Heparan sulfate and heparanase play key roles in mouse β cell survival and autoimmune diabetes

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The autoimmune type 1 diabetes (T1D) that arises spontaneously in NOD mice is considered to be a model of T1D in humans. It is characterized by the invasion of pancreatic islets by mononuclear cells (MNCs), which ultimately leads to destruction of insulin-producing β cells. Although T cell dependent, the molecular mechanisms triggering β cell death have not been fully elucidated. Here, we report that a glycosaminoglycan, heparan sulfate (HS), is expressed at extraordinarily high levels within mouse islets and is essential for β cell survival. In vitro, β cells rapidly lost their HS and died. β Cell death was prevented by HS replacement, a treatment that also rendered the β cells resistant to damage from ROS. In vivo, autoimmune destruction of islets in NOD mice was associated with production of catalytically active heparanase, an HS-degrading enzyme, by islet-infiltrating MNCs and loss of islet HS. Furthermore, in vivo treatment with the heparanase inhibitor PI-88 preserved intraislet HS and protected NOD mice from T1D. Our results identified HS as a critical molecular requirement for islet β cell survival and HS degradation as a mechanism for β cell destruction. Our findings suggest that preservation of islet HS could be a therapeutic strategy for preventing T1D.

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Introduction

The NOD mouse strain spontaneously develops autoimmune type 1 diabetes (T1D) and is recognized as an experimental model for T1D in humans. The disease develops slowly in NOD mice, and the autoimmune pathology initially involves a nondestructive insulitis (NDI), in which mononuclear cells (MNCs) accumulate around the periphery of the islets. Autoimmune destruction of insulin-producing pancreatic β cells and T1D occurs when the insulitis MNCs become destructive and invade the islets (1). The trigger for this conversion is unknown. Although autoimmune diabetes in NOD mice is T cell dependent, it is unclear how β cells are destroyed once autoreactive T lymphocytes have entered the islets. Evidence suggests that CD8+ T cells recognize peptides derived from β cell–specific autoantigens (including proinsulin/insulin, GAD, IGRP, and chromogranin A) in the context of class I MHC molecules on the cell surface and kill the β cells via the perforin/granzyme pathway of cytotoxicity or induce apoptosis by Fas/FasL signaling (2–7). CD4+ T cells activated by autoantigen peptide/class II MHC complexes on intraislet APCs are likely to amplify islet inflammation by producing nonspecific inflammatory mediators, such as cytokines and chemokines. Intraislet APCs activated in the cytokine milieu could also indirectly damage β cells by producing ROS or cytokines that induce endogenous production of free radicals in the β cells (3).

Intervention therapies have been developed to impede the inflammatory response to islets in NOD/Lt mice. mAb treatment targeting CD4+ or CD3+ T cells has been particularly effective in preventing the development of T1D (8, 9). In the case of anti-CD4 mAb therapy, continual treatment was mandatory and induced CD4+ T cell depletion (9). Anti-CD3 mAb therapy rescued NOD mice from T1D, even when treatment was delayed until after T1D onset, and restored self tolerance after only transient T cell depletion (10). Other experimental therapies targeting cytokines including IL-16, IL-21, and TNF inhibited the recruitment of diabetogenic T cells to the pancreas, reduced insulitis, and prevented T1D (11–13). NOD islets in situ produce chemokines, particularly CCL5 (14), that recruit inflammatory cells, which suggests that β cells themselves could contribute to the initiation and expansion of peri-islet insulitis. Blockade of chemokine signaling via transgenic expression of a chemokine-blocking protein or decoy receptor by β cells has markedly decreased insulitis and T1D incidence in NOD mice (15, 16). Despite the development of effective strategies for reducing insulitis and preventing T1D in NOD mice, practical problems have impeded their clinical application. Notably, recent clinical trials have revealed inconsistent improvement in T1D control after anti-CD3 therapy, and long-term protection from disease progression remains an elusive milestone (17).

Intrinsic properties of β cells have been identified that render them particularly vulnerable to inflammatory insult. In addition to their capacity to secrete chemokines that could exacerbate peri-islet inflammation, islet β cells express low levels of free radical scavenger enzymes, potentially increasing their sensitivity to free radical–mediated damage (18). Conversely, the extent to which islets and β cells use intrinsic defense and survival mechanisms for their protection has largely been underexplored. We recently reported that in situ NOD mouse islets are surrounded by a continuous basement membrane (BM) containing the heparan sulfate proteoglycan (HSPG) perlecan (19). HSPGs consist of a core protein to which a number of side-chains of the glycosaminoglycan or complex sugar heparan sulfate (HS) are covalently attached. HS is a linear polysaccharide consisting of repeating disaccharides of glucosamine and glucuronic acid; there are large regions of N-acetylg glucosamine and glucuronic
acid interspersed with small regions containing sulfated glucosamine and iduronic acid residues. Although regions of the HS chains are characteristically sulfated, their chemical structure shows additional heterogeneity, varying in the position of O- and N-sulfation and in epimerization of glucuronic acid to iduronic acid (20, 21). HSPGs in extracellular matrices (ECMs) and on the surface of cells act as reservoirs for growth factors, cytokines, and chemokines. However, when localized in BMs, HSPGs have the additional function of obstructing cell migration (20). This property suggests that the islet BM containing the HSPG perlecan would constitute a primary defense mechanism against invasion by destructive insulitis (DI) MNCs.

To migrate to sites of inflammation, such as peri-islet insulitis in the pancreas, leukocytes need to first extravasate from blood vessels and then produce a panel of degradative enzymes to solubilize the matrix components of the subvascular endothelial BM. Of these, heparanase (encoded by HPSE) plays a particularly critical role because it is the only known mammalian endoglycosidase that can efficiently degrade the HS side-chains of HSPGs (20, 22). In support of this, inhibition of heparanase using sulfated oligosaccharides or modified heparins has been shown to prevent inflammation and limit metastasis of experimental tumors (20, 22). Likewise, we have previously observed that MNC entry into islets, which marks the onset of destructive autoimmunity in NOD mice, correlates with degradation of the islet BM (19).

However, to our knowledge, a role for heparanase in the repertoire of effector molecules produced by insulitis MNCs has not previously been investigated.

In this study, we report that a protective role for HS extends beyond that of the islet BM (19). We observed remarkably high levels of HS in normal islets in situ. While investigating the HS content of pancreatic islet BMs, we identified remarkably high levels of intracellular HS within the islets of conventional mouse strains, young NOD mice, and immunoincompetent NOD/SCID mice. HS is highly expressed within pancreatic islets in situ, and HS loss in vitro correlates with β cell death. (A) Detection of HS in NOD/SCID pancreatic islets in situ by Alcian blue histochemistry and immunohistochemistry using HepSS-1 anti-HS mAb. No background staining with isotype control mouse IgM was observed. Scale bars: 50 μm. (B) Intra-islet HS content (quantified by Image J with Color Deconvolution plugin) showed that the area of intraislet staining with Alcian blue was significantly higher in islets in situ (n = 50) than after isolation (n = 45; P < 0.0001). (C) Isolated BALB/c islet showing loss of intraislet HS by Alcian blue staining. Scale bar: 50 μm. (D) Intraislet HS staining was further reduced in BALB/c islet β cells during culture, as demonstrated by FACS staining using HepSS-1 (anti-HSlo; blue histogram) and 10E4 (anti-HShi; pink histogram) mAbs or an isotype control IgM (green histogram); black histogram denotes islet cell autofluorescence. Loss of β cell viability after culture for 2 days, as measured by Sytox green uptake (dead/dying cells), is shown at right. SSC-W, side scatter width. (E) Loss of intracellular HSlo and HShi and increase of cell surface HShi and HShi was seen in β cells cultured for 2 days. *P = 0.0022; †P < 0.0001; **P = 0.0225; ‡P = 0.0202. (F) Loss of intracellular HS during culture for 2 days, as in E, correlated with a significant increase in percent Sytox green+ β cells (P < 0.0001). Data are mean ± SEM (n = 6).

Results

HS is highly expressed in mouse islets in situ. While investigating the HS content of pancreatic islet BMs, we identified remarkably high levels of intracellular HS within the islets of conventional mouse strains, young NOD mice, and immunoincompetent NOD/SCID mice. In this study, we report that a protective role for HS extends beyond that of the islet BM (19). We observed remarkably high levels of HS in normal islets in situ, primarily as a result of its unusual intracellular localization within β cells. In vitro, we showed that HS functioned as an important survival mechanism for β cells, with loss of β cell HS representing a mechanism of β cell death. Surprisingly, HS replacement provided protection from free radical–mediated cell death. We showed that in vivo, destructive autoimmunity was associated with degradation of islet HS by heparanase produced by insulitis MNCs and was prevented by administration of a heparanase inhibitor. We therefore identified endogenous HS as a β cell target for autoimmune damage, with heparanase as a mediator of β cell autoimmunity, findings we believe to be novel. Our results suggest that heparanase inhibition may be a promising therapeutic strategy for protecting β cells from HS depletion and T1D progression.

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Loss of HS from β cells in vitro correlates with β cell death. Compared with islets in situ in the pancreas, Alcian blue histochemistry showed the HS content of islets was significantly reduced by more than 50% after their isolation in vitro (Figure 1, B and C). After enzymatic dispersion of the islets to single-cell suspensions, flow cytometry analyses showed that isolated β cells stained for both highly sulfated HS (HS\(^{hi}\); identified by the 10E4 mAb) and undersulfated HS (HS\(^{lo}\); identified by the HepSS-1 mAb), but the remaining HS was localized mainly intracellularly, with low levels of HS present on the cell surface (Figure 1D). Furthermore, the majority of isolated islet cells were insulin-positive β cells (Supplemental Figure 2). Cell surface HS was significantly increased 2- to 3-fold, and intracellular HS was significantly decreased to 67% of preculture levels after 2 days of culture (Figure 1, D and E), which suggests that cell surface HS may be replenished from intracellular HS stores. The loss of intracellular HS during culture was accompanied by a substantial (>2-fold) increase in β cell death, as determined by uptake of Sytox green, a DNA dye that stains dead and dying cells (Figure 1, D and F). These findings suggested that β cells may require intracellular HS for their survival.

HS replacement protects isolated β cells from culture-induced and ROS-induced cell death in vitro. (A) Flow cytometry analysis of isolated β cell viability after culture for 2 days in the absence (control) or presence of 50 μg/ml heparin, HS\(^{hi}\), or PI-88. Viability was assessed by Sytox green uptake or by calcein-AM (viable and apoptotic cells) and PI (dead and apoptotic cells) uptake (see Supplemental Table 1). CB, counting beads. Percent cells is shown for the respective regions. (B) Absolute number of β cells and number of dead β cells in 2 day cultures as in A. The number of β cells was calculated using counting beads. See Supplemental Table 1 for statistical analyses of the same day 2 cultures (note that percent Sytox green+ cells approximated percent PI-+ cells). (C) Coculture with HS\(^{hi}\), but not HS\(^{lo}\), protected isolated islet β cells from 2 day culture–induced cell death, as determined by percent Sytox green+ cells and intracellular insulin staining of β cells. GMFR for insulin staining was compared with serum control. *P < 0.05 vs. control. (D) Ability of β cells cultured in the presence or absence of heparin (50 μg/ml) for 1 hour or 1–2 days to resist H\(_2\)O\(_2\)-induced (i.e., ROS) cell death (see Table 1). Data in A and D are representative of 3–5 separate experiments.
was confirmed by uptake of calcein-AM, a fluorescent dye that labels viable and apoptotic cells, and by staining with propidium iodide (PI), a DNA-binding dye that labels dead and apoptotic cells. In the presence of heparin, the highly viable calcein ‘PI’ population of β cells was increased from 25% to 86%, the dead population (calcein ‘PI’) reduced from 18% to 1%, and the apoptotic population (calcein ‘PI’) decreased from 52% to 6% (Figure 2A). HSlo and the HSlo mimetic PI-88 (27, 28), at 50 μg/ml, were just as effective as heparin at promoting β cell survival (Figure 2A and Supplemental Table 1), although only heparin was active at 5 μg/ml (Supplemental Figure 3). All 3 compounds decreased the absolute number of dead β cells by 6- to 14-fold compared with controls, despite comparable total cell numbers in the cultures (Figure 2B).

In contrast to HSlo, treatment with HSlo do not protect the β cells (Figure 2C). The effect of heparin treatment on β cell viability was also independent of the source of heparin (porcine mucosal or bovine lung; Supplemental Figure 4, A and B) and occurred after continuous culture with heparin for either 1 or 2 days (Figure 2D and Supplemental Figure 4C), or after pulsing with heparin for 1 hour followed by culture for 2 days in the absence of heparin (data not shown). Flow cytometry analyses, together with confocal microscopy, showed that β cell protection correlated with the intracellular localization of considerable amounts of FITC-labeled heparin in 86% of β cells (Supplemental Figure 5).

Importantly, heparin-treated β cells became extraordinarily resistant to ROS-induced death, generated by acute treatment of the cells with H2O2. Whereas H2O2 treatment of control β cells after 1 hour of culture significantly increased cell death from 35.5 ± 4.2% to 88.5 ± 1.0%, β cells cultured with 50 μg/ml heparin for 1 or 2 days showed excellent viability in the presence of ROS (10.8 ± 1.1% and 8.0 ± 1.1% cell death, respectively; Table 1 and Figure 2D). Similarly, HSlo and PI-88, but not HSlo (Supplemental Figure 6), also protected β cells from ROS-induced cell death. These results demonstrated that intracellular uptake of HSlo mimetics, such as heparin and PI-88, protected β cells from both culture-induced death and death induced by exogenous ROS. This also suggests that culture-induced death of β cells may be mediated by endogenous ROS.

Pancreatic islets are susceptible to damage by heparanase. Isolated islets with residual intraislet HS were examined for their susceptibility to damage by heparanase, the only known mammalian endoglycosidase that can cleave HS (29). Treatment of isolated islets with 20 μg/ml exogenous human heparanase for 24 hours resulted in peripheral islet damage with increased intraislet cell apoptosis (Supplemental Figure 7). In contrast, the same treatment did not induce apoptosis or death of either B16 melanoma cells or L929 fibroblasts (data not shown). While heparanase was therefore not universally toxic for all types of cells, our data strongly suggest that islets are particularly susceptible to heparanase-mediated damage. To investigate the in vivo relevance of this susceptibility, islets undergoing different levels of autoimmune damage in NOD mice were then examined for HS and heparanase expression. We observed by histochemistry that islets lacking insulitis or with ND1 contained widespread and well-preserved HS (Figure 3A). In contrast, islets damaged by DI showed extensive disruption of islet HS, as detected by Alcian blue staining, particularly in close proximity to infiltrating MNCs (Figure 3A). Likewise, prediabetic NOD/Lt mice and NOD/Lt mice at diabetes onset showed a significant reduction in islet HS content compared with intact adult NOD/SCID islets (Figure 3B). When islets from prediabetic and onset diabetic NOD/Lt mice were isolated with their associated insulitis MNCs and examined by quantitative real-time RT-PCR for Hps transcripts, we observed a 4- to 5-fold increase in transcripts in these islets compared with young NOD islets with low insulitis and a 6- to 8-fold increase compared with NOD/SCID islets lacking insulitis (Figure 3, C and D). This finding correlated with an approximately 28-fold increase in mRNA for the common leukocyte marker Cd45 compared with NOD/SCID islets (Figure 3E), which suggests that the increased Hps mRNA expression was derived from insulitis MNCs. This conclusion was confirmed immunohistochemically, with heparanase protein being strongly expressed by DI MNCs (Figure 3F). Heparanase exists as a 65-kDa proenzyme that is proteolytically cleaved to 8-kDa and 48- to 50-kDa polypeptides that dimerize to form the active enzyme (30–32).

Western blotting analysis revealed substantially higher levels of the 48-kDa active mouse heparanase in late prediabetic (9–10 weeks of age; 6-fold) and onset-diabetic NOD islets (9-fold) compared with NOD/SCID islets, with prediabetic NOD islets from 7- to 8-week-old mice containing mainly proheparanase (compared with late prediabetic and onset-diabetic NOD islets; Figure 3G). This finding suggests that insulitis MNCs gain the ability to generate enzymatically active heparanase at the same time as they become able to mediate destructive disease. We also noted that CBA/H, NOD/SCID, and young NOD islets contained substantial Hps transcript levels, 20- to 33-fold those of normal CBA/H kidneys (Figure 3D). Heparanase protein was also detected by immunohistochemistry — albeit weakly — at or near the cell surface of islet β cells, with Western blotting indicating low levels of the enzyme, in NOD/SCID islets (Figure 3, F and G). These findings suggest that active heparanase produced by invading insulitis MNCs, but not inactive heparanase endogenously produced within islets, contributes to autoimmune islet β cell damage during T1D development.

Inhibition of heparanase protects NOD mice from destructive autoimmune and T1D. The sulfated phosphomanno-oligosaccharide PI-88 (Muparfostat; Progen Pharmaceuticals) is a HS mimetic that is a non-cleavable competitive inhibitor of heparanase (27, 29) and a potent inhibitor of tumor growth (27) and is in clinical trials as an anticaner drug (33). To test whether heparanase plays a direct role in the development of autoimmune diabetes, we treated 10.5- to 11-week-old prediabetic female NOD mice (presumably exhibiting both ND1 and DI) daily with PI-88. Compared with control saline-treated mice, PI-88 treatment significantly delayed development of T1D by

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### Table 1

<table>
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<th>1 day</th>
<th>2 days</th>
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<td>83.5 ± 2.6A</td>
<td>10.8 ± 1.1C</td>
<td>8.9 ± 1.1C</td>
</tr>
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β cells were cultured in the presence or absence of 50 μg/ml heparin for 1 hour, 1 day, or 2 days and then treated or not with 30% H2O2 (as a source of ROS) for 5 minutes. Sytox green uptake was used to assess percent cell death by flow cytometry (n = 4 per group). A P < 0.05 vs. untreated control culture. *P < 0.05 vs. untreated heparin control. **P < 0.05 vs. H2O2-treated control culture.
10 weeks, and the proportion of mice with diabetes at 36 weeks of age was reduced by 50% (Figure 4A). At the end of the experiment, islets in the pancreas from PI-88–treated and saline-treated nondiabetic mice were examined to assess the level of insulitis. Compared with saline-treated controls, PI-88–treated mice showed a significant reduction in islet-associated HS in prediabetic (PD) and onset-diabetic (OD) NOD/Lt mice compared with NOD/SCID (Sc) mice. Y, young 4- to 5-week-old NOD/Lt donors. n = 60–70 islets/group, 6–7 pancreases/group. Data are mean ± SEM. *P < 0.001. (C) Stereomacroscopic view of prediabetic NOD/Lt islets showing a clear, well-defined boundary of an intact islet and islets with insulitis, indicated by a translucent appendage of insulitis MNCs (arrows). (D) Hpsae and (E) Cd45 transcript expression in isolated islets, expressed relative to CBA/H kidney (K) and NOD/SCID islets, respectively. N, normal CBA/H donors. Results are mean ± SD (n = 3 per group). *P < 0.0001. (F) Heparanase immunohistochemistry showed strong expression of heparanase protein by DI MNCs in a diabetes-onset NOD/Lt pancreas (HP130 mAb). Note weak cell surface expression of heparanase (arrows) on NOD/SCID and diabetes-onset NOD/Lt islet cells. Scale bars: 100 μm. (G) Western blot analysis of mouse proheparanase (prohpse; 1453 pAb; 65 kDa) and active heparanase (hpse; HP130 mAb; 48 kDa) in protein extracts from pooled islets (10 donors/group). Heparanase (50 ng) purified from human platelets (Hu Hpse) was included as a 50-kDa standard.

Discussion

We report here that pancreatic β cells have a uniquely abundant intracellular store of HS, which plays a role in β cell survival that we believe to be previously unrecognized. During islet
isolation, approximately 50% of the islet HS content was lost, and during culture of isolated β cells, there was a progressive decline in intracellular HS that correlated with increased β cell apoptosis and death. Surprisingly, HS replacement — achieved by culturing β cells with the highly sulfated HS analog heparin and confirmed by uptake of FITC-labeled heparin — rescued the β cells from apoptosis and dramatically improved β cell survival. Furthermore, the transition from calcine^+PI^+ apoptotic β cells to calcine^+PI^- viable cells was confirmed by loss of staining with 7-amino-actinomycin D (7AAD), a DNA-binding dye conventionally used for identifying apoptotic cells (data not shown). Rescue by HS replacement with heparin, HS\textsubscript{hi}, or PI-88 not only prevented culture-induced cell death, but also rendered the β cells remarkably resistant to death induced by free radicals (i.e., ROS), which suggests that β cell death in vitro may be mediated by excessive levels of free radicals produced endogenously in the β cells. Overall, these findings suggest that in situ, normal levels of endogenous intracellular HS may function, at least in part, to protect β cells from free radical–induced cell injury.

Heparin imported into β cells in vitro and endogenous β cell HS in situ (i.e., in the pancreas) may interact directly with ROS and function as an antioxidant or free radical sink (34–36). The depolymerization of HS by free radical species, for example, could represent an antioxidant mechanism for protecting β cells from free radical damage. In vitro, exposure of rat glomerular BM HS to ROS generated by hypoxanthine/xanthine oxidase reactions results in HS fragmentation or depolymerization, detected by PAGE (34). Similarly, hydroxyl radicals have previously been shown to efficiently depolymerize heparin to low–molecular weight heparins (35). Alternatively, free radicals may be chemically detoxified by removing the anomeric hydrogen of internal sugar residues of HS, as previously reported for polyelectrolyte configurations of glucose, such as phosphorylated or sulfated glucans (36). Unlike depolymerization, a role for HS as a free radical sink would be
expected to better preserve other biological functions of the HS chains. β cells are highly metabolically active and consequentially generate considerable levels of intracellular ROS via disulfide bond formation during insulin biosynthesis in the endoplasmic reticulum or during oxidative phosphorylation or disulfide bond formation for protein folding in mitochondria (37, 38). β cell HS could therefore provide a constitutive mechanism for detoxifying endogenous ROS and thereby compensate for the reported low levels of free radical scavenger enzymes in β cells (18).

We found that inadvertent depletion of intracellular HS during isolation of islets and β cells in vitro rendered β cells particularly vulnerable to damage induced by exogenously delivered ROS and that HS replacement via treatment of β cells with heparin or other HS mimetics in culture provided resistance to ROS. Islets have been reported to undergo matrix detachment-induced apoptosis (anoikis) during their enzyme-mediated isolation (39, 40), and this process in epithelial cells correlates with ROS induction (41). Taken together, these properties suggest a scenario in which, during islet/β cell isolation, islets undergo matrix detachment that substantially elevates endogenous ROS levels in β cells to levels exceeding the protective capabilities of HS, resulting in HS degradation and apoptosis. We further speculate that providing additional exogenous ROS (by acute treatment with H₂O₂) at this time may precipitate β cell apoptosis and death. Like β cells, apoptosis induced in rat kidney glomerular mesangial cells by chronic treatment with H₂O₂ for 24 hours was prevented by pretreatment with heparin or HSPG. Explant-induced apoptosis of glomerular cells was similarly protected by acute treatment with heparin in vitro (42), and we speculate that this early damage may also be attributable to matrix detachment. Furthermore, ultra–low–molecular weight heparin has been reported to prevent glutamate-induced apoptosis in brain cortical cells in vitro and to prevent free radical–induced damage after ischemia/reperfusion injury in rat brain in vivo (43, 44). Our demonstration that HS replacement by heparin rescued β cells from free radical damage and apoptosis in vitro is consistent with these prior reports and with the notion that islets/β cells in situ express exceptionally high levels of endogenous HS in order to execute similar pro-survival functions in vivo. Apoptosis in β cells is attributed to imbalanced expression of pro- and antiapoptotic genes of the Bcl-2 family (45–47), which suggests that HS may also function in regulating gene expression in β cells, as previously reported for other cell types (48, 49). In any case, our findings strongly implicate HS preservation during human islet isolation or HS replacement as strategies for improving islet viability and for reducing the number of islets (and donors) required for clinical islet transplantation as a treatment for established T1D.

In addition to playing a critical role in β cell survival, HS may also perform other homeostatic functions as a reservoir and coreceptor for essential growth factors (50) and has previously been reported to regulate the postnatal development of β cell function (26). Under conditions of autoimmune disease, we demonstrated here that islet HS acts as a critical target for destruction by heparanase. This finding builds on our earlier observation that islets are surrounded by a continuous BM containing the HSPG perlecan, with MNC entry into the islets during destructive autoimmunity being accompanied by degradation of perlecan in the islet BM (19). Thus, DI MNCs produce catalytically active heparanase that — by solubilizing the islet BM HS — allows intra-islet MNC invasion, induces β cell death, and initiates autoimmune diabetes. Paradoxically, we also showed that islet β cells endogenously produced heparanase, which probably has a homeostatic function in regulating the turnover of HS (50) and possibly gene expression (48, 49).

Overall, our data are completely consistent with the idea that T1D is heparanase dependent, since T1D is an autoimmune disease mediated by insulitis MNCs, PI-88 efficiently inhibited heparanase, and PI-88 treatment altered autoimmune–associated pathology in NOD mice. Although the drug can also disrupt interactions between HS and certain growth factors (27), this secondary role is unlikely to enhance β cell survival. In further support of a causal role for heparanase in T1D, we also found that more specific heparanase inhibitors that have minimal growth factor binding and anticoagulant activity were as effective as PI-88 at protecting approximately 50% of NOD mice from developing T1D (data not shown). Heparanase may nevertheless affect T1D development at multiple sites. First, heparanase is likely to function at the level of facilitating MNC migration across the subvascular endothelial BM in the pancreas (20) and, possibly, subsequent passage through the ECM of the pancreas. Secondly, at the level of the islet microenvironment, we propose that the production of catalytically active heparanase by insulitis MNCs and degradation of islet BM HS signals the onset of destructive autoimmunity. Thereafter, migration of insulitis MNCs into the targeted islets results in local intraislet production of active heparanase, progressive loss of β cell HS, β cell apoptosis, loss of insulin production, β cell death, and ultimately T1D. Likewise, PI-88–mediated inhibition of heparanase could therefore inhibit T1D development at each tier of heparanase involvement. However, long-term treatment of NOD mice with PI-88 resulted in a significant increase in the percentage of intact islets (without insulitis) and a significant decrease in the percentage of islets with DI (Figure 4B), suggestive of drug-mediated protection occurring at both vascular and peri-islet sites.

Heparanase has previously been shown to function in disease by regulating BM breakdown and ECM remodeling via HS degradation. These actions facilitate cell migration in inflammation, spread of metastatic tumor cells and their reestablishment as solid tumors at secondary sites, and glomerular BM degeneration in diabetic nephropathy (29, 51, 52). In addition to these more standard functions, our present study identified that heparanase produced by insulitis MNCs plays a direct and fundamental role as an effector mechanism in T1D by degrading β cell HS, a process that leads to β cell death. Our findings unveil a critical role for islet HS in β cell survival and highlight the potential for new therapeutic approaches for rescuing β cell function at the time of T1D onset by HS replacement therapy, as well as for blocking progressive loss of islet-associated HS during disease development with heparanase inhibitors, both of which could be achieved using appropriately designed HS mimetics.

**Methods**

Further information can be found in Supplemental Methods.

**Mice and T1D monitoring.** Specific pathogen–free female NOD/Lt mice (4–5 weeks old) were obtained from the Animal Resource Centre (Perth, Australia), and other strains were obtained from the Animal Services Division of The John Curtin School of Medical Research or The Australian National University Bioscience Facility (Canberra, Australia). Onset of clinical diabetes was determined by measuring urine glucose twice weekly with Multistix reagent strips (Bayer) and confirmed by measuring nonfasting blood glucose levels in tail vein blood using a MediSense glucometer (Abbott Laboratories). Hyperglycemia was defined in our study as 2 consecutive blood glucose readings of 16 mmol/l or greater.
In vivo treatment of prediabetic NOD/Lt mice with PI-88. We treated NOD/Lt female mice at 10.5–11 weeks of age daily for 180 days by i.p. injection with 10 mg/kg PI-88 (provided by Progen Pharmaceuticals) or with 8 μl/g saline diluent.

Islet isolation. Islets were isolated as previously described (53, 54). Briefly, islets were isolated from anesthetized donor mice (Avertin, 0.025 ml/g body weight) by initially perfusing the pancreas in situ via the pancreatic duct with 2.5 mg/ml collagenase P (Roche Diagnostics). The digested pancreas was digested further with an additional 2.5 mg/ml collagenase P in a stationary water bath at 37°C. The digested tissue was then briefly shaken, vortexed lightly, placed on ice to terminate enzymatic digestion, and subsequently washed 3 times. Islets were hand-picked from the digested pancreas tissue with the aid of a dissecting microscope (Kyowa Optical SDZ-P). For isolation of islets from prediabetic and diabetes-onset NOD/Lt donors, an inverted microscope (Olympus CK) was also used to help distinguish islets with insulins from dissociated acinar tissue. We photographed islets showing attached insulins using an inverted microscope (Leica Microsystems).

Preparation and culture of isolated β cells. We dispersed BALB/c mouse islets into single-cell suspensions using the method of Jøsefelsen et al. (55), modified to use 1 mg/ml Dispase II (Roche) in the chelation buffer, and subsequently cultured the islet cells on uncoated plastic. The cells were cultured at 2.5–4 × 10⁶ cells/well in CELLSTAR 96-well plates (Greiner Bio-one) in RPMI 1640 medium (Gibco; Invitrogen), supplemented with 10% fetal calf serum (SAFC Biosciences), in the presence or absence of heparin from bovine lung (Sigma-Aldrich) or porcine intestinal mucosa (Celsus Laboratories), PI-88 (provided by Progen Pharmaceuticals), porcine mucosal HSα (HI-11098; Celsus Laboratories) or HSα (HSO-10595; Celsus Laboratories) at 0.5–50 μg/ml for 1 hour to 2 days in 5% CO₂, 95% air, at 37°C. During culture, the isolated β cells adhered to the hydrophilic tissue culture-treated plastic surface of the wells in the culture plates. β Cells were treated with 30% H₂O₂ (Chem-Supply) for 5 minutes at 37°C as a source of ROS and were used as a positive control for analysis of β cells on death day 0.

Flow cytometry. We assessed cell death by incubating β cells with 31.25 nM Sytox green (Invitrogen) for 15 minutes at 37°C. The region identifying dead cells was defined using H₂O₂-treated cells as positive controls. Cell viability was also analyzed by staining initially with calcein-AM (0.04 μM; Invitrogen) and subsequently with propidium iodide (PI; 2.5 μg/ml; BD Biosciences — Pharmingen), each for 15 minutes at 37°C. Viable cells were identified as calcein−PI−, apoptotic cells as calcein−PI+, and dead cells as calcein−PI+. When PI was replaced with 7AAD (10 μg/ml; Molecular Probes) in the presence of calcein, a similar proportion of apoptotic cells was observed. 10E4 and HepSS-1 mouse mAbs (Seikagaku), sheep anti-mouse Ig-R–phycoerythrin (Chemicon) or goat anti-mouse Ig-R–phycoerythrin (Southern Biotech), BD Cyanofix/Cytoperm Fixation/Permeabilization Kit (BD Biosciences — Pharmingen), and mouse IgM (BD Biosciences — Pharmingen) as isotype control were used for HS staining of β cells, gated for viability using forward and side scatter properties. Flow-Count Fluorospheres (Beckman Coulter) were added (4,000 beads/cell sample) prior to staining to determine cell numbers. The geometric mean fluorescence intensity ratio (GMFR) was calculated by dividing the geometric mean fluorescence intensity of cells stained with primary Ab by that of cells stained with the isotype control. Cells were analyzed using a BD LSRI flow cytometer and CellQuest software (version 6.0, BD Biosciences).

Western blotting. Islets were lysed in 1% CHAPS (3-[3-cholamiddopro pyl]dimethylammonio)-1-propanesulfonate; Sigma-Aldrich) containing 50 mM EDTA, 50 μM E64 (Sigma-Aldrich), 1 mg/ml Pefabloc SC (Roche Diagnostics), and Complete Protease Inhibitor Cocktail Tablets (EDTA-free, at ≥20 recommended concentration; Roche Diagnostics) at pH 8.0. The protein concentration of islet lysates was measured using Bio-Rad protein Dye Reagent (Bio-Rad Labs). Lysates (300 μg/sample) were incubated with concanavalin A–sepharose beads (Pharmacia) for 2 hours, washed, and then boiled in reducing buffer prior to SDS-PAGE. Marker proteins (Precision Plus Protein Standards, Dual Color; Bio-Rad) and 50 ng purified human platelet heparanase (56) were run in parallel. Nitrocellulose membranes were probed with HP130 mAb (Insight Bio- pharmaceuticals) or affinity-purified rabbit anti–recombinant human proheparanase polyclonal Ab 1453 (provided by I. Vlodavsky, Rappaport Faculty of Medicine, Technion, Haifa, Israel) (57) followed by HRP-conjugated rabbit anti-mouse Ig (Dako) or HRP-conjugated swine anti-rabbit Ig (Dako). Immunoreactive proteins were detected using ECL Western Blotting Reagent (Amersham) and a LAS-1000 Chemiluminescence and Fluorescence Imaging System (Fuji Photo Film).

Real-time PCR. For analysis of heparanase, CD45, and ubiquitin conjugating enzyme E2D1 (UBC, endogenous reference gene) transcripts in isolated islets, we used validated TaqMan real-time gene expression assays (Applied Biosystems, catalog nos. Mm00461768_m1, Mm00448463_m1, and Mm00461307_g1). The mean Ct values for UCB between experimental groups were 27.6–29.9. Quantification of changes in transcript expression was calculated using the comparative Ct method as previously described (58), which corrects for any variation between samples in template input by normalization to the endogenous reference gene.

Histology and immunohistochemistry. Pancreas and islet samples were fixed in 10% neutral-buffered formalin, and paraffin sections (4 μm) were prepared. Sections were stained with H&E or with Alcian blue (0.65 M MgCl₂, pH 5.8; Sigma-Aldrich) for selective identification of HS glycosaminoglycans (refs. 23, 59, and Supplemental Figure 1) and counterstained using safranin. For quantitative studies of NOD/Lt islet integrity, H&E-stained semiserial sections of each pancreas specimen (minimum interval of 48 μm between sections analyzed) were examined by 2 observers in a blinded fashion, and the percentage of islets with each descriptor (normal, NDI, DI, or completely destroyed) were quantified and expressed as mean ± SEM of the total islets counted in the pancreas sections. Image J software (version 1.44a; NIH) with Color Deconvolution plugin (60) was used to quantify the intensity of Alcian blue staining. For measurement of the intensity (optical density), images were deconvoluted using the plugin to separate the Alcian blue staining component from the counterstain safranin. The mean optical density of Alcian blue staining was then determined by converting the mean pixel value for islet tissue (including insulitis) to optical density, using the calculation log₂(S/mean pixel value). For analysis of islets from saline- and PI-88–treated NOD/Lt mice, 3 sections per specimen were analyzed for 3 (saline) or 4 (PI-88) specimens, with the minimum interval between sections analyzed being 48 μm. For assessment of area of Alcian blue staining, BALB/c islets in each specimen (n = 50 from 4 pancreas specimens; 1 section/specimen), BALB/c islets after isolation (n = 45 from 3 pooled islet specimens; 1 section/specimen), and NOD/Lt islets in situ at different stages of autoimmune disease and NOD/SCID islets in situ (n = 6–7 pancreas specimens/group; 3 sections/specimen) were analyzed, with the minimum interval between sections analyzed being 64 μm. We localized HS (10E4 or HepSS-1 mAbs) and heparanase (HP-130 mAb) by immunohistochemistry after antigen retrieval using 0.05% pronase protease (Calbiochem) for HS (61) or heat-induced standard antigen retrieval for heparanase (62), using the M.O.M. immunodetection kit (Vector Laboratories). Control sections were incubated with isotype control Ig.

Statistics. For statistical analyses of differences between groups, we used 2-tailed, unpaired Student’s t test, Mann-Whitney test, or ANOVA with Kruskal-Wallis post-test for histological analyses; Mann-Whitney test for real-time PCR analyses; Fisher exact test for cumulative incidence of diabetes; and Student’s t test, Mann-Whitney test, or ANOVA with Bonferroni post-test for flow cytometry data analyses. A P value less than 0.05 was considered statistically significant.
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