Presence of diabetes autoantigens in extracellular vesicles derived from human islets

Craig P. Hasilo1,2,5, Sarita Negi1,2,5, Isabelle Allaëys3, Nathalie Cloutier3, Alissa K. Rutman1,2,5, Marco Gasparrini1,2, Éric Bonneil4, Pierre Thibault4,5, Éric Boilard3,5 & Steven Paraskevas1,2,5

Beta-cell (β-cell) injury is the hallmark of autoimmune diabetes. However, the mechanisms by which autoreactive responses are generated in susceptible individuals are not well understood. Extracellular vesicles (EV) are produced by mammalian cells under normal and stressed physiological states. They are an important part of cellular communication, and may serve a role in antigen processing and presentation. We hypothesized that isolated human islets in culture produce EV that contain diabetes autoantigens (DAA) from these otherwise normal, non-diabetic donors. Here we report the caspase-independent production of EV by human islets in culture, and the characterization of DAA glutamic acid decarboxylase 65 (GAD65) and zinc transporter 8 (ZnT8), as well as the β-cell resident glucose transporter 2 (Glut2), present within the EV.

An underlying hallmark of type 1 diabetes (T1DM) is the direct injury to the insulin-producing β-cells within the islets of Langerhans. Associated with a break in self-tolerance in these individuals, is the development of diabetes-associated autoantibodies (AAb). The major AAb found to correlate with diabetes progression (at least one AAb is detected in 90% of patients1) include those against the 65 kDa form of glutamic acid decarboxylase (GAD65)2, the zinc transporter 8 (ZnT8)3, the tyrosine phosphatase islet antigen-2 (IA-2)4, and insulin/proinsulin5. The detection of at least two AAb in asymptomatic individuals is defined as presymptomatic normoglycemia, a state which carries 5-year and 10-year risks of 44 and 70%, respectively, in progressing to overt diabetes6. Although the development of insulitis, β-cell injury, and AAb have been extensively studied, the factors or signals which initiate this process are not known.

Extracellular vesicles (EV) are produced by most, if not all cells under various stimuli. It is widely accepted that EV are an important part of cellular communication, and may play a role in the development of immune responses. Indeed, a pathophysiological role for EV have been identified in disease states such as cancer7,8, rheumatoid arthritis9, systemic lupus erythematosi10,11, thrombocytopenia12, as well as neurodegenerative13,14 and vascular disease15,16, to name a few. The cargo payload carried by EV varies greatly depending on their size and origin. Traditionally, EV have been classified according to size, with exosomes (Exo) generally representing EV of 150 nm or less, microvesicles (MV) being 100 to 1,000 nm, and apoptotic bodies (AB) ranging from 1,000 to 5,000 nm and often containing nucleic acids, mitochondrial proteins and other subcellular structures17. More recently, however, it has been illustrated that the more encompassing term EV should be used, as it accurately reflects all vesicle types released by cells18, as Exo up to 250 nm and Exo-like vesicles derived from apoptotic cells have been described19,20. EV may contain neo-antigens (Ag) that, when exposed in times of injury, may act as triggers of autoimmunity13.

EV production in the context of T1DM is only beginning to be appreciated with a few preliminary reports in murine and human contexts. Exo from mouse insulinoma cell lines or islet-derived mesenchymal stem cells have immunostimulatory capacity in non-obese diabetic (NOD) mice, eliciting a response in autoreactive T-cells and marginal zone-like B-cells18-20. Human islets have been evaluated for miRNA content in the context of

1Human Islet Transplant Laboratory, McGill University Health Centre, Montréal, Québec, Canada. 2Research Institute of the McGill University Health Centre, Montréal, Québec, Canada. 3Centre de Recherche en Rhumatologie et Immunologie, Centre de Recherche du Centre Hospitalier Universitaire de Québec, Faculté de Médecine de l’Université Laval, Québec, Québec, Canada. 4Institut de Recherche en Immunologie et en Cancérologie, Université de Montréal, Montréal, Québec, Canada. 5Canadian National Transplant Research Program, Edmonton, Alberta, Canada. Correspondence and requests for materials should be addressed to S.P. (email: steven.paraskevas@mcgill.ca)
islet-endothelial cell interactions, and produce GAD65, IA-2 and proinsulin containing-Exo, following cytokine exposure.

Our study focused on determining whether human islet-derived EV harbor known diabetes autoantigens (DAA). The possibility exists that some β-cell components not normally externalized may be released following sub-lethal injury. Here we report the production of EV by normal, nondiabetic human islets following isolation, their characterization using flow cytometry and proteomic-based approaches, and the observation of T1DM autoantigens in these extracellular particles (Fig. 1).

Results

**Human islets produce a heterogeneous EV population in culture.** To evaluate if our human islet preparations were producing EV in culture, we first used Nanoparticle Tracking Analysis (NTA) to quantify the release of particles by islet preparations. Since it has been suggested that culture conditions may result in varying levels of EV secreted into the media, we first investigated EV elution after 24 hours. Particle tracking by NTA revealed concentrations that varied between donor islet preparations. Generally, the islet preparations produced between $4.53 \pm 1.49 \times 10^6$ to $5.61 \pm 0.88 \times 10^7$ particles/µl (Fig. 2a), with a mean size of $193 \pm 21$ nm (Fig. 2b).

To further characterize EV and to confirm their membrane moiety, we analyzed islet conditioned media (ICM) from each islet preparation by flow cytometry, Annexin V (AnnV) and CellTracker Deep Red (CT)
were respectively chosen as markers of membrane-phospholipid moiety (phosphatidylserine, PS) and of cytosol content, as in previous studies. Similar variability was observed by flow cytometry analyses, as EV were detected in quantities ranging from $70.77 \pm 44.86$ to $5,302.36 \pm 1,236.71$ AnnV$^+$CT$^+$EV/$\mu l$ between ICM preparations (Fig. 3a) containing a heterogeneous population of AnnV$^+$CT$^+$EV with a peak density in the 100–1,000 nm range (Fig. 3b). The addition of the detergent Triton X-100 to the ICM resulted in a dose-dependent decrease in AnnV$^+$EV (Supplementary Fig. S1), confirming the signal detected was indeed due to the presence of membrane bound vesicles. We then performed whole mount negative staining with transmission electron microscopy (TEM). Visualization of the fractionated ICM confirmed the presence of membrane vesicles contained within (Fig. 3c,d).

Proteomic analysis of islet-derived EV shows commonality with that of islet lysates. In order to obtain a complete protein profile of vesicles from ICM, protein extracts from 50,000 $\times$ g (50 K) and 200,000 $\times$ g (200 K) serial centrifugations of ICM (ICM-50 K and ICM-200 K, respectively), and islet cell lysates (ICL) were used for analysis by liquid chromatography-mass spectrometry (LC-MS/MS). We identified 182, 211, and 1,622 proteins in ICM-50 K, ICM-200 K, and ICL, respectively (Fig. 4a). Interestingly, all the fractions from ICM showed $\geq 56\%$ homology with ICL indicating that islet-derived vesicles carry islet-specific proteins. To confirm that this data set is unique to islets, we compared our data to the proteins identified in the serially centrifuged 50 K and 200 K fractions generated from conditioned media harvested from a serum-starved human umbilical vein endothelial cell line (HUVEC; ExoCarta and Vesiclepedia databases, unique accession ID: vesiclepedia_560). Indeed, the commonality between the human ICM-50 K and ICM-200 K fractions compared to that of the HUVEC were 31 and 16%, respectively (Fig. 4b,c). These observations suggest that islet-derived EV carry cell-type specific protein signatures. The proteomic data will be made available by submission to ExoCarta and Vesiclepedia.

We performed the Go-term enrichment analysis to evaluate the distribution of identified proteins by putative cellular compartment, biological process and molecular function. Enrichment of cellular component showed that a large number of proteins present in islet derived EV are of exosomal origin (Fig. 5a,b) and carry several proteins commonly associated with EV, such as annexins, heat shock protein 70 kDa and 90 kDa variants (Hsp70 and Hsp90, respectively), integrins, clathrin, Ras-related protein Rab5C, and cluster of differentiation factor 14 (CD14) to name a few (Table 1). The majority of the identified proteins in these vesicles were associated with cytoplasm, membrane and cytoskeleton (Fig. 5a,b). Islet-derived EVs were also enriched in proteins associated
Figure 3. Human islet conditioned media contains a heterogeneous population of AnnV⁺ CellTracker⁺ EV. (a) Bar graph showing the mean levels of each ICM (n = 10) detected by flow cytometry varies in their content of AnnV⁺ Cell Tracker⁺ EV after 24 hours in culture. (b) Representative contour plot (FSC vs SSC) shows the peak density of AnnV⁺ Cell Tracker⁺ EV is within 100 nm to 1,000 nm size based on the detection of 100, 500 and 1,000 nm silica beads. (c,d) Visualization of fractionated ICM by whole mount negative staining TEM confirms the presence of membrane vesicles. Scale bars represent 100 nm. Data are expressed as mean ± SEM of n = 3 independent experiments performed on ICM from ten (n = 10) different donors.

Figure 4. Proteomic analysis reveals homology of islet conditioned media extracellular vesicles with islet cell lysate, and a signature distinct from extracellular vesicles produced by human umbilical vein endothelial cells. Venn diagrams depicting the overlap of genes detected in the islet conditioned media (ICM)-50 K and ICM-200 K fractions with that of the islet cell lysate (ICL) (a). Comparison of homology of genes detected in ICM-50 K compared to that of human umbilical vein endothelial cell (HUVEC)-50 K, as well as in ICM-200 K and HUVEC-200 K (islet and endothelial 50,000 g (b) and 200,000 g (c), respectively).
with lysosomes, mitochondria and nucleolus (Fig. 5a,b). Moreover, enrichment analysis for molecular function identified proteins involved in immune response, heat shock and complement activation were also enriched in our EV (Fig. 5a,c,d, Supplementary Fig. S2a,b).

DAA are present in the EV produced by cultured human islets. To investigate the presence of DAA in human ICM EV, we employed flow cytometry with a small particle analyzer, and gated on individual autoantigens in combination with the EV markers AnnV and CT. This analysis revealed the presence of Ag^+ AnnV^+ CT^+ EV for GAD65 (Fig. 6a), ZnT8 (Fig. 6b), the β-cell facilitative glucose transporter, Glut2 (Fig. 6c), the tetraspanin superfamily member, cluster of differentiation factor 9 (CD9) (Supplementary Fig. S3a), and confirmed the proteomic detection of Hsp70 (Supplementary Fig. S3b), and CD14 (Supplementary Fig. S3c), but not IA-2, nor heat shock cognate 70 kDa variant (Hsc70) in the ICM tested (Table 2). Individual contour plots for each of these biomarkers revealed a peak density of EV within a similar size range to that of AnnV^+ CT^+ EV alone (Supplementary Fig. S4).

Similar to the variability seen with AnnV^+ CT^+ EV alone, we noted a wide variation in Ag^+ AnnV^+ CT^+ EV levels between islet preparations (Fig. 7a–f) and in the total percentage of AnnV^+ CT^+ EV with detected antigens on the surface (Fig. 7g). To ensure that genuine EV were being identified and to verify the specificity of the antibodies used, samples of ICM from different islet preparations (n = 4 donors) were treated with centrifugation at 100,000 × g for 1 hour, 50 mM ethylenediaminetetraacetic acid (EDTA), or 0.3% Triton X-100, with resulting signal depletion. In the case of GAD65, a significantly greater level of GAD65^+ AnnV^+ CT^+ EV/µl was observed between control (non-treated ICM) and ICM centrifuged at 100,000 × g (p = 0.012), ICM with 50 mM EDTA (p = 0.005) and ICM treated with 0.3% Triton X-100 (p = 0.005), indicating that the markers detected did indeed come from the EV present within the ICM (Fig. 6a and Supplementary Fig. S4). The same pattern was observed for each of ZnT8 (Fig. 6b), Glut2 (Fig. 6c), CD9 (Supplementary Fig. S3a), Hsp70 (Supplementary Fig. S3b), and CD14 (Supplementary Fig. S3c), where treatment of the ICM by 100,000 × g centrifugation, EDTA and Triton X-100 all resulted in a significant depletion of Ag^+ AnnV^+ CT^+ EV/µl detected (p < 0.05).

TEM analyses revealed a population of EV ranging in size from 30 to 700 nm (Figs. 3c,d and 6d–f, and Supplementary Fig. S5). Assessment of immunogold double labeling on EV by whole mount negative staining TEM revealed Ag^+ AnnV^+ , Ag^+ AnnV^− , Ag^− AnnV^+ , and Ag^− AnnV^− EV for DAA GAD65 (Fig. 6d and Figure 5. Comparison of proteomic data by enrichment analyses of fractionated human islet conditioned media with FunRich component and pathway analyses. Bar graphs of cellular componments overrepresented in highly abundant proteins are shown for islet conditioned media (ICM)-50 K (a) and ICM-200 K (b). Bar graphs illustrating the biological process overrepresented in highly abundant proteins for ICM-50 K (c) and ICM-200 K (d).
Supplementary Fig. S5a,d) and ZnT8 (Fig. 6e and Supplementary Fig. S5b,e), and β-cell marker Glut2 (Fig. 6f and Supplementary Fig. S5c,f).

To assess the contribution of caspase-dependent apoptosis to islet-derived EV, we cultured islet preparations (n = 3 donors) with 50 μM of the pan-caspase inhibitor carbobenzoxy-valyl-alanyl-aspartyl-[O-methyl]fluoromethylketone (ZVAD) or dimethyl sulfoxide (DMSO; vehicle) and compared EV production to that of islets in culture with no intervention (Normal). No difference was observed on GAD65+AnnV+CT+EV (Supplementary Fig. S6a) or AnnV+CT+EV (Supplementary Fig. S6b) production in the ICM between ZVAD, DMSO and Normal islets (p > 0.05). A time course assessment of EV production revealed that the greatest concentration of both GAD65+AnnV+CT+EV and AnnV+CT+EV were produced at 24 hours (p < 0.05), declined at 48 hours and then stabilized by 72 hours, but this was independent of ZVAD treatment (Supplementary Fig. S6). Furthermore, there was no size-dependent treatment effect observed on EV production (Supplementary Fig. S7).

**Discussion**

In this study, we have demonstrated the production of a heterogeneous population of EV by islets in culture, and characterized human islet-derived EV with respect to DAA and β-cell markers. These EV expose PS on their outer surface, as assessed by AnnV binding, contain intracellular esterases of islet-origin, as assessed using CT, and present DAA GAD65 and ZnT8, and glucose transporter Glut2. We have shown the precise detection of each of these Ag-containing vesicles using flow cytometry, as centrifugation at 100,000 g, the addition of EDTA to chelate free calcium and prevent AnnV binding to PS, and the detergent-based depletion of membrane bound structures via the addition of Triton X-100 all significantly reduced the detection of Ag positive EV. Visualization of these DAA-containing EV was confirmed by whole mount immunogold TEM, showing a morphological diversity of EV in human ICM. A large proportion of vesicles were AnnV negative, as has been reported in the case of platelet-derived microparticles and adipocyte-derived EV. Vesicles that are PS negative may represent a population of EV that are less immunostimulatory and capable of remaining in the circulation for longer durations. The significance of AnnV negative particles is not yet fully understood, and while they may represent an

| Accession No. | Symbol | Name | ICM-EV fraction |
|---------------|--------|------|----------------|
| P07355        | ANXA2  | Annexin A2 | ICM-200 K |
| P09525        | ANXA4  | Annexin A4 | ICM-50 K   |
| P08758        | ANXA5  | Annexin A5 | ICM-50 K   |
| P08133        | ANXA6  | Annexin A6 | ICM-50 K   |
| P20073        | ANXA7  | Annexin A7 | ICM-50 K   |
| P14625        | HSP90B1| Endoplasm | ICM-50 K, ICM-200 K |
| P34932        | HSPA4  | Heat shock 70kDa protein 4 | ICM-200 K |
| P11021        | HSPA5  | 78 kDa glucose-regulated protein | ICM-50 K |
| P11142        | HSPA8  | Heat shock cognate 71 kDa protein | ICM-50 K, ICM-200 K |
| P38646        | HSPA9  | Stress-70 protein, mitochondrial | ICM-50 K |
| P04792        | HSPB1  | Heat shock protein beta-1 | ICM-200 K |
| P04406        | GAPDH  | Glyceraldehyde-3-phosphate dehydrogenase | ICM-200 K |
| P60709        | ACTB   | Actin, beta | ICM-200 K |
| P06733        | ENO1   | Enolase 1, (alpha) | ICM-200 K |
| P14618        | PKM    | Pyruvate kinase, muscle | ICM-50 K, ICM-200 K |
| P68104        | EEF1A1 | Eukaryotic translation elongation factor 1 alpha 1 | ICM-50 K, ICM-200 K |
| P04075        | ALDOA  | Aldolase A, fructose-bisphosphate | ICM-50 K |
| P13639        | EEF2   | Eukaryotic translation elongation factor 2 | ICM-200 K |
| P00558        | PGK1   | Phosphoglycerate kinase 1 | ICM-50 K |
| P62258        | YWHAE  | Tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, epsilon | ICM-50 K |
| Q00610        | CLTC   | Clathrin, heavy chain (Hc) | ICM-200 K |
| P62937        | PPIA   | Peptidylprolyl isomerase A (cyclophilin A) | ICM-50 K |
| P60174        | TP1    | Triosephosphate isomerase 1 | ICM-50 K |
| Q06830        | PRDX1  | Peroxiredoxin 1 | ICM-200 K |
| P21333        | FLNA   | Filamin A, alpha | ICM-50 K, ICM-200 K |
| P01023        | A2M    | Alpha 2-macroglobulin | ICM-50 K, ICM-200 K |
| P51148        | RAB5C  | RAB5C, member RAS oncogene family | ICM-200 K |
| P01275        | GCCG   | Glucagon | ICM-50 K, ICM-200 K |
| P08571        | CD14   | CD 14 molecule | ICM-50 K, ICM-200 K |
| P61278        | SMS    | Somatostatin | ICM-50 K, ICM-200 K |

Table 1. List of proteins commonly associated with extracellular vesicles and islet cells identified by LC-MS/MS within fractionated human ICM.
underappreciated population of EV warranting further investigation, analyses of these EV falls out of the scope of this present study.

While the concentration of EV released into the ICM varied dramatically between islet preparations, as detected by NTA (Fig. 2a) and flow cytometry (Fig. 3a), the majority of the ICM contained EV that were consistent in the profile of size distribution. The discrepancy may be due to the fact that for flow cytometry, we conservatively evaluated EV harboring PS and containing esterases, which are respectively indicators of membrane

Figure 6. Diabetes autoantigens and β-cell markers are present on extracellular vesicles produced by non-diabetic donors. (a–c) Bar graphs of islet conditioned media (ICM) compared to signal depletion by 100,000 g centrifugation, 50 mM EDTA, or 0.3% Triton X-100 for GAD65+/AnnV+/CT+ EV (a), ZnT8+/AnnV+/CT+ EV (b), and Glut2+/AnnV+/CT+ EV (c). (d–f) Micrographs of immunogold double labeled islet-derived EV preparations from fractionated ICM confirms the presence of AnnV (white arrows), GAD65 ((b) red arrows), ZnT8 ((d) black arrow) and Glut2 ((f) blue arrow). Data are expressed as mean ± SEM from n = 4 human islet preparations from different donors. *p < 0.05, repeated measures one-way analysis of variance with Tukey’s post-hoc test performed on raw values prior to conversion to percent of control. Scale bars represent 100 nm.
moiety and cytosol content, whereas NTA counts EV without regard to specific markers. Notably, the particle size distribution was similar between NTA and flow cytometry with a range of 100 to 1,000 nm (Figs 2b and 3b). Furthermore, the visualization of islet-derived EV by whole mount negative staining TEM confirmed that the islet-derived EV were indeed membrane-bound structures (Figs 3c,d and 6d–f and Supplementary Fig. S5). Other groups have similarly reported that the majority of protocols used to isolate EV (or more specifically, Exo) result in diverse populations of EV from endosomal and non-endosomal origin15, 16, 31.

Proteomic evaluation of islet-derived EV reflected an islet-specific protein signature as they shared a well-documented limitation of LC-MS/MS is the inefficient solubilization of transmembrane and membrane-anchored proteins due to their hydrophobicity and poor ionization of peptides35–37. Tryptic digestion at play.

Table 2. List of autoantigens and biomarkers assessed, and the method used for their detection, within the human islet conditioned media.

| Marker     | Category                      | Description                                                                 | Detection          |
|------------|-------------------------------|-----------------------------------------------------------------------------|--------------------|
| GAD65 (GAD2) | Diabetes autoantigen          | Diabetes autoantigen; enzyme that converts glutamate to GABA for storage in secretory vesicles in pancreatic β-cells and GABA-secreting neurons56. | Flow Cytometry, TEM |
| IA-2 (PTPRN)   | Diabetes autoantigen          | Islet antigen-2; protein tyrosine phosphatase family [PTPRN]; enriched in pancreatic β-cells. Membrane bound associated with insulin granules14, 15. | Not detected       |
| ZnT8 (SLC30A8) | Diabetes autoantigen          | Zinc transporter present on the cell surface and localized to the insulin storage granules. Highest levels are within the pancreatic β-cell65. | Flow Cytometry, TEM |
| Glut2 (SLC2A2) | β-cell marker; New-onset diabetes autoantigen | Glucose transporter-2; resident on cell surface for passive diffusion of glucose into cell. High levels on pancreatic β-cells, hepatocytes, kidney proximal tubule cells and cells within the small intestine66. | Flow Cytometry, TEM |
| Hsc70       | Injury marker                 | Heat shock cognate protein 70, membrane associated, maintains GAD65 in close proximity to GABA-containing vesicles67, common EV marker. | LC-MS/MS          |
| Hsp70       | Injury marker                 | Highly stress-inducible injury marker, stabilizes proteins against aggregation, known to be pro-inflammatory66, common marker of EV. | Flow Cytometry, LC-MS/MS |
| CD9         | EV marker                     | Tetraspanin, four transmembrane spanning regions involved in protein scaffold maintenance, modulation of cell adhesion, motility and migration, membrane fusion, and exocytic vesicle formation67. | Flow Cytometry     |
| CD14        | EV Marker                     | Monocyte differentiation antigen. Acts as a GPI-anchored coreceptor for TLR2:TLR6; internalized into the endosomal compartment in a lipid raft-dependent mechanism68. | Flow Cytometry, LC-MS/MS |
| Annexin V (ANXA5) | EV marker                  | Intracellular protein with anchoring domains that binds to calcium and to phosphatidylserine69. | Flow Cytometry, LC-MS/MS, TEM |

A recent study confirms our hypothesis that DAA are contained within EV produced by human islets. This study documents the trafficking of GAD65, IA-2 and proinsulin in Exo produced from the trans-golgi network in a Rab11-controlled endosomal pathway by humans and rat islets in culture22. These DAA containing Exo were taken up by activated dendritic cells, with a concurrent cytokine-induced increase in production of Exo and uptake by dendritic cells under stimulation, thus detailing a potential antigen-presentation model for the break in self-tolerance in T1D. In this study, the authors were able to detect GAD65 and IA-2 in their EV preparations by LC-MS/MS by introducing a pre-purification of their proteomic samples with an additional SDS-PAGE step prior to tryptic digestion. In our study, the presence of albumin in the culture media necessitated that albumin-containing bands be cut out prior to tryptic digestion. In this way, proteins of similar size to the 66.5 kDa albumin may have been excised with the albumin-containing band and therefore not included in the tryptic digests. While this may account for GAD65 not being detected in our LC-MS/MS analyses, other factors may be at play.

A well-documented limitation of LC-MS/MS is the inefficient solubilization of transmembrane and membrane-anchored proteins due to their hydrophobicity and poor ionization of peptides35–37. Tryptic digestion...
is not always sufficient to release peptide fragments contained within the transmembrane regions, hindering the detection of these markers. In the case of GAD65, the NH$_2$-terminal membrane anchoring region has been identified as being responsible for the hydrophobicity of the enzyme. Similarly, ZnT8 belongs to the SLC family of zinc transporters that are predicted to contain 6 transmembrane domains, and Glut2 belongs to the SLC2 family of membrane transporters containing 12 membrane-spanning domains. This may explain why GAD65, ZnT8 and Glut2 were not detected in our LC-MS/MS analyses, despite their identification by flow cytometry and immunogold TEM. The same may hold true for CD9, CD81 and other members of the tetraspanin family that are often detected as common EV markers.

In its physiological state, GAD65 has been localized within β-cells to subcellular MV and Golgi membranes. An accumulation of GAD65 in β-cell Golgi membranes was shown in pancreas biopsies from T1DM patients and in vitro models of endoplasmic reticulum stress. Palmitoylation of GAD65 was the cause of this accumulation and also the proposed mechanism for its immunogenicity once externalized.

The localization of ZnT8 to EV is an intuitive discovery. The storage of insulin within the dense core secretory vesicles necessitates the internalization of high concentrations of zinc for solid hexamer formation of 6 insulin units.

Figure 7. Antigen levels on extracellular vesicles present in islet conditioned media from different donor islet preparations. Column scatter graphs of islet conditioned media (ICM) analyzed by flow cytometry for GAD65$^+$ AnnV$^+$ CT$^+$ EV (a), ZnT8$^+$ AnnV$^+$ CT$^+$ EV (b), Glut2$^+$ AnnV$^+$ CT$^+$ EV (c), CD9$^+$ AnnV$^+$ CT$^+$ EV (d), Hsp70$^+$ AnnV$^+$ CT$^+$ EV (e), and CD14$^+$ AnnV$^+$ CT$^+$ EV (f). Bar graph depicting the percent of AnnV$^+$ CT$^+$ EV that were positive for the antigens described in the above panels (g). Data are expressed as mean ± SEM from n = 10 human islet preparations from different donors, and are representative of n = 3 independent experiments.
EV may be used as biomarkers for the detection of early islet injury and may serve to identify susceptible individuals to the events that lead to a break in self-tolerance. The combination of GAD65, ZnT8 and Glut2 on islet-derived β-cell specific method of antigen presentation that is only beginning to be appreciated as a plausible contribution to the immune response against the β-cells. It is not known if this aspect may change the type and content of EV produced. In that study, islets were allowed to recover for an unspecified period of time, then stimulated by cytokine administration. It is interesting to note that the findings of Cianciaruso et al. did not include the detection of ZnT8 or Glut2. This may be due to several factors including the size-restricted analyses performed on Exo, whereby ZnT8 and Glut2 were localized to Exo-like and larger EV in our study (Fig. 6d–f and Supplementary Fig. S5). Furthermore, in that study, islets were allowed to recover for an unspecified period of time, then stimulated by cytokine administration. It is not known if this aspect may change the type and content of EV produced.

**Methods**

**Consent for research and regulatory approvals.** This study was approved by the Centre for Applied Ethics at the McGill University Health Centre (MUHC). All studies were conducted in accordance with relevant guidelines and procedures on pancreatic islets originating from end-of-life organ donation after research consent was obtained from next-of-kin by the appropriate organ donation personnel at Transplant Québec. Only post-mortem biological samples were used in this study with no patient identifiers.

**Human islet isolation and the collection of islet conditioned media.** Islets of Langerhans were isolated, purified, and cultured at the MUHC Human Islet Transplant Laboratory, using previously described protocols. Briefly, the pancreas was loaded intraductally with cold CIzyme (Collagenase HA; VitaCyte LLP, Indianapolis, IN) and either Neutral Protease (Serva Electrophoresis GmbH, Heidelberg, Germany) or Thermolysin (VitaCyte LLP, Indianapolis, IN) enzymes at 4°C prior to warm dissociation at 37°C in a recirculation loop. Iodixanol-based (Optiprep) continuous density gradient purification was performed after the collection of the digested pancreas, and islet fractions with ≥80% purity, ≥90% viability and a glucose-stimulated insulin secretion ratio of >1.0 were used for these studies. Human ICM (CMRL 1066, 10 mM Heps, 2.5% human serum albumin, 10 µg/ml ciprofloxacin) was harvested from islet preparations (n = 10) after post-isolation culture for up to 72 hours, centrifuged at 1,200 g for 15 min. to discard dead cells and debris, then stored at −80°C until use. Relevant donor characteristics have been detailed in Supplementary Table S1, including the analyses of the islet preparations from each donor that were used to generate the ICM or ICL.

**Nanoparticle Tracking Analyses.** Individual ICM samples, collected at 24 hours after isolation, were analyzed by the Nanosight NS500 system (Nanosight Ltd., Amesbury, UK). Human ICM were measured by NTA to quantify the mean size and concentration of particles. ICM preparations were diluted (1:50) in PBS and analyzed with the Nanoparticle Analysis System & NTA 1.4 Analytical Software. At least 5 recordings, 30 seconds each were obtained at 37°C with a camera shutter speed set to 30.0 ms, a camera level of 14 and detection threshold set to 9.

**Flow cytometry Analyses.** Flow cytometry analyses were performed on aliquots of ICM using a FACSA Flow cytometry analyses were performed on aliquots of ICM using a FACSA Canto II with small particle analyzer, as previously described. Samples of 25 µl ICM (n = 10), collected 24 hours after isolation, were labeled for 30 min. at RT with the following fluorescent-probe conjugated surface markers or antibodies: DAA anti-GAD65-PE (0.25 µg, clone 144, ImmuQuest), anti-IA-2-APC (0.4 µl, aa287–316, LifeSpan Biosciences), and anti-ZnT8-FITC (3 µg, aa263–369, LifeSpan Biosciences, Inc.); anti-Glut2-PE (5 µl, aa263–369, LifeSpan Biosciences, Inc.).
EV Enrichment. Vesicles were enriched from human ICM collected at 24 hours, using a protocol described previously \(^1\). Briefly, aliquots of 30 ml were thawed and fractionated by serial centrifugation at 50,000 \(g\) for 15 min to isolate apoptotic body-like and MV-like EV, followed by 200,000 \(g\) for 18 hours to isolate Exo-like EV, producing the ICM-50 K and ICM-200 K pellets, respectively, and stored at -80 \(^\circ\)C until use.

Mass Spectrometry. For proteomic analyses by LC-MS/MS, EV fractions were isolated from ICM from different donors (n = 5) through sequential centrifugation, as outlined above. Twenty micrograms of protein from fractions of ICM-50 K and ICM-200 K were separated by SDS-PAGE. Tryptic digests were analyzed by LC-MS/MS on an LTQ-Orbitrap Elite and the data were processed using PEAKS 7.0 (Bio-informatics Solutions), and the human Uniprot database, as previously described \(^1\). Processed data was compared to that of the ICL, and proteins identified in the ultracentrifuged 50 K and 200 K fractions derived from control, unconditioned media were subtracted from those detected in the respective fractions of ICM-50 K and ICM-200 K. Data analysis was performed with Scaffold 4.4 with a False Discovery Rate of 1%. Gene Ontology enrichment analysis for cellular component, molecular function and biological pathways were performed using FunRich v3.0, and only the enriched terms with a p-value of <0.01 were considered, as previously described \(^2\).

Whole Mount Negative Staining and Immunogold Transmission Electron Microscopy. Whole mount negative staining and immunogold labeling experiments for TEM were performed on the aforementioned fractionated human ICM samples, as previously described \(^1\,3\,4\). Briefly, a 1:1 dilution of the ICM-50 K or ICM-200 K pellets was created using a buffer of 4% paraformaldehyde in 50 mM HEPES, pH 7.4, and incubated for 1 hour. Aliquots of 10 \(\mu\)l were adsorbed to Formvar carbon-coated copper grids for 20 min. and contrasted for whole mount negative staining electron microscopy or immunolabeling. Immunogold double labeling was performed to confirm the results obtained by flow cytometry. Primary antibodies used were of the same clonality described above. After blocking with 1% bovine casein ovalbumin for 5 min., combinations of either anti-human GAD65, anti-human Glut2 or anti-human ZnT8 (epitope: aa263–369; LifeSpan Biosciences, Inc.) (clones as above) with anti-annexin V (R&D Systems) were incubated for 1 hour. Control samples were run in parallel by incubation for 1 hour with 1% bovine casein ovalbumin alone. Grids were incubated with a goat anti-mouse IgG conjugated to 18 nm colloidal gold (1:20 dilution; Jackson Laboratories) and goat anti-rabbit IgG conjugated to 12 nm colloidal gold (1:20 dilution; Jackson Laboratories) secondary antibodies for 30 min., followed by washing and fixation. Negative staining was performed with 10 \(\mu\)l of uranyl acetate for 40 sec., and then the grids were air dried for at least 30 min. prior to analyses on a FEI Tecnai 12 BioTwin 120 kV TEM. At least 50 fields of view were examined for the presence of gold particles at 13,000 \(\times\) magnification from each of the control and primary antibody-labeled grids. Electron micrograph images were captured on the Advanced Microscopy Techniques (AMT) XR80C CCD Camera System with AMT Image Capture Engine V601 at 68,000 \(\times\), 49,000 \(\times\), or 18,500 \(\times\) magnification.

Effects of Pan-Caspase Inhibition on Human Islet EV Production. To examine the effect of the pan-caspase inhibitor ZVAD-fmk (ZVAD) on EV production, islets were cultured with media supplemented with 50\(\mu\)M ZVAD, DMSO vehicle control (DMSO), or no intervention (Normal) at 37 \(^\circ\)C for up to 72 hours. ICM was collected at 24, 48 and 72 hours, as outlined above, and stored at -80 \(^\circ\)C until analysis by flow cytometry.

Statistical Analyses. All data are expressed as means \(\pm\) SEM derived from at least three independent experiments unless otherwise specified. Statistical significance of results was evaluated on raw values prior to conversion to percent of control, by a repeated measures one-way analysis of variance (ANOVA) test with Tukey’s post-hoc pairwise comparison, or t-test, using SPSS Statistics (Version 21). P values < 0.05 were considered significant.

References
1. Notkins, A. L. & Lernmark, A. Autoimmune type 1 diabetes: resolved and unresolved issues. J Clin Invest 108, 1247–1252 (2001).
2. Baekkeskov, S. et al. Identification of the 64 K autoantigen in insulin-dependent diabetes as the GABA-synthesizing enzyme glutamic acid decarboxylase. Nature 347, 151–156 (1990).
3. Wenzlau, J. M. et al. The cation efflux transporter ZnT8 (Slc30A8) is a major autoantigen in human type 1 diabetes. Proc Natl Acad Sci USA 104, 17040–17045 (2007).
4. Payton, M. A., Hawkes, C. J. & Christie, M. R. Relationship of the 37,000- and 40,000-M(r) tryptic fragments of islet antigens in insulin-dependent diabetes to the protein tyrosine phosphatase-like molecule IA-2 (ICA512). J Clin Invest 96, 1506–1511 (1995).
5. Wenzlau, J. M. & Hutton, J. C. Novel diabetes autoantibodies and prediction of type 1 diabetes. Curr Diab Rep 13, 608–615 (2013).
6. Insel, R. A. et al. Staging presymptomatic type 1 diabetes: a scientific statement of JDFR; the Endocrine Society, and the American Diabetes Association. Diabetes Care 38, 1964–1974 (2015).
7. McKiernan, J. et al. A Novel Urine Exosome Gene Expression Assay to Predict High-grade Prostate Cancer at Initial Biopsy. JAMA Oncol (2016).
8. Hoshino, A. et al. Tumour exosome integrins determine organotropic metastasis. Nature 527, 329–335 (2015).
9. Bolliard, E. et al. Platelets amplify inflammation in arthritis via collagen-dependent microparticle production. Science 327, 580–583 (2010).
10. Nielsen, C. T. et al. Increased IgG on cell-derived plasma microparticles in systemic lupus erythematosus is associated with autoantibodies and complement activation. Arthritis Rheum 64, 1227–1236 (2012).
11. Selly, E. M., Sayed, D., Abdel Aal, R. F., Ahmad, H. M. & Abdou, M. A. Increased circulating red cell microparticles (RMP) and platelet microparticles (PMP) in immune thrombocytopenic purpura. Thromb Res 131, e59–e63 (2013).
12. Tomlinson, P. R. et al. Identification of distinct circulating exosomes in Parkinson’s disease. Am J Clin Transl Neurol 2, 353–361 (2015).
13. Dieude, M. et al. The 20S proteasome core, active within apoptotic exosome-like vesicles, induces autoantibody production and accelerates rejection. Sci Transl Med 7, 318ra200 (2015).
14. Pallet, N. et al. A comprehensive characterization of membrane vesicles released by autophagic human endothelial cells. Proteomics 13, 1108–1120 (2013).
15. Gyorgy, B. et al. Membrane vesicles, current state-of-the-art: emerging role of extracellular vesicles. Cell Mol Life Sci 68, 2667–2688 (2011).
16. Marcus, G. et al. Revealing the diversity of extracellular vesicles using high-dimensional flow cytometry analyses. Scientific reports 6, 35928 (2016).
17. van der Pol, E., Boing, A. N., Gooi, E. L. & Nieuwland, R. Recent developments in the nomenclature, presence, isolation, detection and clinical impact of extracellular vesicles. J Thromb Haemost 14, 48–56 (2016).
18. Bashratyan, R., Sheng, H., Regn, D., Rahman, M. J. & Dai, Y. D. Insulinoma-released exosomes activate autoreactive marginal zone-like B cells that expand endogenously in prediabetic NOD mice. European journal of immunology 43, 2588–2597 (2013).
19. Rahman, M. J., Regn, D., Bashratyan, R. & Dai, Y. D. Exosomes released by islet-derived mesenchymal stem cells trigger autoimmune responses in NOD mice. Diabetes 63, 1008–1020 (2014).
20. Sheng, H. et al. Insulinoma-released exosomes or microparticles are immunostimulatory and can activate autoreactive T cells spontaneously developed in nonobese diabetic mice. J Immunol 187, 1591–1600 (2011).
21. Figliolini, F. et al. Isolation, characterization and potential role in beta cell-endothelium cross-talk of extracellular vesicles released from human pancreatic islets. Plos one 9, e105251 (2014).
22. Cianciaruso, C. et al. Primary Human and Rat beta-Cells Release the Intracellular Autoantigens GAD65, IA-2, and Proinsulin in Exosomes Together With Cytokine-Induced Enhancers of Immunity. Diabetes 66, 460–473 (2017).
23. Korsgren, S. et al. On the etiology of type 1 diabetes: a new animal model signifying a decisive role for bacteria eliciting an acute innate immunity response. Am J Pathol 181, 1735–1748 (2012).
24. Buendia, P. et al. Endothelial microparticles mediate inflammation-induced vascular calcification. FASEB J 29, 173–181 (2015).
25. Di Buduo, C. A. et al. Programmable 3D silk bone marrow niche for platelet generation ex vivo and modeling of megakaryopoiesis pathologies. Blood 123, 2254–2264 (2013).
26. Rousseau, M. et al. Detection and quantification of microparticles from different cellular lineages using flow cytometry. Evaluation of the impact of secreted phoshoase A2 on microparticle assessment. Plos one 10, e0146182 (2015).
27. Arraud, N. et al. Extracellular vesicles from blood plasma: determination of their morphology, size, phenotype and concentration. J Thromb Haemost 12, 614–627 (2014).
28. Connor, D. E., Exner, T., Ma, D. D. & Joseph, J. E. The majority of circulating platelet-derived microparticles fail to bind annexin V, lack phospholipid-dependent procoagulant activity and demonstrate greater expression of glycoprotein Ib. Thromb Haemost 103, 1044–1052 (2010).
29. Connolly, K. D. et al. Characterisation of adipocyte-derived extracellular vesicles released pre- and post-adipogenesis. Journal of extracellular vesicles 4, 29159 (2015).
30. Bolliard, E., Duchez, A. C. & Brisson, A. The diversity of platelet microparticles. Curr Opin Hematol 22, 437–444 (2015).
31. Bobrie, A., Colombo, M., Krumeich, S., Raposo, G. & Thery, C. Diverse subpopulations of vesicles secreted by different intracellular mechanisms are present in exosome preparations obtained by differential ultracentrifugation. Journal of extracellular vesicles 1 (2012).
32. Aikin, R., Rosenberg, L., Parakavas, S. & Maysinger, D. Inhibition of caspase-mediated PARP-1 cleavage results in increased necrosis in isolated islets of Langerhans. J Mol Med (Berl) 82, 389–397 (2004).
33. Barsnes, N. R., Wyllie, S. & Goss, J. A. Inflammation-mediated dysfunction and apoptosis in pancreatic islet transplantation: implications for infrahepatic grafts. J Leukoc Biol 77, 587–597 (2005).
34. Piemonti, L. et al. Alloantibody and autoantibody monitoring predicts islet transplantation outcome in human type 1 diabetes. Diabetes 62, 1656–1664 (2013).
35. Gerl, M. J. et al. Analysis of transmembrane domains and lipid modified peptides with matrix-assisted laser desorption ionization-time-of-flight mass spectrometry. Anal Chem 86, 3722–3726 (2014).
36. Choksawangkarn, W., Edwards, N., Wang, Y., Gutierrez, P. & Fenselau, C. Comparative study of workflows optimized for in-gel, in-solution, and on-filter proteolysis in the analysis of plasma membrane proteins. J Proteome Res 11, 3030–3034 (2012).
37. Castellanos-Serra, L., Ramos, Y. & Huerta, V. An in-gel digestion procedure that facilitates the identification of highly hydrophobic proteins by electrospray ionization-mass spectrometry analysis. Proteomics 5, 2729–2738 (2005).
38. Kanaani, J., Lisson, D., Kash, S. F. & Baeckkesov, S. The hydrophilic isomorph of glutamate decarboxylase, GAD67, is targeted to membranes and nerve terminals independent of dimerization with the hydrophilic membrane-anchored isoform, GAD65, J Biol Chem 274, 37200–37209 (1999).
39. Seve, M., Chimenti, F., Devergnas, S. & Favier, A. In silico identification and expression of SLC30 family genes: an expressed sequence tag data mining strategy for the characterization of zinc transporters’ tissue expression. BMC genomics 5, 32 (2004).
40. Mueckler, M. & Thorens, B. The SLC2 (GLUT) family of membrane transporters. Mol Aspects Med 34, 121–138 (2013).
41. Christgau, S. et al. Membrane anchoring of the autoantigen GAD65 to microvesicles in pancreatic beta-cells by palmitoylation in the NH2-terminal domain. J Cell Biol 118, 309–320 (1992).
42. Kanaani, J. et al. Compartmentalization of GABA synthesis by GAD67 differs between pancreatic beta cells and neurons. Plos one 10, e0117130 (2015).
43. Phelps, E. A. et al. Aberrant Accumulation of the Diabetes Autoantigen GAD65 in Golgi Membranes in Conditions of ER Stress and Autoimmunity. Diabetes 65, 2686–2699 (2016).
44. Sondergaard, L. G. et al. Zinc ions in beta-cells of obese, insulin-resistant, and type 2 diabetic rats traced by autoradiography. APAMS 111, 1147–1154 (2003).
45. Chimienti, F. et al. In vivo expression and functional characterization of the zinc transporter ZnT8 in glucose-induced insulin secretion. J Cell Sci 119, 4199–4206 (2006).
46. Wenzlau, J. M. et al. A common nonsynonymous single nucleotide polymorphism in the SLC30A8 gene determines ZnT8 autoantibody specificity in type 1 diabetes. Diabetes 57, 2693–2697 (2008).
47. Sladek, R. et al. A genome-wide association study identifies novel risk loci for type 2 diabetes. Nature 445, 881–885 (2007).
48. Thorens, B. GLUT2, glucose sensing and glucose homeostasis. Diabetologia 58, 221–232 (2015).
49. Inman, L. R. et al. Autoantibodies to the GLUT-2 glucose transporter of beta cells in insulin-dependent diabetes mellitus of recent onset. Proc Natl Acad Sci USA 90, 1281–1284 (1993).
50. Lakey, J. R. et al. Variables in organ donors that affect the recovery of human islets of Langerhans. Transplantation 61, 1047–1053 (1996).
51. Melzi, R. et al. Donor and isolation variables associated with human islet monocyte chemoattractant protein-1 release. Transplantation 78, 1564–1567 (2004).
52. Lakey, J. R., Rajotte, R. V., Warnock, G. L. & Kneteman, N. M. Human pancreas preservation prior to islet isolation. Cold ischemic tolerance. Transplantation 59, 689–694 (1995).
53. Negi, S. et al. Analysis of beta-cell gene expression reveals inflammatory signaling and evidence of dedifferentiation following human islet isolation and culture. PLoS one 7, e30415 (2012).
54. Paraskevas, S. et al. Modulation of JNK and p38 stress activated protein kinases in isolated islets of Langerhans: insulin as an autocrine survival signal. Ann Surg 233, 124–133 (2001).
55. Paraskevas, S. et al. Activation and expression of ERK, JNK, and p38 MAP-kinases in isolated islets of Langerhans: implications for cultured islet survival. FEBS Lett 455, 203–208 (1999).
56. Paraskevas, S. et al. Apoptosis occurs in freshly isolated human islets under standard culture conditions. Transplant Proc 29, 750–752 (1997).
57. Paraskevas, S., Maysinger, D., Wang, R., Duguid, T. P. & Rosenberg, L. Cell loss in isolated human islets occurs by apoptosis. Pancreas 20, 270–276 (2000).
58. Thomas, D. et al. Proapoptotic Bax is hyperexpressed in isolated human islets compared with antia apoptotic Bcl-2. Transplantation 74, 1489–1496 (2002).
59. Kaddis, J. S., Danobeitia, J. S., Niland, J. C., Stiller, T. & Fernandez, L. A. Multicenter analysis of novel and established variables associated with successful human islet isolation outcomes. Am J Transplant 10, 646–656 (2010).
60. Caballero-Corbalan, J. et al. Vitacyle collagenase HA: a novel enzyme blend for efficient human islet transplantation. Transplantation 88, 1400–1402 (2009).
61. Balmurugan, A. N. et al. A new enzyme mixture to increase the yield and transplant rate of autologous and allogeneic human islet products. Transplantation 93, 693–702 (2012).
62. Pathan, M. et al. FunRich: An open access standalone functional enrichment and interaction network analysis tool. Proteomics 15, 2597–2601 (2015).
63. Thery, C., Amigorena, S., Raposo, G. & Clayton, A. Isolation and characterization of exosomes from cell culture supernatants and biological fluids. Curr Protec Cell Biol Chapter 3, Unit 3 22 (2006).
64. Bonfaccio, E., Lampasona, V., Genovese, S., Ferrari, M. & Bocchi, E. Identification of protein tyrosine phosphatase-like IA2 (islet cell antigen 512) as the insulin-dependent diabetes-related 37/40 K autoantigen and a target of islet-cell antibodies. J Immunol 155, 5419–5426 (1995).
65. Cho, J. A., Lee, Y. S., Kim, S. H., Ko, J. K. & Kim, C. W. MHC independent anti-tumor immune responses induced by Hsp70–antigen 512) as the insulin-dependent diabetes-related 37/40 K autoantigen and a target of islet-cell antibodies. J Immunol 155, 5419–5426 (1995).
66. Mazurov, D., Barbashova, L. & Filatov, A. Tetraspanin protein CD9 interacts with metalloprotease CD10 and enhances its release via autocrine survival signal. Ann Surg 233, 124–133 (2001).
67. Paraskevas, S., Maysinger, D., Wang, R., Duguid, T. P. & Rosenberg, L. Cell loss in isolated human islets occurs by apoptosis. Pancreas 20, 270–276 (2000).
68. Thomas, D. et al. Proapoptotic Bax is hyperexpressed in isolated human islets compared with antia apoptotic Bcl-2. Transplantation 74, 1489–1496 (2002).
69. Kaddis, J. S., Danobeitia, J. S., Niland, J. C., Stiller, T. & Fernandez, L. A. Multicenter analysis of novel and established variables associated with successful human islet isolation outcomes. Am J Transplant 10, 646–656 (2010).
70. Caballero-Corbalan, J. et al. Vitacyle collagenase HA: a novel enzyme blend for efficient human islet transplantation. Transplantation 88, 1400–1402 (2009).
71. Balumurugan, A. N. et al. A new enzyme mixture to increase the yield and transplant rate of autologous and allogeneic human islet products. Transplantation 93, 693–702 (2012).
72. Pathan, M. et al. FunRich: An open access standalone functional enrichment and interaction network analysis tool. Proteomics 15, 2597–2601 (2015).
73. Thery, C., Amigorena, S., Raposo, G. & Clayton, A. Isolation and characterization of exosomes from cell culture supernatants and biological fluids. Curr Protec Cell Biol Chapter 3, Unit 3 22 (2006).
74. Bonfaccio, E., Lampasona, V., Genovese, S., Ferrari, M. & Bocchi, E. Identification of protein tyrosine phosphatase-like IA2 (islet cell antigen 512) as the insulin-dependent diabetes-related 37/40 K autoantigen and a target of islet-cell antibodies. J Immunol 155, 5419–5426 (1995).
75. Cho, J. A., Lee, Y. S., Kim, S. H., Ko, J. K. & Kim, C. W. MHC independent anti-tumor immune responses induced by Hsp70– enriched exosomes generate tumor regression in murine models. Cancer Lett 275, 256–265 (2009).
76. Mazurov, D., Barbashova, L. & Filatov, A. Tetraspanin protein CD9 interacts with metalloprotease CD10 and enhances its release via exosomes. FEBS J 280, 1200–1213 (2013).
77. Zanoni, I. & Granucci, F. Role of CD14 in host protection against infections and in metabolism regulation. Front Cell Infect Microbiol 3, 32 (2013).
78. Bouter, A. et al. Review: Annexin-A5 and cell membrane repair. Placenta 36(Suppl 1), S43–49 (2015).

Acknowledgements
This research was performed with the support of Transplant Quebec, responsible for coordinating the organ donation services and obtaining consent for research. We thank the families of the organ donors for this invaluable contribution. We also thank Marie-Josée Hébert, Mélanie Dieudé and Julie Turgeon (Centre de Recherche du Centre Hospitalier Universitaire de Montréal, Canadian National Transplant Research Program) for their expert technical assistance and resources supporting this project. Funding support for these studies was provided from the following sources: Canadian National Transplantation Research Program (CNTRP) and the McGill University Health Centre Foundation (Royal Victoria Hospital and Montreal General Hospital Foundations). C.P.H. is the recipient of the Astellas CNRTP Training Award, and É.B. is the recipient of a new investigator salary award from the Canadian Institutes of Health Research.

Author Contributions
C.P.H., S.N., S.P., N.C., I.A., É. Boilard, É. Bonneil, and P.T. designed the experiments. Human islet isolations, generation of ICM and ICM fractionation were performed by C.P.H., M.G., A.K.R., S.N., and S.P. NTA was performed by A.K.R., S.N. and C.P.H. Flow cytometry was performed by N.C., I.A., C.P.H. and É. Boilard. LC-MS/MS was performed by É. Bonneil. TEM studies were performed by C.P.H. Data analyses were performed by C.P.H., S.N., A.K.R., N.C., I.A., and É. Boilard. C.P.H., S.N. and S.P. wrote the manuscript. All authors contributed to the editing of the manuscript.

Additional Information
Supplementary information accompanies this paper at doi:10.1038/s41598-017-04977-y

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

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.
