RAGE binds preamyloid IAPP intermediates and mediates pancreatic β cell proteotoxicity

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Islet amyloidosis is characterized by the aberrant accumulation of islet amyloid polypeptide (IAPP) in pancreatic islets, resulting in β cell toxicity, which exacerbates type 2 diabetes and islet transplant failure. It is not fully clear how IAPP induces cellular stress or how IAPP-induced toxicity can be prevented or treated. We recently defined the properties of toxic IAPP species. Here, we have identified a receptor-mediated mechanism of islet amyloidosis–induced proteotoxicity. In human diabetic pancreas and in cellular and mouse models of islet amyloidosis, increased expression of the receptor for advanced glycation endproducts (RAGE) correlated with human IAPP–induced (h-IAPP–induced) β cell and islet inflammation, toxicity, and apoptosis. RAGE selectively bound toxic intermediates, but not nontoxic forms of h-IAPP, including amyloid fibrils. The isolated extracellular ligand–binding domains of soluble RAGE (sRAGE) blocked both h-IAPP toxicity and amyloid formation. Inhibition of the interaction between h-IAPP and RAGE by sRAGE, RAGE-blocking antibodies, or genetic RAGE deletion protected pancreatic islets, β cells, and smooth muscle cells from h-IAPP–induced inflammation and metabolic dysfunction. sRAGE-treated h-IAPP Tg mice were protected from amyloid deposition, loss of β cell area, β cell inflammation, stress, apoptosis, and glucose intolerance. These findings establish RAGE as a mediator of IAPP-induced toxicity and suggest that targeting the IAPP/RAGE axis is a potential strategy to mitigate this source of β cell dysfunction in metabolic disease.

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Introduction

The misfolding and aggregation of soluble proteins into amyloid fibrils plays a key role in a range of human diseases (1–8). Pathological aggregation of the endocrine hormone human islet amyloid polypeptide (h-IAPP, also known as amylin) is a key feature in islet amyloidosis. h-IAPP is cosecreted with insulin and plays an adaptive role in metabolism (6, 13–16), but in T2D, it aggregates by an unknown mechanism and is deposited as pancreatic islet amyloid plaques associated with reduced β cell volume (2, 4, 7, 8, 17, 18). Aggregation of h-IAPP into amyloid fibrils involves 3 observable stages: Pre-amyloid oligomers (or prefibrillar intermediates) formed in the lag phase (LP) (first phase) assemble into amyloid fibrils in the growth phase (GP) (second phase), leading to an equilibrium between amyloid fibrils and residual soluble peptide in the saturation phase (SP) (third phase) (Supplemental Figure 1, A and B; supplemental material available online with this article; https://doi.org/10.1172/JCI85210DS1). Toxic h-IAPP aggregates cause pancreatic β cell and islet dysfunction and death, contributing to T2D, islet transplant failure, and, ultimately, cardiovascular and microvascular complications (19–23).

Islet amyloidosis is characterized by the aberrant accumulation of islet amyloid polypeptide (IAPP) in pancreatic islets, resulting in β cell toxicity, which exacerbates type 2 diabetes and islet transplant failure. It is not fully clear how IAPP induces cellular stress or how IAPP-induced toxicity can be prevented or treated. We recently defined the properties of toxic IAPP species. Here, we have identified a receptor-mediated mechanism of islet amyloidosis–induced proteotoxicity. In human diabetic pancreas and in cellular and mouse models of islet amyloidosis, increased expression of the receptor for advanced glycation endproducts (RAGE) correlated with human IAPP–induced (h-IAPP–induced) β cell and islet inflammation, toxicity, and apoptosis. RAGE selectively bound toxic intermediates, but not nontoxic forms of h-IAPP, including amyloid fibrils. The isolated extracellular ligand–binding domains of soluble RAGE (sRAGE) blocked both h-IAPP toxicity and amyloid formation. Inhibition of the interaction between h-IAPP and RAGE by sRAGE, RAGE-blocking antibodies, or genetic RAGE deletion protected pancreatic islets, β cells, and smooth muscle cells from h-IAPP–induced inflammation and metabolic dysfunction. sRAGE-treated h-IAPP Tg mice were protected from amyloid deposition, loss of β cell area, β cell inflammation, stress, apoptosis, and glucose intolerance. These findings establish RAGE as a mediator of IAPP-induced toxicity and suggest that targeting the IAPP/RAGE axis is a potential strategy to mitigate this source of β cell dysfunction in metabolic disease.

Substantial evidence reveals that both extracellular and intracellular h-IAPP oligomers contribute to islet β cell toxicity. Postmortem histological studies of human subjects with T2D show predominantly extracellular pancreatic islet amyloid deposition (4, 6, 17). Rodents do not develop islet amyloidosis, as rodent IAPP is nonamyloidogenic and nontoxic (17, 24, 25), but a variety of Tg h-IAPP rodent models have been created that form either extracellular or intracellular islet amyloid, or both (26). Mice that overexpress h-IAPP demonstrate intracellular oligomer formation and defects in autophagy and/or ER stress (27–30), while cultured Tg murine islets expressing physiological levels of h-IAPP do not display ER stress during islet amyloidosis (31). Extracellular h-IAPP oligomers are toxic (6, 19, 32–35). Findings that h-IAPP secretion is necessary for β cell proteotoxicity and amyloid formation (32), studies that show extracellular h-IAPP oligomers can be translocated into β cells (36), and receptor-mediated mechanisms of cytotoxicity (33) all support a role for extracellular oligomers in h-IAPP–induced β cell and islet pathogenesis. Here, we focus on islet amyloidosis–induced proteotoxicity by extracellular h-IAPP oligomers.

Amyloidogenic h-IAPP induces β cell apoptosis in vitro, in rodent models in vivo, and in human T2D (10, 19, 26, 34, 35, 37, 38). Multiple mechanisms of toxicity have been proposed, but there is no consensus about how h-IAPP causes β cell damage (6, 17, 39). Local islet inflammation and the production of cytokines and chemokines have been suggested to contribute to toxicity, as have the disruption of cell membranes by h-IAPP aggregates; impairment of the ER–associated protein degradation/ubiquitin/proteasome (ERAD/ubiquitin/proteasome) pathway and ER stress; defects in autophagy; activation of NADPH-oxidase and consequent oxida-
highly effective therapeutic agent to prevent both h-IAPP toxicity and amyloid formation in vivo. These findings highlight RAGE as a therapeutic target for β cell preservation in metabolic disease and provide critical information for the design of inhibitors of h-IAPP/RAGE interactions.

**Results**

RAGE is upregulated in h-IAPP–induced β cell and islet proteotoxicity. We have shown that toxic h-IAPP LP intermediates, but not nontoxic h-IAPP amyloid fibrils, induce the production of ROS in INS-1 β cells and that this event precedes the detection of metabolic dysfunction (19). If RAGE is a mediator of h-IAPP–induced β cell toxicity, then we would expect that toxic h-IAPP LP intermediates would likewise upregulate RAGE expression before detectable loss of β cell viability. This was indeed the case. We prepared solutions of h-IAPP LP intermediates, determined that they were prefibrillar by the amyloid-sensitive dye thioflavin-T and by transmission electron microscopy (TEM) (Figure 1, A and B), and incubated them with rat INS-1 β cells for a short period of time (1 hour), which does not provoke loss of cell viability (Supplemental Figure 3A) (19), as well as a longer incubation time (5 hour), which is required to detect metabolic dysfunction in cellular toxicity assays (Figure 1C). The results demonstrated that exogenous h-IAPP LP intermediates upregulated β cell RAGE protein expression within 1 hour of peptide incubation on cells, preceding the detection of β cell dysfunction (Supplemental Figure 3, A and B). We observed similar results during high-glucose–induced β cell stress. INS-1 β
Amyloid fibrils did not increase RAGE expression (Figure 1, C and D). Treatment of cells with either nontoxic rat IAPP (r-IAPP) or nontoxic h-IAPP elevated RAGE protein levels (Figure 1, C and D). Treatment of incubation resulted in significant cellular toxicity along with β cell stress. These results are consistent with a role for RAGE in detectable glucotoxicity (Supplemental Figure 4, A and B). These cells incubated at a high glucose concentration (16.7 mM) showed an increase in RAGE protein levels compared with cells cultured at a standard glucose concentration (11.7 mM), which also preceded an increase in RAGE protein levels compared with cells cultured at a high glucose concentration (16.7 mM) showed an increase in RAGE protein levels compared with cells cultured at a high glucose concentration (16.7 mM) showed an increase in RAGE protein levels compared with cells cultured at a high glucose concentration (16.7 mM) showed an increase in RAGE protein levels compared with cells cultured at a high glucose concentration (16.7 mM) showed an increase in RAGE protein levels compared with cells cultured at a high glucose concentration (16.7 mM) showed an increase in RAGE protein levels compared with cells cultured at a high glucose concentration (16.7 mM) showed an increase in RAGE protein levels compared with cells cultured at a high glucose concentration (16.7 mM) showed an 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sRAGE is an inhibitor of amyloid formation by h-IAPP. We hypothesized that sRAGE would inhibit h-IAPP amyloid formation, since it selectively binds toxic LP intermediates. We added sRAGE to h-IAPP at different time points over the course of aggregation (Figure 2A) and monitored the effect of sRAGE on h-IAPP secondary structure formation: $t_0$ species (black), ELP species (orange), MLP species (green), LLP species (purple), and SP amyloid fibrils (blue). CD spectra of samples were recorded after a 48-hour incubation. h-IAPP readily formed amyloid by itself in the absence of sRAGE (red), but sRAGE inhibited β-sheet formation when added to h-IAPP at time points before the formation of toxic LP intermediates or when they were present, demonstrating that inhibition by sRAGE was sustained and not transient. The inhibitory effects of sRAGE on h-IAPP β-sheet formation decreased as it was added to h-IAPP at later time points in the LP. Representative TEM images show the morphology of protein species at time points assessed by difference-CD. Amorphous sRAGE alone, h-IAPP SP amyloid fibrils, and the effect of the addition of sRAGE on different h-IAPP kinetic species: ELP species, LLP species, and SP amyloid fibrils. Scale bars: 200 nm. The final peptide concentration in biophysical experiments was 20 μM. Thioflavin-T, CD, and TEM data are representative of 3 to 5 independent experiments. Data in A represent the mean ± SD of 3 to 6 technical replicates per time point. Error bars for some data points are smaller than the size of the symbols. incub, incubation.
tion in each sample was then monitored by thioflavin-T binding, difference-CD, and TEM. Aliquots from the original h-IAPP and sRAGE stock solutions were measured at the same time points, serving as control solutions.

A decrease in temperature increases the length of the LP of amyloid formation, facilitating the identification of distinct events in the LP. Thus, we performed the sRAGE/h-IAPP binding experiments at lower temperatures (15°C) than those conducted in the absence of sRAGE depicted in Figure 2C (25°C) to better resolve the effect of adding sRAGE to different h-IAPP LP species. The data indicate that addition of sRAGE to ELP or MLP species prevented amyloid formation but that sRAGE had no effect when added during the GP or SP (Figure 3A). These data are supported by difference-CD and TEM studies (Figure 3, B–H). The data show that addition of sRAGE to h-IAPP at or before time points of toxic species formation prevented β-sheet formation, as detected by difference-CD (Figures 3B and Supplemental Figure 7, A–F), and amyloid formation, as detected by TEM (Figure 3, C–F). The data also show that sRAGE had less of an effect when it was added later in the LP (Figure 3, B and G). Thus, sRAGE spe-

**Figure 4.** sRAGE inhibits h-IAPP-induced inflammatory gene expression in β cells. qPCR analysis of INS-1 β cells treated (5 hour) with h-IAPP LP intermediates, a 1:1 molar ratio of h-IAPP/sRAGE, sRAGE alone, r-IAPP, or buffer. The final peptide concentration after transferring aliquots into β cell assays was 14 μM. Data represent the mean ± SEM of 3 to 4 independent experiments (3–9 technical replicates per experiment) and are presented as the fold change relative to buffer-treated β cells. *P < 0.05 and ****P ≤ 0.0001, by 1-way ANOVA.
We conducted WB experiments using INS-1 β cells in parallel with qPCR, thiocflavin-T, and cell viability studies and observed that the upregulation of β cell inflammatory markers and proteotoxicity by h-IAPP LP intermediates was RAGE dependent. WB data in Figure 5 show that preamyloid h-IAPP intermediates significantly increased RAGE protein expression and that blocking h-IAPP/sRAGE interactions using sRAGE prevented h-IAPP-induced RAGE upregulation. We obtained similar results using RAGE-expressing primary murine aortic SMCs (Supplemental Figure 8A).

We next tested the effect of blocking cellular h-IAPP/sRAGE interactions on h-IAPP-induced toxicity toward RAGE-expressing murine pancreatic islets, β cells, and SMCs using 3 independent strategies: treatment with sRAGE, treatment with RAGE-blocking antibodies (IgGs) that directly block cellular RAGE/ligand interactions, and genetic deletion of RAGE (Ager−/−). We found that the addition of sRAGE protected pancreatic islet cells, INS-1 β cells, and SMCs from h-IAPP-induced metabolic dysfunction (Figure 6, A and B, and Supplemental Figure 8B). Pretreatment of INS-1 β cells with increasing concentrations of anti–RAGE IgG (blue bars, Figure 6C) inhibited metabolic dysfunction by h-IAPP LP intermediates in a dose-dependent manner, unlike what we observed with control IgG (red bars, Figure 6C). Primary pancreatic islets isolated from Ager−/− mice (blue bars, Figure 6, D and E) showed significant protection from h-IAPP toxicity compared with islets from age-matched RAGE-expressing (Ager+/+) WT mice (red bars, Figure 6, D and E). We obtained similar results when we added h-IAPP LP intermediates to cultured WT (red bars, Supplemental Figure 8C) versus Ager−/− (blue bars, Supplemental Figure 8C) murine aortic SMCs. Together, the results indicate that RAGE plays a key role in mediating h-IAPP toxicity to cells and pancreatic islets.

RAGE is upregulated in hemi-h-IAPP mice in concert with proinflammatory gene expression and β cell stress and apoptosis, and treatment with sRAGE ameliorates islet amyloidosis-induced pathology and metabolic dysfunction. To examine the role of RAGE in h-IAPP-induced β cell toxicity in vivo and assess the potential efficacy of sRAGE as a therapeutic agent for β cell and islet preservation in islet amyloidosis, we used hemizygous Tg mice that express RAGE, overexpress h-IAPP (hemi-h-IAPP mice), and develop β cell degeneration and islet amyloidosis pathology but not diabetes (26, 53). This prediabetic model decouples the role of RAGE in islet amyloidosis–induced β cell perturbation from the multiple other stress-provoking factors that exist in the complex setting of diabetes, including hyperglycemia-mediated formation of other RAGE ligands such as advanced glycation endproducts (AGEs) (49). The slow development of early-stage islet amyloidosis in this model facilitates the gradual accumulation of prefibrillar h-IAPP aggregates, allowing us to test the hypothesis that toxic h-IAPP intermediates cause RAGE-mediated β cell defects.

We treated hemi-h-IAPP and WT mice with sRAGE (100 µg/day) or PBS 6 days a week for 10 months to inhibit in vivo h-IAPP/RAGE interactions. Metabolic characterization confirmed defective β cell function in the hemi-h-IAPP mice at 12 months of age, but no diabetes, as previously reported (26, 53). The data show that PBS-treated hemi-h-IAPP mice (referred to hereafter as Tg/PBS mice) were mildly but significantly glucose intolerant compared with age-matched and weight-matched WT mice, but no hyperglycemia or significant differences in circulating insulin levels were specifically targets species that reside early in the LP and blocks amyloid formation by sequestering them and preventing their further assembly into fibrils.

RAGE significantly contributes to h-IAPP-mediated cellular proinflammatory gene expression and cytotoxicity. Our finding that toxic h-IAPP LP intermediates bound RAGE and upregulated β cell RAGE expression led us to predict that LP intermediates would modulate inflammatory gene expression and cytotoxicity in cultured β cells and muscle cells, as h-IAPP–induced islet cell and cardiomyocyte inflammation have been reported (40, 42, 44, 45, 51). We also expected that sRAGE would compete with cell-surface RAGE for h-IAPP binding, as it targets toxic LP intermediates and is a competitive inhibitor of ligand binding to cell membrane–bound RAGE. To test this, we used INS-1 β cells and primary murine aortic smooth muscle cells (SMCs) as model systems (50, 52). We prepared h-IAPP LP intermediates, a 1:1 molar mixture of h-IAPP/sRAGE, and multiple controls including sRAGE, r-IAPP, and buffer-only solutions. We then characterized each sample and added aliquots to cultured cells for concurrent real-time quantitative PCR (qPCR) and Western blot (WB) studies.

We found that h-IAPP LP intermediates significantly upregulated mRNA transcripts of Ager (the gene encoding RAGE), Ccl2, Cxcl1, Cxcl2, Il1b, Il18, and Tnfa, and that treatment of the β cells with sRAGE during h-IAPP challenge reduced proinflammatory gene expression. h-IAPP LP intermediates exerted no significant effect on the regulation of antiinflammatory Il10. SAGE alone, like r-IAPP and buffer, had no independent effect on the regulation of any of the examined inflammatory mediators (Figure 4).
observed (Figure 7, A–E). Quantitative histological analyses of pancreatic tissue revealed an increase in mean islet diameter in Tg/PBS mice compared with the WT groups (Figure 7F). Immunofluorescence studies of pancreas sections costained for S100B-insulin or AGEs-insulin confirmed the absence of these RAGE ligands in the β cells and islets of both treatment groups of Tg and WT mice, while positive control pancreas sections from diabetic high-fat diet–fed (HFD-fed) WT mice showed AGE and S100B immunoreactivity (Supplemental Figure 9 and Supplemental Figure 10).

Treatment of hemi_h-IAPP mice with sRAGE (referred to hereafter as Tg/sRAGE mice) protected them from islet amyloidosis–induced pre-diabetic defects in β cell function associated with glucose intolerance (Figure 7, A–E), and morphological analyses showed a significantly lower mean islet diameter compared with that observed in islets from Tg/PBS mice (Figure 7F). Collectively, the results confirmed that hemi_h-IAPP mice develop β cell dysfunction and that sRAGE inhibits h-IAPP toxicity and islet pathology in vivo.

qPCR studies in whole pancreas samples revealed a similar induction pattern of inflammatory mediators in Tg/PBS islets compared with the WT groups (Figure 7F). Immunofluorescence studies confirm the upregulation of RAGE immunoreactivity in Tg/PBS islets compared with WT/PBS islets, as well as significant protection of Tg/sRAGE mice from islet amyloid deposition, β cell stress, and apoptosis. Pancreas sections costained for thioflavin-S and insulin indicated modest amyloid severity in Tg/PBS mice (Figures 9, A and B, and Supplemental Table 1), along with a significant decrease in the mean islet β cell area (from 72.4% ± 3.6% in WT/PBS mice to 59.3% ± 2.7% in Tg/PBS mice) (Figure 9C and Supplemental Table 1). In contrast, we found that amyloid deposition in Tg/sRAGE mice-induced upregulation of proinflammatory gene expression, as indicated by significantly lower Cxcl2 and Il1b mRNA transcript levels compared with Tg/PBS mice and a nonsignificant trend toward lower Ager, Cxcl2, and Il18 gene expression. Upregulation of other inflammatory processes was also observed in Tg/sRAGE mice, as indicated by a significant increase in Il10 and Tnfa mRNA transcript levels (Figure 8). We observed no change in Ccl2 levels in Tg/sRAGE mice compared with levels in Tg/PBS mice. These findings confirm in vivo relevance for sRAGE as an inhibitor of islet amyloidosis–induced modulation of inflammatory cytokines and chemokines.

Immunofluorescence studies confirm the upregulation of RAGE immunoreactivity in Tg/PBS islets compared with WT/PBS islets, as well as significant protection of Tg/sRAGE mice from islet amyloid deposition, β cell stress, and apoptosis. Pancreas sections costained for thioflavin-S and insulin indicated modest amyloid severity in Tg/PBS mice (Figures 9, A and B, and Supplemental Table 1), along with a significant decrease in the mean islet β cell area (from 72.4% ± 3.6% in WT/PBS mice to 59.3% ± 2.7% in Tg/PBS mice) (Figure 9C and Supplemental Table 1). In contrast, we found that amyloid deposition in Tg/sRAGE mice.
was markedly reduced and that the β cell area was preserved to a degree comparable to that observed in WT mice (Figures 9, A–C, Supplemental Figure 11A, and Supplemental Table 1). These findings are consistent with our in vitro data demonstrating inhibition of h-IAPP toxicity and amyloid formation by sRAGE.

Mild islet amyloid deposition in Tg/PBS mice was accompanied by a significant increase in islet β cell stress and apoptosis compared with WT/PBS mice, as assessed by costaining with cleaved caspase-3 (CC3) and insulin and triple staining with TUNEL, insulin, and DAPI, respectively (Figure 9, D–G, Supplemental Figure 11, B and C, and Supplemental Table 1). These results are in line with previous studies showing increased β cell apoptosis in Tg mouse models overexpressing h-IAPP (26). Treatment of Tg mice with sRAGE led to a significant reduction in β cell stress and apoptosis compared with that seen in PBS-treated Tg mice (Figure 9, D–G and Supplemental Table 1). We observed no statistically significant differences between sRAGE-treated and PBS-treated WT mice, confirming that sRAGE does not have independent effects, delivers no harm, and offers no additional protection (Figures 9, E and G, Supplemental Figure 11, B and C, and Supplemental Table 1). Pancreas sections from the same mice, costained for RAGE and insulin, showed a significant increase of approximately 1.9-fold in β cell RAGE expression in Tg/PBS versus WT/PBS islets (Figure 10, A and B, and Supplemental Table 1).

We observed a direct inverse relationship between islet β cell area and β cell stress/preapoptosis and apoptosis in Tg/PBS mice (Supplemental Figure 12, A and B, respectively), consistent with previous reports on islet amyloidosis (10). We found that loss of islet β cell area also directly correlated with β cell RAGE immunoreactivity in Tg/PBS mice (Figure 10C). These data link RAGE with islet amyloidosis–induced β cell toxicity and demonstrate that significant β cell apoptosis and a loss of β cell area do not require extensive islet amyloid deposition or diabetes, consistent with reports in other models (54). Together, the in vivo data support a role for h-IAPP-induced, RAGE-mediated islet defects in prediabetes and confirm that sRAGE is an effective pharmacological agent against islet amyloidosis–induced β cell stress and apoptosis.

h-IAPP upregulates β cell RAGE expression in human subjects with T2D and islet amyloidosis. We probed the relevance of our findings to human pancreas specimens bearing significant β cell stress using T2D and nondiabetic (ND) pancreas samples obtained from the Network for Pancreatic Organ Donors with Diabetes (nPOD). We observed thioflavin-S+ islet amyloid plaques in human T2D pancreas sections costained with insulin,
as has been previously reported (10). No amyloid was detected in ND human islets. h-IAPP immunoreactivity in T2D islets colocalized with thioflavin-S+ and insulin− islet areas, as well as thioflavin-S− and insulin+ islet areas, suggesting that the antibody detects different forms of h-IAPP (Supplemental Figure 13). We found that islet amyloid deposition in human T2D was accompanied by a significant increase in islet β cell RAGE immunoreactivity compared with ND human islets (Figure 11 and Supplemental Table 2). Human T2D RAGE+ islet areas colocalized with insulin+ and h-IAPP+ islet areas.

The RAGE staining pattern in human islets appeared different than that observed in murine islets. In both human and murine islets, we detected RAGE expression on the surface of cells. Close examination revealed that RAGE immunoreactivity in murine islets was more intense and punctate and located on insulin+, insulin−, and insulin− islet areas, consistent with the ubiquitous expression of RAGE, and was increased on insulin+ cells in Tg/PBS mice (Figure 10A). RAGE immunoreactivity in human islets colocalized predominantly with insulin+ cells in both diabetic and ND islets (Figure 11A). The ring-like pattern suggests that RAGE expression in human islets is more uniform and covers a larger area of the β cell surface than is seen in murine islets. The greater RAGE/insulin overlap in human diabetic islets may reflect the fact that these human subjects had longstanding T2D, while the mice were not diabetic. Thus, the differences in staining patterns may well be due to the presence of other glycemia-induced RAGE ligands in diabetic human islets.

We confirmed the absence of RAGE ligands (S100B and AGEs) in our murine Tg/PBS islets (Supplemental Figures 9 and 10) and their presence in diabetic human islets compared with ND islets (data not shown). Together, the human data support our in vitro, ex vivo, and in vivo findings and indicate that pancreatic islet amyloidosis is associated with the upregulation of β cell RAGE expression in T2D.

Discussion

Our data provide the first evidence to our knowledge that RAGE selectively binds to a distinct, transient population of toxic prefibrillar h-IAPP intermediates and plays a role in h-IAPP–induced cellular perturbation in pancreatic islet amyloidosis (Figure 12). Several lines of evidence strongly support this conclusion. First, RAGE was upregulated in β cells prior to cellular metabolic dysfunction, coincident with the upregulation of inflammatory mediators in vitro; in isolated murine islets after challenge with toxic h-IAPP intermediates, leading to loss of islet viability ex vivo; in...
a Tg mouse model of early islet amyloidosis with h-IAPP–induced glucose intolerance, loss of islet β cell area, and upregulation of inflammatory pancreatic mRNA transcripts in vivo; and in pancreatic islet β cells from human subjects with islet amyloidosis and T2D. Second, inhibiting RAGE activity by genetic Ager deletion, using blocking IgGs to prevent cellular h-IAPP/RAGE interactions, or sequestering h-IAPP intermediates with sRAGE to neutralize their toxicity protected cells and pancreatic islets from h-IAPP–induced inflammatory gene expression, metabolic dysfunction, and apoptosis. Third, sRAGE is an inhibitor of both h-IAPP toxicity and amyloid formation in vitro and in vivo. Fourth, RAGE binds only to toxic h-IAPP LP intermediates, while nontoxic forms of IAPP (r-IAPP, h-IAPP$^{t_0}$ species, and amyloid fibrils) do not bind RAGE. These observations, together with the findings of other studies that implicate a role for receptor-mediated mechanisms of toxicity, are consistent with extracellular islet amyloid formation but do not exclude the possibility of additional intracellular mechanisms of toxicity (6, 17, 33, 39).

Figure 9. Islet amyloid deposition, β cell stress, and apoptosis are suppressed by sRAGE treatment in hemi_h-IAPP mice. (A) Representative murine islets costained in pancreas for insulin (Ins, red) and thioflavin-S (Thio-S, green) show spatial overlap between insulin– (black) and thioflavin-S–islet areas. Scale bars: 50 μm. Quantitative immunofluorescence image analysis of (B) islet amyloid severity and (C) islet β cell area. (D) Representative murine islets costained in pancreas show colocalization (yellow) of insulin (red) and CC3 (green) immunoreactive islet areas. Scale bars: 50 μm. (E) Quantitative immunofluorescence image analysis of CC3–(stressed/preapoptotic) β cell area. (F) Representative murine islets costained in pancreas show colocalization (magenta) of TUNEL– (red) and DAPI– (blue) cell nuclei in insulin– (green) β cells. Scale bars: 50 μm and 8 μm (enlarged insets of β cell regions in white boxed areas in A, D, and F). (G) Quantitative immunofluorescence image analysis of TUNEL– (apoptotic) β cell area. Data represent the mean ± SEM of 3 mice per group. *P < 0.05, **P ≤ 0.01, ***P ≤ 0.001, and ****P ≤ 0.0001, by 1-way ANOVA.
creas samples was diluted by mRNA extracts from other cell types besides β cells, our data still demonstrate a potent in vivo suppression of multiple inflammatory mediators by sRAGE. Interestingly, treatment of hemi_h-IAPP mice with sRAGE did not lead to lower Ccl2 gene expression, suggesting a possible role for infiltration and/or alternative polarization of monocytes and macrophages in Tg/sRAGE mice. Further studies are needed to determine the role of infiltrating immune cells in the pathophysiology of islet amyloidosis and the pharmacology of sRAGE. Collectively, these results provide another connection between h-IAPP–induced toxicity and inflammation and suggest that amyloidogenic h-IAPP augments inflammation in pancreatic islet amyloidosis via receptor-mediated processes, such as recruitment of monocytes/macrophages and other myeloid cells, and activation of the RAGE pathway in β cells. These considerations are in line with studies supporting the hypothesis that h-IAPP induces toxicity by multiple and complementary mechanisms (17, 39).

Extensive studies have indicated that adaptive IAPP has a physiological role in the regulation of glucose homeostasis via paracrine mechanisms in the pancreatic islets as well as autocrine mechanisms and effects on the central nervous system, and that modulation of IAPP production and/or secretion in pathological settings leads to alterations in glucose metabolism (6). A number of in vivo and in vitro experiments have demonstrated that IAPP has an inhibitory effect on insulin secretion (62, 63) and glucagon secretion (64, 65). Studies have shown that the decoupling of IAPP and insulin gene expression and secretion in pathological conditions promotes IAPP overproduction and impaired glucose tolerance, but not insulin resistance (62, 66–68). The hemi_h-IAPP mouse model used in our studies over secretes h-IAPP, a process...
that is driven by the insulin II promoter, and recapitulates the h-IAPP hypersecretion observed in glucose intolerance. Previous studies have reported increased blood glucose levels in this Tg mouse model compared with levels in WT mice (53). We did not observe hyperglycemia in this model but did detect impaired glucose tolerance in the mice at 12 months of age, along with pancreatic inflammation, high levels of β cell stress and apoptosis, increased islet size, and a significant decrease in total islet β cell area in Tg/PBS versus WT/PBS mice, consistent with β cell compensation in response to increased insulin requirements in mice (69, 70).

While hyperplasia is not a typical characteristic of most h-IAPP Tg mouse strains, and the variation in islet sizes in the mice used in our studies made it difficult to conclusively diagnose hyperplasia, as defined by the Department of Health and Human Services National Toxicology Program (71), the observed increase in the size of Tg/PBS islets compared with WT control islets was statistically significant (P ≤ 0.001). This may be due in part to age and in response to an increased demand for insulin to control impaired glucose tolerance resulting from hypersecretion of h-IAPP. The late onset of observed β cell defects suggests that the alteration in glucose metabolism is a slow process in this model. In this case, the mice may have been able to compensate during the period before the onset of amyloidogenesis and the subsequent increase in apoptosis that was detected later at 12 months, which hindered the adaptive increase in β cell mass (54). Studies have shown that the total pancreatic β cell mass seen in rodents after birth is a balance between β cell replication from existing β cells, β cell production (differentiation) from other sources, and β cell apoptosis, and that the formation and maintenance of adult β cells depend largely on sources independent of β cell duplication (72), as the rate of β cell

Figure 11. RAGE immunoreactivity in human T2D islets colocalizes with insulin+ and h-IAPP+ β cells. (A) Representative images of pancreatic islets from T2D (nPOD ID 6124) and ND (nPOD ID 6011) human subjects. Islets were triple stained for insulin (red), h-IAPP (blue), and RAGE (green) and show colocalization (yellow and cyan) of RAGE+ islet areas with insulin+ and h-IAPP+ β cells. Scale bars: 50 μm and 8 μm (enlarged insets of β cell regions in white boxed areas). (B) Quantitative immunofluorescence image analysis shows a significant increase in the RAGE+ β cell area in T2D compared with ND islets. Data represent the mean ± SEM of 8 T2D subjects and 7 ND subjects. **P < 0.01, by 2-tailed Student’s t test.
replication in rodents declines with age (73). Our in vivo studies were not longitudinal and can only provide a snapshot at the time of sacrifice. Thus, we are limited in our ability to speculate about the rates of β cell proliferation, differentiation, or apoptosis other than to say that the rates of β cell compensation and apoptosis were probably not linear; rather, they progressively accelerated at different rates and different time points over the 12-month period, with advancing glucose intolerance and islet amyloidosis.

Our in vivo studies confirm the physiological relevance of our in vitro and ex vivo findings and are in line with previous reports that this hemi-h-IAPP mouse model develops β cell degeneration and early-stage pancreatic islet amyloidosis in the absence of diabetes (53). This 12-month-old mouse model decouples islet amyloidosis–induced metabolic dysfunction from the additive effects of hyperglycemia and indicates that the onset of islet amyloidosis occurs before the onset of diabetes and is not a consequence of the disease. The data presented in Figure 9, Figure 10, and Supplemental Figure 12, which together demonstrate a direct relationship between the loss of β cell area and β cell stress, apoptosis, and RAGE expression in Tg/PBS mice, and a lack of correlation between these parameters and islet amyloid area, are all normally distributed and statistically significant. The lack of correlation between islet amyloid severity and the extent of β cell stress, apoptosis, and loss of β cell area is in line with previous reports (54) and supports our other in vitro and in vivo findings that toxic preamyloid h-IAPP intermediates, which gradually form and build up during the slow LP of amyloid formation, are more deleterious than the final thioflavin T-positive amyloid fibrils. Our in vivo findings also establish a role for RAGE in islet amyloidosis–induced β cell proteotoxicity. Our human pancreas studies show an association between increased β cell RAGE expression and h-IAPP amyloidosis in human T2D. Together, our in vivo findings highlight the interaction of RAGE and h-IAPP intermediates as a primary target for β cell and islet preservation in metabolic disease.

The administration of sRAGE in vitro and in a murine model of pancreatic islet amyloidosis blocked toxicity, proinflammatory mediators, and amyloid formation; promoted anti-inflammatory processes; and preserved β cell area and overall β cell and islet morphology. We believe that the dual therapeutic action of sRAGE presents a new paradigm for antiamyloidosis agents. First, sRAGE sequesters and neutralizes h-IAPP intermediates, inhibiting their interaction with RAGE and other cell-surface binding sites, which in turn blocks consequent downstream events leading to toxicity. Second, sRAGE prevents the polymerization of toxic intermediates into amyloid fibrils. While fibrillar h-IAPP is not directly cytotoxic, the accumulation of amyloid plaque causes physical disruption in tissue architecture, elasticity, and homeostasis and thus may contribute to cumulative cellular perturbations.

Our studies define the mode of interaction between toxic h-IAPP species and cellular RAGE. Many RAGE ligands are anionic and are thought to bind to the basic patch on the receptor surface. h-IAPP is cationic and hydrophobic and would thus be expected to repel the basic patch. The data presented here, which indicate a close proximity of h-IAPP with the solvent-exposed sRAGE Trp residues, suggest that toxic h-IAPP intermediates probably bind to the hydrophobic patch on RAGE, highlighting this receptor region as an important site for the engagement of pathogenic species and a potential drug target. This is an important consideration for the design of inhibitors of RAGE activation in pancreatic islet amyloidosis, as the inhibitors of the extracellular RAGE domains that target the basic patch may be ineffective at blocking interactions with distinct hydrophobic ligands, such as h-IAPP.

In summary, we have identified a RAGE-mediated cellular mechanism of h-IAPP–induced β cell toxicity in pancreatic islet amyloidosis that precedes the development of hyperglycemia. Further, this work has unveiled an unanticipated dual therapeutic advantage of sRAGE as an inhibitor of both h-IAPP toxicity and amyloid formation itself. We believe these findings fill a critical gap in knowledge and advance our understanding in the field of IAPP biology, as well as across the field of amyloidosis diseases by identifying a new mediator of proteotoxicity that has significant therapeutic implications for β cell preservation and mitigation of consequent metabolic dysfunction.
Methods

Human pancreas studies. Sections of paraffin-embedded or frozen OCT-embedded pancreas specimens previously prepared from de-identified T2D (n = 8) and ND (n = 7) human subjects were obtained from nPOD. Tissue IHC-immunofluorescence (IHC-IF) methods were optimized directly in the control human pancreatic tissue. Clinical and immunohistological data for each subject can be found in Supplemental Table 2.

Mouse studies. Hemizygous Tg mice that overexpress the amyloidogenic h-IAPP–coding sequence under the regulation of the rat insulin II promoter 5’-UTR (The Jackson Laboratory) have been described previously (26, 53). Male mice were fed a normal chow diet and treated with 100 μl i.p. injections of either sRAGE (100 μg/ml) or PBS 6 days a week for 10 months to produce 4 age-matched study groups: Tg/PBS, Tg/sRAGE, WT/PBS, and WT/sRAGE. Body weight, fasting blood glucose, and i.p. glucose tolerance tests (ip-GTTs) were measured over the course of the study, and fasting plasma insulin concentrations in the Supplemental Methods.

IHC-IF. Six sections of formalin-fixed, paraffin-embedded pancreas specimens (4 μm thick and 30–200 μm apart) were labeled for each marker. All murine and human pancreas sections were stained with primary and secondary antibodies for detection of insulin, amyloid, C3C, TUNEL, RAGE, AGEs, S100B, and IAPP. Images were taken using a Leica fluorescence microscope. Quantitative analysis using MetaMorph LASF imaging software was performed by an investigator blinded to the experimental condition. Detailed methods for histological studies can be found in the Supplemental Methods.

Cell culture. The rat insulinoma cell line 832/13 (INS-1 β cells) was provided by Christopher Newgard (Duke University School of Medicine, Durham, North Carolina, USA) (74). Murine vascular SMCs were obtained from the aortas of 10-week-old male C57BL/6 mice (The Jackson Laboratory) or Ager−/− mice (52). Detailed cell culture methods can be found in the Supplemental Methods.

Islet isolation and culture. Pancreatic islets were isolated from anesthetized 12- to 18-week-old C57BL/6 mice (The Jackson Laboratory) or Ager−/− mice (52) according to institutional guidelines, hand purified, and cultured in supplemented RPMI 1640 medium, as described in the Supplemental Methods.

Protein preparation. IAPP peptides were prepared as previously described (19) or purchased from the KECK Foundation at Yale University (New Haven, Connecticut, USA). Human sRAGE was prepared via a baculovirus expression system using Sf9 cells (Clonetech, Invitrogen, Thermo Fisher Scientific) (75), as described in the Supplemental Methods.

Amyloid formation assays. Amyloid formation reactions were initiated as previously described (19) and were monitored using thioflavin-T binding assays. Aliquots were removed from stock solutions at various time points over the course of amyloid formation and further characterized by difference-CD and TEM.

Toxicity assays. INS-1 β cells were seeded at a density of 30,000 cells per well in 96-well plates 24 hours prior to experiments and incubated for 1 or 5 hours with aliquots of peptide solutions or controls. SMCs were seeded at a density of 300,000 cells per well in 6-well plates 36 hours before experiments, switched to serum-free DMEM 12 hours before experiments, and incubated for 10 hours with aliquots from peptide stock solutions or controls. Hand-purified islets of the same size and number (25 islets/well in 96-well plates) were incubated for 3 to 5 hours with peptide solutions or controls. Final peptide concentrations in cellular and islet assays were 14 μM. Cell and islet viability was assessed by Alamar Blue metabolic assays (76) and light microscopy, as previously described (19). Values were calculated relative to those of control cells or islets treated with buffer or r-IAPP. Data from islet experiments were normalized to the total number of islets per well. β Cell viability data represent numerous (>20) experiments carried out at different times by different laboratory members using multiple preparations of peptide stock solutions.

Light microscopy. Changes in cell and islet morphology were examined by light microscopy as an additional method of evaluating viability. Images were captured using an Olympus BX61 light microscope.

RAGE blockade assays. Blocking antibodies were produced at the Pocono Rabbit Farm & Laboratory, Inc. using previously described methods (77). INS-1 β cells were plated at a density of 30,000 cells per well in 96-well plates 24 hours before experiments. Cells were pretreated (2.5 hours) with either rabbit anti–human RAGE IgG or rabbit nonimmune IgG prior to challenge (5 hours) with toxic h-IAPP intermediates (14 μM). Cell viability was measured by Alamar Blue metabolic assays.

RNA isolation and qPCR. Total cellular RNA and tissue RNA were isolated from β cells, SMCs, or pancreas specimens using the RNeasy Plus Mini Kit or RNeasy Fibrous Tissue Mini Kit (QIAGEN). The quality of RNA was determined by measurement of a 260:280 ratio. RNA (1 μg) was reverse transcribed to cDNA with MultiScribe Reverse Transcriptase (Applied Biosystems). Real-time qPCR was performed using the TaqMan method (50°C for 2 min, 95°C for 10 min, and 40 cycles of 95°C for 15 seconds and 60°C for 1 min) with premade Ager, Ccl2, Cxcl2, Cxcl12, Il1b, Il18, Il10, Tfna, and Vcam1 primers (Life Technologies, Thermo Fisher Scientific). Relative mRNA contents were normalized according to the expression of 18S rRNA using the ΔΔCt method. qPCR was performed using an Applied Biosystems 7500 Real-time PCR machine.

WB analysis. Proteins (25–60 μg per lane) isolated from β cells or murine pancreatic islets were separated on a 4%–12% Bis-Tris NuPAGE gel (Bio-Rad) and transferred onto a PVDF membrane. Membranes were probed overnight (4°C) with the following primary antibodies: rabbit anti-RAGE (1:1,000; GeneTex; GTX23611) and anti-mouse GAPDH (1:1,000; Abcam; ab8245), followed by the following secondary antibodies (1 hour, 25°C): IRDye 800EW goat anti–rabbit IgG (1:1,000; Li-Cor; 925-32211) for detection of RAGE, and IRDye 680RD goat anti-mouse IgG (1:25,000; Li-COR; 925-68070) for detection of GAPDH. Protein signals were quantified using the Odyssey Infrared Imaging System (Li-COR; model 9120). RAGE protein signals were normalized to GAPDH protein signals in the same lane.

TEM. Samples (4 μl) were placed on a carbon-coated, 200-mesh copper grid and negatively stained with saturated uranyl acetate (Electron Microscopy Sciences). The samples were imaged with a Philips CM12 or a FEI BioTwinG2 transmission electron microscope.

Thioflavin-T binding assays. Aliquots (100 μl) of peptide solutions were added to 96-well plates containing 8 μl of a 1 mM thioflavin-T solution (Sigma-Aldrich). The final solution conditions contained 15 or 25 μM peptide, 20 mM Tris HCl, and 74 μM thioflavin-T (pH 7.4). Fluorescence was measured using a Beckman Coulter DTX880 plate reader (excitation, 445 nm; emission, 485 nm). Additional information can be found in the Supplemental Methods.
Far UV CD. Far UV CD was performed using an Applied Photo- 
physics spectrophotometer. Aliquots (300 μl) were transferred from 
amyloid formation assays to a 0.1 cm quartz cuvette within a few 
minutes prior to data collection. Spectra were recorded over a range of 
190 to 260 nm, and CD spectra represented the average of 5 repeats. 
Background spectra were subtracted. In difference-CD experiments, 
the spectra for solutions of sRAGE alone, which were collected at 
the same time as respective h-IAPP/sRAGE mixtures, were subtracted 
from the spectra of h-IAPP/sRAGE mixtures. The samples contained 
20 mM tris HCl (pH 7.4).

Trp fluorescence and SPR. sRAGE Trp fluorescence (280 nm excita-
tion; 350 nm emission) was measured using a Photon Technology Inter-
national instrument. The signal was an average of 20 reads over 20 sec-
onds (2.5 mm bandwidth and 1-second integration time). In SPR studies, 
sRAGE was immobilized on the sensor chip, and the interaction of dif-
f erent h-IAPP species with sRAGE was measured using a GE Healthcare 
SPR instrument. The final peptide concentrations were 20 μM h-IAPP 
or r-IAPP and 20 μM sRAGE in 20 mM Tris-HCl (pH 7.4, 25°C).

Statistics. Data analysis was carried out in collaboration with 
biostatisticians using appropriate statistical analysis methods deter-
mined by the distribution type and sample size of the data sets. Nor-
mally distributed data, as judged by the Shapiro-Wilk normality test, 
are presented as either mean ± SEM or mean ± SD of n independent 
experiments (as reported in each figure legend), as appropriate. Data 
that did not pass the Shapiro-Wilk normality test [Figure 8 (SI Figure panel 
only) and Supplemental Figure 4] are presented as mean ± SD. For 
multiple-group comparisons, normally distributed data were ana-
lyzed by one-way or two-way ANOVA, as appropriate, followed by 
post hoc Tukey HSD for pairwise comparisons, or post hoc paired 
t-test for a pre-selected set of group comparisons. For two-group com-
parisons, normally distributed data were first analyzed by the F-test 
to compare variances between two groups, followed by two sample 
tailed t-test with equal or unequal variance. Data that did not pass 
the Shapiro-Wilk normality test were assessed by the Kruskal-Wallis 
rank sum test followed by Dunn’s post hoc group comparison test. 
Data were corrected for multiple comparisons using the Benjamini-
Hochberg multiple comparison correction; an adjusted P value of 
<0.05 was considered significant. The significance of the linear 
regression analyses in correlation studies was determined by the 
Wald test; a P value of <0.05 was considered significant. For paired 
group comparisons, ANOVA with post hoc test results of P < 0.1 but 
> 0.05 were considered non-significant trends.

Study approval. All animal procedures were approved by the IAUC of 
New York University and performed in accordance with NIH animal care 
guidelines. All deidentified human studies were IRB exempt, as determined by the NYU School of Medicine IRB. nPOD is a research organization based in Gainesville, FL, that operates under the approval of the University of Florida’s IRB. This approval allows nPOD to work with Organ Procurement Organizations (OPOs) to obtain tissue from organ donors in the U.S. Adults who have registered while alive to become an organ donor upon their passing are considered to have granted consent for organ recovery for transplant and research. If the donor is a minor child, or is an adult who has not independently registered, the OPO approaches the legal next-of-kin for consent. The 58 OPOs in the U.S. operate regionally, and each have a version of a patient consent form. OPOs within the U.S. are members of the Organ Procurement and Transplantation Network (OPTN) and are overseen by the United Network for Organ Sharing (UNOS).

Author contributions
AA designed the research, conducted experiments, analyzed data, and wrote the manuscript. PC prepared critical reagents and assisted with experiments. AP, JZ, MH, JD, SAP, and JL assisted in vitro experiments. RR and FS assisted with in vivo experiments. HK and HL analyzed data. DPR and AMS designed and directed research, analyzed data, and wrote the manuscript.

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