **NaHCO₃** | **TX100/SDS** |
|------------------------|-------------|-------------|-------------|---------------|
| **Proteases & inhibitors** | | | | |
| Legumain 4/5 | BN1106_s1861B000097 | 7 | 12 | 0 |
| Cathepsin B | BN1106_s1772B000188 | 0 | 13 | 0 |
| Cathepsin A | BN1106_s1241B000264 | 3 | 7 | 0 |
| Cathepsin B6/8 | BN1106_s793B000177 | 2 | 5 | 0 |
| Cathepsin B fragment | BN1106_s8462B000006 | 0 | 4 | 0 |
| Cathepsin B4/5/7 | BN1106_s13444B000002 | 0 | 6 | 0 |
| Cathepsin L1 | BN1106_s8490B000026 | 0 | 5 | 0 |
| Lyosomal Pro-X carboxypeptidase | BN1106_s1620B000120 | 0 | 4 | 0 |
| M17 Leucine aminopeptidase 2 | BN1106_s6178B000566 | 0 | 6 | 0 |
| Protease subunit alpha 6 | BN1106_s1579B000119 | 0 | 4 | 0 |
| Protease subunit alpha type | BN1106_s4981B000066 | 0 | 6 | 0 |
| Protease subunit alpha type | BN1106_s3658B0000104 | 0 | 4 | 0 |
| Lysozyme pro-Xaa carboxypeptidase | BN1106_s8490B000026 | 0 | 5 | 0 |
| Lysosomal pro-Xaa carboxypeptidase | BN1106_s3516B000032 | 0 | 3 | 0 |
| Protease subunit alpha 6 | BN1106_s1259B000205 | 0 | 3 | 0 |
| M17 Leucine aminopeptidase 1 | BN1106_s7079B000034 | 0 | 2 | 0 |
| Protease subunit beta 2 | BN1106_s2284B000154 | 0 | 4 | 0 |
| Protease subunit alpha type | BN1106_s1628B000138 | 0 | 2 | 0 |
| **Pumps & transporters** | | | | |
| ABCB1 | BN1106_s274B0000296 | 4 | 2 | 3 |
| MDR p-glycoprotein 1 | BN1106_s2471B000098 | 0 | 0 | 3 |
| Glucose-transporter | BN1106_s584B0000350 | 0 | 0 | 2 |
| Phospholipid-translocating ATPase IIB | BN1106_s435B0000302 | 0 | 0 | 3 |
| ATP synthase subunit alpha | BN1106_s4332B000087 | 0 | 2 | 0 |
| **Metabolism** | | | | |
| Propionyl-CoA carboxylase alpha | BN1106_s1252B0000359 | 17 | 3 | 0 |
| Glutamate dehydrogenase 1 NAD(P)+ | BN1106_s5768B000030 | 5 | 6 | 0 |
| Aldolase | BN1106_s446B0000665 | 2 | 2 | 0 |
| GAPDH | BN1106_s5174B000030 | 5 | 0 | 0 |
| Glutamine synthetase | BN1106_s2400B000186 | 3 | 3 | 0 |
| Propionyl-CoA carboxylase beta | BN1106_s1551B0000468 | 0 | 3 | 0 |
| Malic enzyme | BN1106_s333B0000362 | 6 | 0 | 0 |
| Hexokinase A | BN1106_s175B000070 | 0 | 0 | 2 |
| Retinol dehydrogenase | BN1106_s2277B000049 | 0 | 2 | 0 |
| Malate dehydrogenase | BN1106_s1459B0000183 | 3 | 0 | 0 |
| **Cytoskeleton** | | | | |
| Actin 5C | BN1106_s2007B000133 | 11 | 4 | 0 |
| Ezrin | BN1106_s1300B000145 | 2 | 0 | 2 |
| Fimbrin | BN1106_s1403B000129 | 7 | 2 | 0 |
| Alpha actinin | BN1106_s4069B000247 | 4 | 0 | 0 |
| Gelsolin | BN1106_s2349B000188 | 3 | 2 | 0 |
| Beta tubulin 1 | BN1106_s55B000372 | 3 | 0 | 0 |
| **Exosome biogenesis / vesicle trafficking** | | | | |
| Ubiquitin-60S ribosomal protein L40 | BN1106_s5657B000103 | 2 | 0 | 2 |
| Hsp70 | BN1106_s309B0000234 | 9 | 2 | 3 |
| Chaperonin containing tcp1 | BN1106_s1242B000159 | 2 | 0 | 0 |
| 14–3–3 protein | BN1106_s3904B000042 | 0 | 0 | 2 |
| Annexin | BN1106_s819B0000364 | 8 | 4 | 0 |
| Tetraspanin-CD63 receptor | BN1106_s1657B000161 | 0 | 5 | 4 |
| Acid sphingomyelinase | BN1106_s1285B000159 | 2 | 6 | 0 |
| Annexin | BN1106_s945B0000218 | 2 | 9 | 0 |
| Myoferlin | BN1106_s3585B000136 | 0 | 0 | 11 |
| Tetraspanin-1 | BN1106_s1657B000161 | 0 | 5 | 4 |
| ALIX | BN1106_s2963B000136 | 0 | 0 | 6 |
| IST1 | BN1106_s3747B000112 | 2 | 0 | 7 |
| Otoferlin | BN1106_s2361B000048 | 0 | 0 | 7 |
| Vacular assembly/sorting protein D1D2-like protein | BN1106_s2316B000077 | 0 | 0 | 5 |
| ALIX | BN1106_s1871B0000313 | 0 | 0 | 4 |
| Charged multivesicular body protein | BN1106_s2655B0000264 | 0 | 0 | 4 |
| Vacular protein sorting-associated protein 4B | BN1106_s1437B0000141 | 0 | 0 | 4 |
surface. These included propionyl-CoA carboxylase, glutamate dehydrogenase, glutamine synthase, malic enzyme, and malate dehydrogenase and GAPDH whereas other enzymes were found associated with the EV membrane or within the EV lumen. A small number of host-derived proteins were also identified including immunoglobulins, casein, and acylaminoacid-releasing enzyme (supplemental Table S3).

Comparison of our data sets for adult fluke total secretome and the exosome-like EVs with the tegumental proteome reported by Wilson et al. (6) revealed that there is an overlap in protein expression between these samples (Fig. 3). Of the 180 proteins identified in the adult fluke EVs, 35 and 22 proteins were shared with the total secretome and tegument, respectively. A total of nine proteins were shared by the total secretome and tegument, whereas only nine proteins were found in all three samples. supplemental Fig. 2 shows the composition of the total secretome and EV proteome data sets grouped according to function.

Transcriptome Profiling Reveals that Fasciola EV Biogenesis Pathways are Constitutively Expressed While Cargo Molecules are Developmentally Regulated—A significant number of the proteins identified have roles in the endosomal pathway, or in EV biogenesis, and provide insight into the mechanism used by F. hepatica to generate EVs. These include various components of the ESCRT-dependent exosome biogenesis pathway (e.g. a number of charged multivesicular body proteins, the AAA-ATPase Vps4, IST1, syntenin and ALIX), and the ceramide-dependent pathway (e.g. sphingomyelinase). Several members of the Rab GTPase family, such as Rab8a, Rab11b, and Rab27 that have roles in EV biogenesis and release were also found. Also identified were SNAREs (synaptotagmin and VAMP7) as well as myoferlin and otoferlin that mediate vesicle trafficking and fusion. A cartoon summarizing the proteome of the exosome-like EVs from adult fluke is shown in Fig. 4 and the full mass spectrometry report is shown in supplemental Table S3.

To gain insight into the putative mechanism(s) of EV biogenesis in F. hepatica we conducted an extensive search of the literature for proteins implicated in this process via functional studies, largely in mammalian cells. A total of 102 pro-
### TABLE II
Top 50 proteins identified within the lumen of EVs released by adult *F. hepatica*

| Protein ID Identifier | Unique Peptides |
|-----------------------|-----------------|
| **Proteases**         |                 |
| Legumain 4/5           | BN1106_s1861B000097  | 14 |
| Cathepsin A            | BN1106_s1241B0000264 | 11 |
| Cathepsin B            | BN1106_s1772B000188  | 8  |
| M17 Leucine aminopeptidase 2 | BN1106_s617B0000566 | 8  |
| Cathepsin B6/8         | BN1106_s733B000177  | 7  |
| Cathepsin L1           | BN1106_s6490B000026 | 7  |
| 20S proteasome subunit alpha 6 | BN1106_s1579B0000119 | 7  |
| Cathepsin B4/5/7       | BN1106_s1344B000002 | 6  |
| Lysosomal Pro-X carboxypeptidase | BN1106_s1620B000120 | 6  |
| Proteasome subunit beta type-5 | BN1106_s1639B0000051 | 6  |
| 20S proteasome subunit alpha 3 | BN1106_s7079B000034 | 5  |
| M17 Leucine aminopeptidase 1 | BN1106_s7079B0000034 | 5  |
| **Membrane structure** |                 |
| Annexin                | BN1106_s819B0000364 | 17 |
| Myoferlin              | BN1106_s3585B000136 | 14 |
| Tetraspanin-CD63 receptor | BN1106_s1657B000161 | 8  |
| Annexin                | BN1106_s945B000218  | 8  |
| Tetraspanin-1          | BN1106_s915B000136  | 8  |
| Otoferlin              | BN1106_s3261B000048 | 8  |
| T-cell immunomodulatory protein | BN1106_s92B0000560 | 7  |
| Annexin                | BN1106_s3266B000046 | 6  |
| **EV biogenesis**      |                 |
| ALIX                   | BN1106_s2963B000136 | 9  |
| Acid sphingomyelinase  | BN1106_s1285B000159 | 8  |
| IST1                   | BN1106_s3747B000012 | 7  |
| Vacuolar protein sorting-associated protein 4B | BN1106_s1437B000014 | 7  |
| ALIX                   | BN1106_s1871B0000313 | 6  |
| Rab8                   | BN1106_s2588B000276  | 6  |
| Rab11b                 | BN1106_s844B0000259 | 6  |
| **Channels and transporters** |                 |
| ABCB1                  | BN1106_s274B0000296 | 30 |
| Na⁺/K⁺ transporting ATPase subunit alpha | BN1106_s521B0000167 | 12 |
| Niemann-Pick disease type C1 protein | BN1106_s1498B000025 | 10 |
| MDR p-glycoprotein 1   | BN1106_s2471B000098 | 8  |
| vATPase subunit I      | BN1106_s1877B000008 | 8  |
| Inositol transporter   | BN1106_s3611B000052 | 7  |
| **Enzymes**            |                 |
| Phosphodiesterase-nucleotide pyrophosphatase | BN1106_s1985B000403 | 9  |
| Aldolase               | BN1106_s446B0000065 | 9  |
| Peroxiredoxin          | BN1106_s1614B000280 | 5  |
| Enolase                | BN1106_s3227B0000227 | 8  |
| Glutamate dehydrogenase 1 NAD(P)⁺ | BN1106_s5767B000030 | 7  |
| Beta-1,4-galactosyltransferase 2 | BN1106_s741B0000214 | 6  |
| **Uncharacterized/other** |                 |
| Major vault protein    | BN1106_s7273B000042 | 15 |
| Uncharacterized 5C     | BN1106_s2907B000133 | 10 |
| Uncharacterized        | BN1106_s3001B000132 | 9  |
| DM9 domain-containing protein | BN1106_s5689B000026 | 7  |
| Uncharacterized        | BN1106_s216B000184  | 7  |
| Uncharacterized        | BN1106_s6821B000024 | 6  |
| Ferritin               | BN1106_s3950B000041 | 6  |
| Ezrin                  | BN1106_s1300B000145 | 6  |
| 14–3-3 protein         | BN1106_s3904B000042 | 5  |
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Fig. 3. Expression of proteins in the tegument, total secretome and EVs of adult F. hepatica. Venn diagram comparing the proteins identified in the total secretome and extracellular vesicles (EVs) released by adult F. hepatica (this study), and those found in tegmental fractions (tegument) by Wilson et al. (6).

...teins were identified and categorized according to their role in vesicle formation including: components of the endosomal sorting complexes required for transport (ESCRT) pathway, components of the ceramide pathway, roles in EV release (Rabs and SNARES), RNA/protein/lipid sorting and membrane fusion/repair (supplemental Table 4). These proteins were used as queries to interrogate the newly available F. hepatica genome (11). All components of the ESCRT machinery, except for the charged multivesicular body protein 6 (CHMP6) from the ESCRT-III complex, were present in the F. hepatica genome. The ESCRT pathway was also well represented at the proteome level with up to ten putative members identified during the MS analysis of adult fluke exosome-like EVs; most of them were components of the ESCRT-III and Vps4 complexes that have roles in multivesicular body (MVB) formation and the final membrane budding/abscission steps during the latter stages of EV biogenesis (39). BLAST results also showed that many components of ESCRT-independent pathways for exosome release are present in the fluke genome but less widely represented at the proteome level. The most significant matches of the MS data include acid sphingomyelinase and tetraspanin CD63, that are both involved in exosome biogenesis in mammalian cells.

Having identified putative homologs of the EV biogenesis pathway members in F. hepatica, we then examined their relative expression levels across the intramammalian lifecycle stages of F. hepatica. Transcriptome data sets for Fasciola NEJs at 1 h, 3 h, and 24 h time-points postexcystment were used as well as 21-day old immature liver-stage flukes and adult flukes from the bile duct (11). Relative transcript expression levels (fold change on a log2 scale) are shown as a heat-map in Fig. 5. Taking this broad view, it is apparent that, with a few exceptions, putative EV biogenesis pathway members in F. hepatica are constitutively expressed (albeit at low levels), and do not differ greatly in expression levels as the parasite migrates from the small intestine, through the liver and into the bile ducts. However, when we performed the same analysis of the transcripts encoding the remaining proteins identified during our proteomics analysis of adult fluke exosome-like EVs (i.e. those not implicated in EV biogenesis and thus termed putative cargo molecules), we found that most were expressed at higher levels and were developmentally regulated. As can be seen in the heat-map shown in Fig. 6, the transcripts can be broadly segregated into two groups: (1) those with low levels of expression in the NEJ stage that increase in the 21-day old and adult flukes, and (2) those with high levels of expression in the NEJ stage that decrease in the 21-day old and adult flukes. Group one members included several proteases such as legumain 4/5, Pro-Xaa carboxypeptidase, cathepsin B4/5/7 and members of cathepsin L clades 1, 2, and 5. Others included saposin-like and CD59-like proteins, a calcium-binding protein, and a range of highly expressed, but uncharacterized, proteins. Notable group two members included thioredoxin, peroxiredoxin, cathepsin B2, various protease subunits and metabolic enzymes such as enolase, malate dehydrogenase, and propionyl CoA carboxylase.

DISCUSSION

The recent availability of new -omics resources for F. hepatica permitted us to conduct the most comprehensive analysis of any helminth secretome to date. We have discovered that adult F. hepatica secretions are more complex than previously thought, and contain at least two subpopulations of EVs that differ according to size, cargo and mechanism of release from the parasite. In doing so we have challenged the classic view that soluble proteins are the sole mediators of the host-parasite interaction and identified new mechanisms by which parasite-derived molecules can interact with host cells. These mechanisms can be classified under various headings: (1) Extracellular interaction with host macromolecules. In terms of protein content, soluble molecules made the greatest contribution to the adult fluke total secretome (87.8%). This fraction is dominated by members of the cathepsin L cysteine peptidase family which represent >80% of the total protein secreted (8–10). Fasciola cathepsin Ls perform a variety of roles, mostly associated with extracellular degradation of host macromolecules such as collagen to aid parasite migration through host tissues (36, 37), hemoglobin as a source of nutrition (30) and immunoglobulins to avoid expulsion by the host immune system (40). The considerable expenditure devoted to secretion of soluble proteins by F. hepatica clearly shows that these tasks are critical for completion of the parasite’s life-cycle. (2) Endocytosis of soluble parasite molecules. We have shown that cathepsin L1 and helminth defense molecule (HDM), which are both present as soluble proteins within F. hepatica secretions, are internalized via the endolysosomal pathway and modulate host macrophage function by degradation of toll-like receptor 3 and inhibition of vacuolar-ATPase respectively (26, 41). The receptors required for uptake of these molecules are currently unknown however...
HDM binds to cholesterol within lipid rafts on the cell surface that may initiate internalization (26). Internalization of parasite EVs. Although we estimate that the 15 K and 120 K EV pellets hold only 8.6% and 3.6% of the total protein secreted by adult *F. hepatica*, respectively, they contain potent immunomodulators/virulence factors (see below) that would exert greater effects on host cell function than their abundance might suggest. EVs secreted by adult flukes will drain, via the bile ducts, into the host gut and we have shown that fluke exosome-like EVs are internalized by host intestinal epithelial cells (IECs) *in vitro* (12). Although traditionally regarded as a mere physical barrier to infection, IECs are now known to express pattern-recognition receptors that enable them to detect a range of microbial ligands and transmit signals to underlying gut-associated lymphoid tissues (42). Thus, parasite-derived EVs could maintain a systemic Th2/regulatory environment that is permissive to fluke survival and reproduction via modulation of IEC immune surveillance.

We, and others, have shown that several proteins released by *F. hepatica* have the ability to modulate host immune cell function (43). In this study we have found the majority of these immunomodulators are packaged within the exosome-like EVs secreted by adult fluke. Although *F. hepatica* peroxiredoxin (Prx) actively drives host Th2 immune responses via stimulation of M2 macrophages (44, 45), most of the other fluke immunomodulators identified so far (despite displaying varied structures and biochemical activities), indirectly promote Th2 responses by suppressing the development of Th1/Th17-associated inflammation. These include *Fasciola* helminth defense molecule (HDM), cathepsin L1 (FhCL1), fatty acid-binding protein (FABP), kunitz-type molecule (Fh-KTM) and Sigma class glutathione transferase (GST) (23, 41, 46–48). The role of *F. hepatica* Prx in switching the host immune response to a Th2 phenotype during the early stages of infection has been well documented (44, 45). This is in agreement with our transcriptome profiling of EV cargo molecules (Group 2, Fig. 6). However, our transcriptomics and proteomics profiling experiments have shown that the remaining immunomodulators (FhCL1, HDM, FABP, Fh-KTM, and GST sigma) are constitutively expressed across the intramamalian life-cycle stages of the fluke and are actually up-regulated as the flukes migrate through the liver, enter the bile ducts and mature into adult worms (10,11; Group 1, Fig. 6). These observations suggest that the latter molecules act to maintain the Th2/regulatory phenotype seen during chronic infection. Thus, EV cargo molecules, as well as components of the soluble secretome of adult *F. hepatica* (10) are developmentally regulated to facilitate the parasites migration through host tissue and to counteract host immune attack. Work is ongoing in our laboratory to characterize the EVs secreted by the NEJ stage.

We have also identified potential immunomodulators not previously thought to be secreted by adult *F. hepatica*, the most notable being members of the cathepsin B family. Several cathepsin B transcripts were previously identified in adult flukes, but these enzymes were not detected in the adult fluke secretome and therefore it was suggested that they have intracellular house-keeping roles (10). However, we have found that the small exosome-like EVs contain at least four developmentally regulated cathepsin B peptidases. This finding establishes that both cathepsin L and B peptidases are secreted by adult flukes, albeit via different mechanisms, the former via the gut and the latter via the tegument. Studies in *Schistosoma mansoni* have shown that a cathepsin B,
SmCB2, is located in the tegument of the adult parasites (49). Also identified, within the EV lumen, were a number of leaderless proteins (including Prx, heat shock proteins and annexins) that are homologs of host damage-associated molecular pattern molecules (DAMPs). DAMPs are leaderless cytosolic proteins that are released into the extracellular environment via atypical secretory pathways in response to tissue damage (50). They interact with pattern recognition receptors expressed by immune cells and thus set off an “alarm” system which often leads to a wound-healing scenario (50). Because of their abundance within helminth secretions, we previously speculated that DAMP homologs can direct the

Fig. 5. Expression of transcripts encoding putative EV biogenesis pathway members across the intramamalian lifecycle stages of *F. hepatica*. Heatmaps are colored on a log2 scale with up-regulation in red and down-regulation in blue relative to metacercariae, depicting log fold change between developmental stages, grouped by hierarchical clustering. The *F. hepatica* life-cycle stages analyzed are: newly excysted juveniles (NEJ; 1-, 3- and 24-h postexcystment), 21-day old immature liver-stage flukes (Juv_21d), and adult flukes from the bile duct.
host immune response (51) and their packaging into EVs provides an alternative route for their non-canonical secretion. Although the mechanism for targeting cytosolic proteins into EVs is not fully understood, ATP-binding cassette (ABC) transporters are responsible for packaging some leaderless proteins (e.g., FGF1, IL-1β, and Hsp70) into EVs destined for secretion (52–54). We identified two ABC transporters associated with the *F. hepatica* EV membrane.

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**Fig. 6. Expression of transcripts encoding putative EV cargo proteins across the intramammalian life cycle stages of *F. hepatica*.** Heatmaps are colored on a log2 scale with up-regulation in red and down-regulation in blue relative to metacercariae, depicting log fold change between developmental stages, grouped by hierarchical clustering. The *F. hepatica* life cycle stages analyzed are: newly excysted juveniles (NEJ, 1-, 3- and 24-h postexcystment), 21-day-old immature liver-stage flukes (Juv_21d), and adult flukes from the bile duct.
(ABCB1 and p-glycoprotein 1) that may perform similar packaging roles in the parasite.

The recently available *F. hepatica* genome (11) has allowed us, for the first time, to identify pathways involved in EV biogenesis in a helminth parasite. Exosomes arise from the endosomal system and are formed by inward budding of the multi-vesicular body (MVB) membrane that allows capture of cytoplasmic cargo. This step is thought to occur via two different routes. Endosomal sorting complexes required for transport (ESCRT) 0, I, II, and III, together with syntenin-ALIX complexes, capture ubiquitinated proteins, whereas in the ESCRT-independent pathway tetraspanins (e.g. CD63) and various lipid-related enzymes (notably sphingomyelinase) can also drive vesicle formation (55, 56). The final abscission of the budding exosomes into the MVB lumen requires the AAA-ATPase VPS4 (57). The exosomes are then secreted into the extracellular space following the fusion of the MVB with the plasma membrane, which is dependent on small GTPases including Rab8b, Rab11, Rab27a, and Rab27b and the Rab effectors otoferlin and synaptotagmin-like protein (58–61).

We identified *F. hepatica* homologs of all these key regulators of exosome biogenesis during our proteomics analysis of the 120 K EV pellet (summarized in Fig. 4). However, it is significant that although the ESCRT pathway was well represented at the proteome level (10 members were identified in the MS analysis), the only representatives of the lipid-related/ESCRT-independent pathway were sphingomyelinase and tetraspanin CD63 (supplemental Table S4). Based on these findings we propose that exosome biogenesis in adult fluke primarily occurs via ESCRT-independent MVB formation in the tegumental syncytium with the exosomes being shed from the apical plasma membrane (although functional studies such as RNAi are needed to confirm this). In support of this proposal, MVBs have been described during ultrastructural examination of the tegumental syncytia of *F. hepatica* and the related trematode *E. caproni* (6, 12). Furthermore, our comparative proteomics analysis revealed that 42% of the proteins identified in the *F. hepatica* tegumental extracts by Wilson et al. (6) were also packaged into the 120 K EVs; these contained a variety of proteins (e.g. tegumental antigen 1, DM9-containing protein, annexin, and tetraspanin) that localize primarily within the tegument of *F. hepatica* and related trematodes (6, 38, 62–64). Determining the specific roles of individual pathway members during exosome biogenesis in liver fluke needs further investigation, perhaps by RNAi-mediated gene silencing which is functional in this parasite (65).

Targeting key regulators of EV biogenesis (including sphingomyelinase, ALIX and Rab GTPases) using chemical inhibitors or RNAi has been shown to significantly reduce the number of EVs released by mammalian cells. Indeed, suppressing EV production to inhibit their effect in disease is an emerging therapeutic strategy that has yielded impressive results, notably in cancer therapy (reviewed by 14). We have identified *F. hepatica* homologs of several key regulators of EV biogenesis that could be targeted for parasite control if specific inhibitors can be found. Although technically challenging, selective inhibition of parasite EV biogenesis would prevent the delivery of a range of immunomodulators to host cells and, conceivably, tip the host immune balance in favor of Th1-driven parasite elimination. An alternative anti-parasite control strategy would be to block the uptake of parasite-derived EVs by host cells. Although direct fusion of EVs with the plasma membrane of target cells has been reported (66), uptake mechanisms akin to receptor-mediated endocytosis (dependent on the complement of proteins expressed on the vesicle surface) appear to be the dominant route of entry into recipient cells (67). Many of the proteins we identified on the external surface of *F. hepatica* EVs are metabolic enzymes, or other housekeeping proteins, with reported “moonlighting” roles in other helminths and microbial pathogens (68). These include malic enzyme, enolase, GAPDH, aldolase and annexin which act as parasite adhesion molecules via interaction with plasminogen and/or fibronectin on the host cell surface (69–74) and could be targets for blocking antibodies. Recently, it was shown that antibodies against *O. viverrini* tetraspanin blocked uptake of parasite EVs by host cholangiocytes, providing proof-of-concept, at least mechanistically, for this approach as an anti-parasite therapy (75).

In this report we have performed the first comparative analysis of the soluble and vesicular secretions of a helminth parasite. This was made possible by integrating the recently available genome, transcriptome, and proteome data sets for this pathogen (11). Our study underscores the importance of investigating the role of EVs during acute and chronic fluke infection and presents new opportunities to disrupt EV biogenesis or interaction with target cells toward parasite control.

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* This article contains supplemental Figs. S1, S2 and Tables S1 to S4.

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