Beta cells transfer vesicles containing insulin to phagocytes for presentation to T cells

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Beta cells from nondiabetic mice transfer secretory vesicles to phagocytic cells. The passage was shown in culture studies where the transfer was probed with CD4 T cells reactive to insulin peptides. Two sets of vesicles were transferred, one containing insulin and another containing catabolites of insulin. The passage required live beta cells in a close cell contact interaction with the phagocytes. It was increased by high glucose concentration and required mobilization of intracellular Ca2++. Live images of beta cell–phagocyte interactions documented the intimacy of the membrane contact and the passage of the granules. The passage was found in beta cells isolated from islets of young nondiabetic diabetic (NOD) mice and nondiabetic mice as well as from nondiabetic humans. Ultrastructural analysis showed inrasislet phagocytes containing vesicles having the distinct morphology of dense-core granules. These findings document a process whereby the contents of secretory granules become available to the immune system.

autoimmune diabetes | autoimmune immunity | insulin reactivity | insulin-reactive T cells

A n important issue in understanding the initiation of the autoimmune diabetic process is the manner in which the products of the secretory granule, in particular insulin, are handled by the immune system and made available to the autoreactive T cells. Autoreactivity to insulin is evident in type 1 diabetes and is a major component of the spontaneous diabetes of the NOD mouse (reviewed in refs. 1, 2). In the nondiabetic diabetic (NOD) mouse, CD4 T cells recognizing insulin peptides are found among the T cells that infiltrate the islets of Langerhans early in the autoimmune process. Such T cells induce disease when isolated and injected into nondiabetic mice (3–5). Altogether, a persuasive case has been made for insulin autoreactivity as driving forward diabetic autoimmunity (6–10). Our previous investigations showed that the islets of Langerhans of NOD mice contained resident phagocytes represented by two lineages: typical macrophages composed the majority, whereas a minority was made up of a subset of dendritic cells (DCs) expressing the CD103 integrin—that is, the CD103+/CD8α− DC lineage under the control of the Batf3 transcription factor (11). Islet phagocytes were found to be in intimate contact with both beta cells and the vessel walls, throwing small projections into the vessel lumen (12). Importantly, when isolated they were strong antigen-presenting cells (APCs), particularly of insulin epitopes (5, 13). Furthermore, an initial ultrastructural analysis showed dense-core secretory granules inside vacuoles of the islet phagocytes residing in the islets (13), and by immunofluorescence islet phagocytes were shown to contain products from the beta cells (5, 13, 14). Direct evidence of insulin peptides within the beta cells and in islet phagocytes was obtained using a monoclonal antibody that was exclusively reactive with an insulin B chain peptide segment and not with native insulin (5).

The passage of insulin to APCs took place even in nondiabetic mice; for a telling example, it was evident in NOD.Rag1−/− mice that have no lymphocytes and do not develop diabetes. In brief, inflammation was not a requisite for the acquisition of secretory granule antigens by the APCs. A structural study in active diabetic rats also showed granules inside monocyte-like cells (15). In toto, these results indicated that the products of the secretory granule had been passed to the APCs in vivo. These findings point to a scenario in which beta cells donate the antigenic epitopes that trigger autoreactive T cells to the local phagocytes as an initial stage before the sensitization of the draining pancreatic node (16–21; as reviewed in ref. 22).

This study further characterizes the interactions between beta cells and APCs probing with CD4 T cells to insulin. It examines the conditions that generate the insulin peptides recognized by diverse CD4 T cells and documents the passage of vesicles containing the peptides to APCs in a contact reaction dependent on live beta cells. Importantly, it documents that such peptides are also generated from human beta cells, further highlighting the importance of insulin immunoreactivity in T1D.

Results and Discussion

A Functional Assay to Examine the Passage of Insulin Epitopes to Phagocytes. We previously characterized two distinct sets of insulin-reactive CD4 T cells (5, 8, 23). One set recognizes a peptide resulting from insulin processing by APCs and encompasses residues 13–21 of the B chain. In NOD diabetes, these T cells are poorly represented, most likely because of thymic deleltional effects. We have referred to them as “type A,” represented here by the ITF-3 clone. The second set recognizes primarily the 12–20 segment of the insulin B chain, a one-amino acid shift of the binding register. This peptide is not expressed after the processing of insulin by APCs but is found when APCs react with denatured insulin or insulin peptides. These clones are relatively abundant in NOD diabetes. We have referred to these T cells as “type B,” herein represented by the 8F10 clone. Therefore, comparing the response of these two T cells is useful, in that the T cells directed to

Significance

This report documents that beta cells from islets of Langerhans normally transfer some of their secretory granules to resident phagocytes. The transfer involves a close contact interaction between live beta cells and phagocytes, increases upon glucose stimulation, and requires mobilization of intracellular Ca2++. In autoimmune diabetes, the CD4 T cells to various peptides of the insulin B chain recognize the transferred antigens in the phagocytes represented in islets by macrophages and a subset of dendritic cells. We have identified a process whereby antigens become available for recognition by autoreactive T cells in type 1 diabetes.


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the 12–20 segment probe the presentation of denatured insulin or the insulin catabolites but not the native insulin molecule.

In an initial report, we had stained whole isolated islets with antibodies to insulin and also to the insulin B9–23 peptide (5). The antibody to B9–23 identified positive granules in beta cells that were larger than the insulin granules and stained positive with anti-Lamp2 antibodies, suggesting that B9–23 granules may compose a distinct pool of granules (figure 3 in ref. 5). The intraislet APCs contained B9–23-positive granules at an average of about 10 per cell. To quantitate our previous findings using flow cytometry, islet cells were isolated and examined using antibodies to insulin and the B9–23 peptide (5). We found insulin and B9–23 reactivity both in beta cells (Fig. L4) and in CD45+ CD11c+ F4/80+ macrophages (Fig. 1B), importantly from islets of nondiabetic NOD Rag1−/− mice. Similar results were found in islets from 4–6-wk-old NOD mice or C57BL/6 mice. In a different manipulation, beta cells were isolated from NOD Rag1−/− mice and subjected to subcellular fractionation of the secretory granules using differential centrifugation of vesicles (24, 25). The granules were then added to CD11c+ DCs isolated from FMS-like tyrosine kinase 3 ligand (FLT-3L)-treated mice, and presentation to the two T cells was examined. (The DCs comprised two sets, a SIRPα+ and a CD24+, the two phagocytes in the islets are a macrophage-SIRPα+ and the CD103+DC. In this report, we do not differentiate between the two sets.) The 25,000 × g fraction that contains the mature secretory granules stimulated the IIT-3 T cells that recognized insulin epitopes and had a much smaller reactivity to SF10 that only recognizes peptides or denatured insulin. The reverse was found for the 5,000 × g fraction that stimulated strongly the SF10 (Fig. 1C). In sum, beta cells contain vesicles that are immunoreactive to B chain-specific antibody; the intraislet APCs likewise show positive reactivity with both B chain peptide as well as insulin-specific antibodies.

A culture assay was developed to examine the transfer of insulin immunogenic material from beta cells to phagocytes. Endocrine cells harvested from isolated islets were placed in culture in different media from short time periods of 1–3 h to overnight, after which DCs were added for several hours. (We refer to the endocrine cells as beta cells, as we are probing only insulin transfer.) Lastly, the presence in the DCs of the peptide bound to

![Image](image.png)

Fig. 1. Beta cells transfer immunogenic insulin to phagocytes. Flow cytometry plots of isolated endocrine cells (A) or intraislet macrophages (B) stained with antibodies reactive to insulin and to the B9–23 peptide. Cells were gated as indicated in the panels. Red histograms correspond to the isotype control antibody staining. Blue histograms are either antiinsulin or anti-B9–23 peptide staining. Numbers in each plot indicate the mean fluorescence intensity for their respective histogram. (C) Secretory granules were isolated by differential centrifugation from beta cells isolated from NOD Rag1−/− mice and offered to spleen DCs, and the response of the SF10 T cell or IIT-3 T cell was then assayed. Shown are the responses to the 5,000 and 25,000 × g fractions (in 5K and 25K, respectively) and as a control to the B9–23 peptide. (D) The characterization of the two CD4 T cells to insulin (8). The FLT-3L DCs were incubated with insulin or with the B9–23 peptide, each at 10 μM. SF10 reacts with peptides B9–23 or B12–20 (sequence shown below the graph) but not with insulin or B12–21. IIT-3 reacts with insulin and peptides B9–23 and B13–21. (E) A representative assay (of n > 25). Indicated are the cells used in the assay. Beta cells were from 6-wk-old NOD Rag1−/− mice; the APCs were DCs obtained from the spleen of mice previously injected with FLT-3L. Background response of the T cells never exceeds 150 cpm. (F) Summary of the first series of experiments. The explanation is in the text. (G) Antibody to I-Aβ7 inhibits the transfer. The culture included the presence or absence of 10 μg/mL of the antibody Ag2.42.67 specific for I-Aβ7. (H) Same as in A but testing islets from B6 mice. Shown is a representative experiment of two experiments. (I) As in A but testing human islets. The results are pooled from two experiments.
the I-A\(^{B}\) class II MHC molecule was probed using either of the two insulin-reactive CD4 T cells.

**Testing Beta Cells from Multiple Sources.** Fig. 1D shows the specificity of the T cells used in these experiments: The CD4 T-cell 8F10 cells were obtained from NOD.

By streptozotocin before the addition of the DCs (Fig. 2), there was a complete lack of presentation when all beta cells were killed responsively for the transfer of peptides to the DCs. Indeed, there were no responses when all beta cells were killed—expected, in the absence of additional DCs, the T cells never responded to beta cells, because they lack expression of MHC-II molecules (Fig. 1E).

Consistently, the addition of DCs after the first 24 h of culture of beta cells in media containing 5 mM glucose resulted in a variable low level of presentation (low glucose in the figures). (Identical results were also found with 2.5 mM glucose in the media.) Both T cells reacted on average about 10-fold higher overall than the background stimulation (Fig. 1E and F). Culturing beta cells with 25 mM glucose resulted in a significant increase in insulin presentation by DCs. The response of SF10 (i.e., to B12–20 peptide) increased by 352% (n = 13 experiments); with ITT-3 (i.e., to B13–21 peptide), there was a similar increase (351%, n = 8 experiments) (Fig. 1E and F). The age of the mice did not influence the transfer. In various experiments, beta cells from 4–16-wk-old NOD.Rag1\(^{−/−}\) mice presented equally. Similar results were obtained examining beta cells from young prediabetic NOD mice 6, 8, and 14 wk of age. Not shown are results indicating similar findings with bone marrow-derived macrophages as the capture APCs.

The presentation of insulin peptides, as expected, depended on the interaction with class II MHC molecules and was inhibited by addition of a blocking monoclonal antibody to the I-A\(^{B}\) protein (Fig. 1G). The specificity of the insulin reactivity was shown by testing islets from mice that express only insulin-2 with a mutation at Tyr-16 of the beta chain (6). This is a nonimmunogenic insulin known to be nonreactive with either type A or type B T cells (6–8, 23); their beta cells did not elicit a positive response. (The results were as follows. The mean cpm response of T cells alone, T cells plus beta cells and DCs, and the control of beta cells plus DCs plus B3–23 peptides were for SF10, 58, 47, and 38,610, respectively, and for ITT-3, 35, 46, and 41,991, respectively.)

C57BL/6 mice do not develop spontaneous diabetes. Their beta cells incubated with DCs from NOD mice transferred the immunogenic peptides under low glucose media, whereas the transfer was potentiated by incubating them in high glucose media (Fig. 1H). In brief, there was no difference in the generation of peptides between beta cells of a different MHC haplotype or diabetes-susceptible status. The same transfer took place testing human islet preparations obtained from two recently deceased nondiabetic patients. The segment of the insulin B chain that stimulates the murine 12–20- and 13–21-reactive CD4 T cells is identical between human and mouse (Fig. 1D). As expected, in both situations, the DCs and the CD4 T cells were from NOD mice—that is, syngeneic—based on MHC restriction rules. Clearly two events are taking place: One is the transfer of the antigenic material from the beta cells, and the second is the DC–T cell interaction, the latter requiring the diabetes-susceptible MHC molecule.

**Nature of the Transfer.** Phagocytes could take up extracellular insulin or insulin peptides and then present them to the T cells. Separation of the beta cells from the DC by a cell-impermeable membrane abolished presentation, indicating that close cell contact was required (Fig. 2A). About 30% of beta cells invariably died after the first 24 h of culture, but the dead beta cells were not responsible for the transfer of peptides to the DCs. Indeed, there was a complete lack of presentation when all beta cells were killed by streptozotocin before the addition of the DCs (Fig. 2B).

Presentation required the processing of the granules by the DCs. DCs were pretreated with chloroquine, the drug was washed after 1 h, and then the DCs were added to the culture for a short period (because the effects of the drug are reversible). There was significant inhibition by the chloroquine pretreatment, indicating that the insulin vesicle had to be taken to acidic compartments of the phagocyte in order for their contents to be presented to the T cells (Fig. 2C). The control manipulation of testing APCs pulsed with insulin proteins or peptides established, as expected, that presentation of the free peptide was not affected by chloroquine, whereas the presentation of insulin was completely inhibited (Fig. 2D). Presentation of the peptide-MHC complex from the transferred vesicles, however, was short. Following a 1-h period of incubation of beta cells with DCs, the DCs were separated and T cells were added at various times. There was a progressive drop in presentation so that by 1–2 h there was a 50–75% decrease in the response (Fig. 2E). In sum, the transfer granules need to enter an acidic compartment of the APCs to release their contents of insulin or insulin peptide, but their time available for immune recognition is relatively short.

The previous findings indicate a close membrane-to-membrane interaction between beta cells and phagocytes that results in the transfer of the granules containing insulin or insulin products. We therefore investigated the potential role of Ca\(^{2+}\) in the mobilization of granules. We first examined if thapsigargin, a well-established inhibitor of the sarco/endoplasmic reticulum (ER) Ca\(^{2+}\) ATPase, may modulate the transfer of the granules. Antigen presentation was significantly increased by thapsigargin treatment of beta cells in low glucose media and was further enhanced in high glucose media (Fig. 3A). (In 10 different experiments, addition of thapsigargin increased by 211% over that of high glucose media (Fig. 3A).)

Both thapsigargin and high glucose increase cytoplasmic Ca\(^{2+}\) levels by depleting ER Ca\(^{2+}\) (26), raising the possibility that Ca\(^{2+}\) leakage from the ER, not the Ca\(^{2+}\) influx from the extracellular space, increases the transfer of granules. To test this idea, the assay was changed to study the requirements for extracellular or intracellular Ca\(^{2+}\). Beta cells were incubated with DCs for a limited time of 1 h, after which the DCs were separated from the beta cells: The content of immunoreactive insulin was then probed on the isolated DCs with the T cells. At 5 mM glucose concentration, the transfer was potentiated by the absence of Ca\(^{2+}\) in the transfer of the granules. We first examined if thapsigargin, a well-established inhibitor of the sarco/endoplasmic reticulum (ER) Ca\(^{2+}\) ATPase, may modulate the transfer of the granules. Antigen presentation was significantly increased by thapsigargin treatment of beta cells in low glucose media and was further enhanced in high glucose media (Fig. 3A).

In contrast, chelation of intracellular Ca\(^{2+}\) using 1,2-Bis(aminophenoxy)ethane-N,N,N',N'-tetraacetic acid tetrakis(acetoxyethyl ester) (BAPTA) resulted in complete inhibition, either at low or high glucose concentrations (Fig. 3C and D). The effect of BAPTA chelation was on the beta cells. The isolated DCs were removed and then pulsed with insulin presented to the T cells. This meant that insulin had been transferred in the beta cells and DCs plus BAPTA culture, it would have been presented when the DCs were separated. This conclusion was confirmed in the imaging experiments described below. Collectively, these results suggest that the transfer of granules required an increase in intracellular Ca\(^{2+}\) levels caused by ER Ca\(^{2+}\) depletion.

ER Ca\(^{2+}\) depletion is known to induce ER stress and activate the unfolded protein response. We therefore considered the possibility that the unfolded protein response may be involved in the transfer of granules. In our culture assay, during the several hours of culture of DCs with beta cells, we did not find up-regulation in the expression of any of the ER stress sensors: We evaluated a panel of canonical ER stress markers that are highly transcriptionally up-regulated during ER stress responses (27, 28) (Fig. S1 shows one representative result). In data not presented, the addition to the culture of tunicamycin, a drug known to increase the ER stress response, did not enhance presentation of insulin epitopes to T cells. (In unrelated experiments, the addition to beta cells of each of four cytokines—IFN-alpha A,
IFN-gamma, tumor necrosis factor, and interleukin-1 beta—had no effect on the transfer to either of the two CD4 T cells.)

**Live Imaging of Beta Cells and APCs and Electron Microscopy Examination of Islets.** The passage of granules was also documented by live images of the insulinoma line NIT expressing granules bearing ZnT8 bound to green fluorescent protein (GFP) (29). We had reported in a previous study on interactions between the insulinoma cell line NIT with DCs and T cells directed to ZnT8 the islet-specific membrane transport for Zn ions (29). DCs were labeled with CellTrace Violet (false colored red) and added to NIT-ZnT8-GFP cells. Images showed the DCs contacting the NIT; There was a striking and progressive enrichment of GFP+ granules to the contact area (Fig. 4A and Movie S1) in a synapse-like structure. After a contact period, some of the GFP+ granules appeared in the DCs, which eventually detached from the beta cells. The time of contact ranged between 20 and 30 min. Such images were found invariably in all DC–NIT interactions. A 3D reconstruction shown in Fig. S2 documents the contact area and passage of the granules to the DCs. DCs were examined after contact with NIT-ZnT8-GFP treated with BAPTA. There were few granules in the contact area and limited transfer of them into the DCs (Fig. 4A, Movie S2, and Fig. S2). The contact time in the NIT treated with BAPTA was considerably shorter—just a few minutes. These results paralleled those made in the transfer assay described above (Fig. 1E and F). A similar image of granule transfer was made in the single experiment done with bone marrow-derived macrophages (Movie S3).

Islets from young NOD mice including those from NOD.Rag1−/− mice were examined by electron microscopy. The intraislet phagocytes were found in close contact with beta cells and always next to blood vessels (Fig. 4B). All of the phagocytes had cytolsosomal vesicles, many of which contained material identical to beta cell secretory vesicles—that is, with a central electron-dense material, a light surrounding the zone, and a thin membrane; others had vesicles with irregular electron-dense material (Fig. 4C). The structural features indicate that the entire secretory vesicles from beta cells have been taken into the phagocytes. Fig. 4B and C is from an NOD 8 wk of age; Fig. 4D and E is from 14-wk-old NOD. Rag1−/− mice.

**Summary.** We show here that beta cells pass the contents of their secretory granules to phagocytes. The presentation assays indicated that the passage involved close interaction between live cells that took place under conditions of low glucose. The passage increased by addition of a high glucose concentration or the drug thapsigargin, both of which increase cytosolic Ca2+. The passage took place even in beta cells from mice not undergoing an immunological intervention such as those from NOD.Rag1−/− mice or C57BL/6 mice; it also took place with human islets obtained from nondiabetic subjects. The passage of granules and/or insulin antigens to phagocytes was also documented in vivo by ultrastructural analysis and by live imaging of beta cells and islet APCs (Fig. 4) (13,14). In toto, these findings point to a scenario in which secretory granules are being transferred in a constitutive process to the islet resident phagocytes.

The biological relevance of the beta cell–phagocyte interaction outside the context of an autoimmune reaction needs to be evaluated because it may be key in maintaining islet homeostasis. Secretory granules contain a number of bioactive molecules that could well modulate the biology of the phagocytes; phagocytes including tissue macrophages are known to release cytokines and chemokines. In the absence of macrophages, islets are of much smaller size, suggesting a trophic effect of macrophages on islets (12, 30, 31). We envision a symbiotic interaction between both cells: Experiments in progress indicate that islet macrophages are in an activated state (11, 31).

The nature of the passage of vesicles during the beta cell–APC interaction is of particular interest. In contrast to the release of insulin granule content taking place under glucose-regulated secretion, the passage involved the whole secretory granule, as was evident from three different results. The ultrastructural analysis showed images compatible with intact granules inside the phagocytes, without any evidence of death of beta cells or inflammation within the islets. The imaging studies pointed to a flow of ZnT8-GFP+ granules to areas of contact with the APCs, and a rapid flow of the granules into them. The immunological assays probed with the T cells to insulin required a chloroquine-sensitive stage in the APCs, indicating a catabolic process required to release the granule contents. Had free denatured insulin or B chain peptides...
passed to the APCs, there would not have been a requirement for a chloroquine-sensitive step (Fig. 2C). This finding is made more evident by the reactivity of the 8F10 T cell that recognizes catabolites of the insulin B chain. APCs that take up soluble denatured insulin or B chain peptide do not require intracellular catabolism, as is well established for diverse protein antigens. These findings are pointing thus to a close membrane interaction between the intraislet phagocytes that stimulated the movement of granules and its passage from one cell to another.

A number of the vesicles passed to the APCs must have contained insulin catabolites, some in the form of denatured insulin chains or peptides from the insulin B chain. We base this conclusion on the fact that we probed with a T cell that recognizes only insulin B chain catabolites offered exogenously to an APC. This T cell does not react with insulin epitopes after the processing of insulin by an APC (5, 6) (Fig. 1C). Moreover, the monoclonal antibody to insulin B chain peptide identified positive material inside the beta cell as well as in the islet APC. The importance of this finding is that the antibody exclusively reacts with the B chain peptide and not with native insulin, indicating that some intracellular vesicles contain degradative products of insulin. Extensive studies have unequivocally shown that beta cells contain vesicles bearing insulin catabolites, in a pathway referred to as crinophagy (32–36). It indicates an intracellular degradation of excess insulin granules, as a mode of controlling the homeostasis of insulin granule content (32, 37). The impression one obtains from these findings is that the process of generation of the peptide-positive—crinophagic—granules is taking place also constitutively, albeit increased by high glucose. We expect that as active diabetogenesis ensues this process may be modulated, such as a result of the stress conditions that accompanies diabetogenesis (38–40).

The issue from the perspective of autoimmunity is that the insulin catabolic products constitute the immunogens that select for autoreactive T cells that bypass insulin processing and escape thymus regulation (5, 23).

We suggest that the findings made here represent the first stage by which immunogenic autoantigens of beta cells are made available to the immune system. We previously reported that insulin-reactive T cells, such as the 8F10 used here, entered islets to contact the intraislet APCs, not requiring a prior activation
state in pancreatic lymph nodes (23). Subsequently, presentation of antigens in the pancreatic node represented an active amplification step for diabetogenesis. It is likely that the node is supplied by the movement of islet phagocytes. Recently we reported the seminal role of the CD103+ DCs in such a process (11).

Materials and Methods

Islet Isolation, T-Cell Assays. Islets were isolated following standard procedures described before (11). In brief, collagenase was injected into the pancreas via the common bile duct, after which islets were selected under microscopy with a pipettor. The islet cells were dispersed nonenzymatically and plated in a 12-well tissue culture plate at a density of 2.5 × 10^5 cells/mL in either Dulbecco’s minimal essential media with 10% (vol/vol) fetal calf sera containing either 25 mM or 5 mM glucose (referred to as high or low glucose, respectively). Depending on the experiment, cells were cultured for a few hours or overnight at 37 °C. Islets were obtained in most experiments from NOD.Rag1−/− mice of either sex 4–6 wk of age. Islets also were obtained from NOD mice 4–16 wk of age and C57BL/6 mice 8 wk of age. Two human islets were examined, both obtained from Prodo Laboratories. The patients were a 43-y-old male and a 34-y-old female, neither of whom had a history of diabetes. In most assays, 2 × 10^4 beta cells were incubated with 5 × 10^4 CD11c+ DCs in a total volume of 150 μL of a 96-well tissue culture plate for several hours (from 1 to 24 h depending on the nature of the experiment). T cells were then added for 24 h, and their response was assayed.

We used two reporter hybridoma T-cell lines: IIT-3, which recognizes INS protein and the INS B:13–21 peptide, and 8F10, which recognizes INS B:12–20. The production of IL-2 was measured by the stimulation of the IL-2–dependent line CTLL-2 (5). DCs were harvested from spleens of mice killed 10 d after injections with FLT-3L (10 μg for 3 consecutive days). The spleen cells were positively selected for CD11c+ cells using MACS beads (Miltenyi Biotec).

The following changes in the transfer assay of beta cells–APCs were made. To examine (i) the effects of anti–I-Aα7 antibodies, beta cells and DCs were cultured for a day, after which the anti–I-Aα7 antibody AG2.42.7 at 10 μg/mL and the T cells were added for another 24 h. To examine (ii) whether the transfer process involves cell-to-cell contact, a transwell system was used in which beta cells and DCs were separated via a 96-well transwell plate containing a 0.4-μm polycarbonate membrane (Corning). In control wells, beta cells were incubated with DCs in the lower chamber of the transwell. To examine (iii) the effects of beta cell viability, the beta cells were cultured with streptozocin (50 mM) first for 12 h before addition of the DCs. To examine (iv) the role of acidification of vesicles in the DCs, the DCs were

Fig. 4. Imaging and electron microscopy. (A) DCs labeled with CellTrace Violet (red) were added to NIT GFP-ZnT8-Insulinomas. Left panel (Movie S1) shows the contact area of NIT insulinomas in green with the DCs. Note the accumulation of GFP+ granules at the contact area. No movement of granules was observed in BAPTA/AM-treated NIT cells (Right panel and Movie S2). (B–E) Electron micrographs. B represents an islet from an 8-wk-old female NOD; C–E are islets taken from NOD.Rag1−/− mice at 14 wk of age. The arrow in B indicates a vessel. In the enlarged area, one of the dense-core granules is indicated by an arrow. In panel D a phagocyte is shown in between beta cells, and an arrow points to a dense-core granule. Panel E shows a portion of a phagocyte with endocytosed material in the form of vesicles containing an electron-dense core, with others containing amorphous content.
collected from mice injected with FLT-3L as described previously. The DCs examine (EGTA for 1 h, after which the DCs were isolated and cultured with the T cells. To carbonate, and 2 mML-glutamine. Before the examination, we replaced the imaging two-photon microscopy, we used the NIT-ZnT8-GFP insulinomas fixed in Karnovsky fixative, and processed by standard procedures. For live studies, we acquired 18 z-sections (1.25 μm apart) every minute using a customized SP8 two-photon microscope (Leica) equipped with a 25×0.95 N.A. water-dipping objective and a Mai Tai HP Deepsee Laser (Spectra-Physics) tuned by a pulse-picker. Transferrine emission was separated by high-efficiency custom dichroic mirrors (Semrock) and directed to supersonic external detectors. CellTrace Violet (Molecular Probes) was detected below 484 nm and GFP between 509 and 562 nm. Image sequences were then processed with Fiji (Huang et al., 2019) and Adobe Premiere software. The 3D reconstructions and sectioning were performed with Imaris software (Bitplane). For experiments studying the effects of BAFTA-AM on granule transfer, the above was modified slightly. NI T cell growth media was removed and replaced with fresh growth media containing 10 μM BAFTA-AM. The cells were incubated at 37 °C for 2 h. BAFTA containing media was removed from the dish and replaced with fresh growth media for 15 min at 37 °C. The DCs were then added as detailed above.

In experiments examining for EBS sensors, total RNA was isolated with Ambion RNAqueous-Micr o Kit (Life Technologies). cDNA was obtained from total RNA using First Strand Synthesis Protocol with Reverse Transcriptase (New England BioLabs). TaqMan PCR was performed using TaqMan Fast Universal PCR Master Mix. Primers for quantitative RT-PCR were designed using the PrimerTime predesigned quantitative PCR assays (IDT DNA). PrimerTime primers used S-nuclease detection. StepOne 2.1 software was used to perform quality control and the relative expression quantification. We examined an Hsp70 family member (Hsp, Hpa5), the ER-calcium regulator Wolfram syndrome 1 protein (WFS1), the transcription factor CEBP-homologous protein (CHOP, Ddx5), the activating transcription factor 4 (ATF4), and the growth arrest and DNA damage-inducible protein-34 (GADD34, Ppp1r15a).

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Fig. S1. ERS markers evaluation. Beta cells from NOD RAG1<sup>−/−</sup> mice were incubated in high or low glucose media for 1 h, 4 h, and 24 h. cDNA was synthesized from extracted mRNA. Hspa5 (BIP), Wfs1, Ddit3 (CHOP), Ppp1r15a (GADD34), and Atf4 were amplified with specific primers by quantitative RT-PCR. The fold change in gene expression was calculated using $2^{-\Delta\Delta CT}$. Bars are mean ± SD of biological duplicates. Thapsigargin (Tg), 0.1 μM, was a positive control for ERS induction. Results are representative of three experiments.

Fig. S2. (A and B) Granule accumulation in insulinomas at the contact side of DCs ("synapse"). A is a 3D reconstruction and computational sectioning of DCs at the beginning of synapse formation (Left) and at the point of a mature synapse (Right) (related to Fig. 4 and Movie S1). Green particles increase dramatically inside the DCs during the process of synapse generation (white arrows). B is a quantitation of synapse per DC at 30 min.
**Movie S1.** Contact of DCs with NIT-ZnT8-GFP. Note the granule accumulation in the contact area and flow of granules into DCs. 

**Movie S2.** Same as Movie S1, but the NIT was treated with BAPTA/AM, with less granule accumulation in the contact area and no flow into DCs.
Movie S3. Same as Movie S1 but using cultured bone marrow-derived macrophages instead of DCs as the APCs.