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Glucose-dependent partitioning of arginine to the urea cycle protects β -cells from inflammation

Accalia Fu^{1,2}, Juan Carlos Alvarez-Perez³, Daina Avizonis⁴, Tatsuya Kin⁵, Scott B. Ficarro^{1,6,7,8}, Dong Wook Choi^{1,2}, Esra Karakose³, Mehmet G. Badur⁹, Lindsay Evans¹, Carolina Rosselot³, Gaelle Bridon⁴, Gregory H. Bird^{10,11}, Hyuk-Soo Seo^{1,12}, Sirano Dhe-Paganon^{1,12}, Jurre J. Kamphorst⁹, Andrew F. Stewart³, A. M. James Shapiro⁵, Jarrod A. Marto^{1,6,7,8}, Loren D. Walensky^{10,11}, Russell G. Jones¹³, Adolfo Garcia-Ocana³, Nika N. Danial^{1,14,15,*}

¹Department of Cancer Biology, Dana-Farber Cancer Institute, Boston, MA, USA

²Department of Cell Biology, Harvard Medical School, Boston, MA, USA

³Diabetes, Obesity and Metabolism Institute, Department of Medicine, Division of Endocrinology, Diabetes and Bone Disease, Icahn School of Medicine at Mount Sinai, New York, NY, USA

⁴Rosalind and Morris Goodman Cancer Center Metabolomics Core, Montreal, Canada

⁵Clinical Islet Transplant Program, Department of Surgery, University of Alberta, Edmonton, Canada

⁶Blais Proteomics Center, Dana-Farber Cancer Institute, Boston, MA, USA

⁷Department of Oncologic Pathology, Dana-Farber Cancer Institute, Boston, MA, USA

⁸Department of Pathology, Brigham and Women's Hospital, Harvard Medical School, Boston, MA, USA

Data availability

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^{*}Correspondence and requests for materials should be addressed to Nika N. Danial (nika_danial@dfci.harvard.edu). Author contributions

A.F. and N.N.D. designed experiments, performed and analyzed metabolomics, biochemical, and cell-based studies. T.K. and A.M.J.S. provided human donor islets. D.A., M.G.B., G. B., J.J.K. and R.G.J. provided expertise in metabolomics. A.G.-O. and A.F.S. provided expertise in β -cell biology. J.C.A.-P., C.R. and A.G.-O. carried out human islet xenotransplantation studies, and together with E.K purified human β -cells. E.K. and A.F.S performed RNAseq analysis on purified human β -cells. S.B.F. and J.A.M. carried out proteomic analysis on purified human β -cells. G.H.B. and L.D.W. designed and synthesized stapled peptides. H-S.S. and S.D. performed ITC analyses. D.W.C performed peptide pull down studies. L.E. provided technical support with mouse islet isolation and *in vitro* studies. A.F. and N.N.D. wrote the manuscript, which was reviewed by all authors.

Competing interests

The authors declare the following competing interests: J.A.M. serves on the SAB of 908 Devices, and has received sponsored research support from AstraZeneca and Vertex. L.D.W. is a scientific co-founder and shareholder in Aileron Therapeutics. R.G.J. is a scientific advisory board member for Immunomet Therapeutics and consultant for Agios Pharmaceuticals.

The immunofluorescence data to support the conclusions of this study are available at https://figshare.com/ DOI 10.6084/ m9.figshare.11956506. Uncropped western blots can be downloaded with this manuscript. Proteomic data in Fig. 3a are available in Supplementary Dataset 1, and native mass spectrometry data files are available for download from the MassIVE archive at the University of California, San Diego (ftp://massive.ucsd.edu/MSV000085082/). RNAseq data in Fig. 3b are available in DOI:10.1038/ s41467-017-00992-9. RNAseq data in ED Fig. 4 were obtained with permission from Emmanouil Dermitzakis (Accession EGAS00001000442; http://www.ebi.ac.uk/ega/³³.

All details on experimental design can be found in the Reporting Summary at https://www.nature.com/documents/nr-reporting-summary.pdf

¹⁰Department of Pediatric Oncology, Dana-Farber Cancer Institute, Boston, MA, USA

¹¹Linde Program in Cancer Chemical Biology, Dana-Farber Cancer Institute, Boston, MA, USA

¹²Department of Biological Chemistry and Molecular Pharmacology, Harvard Medical School, Harvard Medical School, Boston, USA

¹³Metabolic and Nutritional Programming, Center for Cancer and Cell Biology, Van Andel Institute, Grand Rapids, MI, USA

¹⁴Department of Medical Oncology, Dana-Farber Cancer Institute, Boston, MA, USA

¹⁵Department of Medicine, Harvard Medical School, Boston, MA, USA

Abstract

Chronic inflammation is linked to diverse disease processes, but the intrinsic mechanisms that determine cellular sensitivity to inflammation are incompletely understood. Here, we show the contribution of glucose metabolism to inflammation-induced changes in the survival of pancreatic islet β -cells. Using metabolomics, biochemical and functional analyses, we investigate the protective versus non-protective effects of glucose in the presence of pro-inflammatory cytokines. When protective, glucose metabolism augments anaplerotic input into the TCA cycle via pyruvate carboxylase (PC) activity, leading to increased aspartate levels. This metabolic mechanism supports the argininosuccinate shunt, which fuels ureagenesis from arginine and conversely diminishes arginine utilization for production of nitric oxide (NO), a chief mediator of inflammatory cytotoxicity. Activation of the PC-urea cycle axis is sufficient to suppress NO synthesis and shield cells from death in the context of inflammation and other stress paradigms. Overall, these studies uncover a previously unappreciated link between glucose metabolism and arginine-utilizing pathways via PC-directed ureagenesis as a protective mechanism.

Introduction

Glucose imparts protective or detrimental effects in a range of cell types depending on the extent and duration of the increase in glucose flux^{1–4}. An example of this is the intricate context- and dose-dependent modulatory effects of glucose on the survival of insulin producing β -cells. In particular, prolonged exposure to high glucose impairs β -cells survival and function cooperatively with inflammation in diabetes and obesity^{1,5–7}. While important advances have been made in understanding the immune cell component of inflammation in these pathologies^{6,8–10}, the cell-intrinsic biochemical connection between glucose metabolism and the response of target cells to inflammation has not been defined.

Results

The effect of glucose metabolism on the survival of human islets undergoing inflammation

To dissect the molecular effectors of glucose metabolism that control β -cell viability in the context of inflammation, we treated human donor islets with a combination of proinflammatory cytokines (TNF- α , IL-1 β , and IFN γ), shown to mimic β -cell inflammation in

diabetes^{6,7}. In this system, we examined how increased glucose metabolism via activation of glucokinase (GK, hexokinase IV), the hexokinase isoform expressed in these cells¹¹, influences inflammation-induced cell death and whether the protective versus non-protective effects of glucose can be distinguished. We have previously shown that GK activation via phosphorylation of the GK-binding protein BAD preserves β -cell survival in response to a variety of stress signals, including inflammatory cytokines¹². This prompted investigation whether other established modes of GK activation could be similarly protective. GK activation at its allosteric site by gain-of-function mutations identified in humans with hyperinsulinemic hypoglycemia (e.g., GK Y214C¹³), or by small molecule allosteric GK activators¹¹ (GKAs, e.g., RO0281675¹⁴), significantly augments the enzyme's affinity for glucose^{11,15}, but does not protect human islets from cytokine-induced death (Fig. 1a-c and Extended Data Fig. 1a-c). In contrast, GK activation near its active site by mimicking BAD phosphorylation using either the phospho-mimic mutant of BAD within its BCL-2 homology 3 (BH3) a-helix (BAD SD), or hydrocarbon stapled peptides modeled after the phospho-BAD BH3 helix (BAD SAHB_A SD)^{12,16-18}, maintains GK's native affinity for glucose18 and spares human islets from inflammation-induced death (Fig. 1a-c and Extended Data Fig. 1a-e). This protective effect requires the GK-activating capacity of phospho-BAD because a BAD BH3 mutant harboring triple-alanine substitutions that does not bind or activate GK, BAD AAA^{12,19}, or the corresponding BAD BH3 stapled peptide (BAD SAHB_A AAA)^{12,16} is not protective under similar settings (Fig. 1b-c and Extended Data Fig. 1a, c-d). Importantly, the pro-survival benefit of BAD SAHB_A SD in human islets is abolished upon GK knockdown (Fig. 1d and Extended Data Fig. 1f), indicating that protection by phospho-BAD mimicry is GK-dependent and on-target. These two independent modes of GK activation (allosteric site activation and phospho-BAD mimicry) essentially establish a system in which we can model the protective versus non-protective outcomes of increased glucose metabolism in order to identify the specific metabolic pathways downstream of glucose that determine cell survival during inflammation (Fig. 1a).

We next tested this modeling of protective versus non-protective glucose metabolism in a few additional settings. We found that the protective effect of phospho-BAD mimicry is also observed in type 2 diabetes (T2D) donor islets treated with inflammatory cytokines, and significantly contrasts with the non-protective outcome of GK activation at its allosteric site (Fig. 1e). Because human islets are comprised of ~40–60% β-cells out of the total of multiple islet cell types, we further verified that the above findings in whole islets demonstrate genuine β -cell survival differences by quantifying viability of cells positive for Newport green (NPG), which stains insulin granules only found in β -cells (Fig. 1f). In addition, the contrasting effects of these GK-activating modalities on sensitivity/resistance to inflammation-induced death is reproducible in the INS-1 β -cell line (Fig. 1g).

Changes in arginine metabolism differentiates protective and non-protective GK activation

The above data indicate that we can effectively model non-protective versus protective outcomes of increased glucose metabolism in β -cells using complimentary genetic and pharmacologic GK-activating tools (Fig. 1a). We then used this system as a discovery platform to learn about the metabolic mechanism whereby glucose determines cell survival during inflammation. To this end, we undertook non-targeted metabolomics to compare the

broad metabolite signatures of human donor islets in the absence (PBS) or presence of inflammatory cytokines following treatment with GKA RO0281675 or BAD SAHB_A SD, hereon referred to as non-protective or protective GK activation/glucose metabolism, respectively. The largest differences across conditions were observed in the presence of cytokines (Fig. 1h and Extended Data Fig. 2), and subsequent topological analyses using the Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway database identified nitrogen (P value of 9.3×10^{-9}), arginine and ornithine metabolism (P value of 5.2×10^{-8}) among the top pathways significantly differentiating the metabolic profiles of protective glucose metabolism via BAD SAHBA SD- from vehicle (DMSO), and from RO0281675-treated islets (nitrogen; P value of 9.4×10^{-15} , arginine and ornithine metabolism; P value of $6.6 \times$ 10⁻⁸) (Fig. 1i). Secondary validation using targeted LC-MS/MS analysis of human islets from five additional donors confirmed these findings. This prompted focused examination of arginine-utilizing pathways (Fig. 2a), which revealed a remarkable difference in urea and NO accumulation in that protective glucose metabolism augments urea levels while diminishing NO in the presence of cytokines (Fig. 2b and Extended Data Fig. 3a). These differences were further recapitulated with a second, structurally distinct allosteric GKA, GKA50²⁰ (Fig. 2c and Extended Data Fig. 3b). Importantly, *GK* depletion in human islets abolished the effects of phospho-BAD mimicry on urea and NO, underscoring GK's role in this setting and pointing to a link between glucose metabolism and arginine-utilizing pathways (Fig. 2d). The differential effect of protective versus non-protective GK activation on urea and NO levels also holds in the INS-1 β-cell line (Fig. 2e). Importantly, the ureaenhancing and NO-reducing outcomes of protective glucose metabolism are also evident in human islets expressing the full-length BAD SD protein in comparison to GK Y214C gainof-function mutant at the allosteric site (Fig. 2f and Extended Data Fig. 3c). Thus, the metabolic features distinguishing non-protective versus protective glucose metabolism are not limited to pharmacologic GK activation and are identical when using genetic tools to activate GK through these two distinct mechanisms (Fig. 1a). Of note, islets expressing the non-GK activating full-length BAD AAA mutant^{12,19} showed comparable levels of urea and NO to vector control (VC)- or GK Y214C-expressing islets (Fig. 2f and Extended Data Fig. 3c). Similarly, the BAD BH3 stapled peptides modeled after the BAD AAA mutant (BAD SAHBA AAA) did not enhance urea or reduce NO levels in the presence of inflammatory cytokines compared with vehicle or RO0281675 (Fig. 2e). These important controls underscore the specificity of both the genetic and chemical tools used to experimentally model and distinguish between features of protective versus non-protective glucose metabolism.

Protective glucose metabolism directs arginine to the urea cycle in human islets

Ureagenesis has not been extensively examined in β -cells. In our studies, the urea levels produced by islets (8–20 µM) are three orders of magnitude lower than normal circulating urea levels in humans (2–7 mM), or pathologic concentrations (~20 mM) shown to impair β -cell function as in chronic kidney disease²¹. We confirmed the expression of many urea cycle enzymes and associated pathways in this cell type by subjecting FACS-purified human β -cells to proteomic and RNAseq analyses (Fig. 3a–b, Extended Data Fig. 4 and Supplementary Dataset 1), as well as immunostaining fresh islet preparations for these enzymes within the insulin positive β -cell compartment (Fig. 3c). To biochemically quantify

the reactions relevant for the synthesis of urea and NO via arginase (ARG) and nitric oxide synthase (NOS) enzymes, respectively, we used the ¹⁵N₂[guanidino]-L-arginine tracer. Labelling of human islets revealed protective glucose metabolism is associated with higher enrichment of ¹⁵N₂ in urea (M+2) and lower citrulline (M+1) synthesis indicative of diminished NO synthesis (BAD SAHB_A SD, Fig. 3d–e). In contrast, ureagenesis was not altered in control samples (Veh and BAD SAHB_A AAA, Fig. 3d), or following non-protective GK activation (RO0281675, Fig. 3d). We did not detect any differences in the expression of iNOS or ARG2, the main NOS and ARG isoforms in human β -cells²². As such, differential urea and NO synthesis in this setting is more likely due to substrate-driven differences in enzyme activity. Importantly, treatment with the arginase inhibitor ABH (2(S)-amino-6-boronohexanoic acid) abrogates the protective effect of GK activation in islets undergoing inflammatory stress (Fig. 3f). Overall, these data indicate that protective glucose metabolism engages ureagenesis in the face of inflammation, which is commensurate with diminished NO accumulation.

Protective glucose metabolism links aspartate to the urea cycle to counter inflammation

The above findings prompted investigation of the specific link between glucose and arginine metabolism that could explain how protective glucose metabolism would lead to increased urea cycle activity. Aspartate is required for the production of the rate-limiting urea cycle substrate argininosuccinate, and is a potential metabolic node for modulation by glucose. Glucose carbons can contribute to aspartate pools via the TCA cycle metabolite oxaloacetate (OAA) (Fig. 4a). Thus, aspartate may link glucose to the urea cycle. We found that total aspartate levels were reduced in human islets treated with inflammatory cytokines and were further exacerbated by non-protective GK activation via RO0281675 (Fig. 4b). In contrast, aspartate pools remained stable in BAD SAHBA SD-treated islets in the absence or presence of cytokines (Fig. 4b). ¹³C₆-glucose tracer analyses provided direct evidence that protective GK activation supports a higher incorporation of glucose carbons into aspartate in the presence of inflammatory cytokines, in contrast to non-protective GK activation at the allosteric site (Fig. 4c and Extended Data Fig. 5a). A role for aspartate in glucose-stimulated ureagenesis stipulates that it would contribute to argininosuccinate synthesis, which would in turn depend on the activity of GOT1/2 aspartate aminotransferases catalyzing aspartate synthesis from glucose-derived OAA (Fig. 4a). Indeed, knockdown of either GOT1 or GOT2 using multiple different shRNAs diminished urea levels along with aspartate (Fig. 4d and Extended Data Fig. 5b-d). GOT1/2 depletion also led to elevated NO levels and reduced the viability of BAD SAHB_A SD-treated islets (Fig. 4d-e and Extended Data Fig. 5d). These results point to a protective ureagenic outcome of GOT1/2 activity, congruent with the idea that aspartate produced by these aminotransferases can support argininosuccinate (AS) synthesis to fuel the urea cycle. Consistent with this notion, AS supplementation is sufficient to increase urea levels and rescue the viability of human islets undergoing inflammation (Fig. 4f-g). Taken together, our data provide support for a previously unappreciated link between glucose and the urea cycle, and suggest that this metabolic mechanism of preserving aspartate pools reduces NO synthesis and shields cells from the cytotoxic effects of inflammation.

Pyruvate carboxylase mediates the protective outcomes of glucose and arginine metabolism

We next investigated the source of the increased aspartate following protective GK activation by determining which other metabolites contained isotopically-labelled carbon from ${}^{13}C_{6}$ glucose in islets undergoing inflammatory stress. Malate M+3 was increased, whereas citrate M+2 was unchanged (BAD SAHB_A SD, Fig. 5a and Extended Data Fig. 6a). We interpret such labeling patterns as an increase in PC activity (M+3 malate), no change in pyruvate dehydrogenase (PDH) activity (M+2 citrate), and higher PC to PDH activity ratio (Extended Data Fig. 6b). Independent biochemical measurements provided corroborative evidence that PC activity is higher in cytokine-treated human islets following protective versus nonprotective GK activation (Extended Data Fig. 6c). We next tested whether PC is relevant for the protective partitioning of arginine to the urea cycle away from utilization by NOS. PC knockdown in human islets undergoing inflammatory stress blunts the urea-enhancing and NO-reducing capacity of protective glucose metabolism concomitant with a significant reduction in aspartate levels (Fig. 5b-c and Extended Data Fig. 6d, lanes 1 vs 3). Moreover, on-target, shRNA-mediated depletion of PC or chemical inhibition of this enzyme using phenyl acetic acid (PAA) blocked the pro-survival benefits of GK activation (Fig. 5d-e and Extended Data Fig. 6d). Remarkably, the requirement for PC in augmenting urea and promoting survival in this setting is not recapitulated by glutaminase (GLS), another anaplerotic enzyme that could serve as a potential source of aspartate for argininosuccinate synthesis (Fig. 4a). Specifically, chemical inhibition of GLS did not interfere with urea levels nor the pro-survival effects of protective glucose metabolism (Fig. 5f and Extended Data Fig. 6e). Taken together, these findings are intriguing from both mechanistic and conceptual standpoints in that they establish a previously unexplored link between mitochondrial pyruvate handling via PC and arginine utilization through the ARG reaction. This mechanism restrains inflammation-induced augmentation of NO production and reveals an anti-inflammatory role for glucose metabolism via regulation of the urea cycle.

PC-dependent effects of protective glucose metabolism on islets and diabetes reversal

The prominence of PC in the above in vitro assays warranted investigation of its relevance in long-term β -cell protective outcomes of GK activation *in vivo*. To this end, we used a model of marginal mass islet transplantation. Islet transplantation in both human subjects and rodent models can have limited efficacy due, in part, to islet graft inflammation. Specifically, local production of cytokines at the site of engraftment and the presence of cytokines in the sera of islet transplant recipients have been reported²³. We found that protective activation of GK in human donor islets prior to transplantation reduced β -cell death in islet grafts as early as day 2 post transplantation and reversed hyperglycemia in diabetic mice over time, while non-protective GK activation or control treatments were ineffective (BAD SAHBA SD versus RO0281675, BAD SAHBA AAA or vehicle control, Fig. 6a,b, Table 1 and Supplementary Tables 2 and 3). The superior engraftment and physiologic functionality of protected islets was also evident from their ability to replenish insulin levels upon engraftment in vivo and from improved glucose clearance during a glucose tolerance test (GTT) (Fig. 6c-d). Importantly, chemical inhibition of PC with PAA in BAD SAHBA SDtreated donor islets blocked the efficiency of islet engraftment and systemic improvement in glucose homeostasis (Fig. 6a-d, Table 1, Supplementary Data Tables 2 and 3 Online). These

data underscore the functional relevance of PC as a downstream mediator of protective glucose metabolism in islets.

PC-directed ureagenesis protects human islets from inflammation and glucotoxicity

While we originally arrived at the connection between PC and arginine metabolism by modeling glucoprotection or lack thereof through different modes of GK activation, additional studies indicated that this metabolic link is itself a protective mechanism and is not restricted to experimental modulation of GK. Specifically, overexpression of PC in human islets augments urea, diminishes NO, and protects from inflammation-induced death independent of any genetic or pharmacologic modulation of GK, and effectively phenocopies the outcome of ARG2 overexpression on these same parameters (Fig. 7a-b and Extended Data Fig. 7a). Conversely, PC depletion alone is sufficient to reduce ureagenesis from arginine and enhance NO levels in cytokine-treated human islets (Fig. 7c and Extended Data Fig. 7b). The ability of PC to fuel ureagenesis suggests that this TCA cycle enzyme prevents the aspartate-argininosuccinate shunt from becoming limited for urea cycle activity in the presence of pro-inflammatory cytokines. Consistent with this notion, AS supplementation of PC-depleted cells rescues urea levels and protects against inflammation toxicity (Fig. 7d,e). This suggests that supplying the urea cycle with substrate is sufficient to enhance ureagenesis when flux through PC is constricted and PC-derived aspartate is limiting. Taken together, the above combination of gain- and loss-of-function studies and metabolite supplementation experiments indicate that arginine utilization through the ARG and NOS routes is acutely modulated by PC activity, and are consistent with a link between mitochondrial pyruvate handling and the urea cycle via the aspartate-argininosuccinate shunt.

In parallel studies, we also examined whether the protective outcome of the PC-ARG pathway is restricted to inflammatory stress or has broader significance in the context of other physiologically relevant islet stress paradigms. Chronic exposure to high glucose (glucotoxicity) is another stress paradigm, that along with inflammation, is known to impair β -cell survival in the diabetic milieu²⁴. To our knowledge, the relevance of arginine metabolism in the context of glucotoxicity has not been examined. We found that chronic exposure of human islets to 33 mM glucose reduces urea and enhances NO levels compared to normal islet culture conditions at 5.8 mM glucose (Fig. 7f–g and VC in Fig. 7h). Remarkably, overexpression of PC or ARG2 reversed these effects and was sufficient to protect islet viability in this setting (Fig. 7g–i). Thus, similar to inflammation, glucotoxicity is associated with diminished urea levels and NO accumulation, which can be countered by PC activity and routing of arginine to the urea cycle (Fig. 7g).

Discussion

Our studies identify urea cycle activation as part of a cell-intrinsic, anti-inflammatory pathway that can be fueled by glucose via PC and the argininosuccinate shunt. This glucosedriven protective mechanism ultimately shifts the direction of arginine metabolism to ureagenesis, sparing cells from damaging NO accumulation. While diminishing NO production is known to mitigate cell death in the context of inflammation, we are not aware

of previous studies showing that glucose can influence this process by controlling arginine utilization. This further highlights the relevance of our findings from a conceptual standpoint. Moreover, we provide multiple gain- and loss-of-function studies that position PC as a mechanistic link mediating this previously unappreciated connection between glucose and arginine metabolism. The ureagenic effect of PC is consistent with the notion that its activity can promote the availability of aspartate to feed the argininosuccinate shunt in support of the urea cycle. Our findings also suggest that altered urea cycle activity and an imbalance between arginine partitioning to urea versus NO synthesis is not limited to inflammation and may be a general metabolic feature of cellular stress that can be countered by PC-directed ureagenesis (Fig. 7g).

PC is known to play an important role in anaplerotic regulation of the TCA cycle and makes significant contribution to aspartate synthesis in multiple cell types^{25,26}. In β -cells, PC-dependent trans-mitochondrial pyruvate cycles produce metabolic signals that regulate insulin secretion^{27–30}. To our knowledge, however, this is the first direct demonstration that PC regulates arginine utilization for ureagenesis while suppressing its use for NO synthesis, and that this elicits a genuine pro-survival effect. Our findings also provide mechanistic insights relevant to inborn errors of metabolism involving PC deficiency where a correlation between decreased circulating urea levels, hyperammonemia and lower PC activity has been reported but not fully understood³¹. Furthermore, given that PC activity is altered in a number of pathologic conditions such as diabetes, cancer, and certain infectious diseases³², our findings warrant investigation into whether and how changes in ureagenesis, NO synthesis and nitrogen metabolism are linked to the disease outcomes in these settings.

Methods

Human islets

Primary human islets freshly isolated from deceased donors were obtained from the Alberta Islet Distribution Program (University of Alberta) or provided by the Integrated Islet Distribution Program (IIDP) at City of Hope (http://iidp.coh.org/) and Prodo Labs (https:// prodolabs.com/human-islets-for-research/). All donor material was obtained with informed consent and de-identified patient information, qualifying all human islet distribution centers as IRB-exempt and thus exempt from requiring approval by the office for human research studies at Dana-Farber Cancer Institute.

A total of 109 human donors were used in this investigation with mean age of 49 ranging from 17–75 years of age, 54% and 46% of which were males and females, respectively. The average BMI of donors was 26.7 (range 19.1–38.5), and baseline islet viability was 87.6% (range 68–98%). Donors obtained from IIDP are from the following Research Resource Identifiers: RRID:SAMN08769195 (UPenn), RRID:SAMN08769390 (UWisconsin), RRID:SAMN08769393 (Sharp/Lacy), RRID:SAMN08769822 (UWisconsin), RRID:SAMN08769830 (Sharp/Lacy), RRID:SAMN08769822 (UWisconsin), RRID:SAMN08773762 (UWisconsin), RRID:SAMN08773728 (UMiami), RRID:SAMN08773856 (UMiami), RRID:SAMN08773865 (Sharp/Lacy), RRID:SAMN08769188 (Sharp/Lacy), RRID:SAMN08769207 (SC-ICRC), RRID:SAMN08769084 (Sharp/Lacy), RRID:SAMN08769130 (UWisconsin),

RRID:SAMN08769124 (SC-ICRC), RRID:SAMN08769080 (Sharp/Lacy),
RRID:SAMN08769078 (UMiami), RRID:SAMN08769070 (SC-ICRC),
RRID:SAMN08769066 (Sharp/Lacy), RRID:SAMN08768765 (UPenn),
RRID:SAMN08768788 (UPenn), RRID:SAMN08768781 (UPenn), RRID:SAMN08611143 (UWisconsin), RRID:SAMN08616281 (Sharp/Lacy), RRID:SAMN08685106 (SC-ICRC),
RRID:SAMN08848150 (SC-ICRC), RRID:SAMN09768368 (UMiami),
RRID:SAMN10079665 (UWisconsin).

Type 2 diabetic donor islets used in Fig. 1e were from two donors; a 46 yr old female with BMI of 36 (RRID:SAMN08768788, UPenn), and a 45 year old male with a BMI of 27 (RRID:SAMN08616281, Sharp/Lacy). Both donors had a family history of diabetes and obesity.

Islet culture and cytokine treatment

Human islets were maintained in PIM(S) media (Prodo Labs) with 5% human A/B serum (Gemini BioProducts) at a density of 1000 islets per 10 ml as previously described³⁴. Primary mouse islets were isolated from 10 week-old C57B6 wild type mice and cultured as previously described¹². Control-treated mice were littermates with experimental treatments. Only male mice were used in this study. The cocktail of inflammatory cytokines (R&D Systems) used to treat human islets consisted of 10 ng/ml TNF- α , 10 ng/ml IL-1 β , and 100 ng/ml IFN γ , and that used to treat mouse islets or INS-1 cells consisted of 20 ng/ml TNF- α , 40 ng/ml IL-1 β , and 10 ng/ml IFN γ . These cytokine concentrations are consistent with published reports in human and mouse islets 35,36 . For argininosuccinate (AS) supplementation experiments (Fig 4f-g and Fig 7d-e), PIMS media was supplemented with argininosuccinic acid disodium salt hydrate (Sigma Aldrich) at 1 mM. This concentration was selected based on dose response studies to determine AS concentrations that effectively increase urea levels in human islets (10-200 µM AS had no effect on urea levels). Of note physiological range of AS concentrations in humans is currently unknown and current methods can only detect serum levels of AS in human subjects with ASL deficiency³⁷. For chronic high glucose (glucotoxic) culture conditions (Fig. 7f, h-i), islets were cultured in 33 mM glucose for 72 h compared to 5.8 mM glucose, which is the glucose concentration in PIM(S) media used for normal culture conditions.

SAHB synthesis and treatment

Peptide synthesis, olefin metathesis, fluorescein isothiocyanate (FITC) derivatization, reverse-phase high-performance liquid chromatography purification, and amino acid analysis were performed as previously described^{12,16}. SAHBs modeled after human and mouse BAD BH3 domain variants were used for treatment of human and mouse islets, respectively. BAD SAHB_A SD is modeled after phospho-mimic S118 and S155 in the human and mouse BAD sequence, respectively^{12,16,18}. BAD SAHB_A AAA is modeled after a triple Ala mutant^{12,19} in the BAD BH3 domain that modifies L151, S155 and D156 in the mouse BH3 domain and their corresponding residues in the human BAD BH3 sequence. The BAD AAA BH3 variant does not activate GK^{12,19} (ED Fig. 1c). Neither BAD SAHB_A SD nor BAD SAHB_A AAA binds to BCL-2/BCL-W/BCL-X_L¹⁶. For survival assays and human donor islet treatments, islets were treated with SAHBs for 5 h with 5 μ M of the indicated

SAHBs in 0.5% DMSO in uptake medium, which consisted of RPMI for mouse islets or PIM(S) for human islets, pH 6.2, which is the optimal pH for cellular uptake of BAD stapled peptides¹². Control islets were treated with vehicle composed of 0.5% DMSO in uptake medium. Islets were then washed and let to recover in complete physiological medium (pH 7.4) prior to any further treatments or assays.

Treatment with GKAs and pharmacologic inhibitors

RO0281675 (Axon Medchem) and GKA50 (Tocris) were dissolved in DMSO and used at 5 μ M in 0.5% DMSO. The following reagents were diluted in water: ABH (2(S)-Amino-6-boronohexanoic acid hydrochloride, 6-Borono-L-norleucine ammonium salt (Santa Cruz) used at 50 μ M, BPTES (Bis-2-(5-phenylacetamido-1,3,4-thiadiazol-2-yl) ethyl sulfide, Sigma) used at 500 nM, and PAA (phenyl acetic acid, Sigma) used at 1 mM.

Quantification of NO and cell survival by flow cytometry

NO levels were measured 24 h after cytokine treatment. Briefly, 50 islets per replicate were gently dispersed in accutase (Millipore Sigma) at 37°C, rinsed in 1 ml PBS, and subsequently stained at 37°C for 30 min with DAF-FM (Molecular Probes) for NO levels. Mean FITC signal intensities of DAF-FM stained cells were normalized to the PBS vehicle control set to 1.0 within a given experiment. Cell survival/death was quantified 48 h after cytokine treatment by staining islets with AnnexinV/7AAD (BD Biosciences) as previously described¹². Cells were monitored for PE/APC (585 nm/660 nm) or FITC/DCF (530 nm/488 nm) fluorescence using a BD FACSCanto II flow cytometer (BD Biosciences) and FACSDiva software (BD Biosciences). Positive cells were identified by comparison to unstained cells for all wavelengths at least 2 s.d. outside the means of unstained samples. Please see Supplementary method online file for gating strategies.

Genetic manipulation of islets

All infections were carried out 24 h prior to SAHB/cytokine treatment for a total of 48 h (when measuring NO levels) or 72 h (when measuring cell viability). GK Y214C mutant (pLenti-C-Myc-DDK, Origene), PC (pDNR223, Harvard Plasmid database) and ARG2 (pENTR223, Harvard Plasmid database) were first recombined into pLenti7.3 using the LR Clonase kit (Invitrogen). Lentiviruses were then packaged according to TRC Sigma protocol in 293T cells. Sigma TRC collection shRNA sequences for GK, GOT1, GOT2 and PC are as follows; human GKsh = 5'-

CCGGCGAGGACGTAATGCGCATCACCTCGAGGTGATGCGCATTACGTCCTCGTTTT TTG-3', human GOT1sh#1= 5'-

CCGGGCGTTGGTACAATGGAACAAACTCGAGTTTGTTCCATTGTACCAACGCTTTT TG-3', GOT1sh#2 = 5'-

CCGGGGCTAATGACAATAGCCTAAATCTCGAGATTTAGGCTATTGTCATTAGCTTTTT G-3', human GOT2 sh#1 = 5'-

CCGGGCCTTCACTATGGTCTGCAAACTCGAGTTTGCAGACCATAGTGAAGGCTTTT TG-3,GOT2sh#2 = 5'-

CCGGGGCTACAAGGTTATCGGTATTACTCGAGTAATACCGATAACCTTGTAGCTTTTT G-3', GOT2sh#3 = 5'-

CCGGCGAGATGTCTTTCTGCCCAAACTCGAGTTTGGGCAGAAAGACATCTCGTTT

TTG-3', mouse PCsh#1 = 5'-CCGGCCCTTCAGCTATTTGTCCTTTCTCGAGAAAGGACAAATAGCTGAAGGGTTTT G-3', and mouse PCsh#2 = 5'-CCGGGCACTACTTCATCGAGGTCAACTCGAGTTGACCTCGATGAAGTAGTGCTTTT TG-3', human PCsh#1 = 5'-CCGGCATGTTCATCTCTTGCCAAATCTCGAGATTTGGCAAGAGATGAACATGTTTT TG-3' and human PCsh#2 = 5'-CCGGATGGGCATCCGCCTGGATAATCTCGAGATTATCCAGGCGGATGCCCATTTTT TTG-3'. For lentiviral expression of full length BAD BH3 mutants (Fig. 1b, Fig. 2f, ED Fig. 1a, ED Fig. 3c and ED Fig. 6c), the cDNAs corresponding to the BAD phosphomimic mutant (S155D) and the phospho-deficient, non-GK activating BAD AAA mutant (described above) were cloned into the pCDH vector using the services of RxBiosciences. Lentiviralmediated knockdown and overexpression in human islets was performed as previously described³⁴. For all genetic manipulations, corresponding changes in protein expression were verified by western blot analysis using the following antibodies and dilutions; GOT1 (Sino Biological 14196-T52-50, 1:1000), GOT2 (My Bio Source MBS769801, 1:2000), glucokinase (Santa Cruz, 1:1000), pyruvate carboxylase (PCB, Santa Cruz, 1:1000), ARG2 (Life technologies, 1:1000), V5 (Cell signaling, 1:1000), MYC-tag (Cell Signaling, 1:1000), BAD (Abcam, 1:3000), and actin (Sigma, 1:20000).

Biotin-SAHB peptide pulldown assays

INS-1 cells (1×10^7) cultured in 10 cm plates were lysed on ice for 20 min in buffer containing 20 mM Tris-HCl pH 8.0, 137 mM NaCl, 2 mM MgCl2, 1 mM EDTA, 0.1% CHAPS, protease and phosphatase inhibitors (Sigma-Aldrich). Protein concentrations of spin-clarified lysates were determined using the BCA Protein assay kit (Thermo Fisher Scientific). Biotin pulldown assays were adapted from Escudero *et al.*³⁸. Briefly, 200 µg of lysates were pre-incubated with 25 µM of biotinylated SAHB peptides (ED Fig. 1d) or DMSO for 2 h at 4°C. Biotinylated peptides were then captured with the addition of 30 µl of streptavidin agarose beads for 2 h at 4°C. Beads were centrifuged at 3000 rpm for 30 sec, washed, and eluted by boiling in 2X LDS sample buffer (Thermo Fisher Scientific). Eluates were fractionated on SDS-PAGE and subjected to western blotting with antibodies against GK and PC.

Isothermal titration calorimetry (ITC) measurements

All calorimetric experiments were carried out using an Affinity ITC from TA Instruments (New Castle, DE) equipped with auto sampler in a buffer containing 20 mM HEPES pH 7.5, 150 mM NaCl, and 0.5 mM TCEP, with 5% DMSO at 25 °C. Recombinant GK protein solution (40 μ M) or buffer (as control) in the calorimetric cell was titrated by injecting 3 μ L of 300 μ M BAD BH3 SAHB_A SD or the corresponding unstapled peptide solution in 200 sec intervals with stirring speed at 75 rpm. Resulting isotherm was subtracted against buffer run and fitted with a single site model to yield thermodynamic parameters of H, S, stoichiometry, and Kd using NanoAnalyze software (TA instruments).

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Untargeted metabolomics

These analyses were performed using the services provided by General Metabolics, LLC. Human donor islets were treated as indicated above, rinsed in 150 mM ammonium formate and stored at -80° C. Polar metabolites from 150 human islets per replicate were extracted by hot 70% ethanol at 75°C for 3 min, vortexed, placed on ice, and spin clarified. Extracts were analyzed by non-targeted, flow-injection metabolomics on an Agilent 6550 Quadrupole Time-of-flight mass spectrometer (Agilent) using negative ionization as previously reported³⁹. Briefly, 1.5 µl was injected using a MPS3 autosampler (Gerstel). The mobile phase consisted of isopropanol/water (60:40, v/v) 1 mM ammonium fluoride with a flow rate of 150 µl per min. For online mass axis correction, homotaurine and hexakis(1H,1H,3Htetrafluoropropoxy)phosphazine (HP-0921, Agilent Technologies) were added to the mobile phase. Mass spectra were recorded in profile mode from m/z 50 to 1000 with a frequency of 1.4 spectra/s for 0.48 min using the highest resolving power (4 GHz HiRes). Source temperature was set to 325°C, with 5 L per min drying gas and a nebulizer pressure of 30 psig. Ions were putatively annotated by matching their measured mass with that of the compounds listed in KEGG database for *Homo sapiens*, allowing a tolerance of 0.001 Da, only deprotonated ions (without adducts) were included and duplicate matches were retained. Log fold change in metabolite signal intensities normalized to the median are presented for all conditions in a heatmap (ED Fig. 2), where each column represents individual replicates of pooled extracts from 5 independent human donors, with a total of 8 replicates. Data were analyzed as described below under Metabolomics data and statistical analyses.

Steady state urea quantification by GC-MS

Polar metabolites were extracted in 100 µl of 80% methanol (GC-grade, Thermo Optima) added to 100 µl of spent media from human or mouse islets (60 islets per 2ml) and lysed by sonication. Extracts were spin-clarified, and supernatants were collected and dried by vacuum centrifugation at 4°C (Labconco). Dried supernatant pellets were resuspended in 10 mg per ml of methoxylamine in pyridine (Sigma), incubated for 30 min at 37°C, and derivatized by silylation with 70 µl of N-Methyl-N-tert-butyldimethylsilyltrifluoroacetamide (MTBSTFA, Sigma) for 1h at 70°C. After spin clarifying, 1 µl of derivatized sample was injected in splitless mode into an Agilent 5977B/7890B GC-MS with a DB-5ms capillary column with DuraGuard (Agilent Technologies), using chromatographic methods as previously described⁴⁰. Data are shown either as concentrations (ng/ml) using a urea standard (Sigma) calibration curve or as values relative to PBS-treated controls. Between sample differences were corrected for by a factor of median chromatographic peak area.

Arginine tracer studies

Human islets (150 in 2 ml media) were incubated in ${}^{15}N_2$ (guanidino)-L-arginine (Cambridge Isotopes, 200 mg/L) for 72 h in RPMI media containing 5.5 mM glucose with 5% human A/B serum. At this time point, endogenous arginine pools were saturated with the label. Cytokine treatment was performed 24 h prior to harvesting. For quantification of M+n urea, media was collected (200 µl per replicate) and polar metabolites were extracted in 800 µl of 100% methanol, spin-clarified and dried by vacuum centrifugation at 4°C

(Labconco). Derivatization, instrumentation and data collection was the same as described above for steady state urea measurement using GC-MS. 1-1000 ng ¹⁵N₂-urea (Cambridge Isotopes) standard was assayed in the same media in parallel to generate a concentration curve for direct quantification of ¹⁵N₂-urea in experimental samples (retention time 13.6 min, target ions 213/215 and qualifier ions 233/235 for m+0/m+2 urea). Mass isotopomer distributions (MIDs) were calculated from proportional ion counts with corrections for natural abundances, individual chromatogram data were normalized to tracer uptake and subsequently compared to the amount in vehicle PBS-treated control set to 1.0. For quantification of M+n arginine and citrulline, islets were washed with cold 150 mM ammonium formate solution pH of 7.4 and extracted with 600 µL of 31.6% MeOH/36.3% ACN in $H_2O(v/v)$. Cells were lysed and homogenized by bead-beating for 2 min at 30Hz using 6.2 mm ceramic beads per sample (TissueLyser II, Qiagen). Cellular extracts were partitioned into aqueous and organic layers following dimethyl chloride treatment and centrifugation. The aqueous supernatants were dried as above, subsequently re-suspended in 30 µL of chilled H₂O, and clarified by centrifugation at 1°C. Each sample was analyzed three times using 5 µl injections in an Agilent 6540 UHD Accurate-Mass Q-TOF mass spectrometer (Agilent Technologies) equipped with a 1290 Infinity ultra-performance LC system (Agilent Technologies). Analyte ionization was accomplished using ESI in positive mode. The source operating conditions were set at 325°C and 9 L per min for gas temperature and flow respectively, nebulizer pressure was set at 40 psi and capillary voltage was set a 4.0 kV. Reference masses 121.0509 and 922.0099 were introduced into the source through a secondary spray nozzle to ensure accurate mass. MS data were acquired in full scan mode mass range: m/z 100–1000; scan time: 1.4s; data collection: centroid and profile modes. Retention times, accurate masses, and MS/MS (10, 20, 30, 40 V) for each compound were confirmed against authentic standards; ¹⁵N₂-arginine and ¹⁵N₁-citrulline (Cambridge Isotopes). Unlabeled islet extracts and quality control NT2196 cell extracts were used to confirm that signals were free of interfering ions in the mass range of interest. Chromatographic separation was performed using an Intrada Amino Acid column 3 µm, 3.0×150mm (Imtakt Corp). The chromatographic gradient started at 100% mobile phase B (0.3% formic acid in ACN) with a 3 min gradient to 27% mobile phase A (100 mM ammonium formate in 20% ACN / 80% water) followed with a 19.5 min gradient to 100% A at a flow rate of 0.6 ml/min. This was followed by a 5.5 min hold time at 100% mobile phase A and a subsequent re-equilibration time (7 min) before next injection. The column temperature was maintained at 10°C. Retention times for arginine and citrulline were 19.6 min and 8.5 min, respectively. Data are reported as ¹⁵N-enrichment, which is the corrected ¹⁵N isotopomer abundance normalized to the total abundance of a given metabolite, which were also normalized to tracer uptake and subsequently compared to the amount in vehicle PBS-treated control set to 1.0.

Glucose tracer studies

Mouse islets were incubated with 11 mM of uniformly labeled ¹³C₆-glucose (Cambridge Isotopes) in RPM1640 (Sigma Aldrich) for 24 h with simultaneous PBS or cytokines treatment. Polar metabolites from 150 islets were extracted in 80% methanol, derivatized, and detected with the same methods as above for steady state urea measurement using GC-MS. MID analysis was performed using a previously described algorithm⁴¹ and is described

below under metabolomics data analysis. Data are reported as ¹³C isotopomer enrichment compared to the total abundance of a given metabolite and normalized to vehicle PBS-treated control set to 1.0 (Fig. 4c, 5a, and ED Fig. 6a–b) and as ¹³C fractional labeling not normalized to vehicle PBS in ED Fig. 5a.

Metabolomics data and corresponding statistical analysis

All data except the untargeted metabolomics analysis were quantified by integrating the area under the curve of each metabolite and its isotopologue using MassHunter Quant (Agilent Technologies). Each metabolite's accurate mass ion and subsequent isotopic ions were extracted (EIC) using a 10 ppm window. In the case of isotopic labeling with ¹³C₆-glucose or ¹⁵N₂-arginine, corrections for natural enrichments were generated for each metabolite (C_{total}^{-1} or N_{total}^{-1}) using a previously developed algorithm⁴². MIDs of the indicated metabolites were calculated as previously described⁴² and expressed as percent of the untreated PBS control set at 1.0. Pathway enrichment, principal component analysis (PCA), fold change and ANOVA statistical analysis were performed using Metaboanalyst software (www.metaboanalyst.ca). Pathway analysis was performed comparing the log2 fold change data between two conditions per analysis, BAD SAHB_A SD versus vehicle, and BAD SAHB_A SD versus RO0281675, all in the presence of cytokines (Fig. 1i).

PC enzymatic assay

PC activity was measured in 150 human lysed by sonication in 100 μ l MOPS lysis buffer with protease inhibitors cocktail (Roche) and incubated with NaH¹⁴CO₃, as previously described⁴³. Briefly, 25 μ l of lysate per reaction replicate or control lysate without substrate (pyruvate) containing NaH¹⁴CO₃, were maintained at 37°C for 60 min and stopped with 10% TCA, according to previously described methods⁴³. Reactions were loaded onto P81 filter paper and volatile non-incorporated NaH¹⁴CO₃ was removed and ¹⁴CO₂ incorporation was measured by liquid scintillation. PC activity is reported as nmol ¹⁴CO₂ calculated from total ¹⁴C counts after subtracting ¹⁴C in control samples without pyruvate, normalized to protein content.

FACS purification of human β-cells for proteomic and RNAseq analyses

Human islets obtained from Prodo Labs were dispersed using Accutase (MT25058CI, Thermo Fisher Scientific) and infected with an adenovirus expressing the bright green fluorescent protein, ZsGreen (Clontech, Mountain View CA) under control of the rat insulin-1 promoter (RIP1) and a mini-CMV enhancer. The RIP1-miniCMV promoter included 177 bases of the hCMV IE-1 promoter ClaI-SpeI fragment ligated to 438 bases of the RIP1 promoter^{44,45}. Following transduction with the Ad.RIP-ZsGreen adenovirus for two hours, RPMI1640 medium containing 10% FBS was added to terminate adenovirus infection. Cells were collected 96 h after infection and loaded onto an Aria II cell sorter (BD biosciences), the live ZsGreen⁺ cell (β -cells) FITC-positive population was collected. The β cell fraction was confirmed to be >92% pure by immunolabeling of sorted cells with insulin, and independently, by qRT-PCR and by RNAseq^{44,45}. This procedure was repeated several times using islets from n=3 donors and sorted β -cells were pelleted and frozen at -80°C until enough starting materials for proteomic analyses were obtained. The donors were non-

diabetic with BMIs of 26.3, 28.8, and 27 and ages 48, 61 and 63, respectively. Please see Supplementary method for gating strategy.

Multidimensional nanoflow-LC-MS/MS proteomic analysis of FACS-purified human β-cells

Lysis buffer (1% SDS, 50 mM HEPES, with protease and phosphatase inhibitors) was added to frozen β -cell pellets (purified and collected as above), and the mixture was heated to 95°C for 5 min. After treating with 25U benzonase (Novagen, Thermo Fisher Scientific) for 30 min at 37°C, proteins were reduced with 10 mM DTT for 30 min at 56°C, alkylated with 22.5 mM iodoacetamide for 30 min in the dark, and captured on magnetic beads as described⁴⁶. After 3 washes ($2 \times 70\%$ ethanol and $1 \times 100\%$ ethanol), beads were reconstituted with 100 mM ammonium bicarbonate and proteins digested overnight with trypsin at 37°C. Peptides were re-captured on the magnetic beads by adding acetonitrile to a final concentration of 95%⁴⁶. After washing with acetonitrile, peptides were eluted in 2% DMSO with 50 mM ammonium formate, pH 10, and analyzed by DEEP-SEQ mass spectrometry⁴⁷ at a depth of 34 fractions using a NanoAcquity UPLC system (Waters, Milford, MA) coupled to a timsTOF Pro ion mobility mass spectrometer (Bruker). Peptide fractions were eluted from the first dimension, reversed phase column (5 cm \times 150 µm packed with 5 µm XBridge C18, Waters) and second dimension anion exchange column (5 $cm \times 150 \mu m$ I.D. fused silica packed with 10 μm POROS 10HQ, Applied Biosystems) using injections of acetonitrile or ammonium formate at pH 10.0. Peptide fractions were diluted in-line with solvent A (0.2 M acetic acid in water) to reduce pH and organic concentration and facilitate retention on the final dimension precolumn (5 cm \times 100 µm I.D. fused silica packed with 7 µm Symmetry C18, Waters). Peptides were eluted with an HPLC gradient (5-35% B in 120 minutes, A=0.2M acetic acid in water, B=0.2M acetic acid in acetonitrile), resolved on an analytical column (30 µm I.D. × 50 cm Monitor C18, Orochem), and introduced to the mass spectrometer by electrospray ionization using a captive spray ion source (spray voltage = 1.6 kV). The mass spectrometer collected ion mobility MS spectra over a mass range of m/z 100-1700 and 1/k0 of 0.6 to 1.6, and then performed 10 cycles of PASEF MS/MS with a target intensity of 20k and a threshold of 250. Active exclusion was enabled with a release time of 0.4 min. MS/MS spectra were matched to peptide sequences from human proteins in the Uniprot database (downloaded 9/4/2018) using PEAKS Studio 10.0 software⁴⁸. Search parameters specified precursor and product ion tolerances of 20 ppm and 50 mmu, respectively, as well as fixed carbamidomethylation and variable oxidation of methionine. Search results were filtered to a false discovery rate of 1% and exported to csv. After mapping peptide sequences to unique gene identifiers using the Pep2gene tool⁴⁹, multiplierz scripts⁵⁰ were used to provide an estimate of relative protein expression⁵¹ based on the three most abundant peptides detected for each gene product. Data in Fig. 3a are the distribution of the relative abundance levels of all proteins detected except those that had less than 3 unique peptides identified. The complete raw dataset with accession numbers and signal intensities can be found in online Supplementary Dataset 1. Native mass spectrometry data files are available for download from the MassIVE archive at the University of California, San Diego (ftp://massive.ucsd.edu/MSV000085082/).

RNAseq expression analysis of FACS-purified human β-cells

The RNAseq data in Fig. 3b are taken from a previously published dataset from normal nondiabetic donors⁴⁴. Briefly, RNA was prepared from freshly FACS-purified human β-cell using the RNeasy Micro kit (Qiagen). RNA yields were 300-500 ng from each FACSpurification, and RNA integrity numbers were between 9.5 and 10.0. PolyA⁺ mRNA from sorted β -cells was purified with oligo dT magnetic beads. The polyA⁺ RNA from β -cells was then fragmented in the presence of divalent cations at 94°C. The fragmented RNA was converted into double stranded cDNA. After polishing the ends of the cDNA, the 3' ends were adenylated. Lastly, Illumina-supplied universal adapters were ligated to the cDNA fragments. The adaptor ligated DNA was size selected to get an average of 250 bp insert size using AmpPure beads and amplified by 15 cycle PCR. The PCR DNA was then purified using AmpPure beads to get the final seq library ready for sequencing. The insert size and DNA concentration of the seq library was determined on Bioanalyzer (Agilent Technologies) and Qubit (Thermo Fisher Scientific), respectively. A pool of 10 barcoded RNAseq libraries was layered on two of the eight lanes of the Illumina flow cell at appropriate concentration and bridge amplified to yield ~25-35 million raw clusters. The DNA reads on the flow cell were then sequenced on HiSeq 2000 using a 100 bp paired end recipe. Results are expressed as Log base 2-transformed signals for millions of counts (reads) per million bases (CPM) (Log2CPM, Fig. 3b), and were calculated as previously described⁴⁴. RNA Enrichments in ED Fig. 4 were calculated using a transcriptome dataset for sorted β -cells, whole islet and negative-sorted islet cells generously provided by Emmanouil Dermitzakis (Accession EGAS00001000442; http://www.ebi.ac.uk/ega/)³³ and corresponding RPKM (reads per kilobase per million mapped reads) values were analyzed following the methodology described in Wang et al.⁴⁴. RNA enrichments were calculated by dividing individual RPKM values for each gene from sorted β -cells and non-sorted islet cells by the whole islet RPKM values, where whole islet values are set to 1.

Immunostaining of human islets to assess expression of urea cycle enzymes in β-cells

Human islets from 3 donors were dispersed into single cells on 12 mm diameter round coverslips (Electron Microscopy Sciences) coated with polyD-lysine (Sigma) and cultured overnight in 48 well plates. Cells were then fixed in 3.7% formaldehyde solution (Sigma) in media containing 1 µg/ml Hoechst (BD Pharmogen) at 37°C for 15 min, rinsed in PBS containing 75 mg/ml glycine, treated with 0.1% Triton-X-100 for antigen retrieval, and blocked in 10% FBS (Gibco) for 1 h. Immunostaining was performed overnight at 4°C for each of the following antibodies individually at 1:200 dilution: ARG2 (ABClonal), ASL (Sigma), ASS1 (Thermo Fisher Scientific), CPS1 (ABClonal), NAGS (ABClonal), DDAH1 (Sino Biological Inc), and DDAH2 (ABClonal). Coverslips were rinsed three times in PBS and stained with secondary antibodies against rabbit and/or mouse conjugated to Alexafluor488 (Life Technologies) at 1:500 dilution. Samples without primary antibody were used as negative controls to account for background fluorescence of secondary Alexfluor488 antibodies. All coverslips were then co-stained with an antibody to insulin (LSBio) at 1:150 dilution for 1 h at room temperature, rinsed and stained with a secondary guinea-pig antibody to Fluor568 (Thermo Fisher Scientific) at 1:500 dilution, and mounted onto microglass with ProLong Diamond Antifade mountant (Thermo Fisher Scientific). Imaging was performed on a Nikon Ti w/ Perfect Focus & Spinning Disk Confocal, with

488, 445 and 561 nm lasers, a Hamamatsu ORCA-R2 cooled CCD camera and MetaMorph image acquisition software with 60 and 100X objective oil-lens. Quantitation was performed on 5 images per antibody with 3–5 cells per image in Fiji software (https://imagej.net) by generating a threshold selection mask in the 561 nm channel image to create an insulin-positive region of interest, which was then mapped onto the corresponding 488 image within which the intensity of the 488 nm signal was measured.

Marginal human islet mass xenotransplantation

Human islets from 14 donors were treated overnight with vehicle (1% DMSO), 5 μ M of the indicated BAD SAHB_A or RO0281675, or were co-treated with 5 μ M BAD SAHB_A SD and 1 mM PAA in 1% DMSO in RPMI SAHB uptake medium containing 11 mM glucose, 10% serum and 0.1% Tween 80 (pH 6.0) at 37 °C or co-treated with 5 µM BAD SAHB_A SD and 1 mM PAA. Islets were then washed and incubated in fresh media for 24 h before transplantation. The presence of SAHBA was verified up to 4 days post uptake in human islets. Donors ranged in age from 21 to 64 years (mean 44); 6 were female and 8 were male. Mean BMI was 25 (range 20.8–32.9) and islet purity ranged from 85% to 90%. Diabetes was induced by single intraperitoneal (i.p.) injection of STZ (S0130, Sigma) at 200 mg/kg body weight in 200 µL total volume in a total of 22 10-12 wk old male NOD-SCID mice (001303, Jackson Labs), with n=3-6 mice per condition. STZ-treated mice were subjected to transplantation surgery 24–48 h after their blood glucose levels reached 300 mg/dl. Mice were anesthetized, and a small incision was made in the skin overlying the kidney to expose the peritoneum. The kidney was then exposed by a small incision in the peritoneum and kept moist throughout the procedure. Using a 23 g needle, a small nick was made in the kidney capsule and 500 human islets pre-treated as indicated above were glided through the nick using PE50 tubing. The tubing was removed gently, and the nick cauterized. The area was then sutured, and skin staples applied. Transplant recipients were monitored daily. Blood glucose, insulin measurements and i.p. GTT were performed as previously described¹². All animal studies were performed in compliance with, and with approval of, the Icahn School of Medicine at Mount Sinai Institutional Animal Care and Use Committee.

Quantitation of β-cell death in islet grafts

Kidneys were excised from transplanted mice, fixed in Bouin's solution, paraffin-embedded, sectioned, and stained with an antibody to insulin (A0564, DAKO), TUNEL (G3250 DeadEnd Fluorometric TUNEL system, Promega) and DAPI (D21490, Invitrogen). Cells doubly positive for insulin and TUNEL staining were quantified and compared to total number of insulin positive cells using fluorescence microscopy. An average of 1166±163 cells were counted per condition and are pooled means of a total of 32 individual mouse grafts with Veh n = 6, RO0281675 n = 5, BAD SAHB_A SD n = 6, BAD SAHB_A SD + PAA n = 4 and BAD SAHB_A AAA n = 4 mice and reported as % β-cell death representing the % TUNEL positive cells normalized to vehicle set to 100%. Imaging was performed at 20X magnification with a CARL ZEISS AXIO Observer Z1 Inverted Microscope using ZEN 2 core imaging Software for Zeiss microscope.

Statistical analyses

One or two-way ANOVA with Tukey's test for multiple means comparisons and student's ttest were used in GraphPad for calculating statistical significance with q-value adjustments for a false discovery rate (FDR) of 1%. Outliers were rarely removed and were determined using two-sided Grubbs with alpha set to 0.05. In the case that an outlier was observed, the data were further inspected with the ROUT method at FDR set to 1% to check for multiple outliers.

Extended Data



Extended Data Fig. 1. Characterization of GK-modulating tools used in this study. a, Western blots showing expression levels of full length MYC-tagged GK Y214C and BAD BH3 mutant proteins in islets used in Fig. 1b and Fig. 2f. Blots are representative of n=2 independent experiments with similar results.

b, Viability of human islets treated with increasing doses of RO0281675 or BAD SAHB_A SD and exposed to cytokines as in Fig. 1c. Based on these dose response studies, we elected to use RO0281675 at 3 μ M and BAD SAHB_A SD at 5 μ M throughout all studies. Data are means \pm s.d. of 3 technical replicates of islet cultures from one human donor. **c**, GK activity in human islets treated with vehicle (DMSO), RO0281675, BAD SAHB_A SD, BAD SAHB_A AAA or a stapled peptide modeled after the BH3 domain of a related BCL-2 family protein (BIM SAHB_A). Data are means \pm s.d. with n=4 (veh and BAD SAHB_A SD) or n=3 (RO0281675, BAD SAHB_A AAA or BIM SAHB_A) technical replicates of islet

cultures from one donor.

d, Specific target engagement by BAD SAHB_A SD as assessed by the capture of GK with biotinylated BAD SAHB_A SD but not BAD SAHB_A AAA or BIM SAHB_A in INS-1 protein lysates. Western blot with the anti-PC antibody serves as negative control for GK. Input denotes INS-1 lysates not incubated with any stapled peptides or vehicle. Representative experiment is shown out of n=2 experiments with similar results.

e, Isothermal titration calorimetry (ITC) measurements showing the binding of recombinant human GK to BAD SAHB_A SD in a 1:1 stoichiometry with binding affinity (dissociation constant, *K*d) of ~580 nM (left). ITC using the corresponding unstapled peptide is shown for comparison with a log higher *K*d (right). Data are representative of n=3 independent ITC experiments with similar results.

f, Western blots showing efficiency of GK knockdown in islets used in Fig. 1d and Fig. 2d. Blots are representative of n=2 independent experiments with similar results.



Extended Data Fig. 2. Untargeted metabolomics analysis of human islets undergoing inflammation stress.

Heatmap presentation of LC-MS untargeted metabolomics analysis of human islets showing PBS and cytokine conditions corresponding to Fig. 1h and i. Data are transformed into log fold change for heatmap presentation with 8 technical replicates of total ion counts shown for islets pooled from n=5 human donors.

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Extended Data Fig. 3. Altered arginine metabolism in the context of protective vs non-protective glucose metabolism.

Cytokines

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PBS

Cytokines

a, Urea and NO levels in human islets treated with the indicated compounds and exposed to cytokines (Fig. 2b), expanded to show the PBS data. PBS urea data are from n=5 (veh), n=3 (RO0281675) and n=4 (BAD SAHB_A SD) human donors. Cytokine urea data are from n=10 (veh), n=7 (RO0281675), and n=12 (BAD SAHB_A SD) donors. PBS NO data are from n=8 (veh, RO0281675), and n=9 (BAD SAHB_A SD) donors. Cytokine NO data are from n=9 (veh, RO0281675) and n=8 (BAD SAHB_A SD) donors.

b, Viability of human islets treated with vehicle (DMSO), the allosteric GK activator (GKA50) or BAD SAHB_A SD and exposed to inflammatory cytokines as in Fig. 2c, n=4 donors.

c, Urea and NO levels in human islets expressing the indicated GK and BAD mutants and treated with cytokines (Fig 2f), expanded to show the PBS data. Urea data for PBS and cytokine conditions are from n = 6 (VC) and n = 7 (GK Y214C, BAD SD and BAD AAA)

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PBS

independent experiments using islet cultures from 2 donors. PBS NO data are from n = 4 (VC), n = 2 (GK Y214C) and n = 4 (BAD SD and BAD AAA) independent experiments using islet cultures from 2 donors. Cytokine NO data are from n = 4 (VC, GK Y214C, BAD SD and BAD AAA) independent experiments using islet cultures from 2 donors. Statistical analyses in (a) and (c) are two-way ANOVA and one-way ANOVA in (b), both with Tukey adjustment for multiple comparisons.

Gene name	β-cell	Whole islet	Non-β-cell
Urea cycle and related pathways			
ARG2	6.39	1	0.87
ASL	6.10	1	0.61
ASS1	5.05	1	1.03
CPS1	1.72	1	0.45
GOT1	5.09	1	1.29
GOT2	2.68	1	0.81
MDH1	10.09	1	1.02
MDH2	6.74	1	0.90
NAGS	3.16	1	1.60
OAT	10.22	1	1.64
PC	4.87	1	0.88
SLC25A12	9.97	1	1.46
SLC25A13	7.10	1	0.99
SLC25A15	1.90	1	0.68
Hormones			
INS	1.48	1	0.09
GCG	0.17	1	4.46
β-cell genes			
GCK	1.60	1	1.00
SLC2A2	3.47	1	0.28
MAFA	2.65	1	0.21
α -cell and non- β -cell genes			
ARX	0.13	1	1.86
DNMT1	0.76	1	0.93
LDHA	0.45	1	0.60

RNAseq analysis of urea cycle enzymes and related pathways in sorted β -cells and negative-sorted islet cells relative to whole islets

Extended Data Fig. 4. Expression of urea cycle enzymes and related pathways in FACS-purified human β -cells subjected to transcriptomic analyses.

RNAseq analysis of urea cycle enzymes and related pathways in sorted human β -cells and negative-sorted islet cells relative to whole islets. Read counts as RPKM (reads per kilobase per million mapped reads) are normalized to whole islet PKRM to assess enrichment. All urea cycle related enzymes and transporters are enriched (>1) in the β -cell fraction compared to whole islets and the negative-sorted cells.



Extended Data Fig. 5. Increased generation of aspartate from glucose following protective GK activation.

a, ¹³C fractional labelling of aspartate from ${}^{13}C_6$ glucose. Data are shown as non-normalized to vehicle PBS and display the fraction of each M+*n* mass isotopomer out of the total pool of aspartate for each condition. For clarity, statistical comparisons are only shown for each M +*n* of a given condition (RO0281675, BAD SAHB_A SD and BAD SAHB_A AAA) compared to the corresponding M+*n* of vehicle control. Data are pooled means from n=6 (Veh), n=5 (RO0281675), and n=6 (BAD SAHB_A SD, BAD SAHB_A AAA) independent mouse islet isolations and experiments.

b, Western blot analysis of GOT1/2 knockdown efficiency using multiple independent hairpins for data shown in Fig. 4d–e and ED Fig. 5c–d. Blots are representative of n=2 independent experiments with similar results.

c–d, Aspartate (**c**), urea and NO (**d**) levels in human islets from the same experiments shown in Fig. 4d–e, displaying the complete set of data on all hairpins tested. Aspartate data are

from n = 4 human donors for shCtrl samples and n = 3 donors for knockdown samples. Urea and NO data are from n = 4 and n = 3 donors, respectively.

Statistical analyses in (a) are two-way ANOVA showing p-value comparisons for each condition to Veh, and one-way ANOVA in (c–d), both with Tukey adjustment for multiple comparisons.

□ Veh □ RO0281675 □ BAD SAHB_A SD ■ BAD SAHB_A AAA





a–b, PDH (**a**) and the ratio of PC/PDH (**b**) activity in mouse islets labeled with ${}^{13}C_6$ glucose, measured as M+2 citrate and the ratio of M+3 malate to M+2 citrate, respectively. Data are from analogous glucose tracer studies as in Fig 5a, showing n=8 (Veh), n=5 (RO0281675, BAD SAHB_A AAA) and n=6 (BAD SAHB_A SD) independent experiments for PDH, and n=8 (Veh, BAD SAHB_A AAA), n=5 (RO0281675) and n=6 (BAD SAHB_A SD) independent experiments for PC/PDH. Statistical analyses were performed using one-way ANOVA with Tukey adjustment for multiple comparisons.

c, PC activity in human islets treated with inflammatory cytokines in the context of protective vs non-protective glucose metabolism. Enzyme activity was measured as nmol 14 CO₂ generated from NaH¹⁴CO₃, n=2 human donors in duplicate.

d, Validation of on-target *PC* knockdown and expression level of V5-tagged human PC (hPC) cDNA used to rescue PC expression in human islets treated with a 3'UTR-targeted shRNA against *PC* in experiments corresponding to Fig. 5d. Blots are representative of n=2 independent experiments with similar results.

e, The GLS inhibitor BPTES (Bis-2-(5-phenylacetamido-1,3,4-thiadiazol-2-yl)ethyl sulfide) does not affect islet urea levels at concentrations that reduce the ratio of glutamate/glutamine (glu/gln, a readout of GLS activity), n=2 human donors.



Extended Data Fig. 7. Validation of ARG2 and PC overexpression and knockdown. a, Western blot analysis of ARG2 and PC expression levels in human islets corresponding to experiments shown in Fig. 7a–b and h–i. Blots are representative of n=2 independent experiments with similar results.

b, Western blot analysis of *PC* knockdown efficiency in experiments corresponding to Fig. 7c–e. Blots are representative of n=3 independent experiments with similar results.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1 |. Protective vs non-protective glucose metabolism in human islets undergoing inflammation and attendant metabolite signatures.

a, Schematic summary showing modelling of protective versus non-protective glucose metabolism using GK-targeted genetic and pharmacologic tools. GOF denotes gain-of-function.

b, Viability of human islets expressing vector control (VC), GK Y214C, BAD SD, or BAD AAA following 48 h treatment with a cocktail of inflammatory cytokines (TNF- α , IL-1 β , and IFN γ). Values are normalized to VC PBS control treatment. Data are from n=3 human donors each with 2 replicates.

c, Viability of human islets treated with vehicle (Veh, DMSO), RO0281675, BAD SAHB_A SD or BAD SAHB_A AAA and exposed to cytokines for 48 h as in (b), n=6 donors. **d**, Viability of human islets subjected to *GK* knockdown and treated with vehicle or BAD

SAHB_A SD in the presence of inflammatory cytokines. Data are from n=5 independent experiments using islet cultures from 2 donors.

e, Viability of T2D donor islets treated as in (c). Data are from n=2 independent experiments using islet cultures from 2 donors.

f, Viability of β -cells within human islets treated as in (c) and visualized by co-staining with Newport Green (NPG) and AnnexinV/7AAD. Data are means \pm s.d. from n=4 technical replicates of islets cultures from one donor.

g, Viability of INS-1 β -cells treated as in (c). Data are means \pm s.d. from n=4 technical replicates.

h, Principal component analysis (PCA) of LC-MS untargeted metabolomics of human islets treated as in (c) for 24 h, n=5 donors pooled and split into 8 replicates for metabolomics analysis.

i, Pathway analysis displayed as bar plot showing pathway -log p-values, revealing nitrogen, arginine and ornithine metabolism as the top pathways changed in vehicle control versus BAD SAHB_A SD or in RO0281675 versus BAD SAHB_A SD comparisons, n=5 donors. For RO0281675 vs. BAD SAHB_A SD comparisons, arginine and ornithine metabolism is not displayed but is statistically enriched with a *P*-value of 1.21×10^{-8} .

Data in **b–d and i** are means \pm s.e.m. with statistical analyses on means from independent experiments using one-way ANOVA with Tukey adjustment for multiple comparisons.



Figure 2. Differential modulation of arginine metabolism by protective vs non-protective GK activation in human islets.

a, Schematic of arginine usage for urea and NO synthesis.

b, Quantification of urea and NO levels in human islets treated with the indicated compounds and exposed to cytokines for 24 h. Values are shown relative to vehicle control PBS samples. Data for urea are from n=10 (Veh), n=7 (RO0281675), and n=12 (BAD SAHB_A SD) human donors. Data for NO are from n=9 (Veh and RO0281675) and n=8 (BAD SAHB_A SD) human donors.

c, Quantification of urea and NO in human islets treated with the indicated compounds and exposed to inflammatory cytokines as in (b). Data for urea are from n=3 donors. Data for NO are from n=2 (veh and GKA50) and n=3 (BAD SAHB_A SD) donors.

d, Data for urea are from n = 4 independent experiments using islet cultures from 2 donors. Data for NO are from n = 4 donors.

e, Urea and NO levels in INS-1 cells treated as in (b) measured at 24 h. Data are means \pm s.d. of n=4 technical replicates.

f, Urea and NO levels in human islet expressing the indicated GK and BAD mutants and treated with cytokines as in (b). Data for urea are from 6 (VC) and 7 (GK Y214C, BAD SD and BAD AAA) replicates using islet cultures from 2 donors performed over n=4 independent experiments. Data for NO are from n=4 independent experiments using islet cultures from 2 donors.

Data in **b–d and f** are means \pm s.e.m. with statistical analyses on means from independent experiments using one-way ANOVA with Tukey (b–d) adjustment for multiple comparisons and Fisher's exact test (f).



Figure 3 |. Protective glucose metabolism directs arginine to the urea cycle away from NO synthesis in islets undergoing inflammation.

a, Histogram of relative abundance of 3160 proteins detected (out of 5399 total, see Supplementary Dataset 1) by LC–MS/MS in 8.7×10^4 purified human ²-cells from n = 3 donors. Green lines indicate gene products related to the urea cycle, argininosuccinate and aspartate metabolism detected at the protein level.

b, Expression levels of genes related to the urea cycle, pyruvate metabolism and arigninosuccinate shunt based on RNAseq analysis of FACS-sorted human β -cells from n=3 human donors, data are in Log2CPM.

c, Co-immunostaining of insulin and individual metabolic enzymes related to the urea cycle in dispersed human islet cells from one donor representing similar results obtained from two additional donors. Representative images are shown (left), scale bar is 10 microns. Mean fluorescence intensity (FI) within the insulin positive region of interest (ROI) was calculated from 5 images per antibody (right). Neg denotes negative control for background Alexa

Fluor 488 signal with insulin co-stain. Statistical analyses are student's t-tests of each enzyme compared to Neg.

Enzyme abbreviations in **a**–**c** are ARG2, arginase 2; ASL, argininosuccinate lyase; ASS1, argininosuccinate synthase 1; CPS1, carbamoyl-phosphate synthase 1; DDAH 1, dimethylarginine dimethylaminohydrolase 1; DDAH 2, dimethylarginine dimethylaminohydrolase 2; GOT1, aspartate aminotransferase 1; GOT2, aspartate aminotransferase 2; MDH1, malate dehydrogenase 1; MDH2, malate dehydrogenase 2; NAGS, N-acetyl-glutamate synthase; OAT, ornithine aminotransferase; PC, pyruvate carboxylase; SLC25A12, solute carrier family 25 member 12/calcium-binding mitochondrial carrier protein Aralar 1; SLC25A13, solute carrier family 25 member 13/calcium-binding mitochondrial carrier protein Aralar 2; SLC25A15, solute carrier family 25 member 15/

d–e, Cytokine-induced changes in the partitioning of ${}^{15}N_2$ -L-arginine to urea (**d**) and citrulline/NO (**e**) synthesis in human islets comparing protective vs non-protective glucose metabolism as modeled by BAD SAHB_A SD vs RO0281675 treatment, respectively, n=4 donors.

f, Chemical inhibition of arginase via ABH interferes with the protective effect of BAD $SAHB_A SD$ in human islets undergoing inflammation, n=5 donors.

Data are means \pm s.e.m. with one-way (d,e) and two-way (f) ANOVA statistical tests with Tukey adjustment for multiple comparisons.

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Figure 4 |. Protective glucose metabolism links aspartate to the urea cycle to counter inflammation.

a, Schematic of the TCA and urea cycles and their connection via the aspartateargininosuccinate shunt. Enzymes of interest are marked in red and their corresponding inhibitors in blue.

b, Total aspartate levels in human islets treated with the indicated compounds and cultured in the absence or presence of inflammatory cytokines. Data are from the untargeted metabolomics analysis in Fig. 1h, n=5 human donors pooled and split into 8 replicates. **c**, Contribution of glucose to total aspartate pools in mouse islets labeled with ${}^{13}C_6$ glucose and treated with inflammatory cytokines in the context of protective vs non-protective glucose metabolism. Data are from n=5 (Veh, RO0281675), n=6 (BAD SAHB_A SD) and n=4 (BAD SAHB_A AAA) independent mouse islet isolations and experiments. See Fig. ED 5a for isotopologue distribution of aspartate in an analogous labelling experiment.

d–e, Quantification of urea and NO (**d**), and viability (**e**) in human islets subjected to shRNA-mediated *GOT1* (*G1*) or *GOT2* (*G2*) depletion and treated with cytokines in the context of protective vs non-protective glucose metabolism, n=4 donors for urea, n=3 donors for NO, and n=5 donors for viability measurements. Data for one hairpin per gene are displayed (sh#1 for *GOT1* and sh#2 for *GOT2*) from the full data set of multiple hairpins, see ED Fig. 5b–d.

f–g, Quantification of urea levels (**f**) and viability (**g**) in human islets supplemented with argininosuccinate (AS) in the presence of inflammatory cytokines, H₂O serves as vehicle control for AS. Data are from 6 independent experiments from n=3 donors. Data are means \pm s.e.m. with statistical analyses on means from independent experiments using one-way ANOVA with Tukey adjustment for multiple comparisons.



Figure 5 |. Pyruvate carboxylase supports aspartate and ureagenesis, and is required for the protective effects of glucose metabolism.

a, M+3 malate levels in mouse islets labeled with ${}^{13}C_6$ glucose and treated as in Fig. 4c. Data are pooled means from n=5 (Veh, RO0281675), n=6 (BAD SAHB_A SD) and n=5 (BAD SAHB_A AAA) independent experiments.

b–c, The effect of *PC* knockdown on urea and NO (**b**) and aspartate levels (**c**) in human islets treated with cytokines in the context of protective glucose metabolism, n=3 human donors in **b**, and n=3 independent experiments from one donor in **c**.

d, The effect of *PC* knockdown on viability of human islets treated as in (b). On-target effects of knockdown were validated by rescue with an shRNA-resistant human PC cDNA compared to vector control (VC), n=3 donors.

e, The effect of PC inhibition by PAA on the viability of human islets treated with cytokines in the context of protective glucose metabolism, n=4 donors.

f, GLS activity is not required for the protective effects of glucose metabolism. Viability of BAD SAHB_A SD-treated human islets exposed to inflammatory cytokines in the absence or presence of BPTES for 48 h, n=3 donors.

Data are means \pm s.e.m. from independent experiments with statistical analyses using oneway ANOVA with Tukey adjustment for multiple comparisons.

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Figure 6 |. Pyruvate carboxylase is required for the protective effects of glucose metabolism in human islets *in vivo* and their capacity to reverse diabetes in mice.

a, Schematic of marginal mass islet transplantation in diabetic NOD-SCID mice using human donor islets pre-treated with the indicated single or double combination of compounds for 24 h prior to transplantation. Islet grafts were excised at day 2 post transplantation for quantification of β -cell death as % insulin and TUNEL double positive cells with t-test statistic. Representative images are shown, scale bars are 10 microns. Data are from n=6 (Veh, BAD SAHB_A SD), n=5 (RO0281675) and n=4 (BAD SAHB_A SD + PAA, BAD SAHB_A AAA) mice.

b, Blood glucose levels of mice treated as in (a) measured up to 53 days post transplantation. Nephrectomy was performed at day 50 to excise grafts and show the requirement of protected donor islets for improving blood glucose. Statistical comparisons are provided in Table 1. Data are from n=6 (Veh, BAD SAHB_A SD), n=3 (RO0281675, BAD SAHB_A SD + PAA) and n=4 (BAD SAHB_A AAA) mice.

c, Human insulin levels in the sera of mice treated as in (a) on day 49 post transplantation. Data are from n=6 (Veh, BAD SAHB_A SD), n=3 (RO0281675, BAD SAHB_A SD + PAA) and n=4 (BAD SAHB_A AAA) mice.

d, Mean blood glucose levels during an intraperitoneal glucose tolerance test (GTT) and corresponding area under the curve (AUC) in mice treated as in (a) on day 42 post transplantation. Number of mice per condition is as in (c).

Data are means \pm s.e.m. from independent experiments with statistical analyses using oneway ANOVA with Tukey adjustment for multiple comparisons.



Figure 7 |. Pyruvate carboxylase is sufficient to promote ureagenesis, restrict NO, and protect human islets from inflammation and glucotoxicity.

a–b, Quantification of urea and NO levels (**a**), and viability (**b**) in human islets expressing PC or ARG2 in the presence of inflammatory cytokines, n=4 human donors for urea and n=3 donors for NO and viability measurements.

c, Contribution of arginine to urea synthesis and quantification of NO levels in *PC*-depleted human islets that were labeled with ${}^{15}N_2$ -arginine and treated with cytokines, n=4 donors. **d–e**, Quantification of urea levels (**d**) and viability (**e**) in human islets subjected to *PC* knockdown and exposed to cytokines in the absence (H₂O) or presence of argininosuccinate (AS). Data are means ± s.e.m of n=4 independent experiments from 2 donors.

f, Urea concentrations in human islets cultured in normal (N) media with 5.8 mM glucose compared to glucotoxic (GT) conditions as in 72 h culture in 33 mM glucose, n=13 donors for (N) and n=8 donors for (GT), statistical analysis was performed using student's t-tests.

g, Schematic summary of results showing protective vs non-protective glucose metabolism and PC-directed urea cycle activation as a mechanism that shields cells from stress-induced augmentation in NO and cytotoxicity similar to ARG overexpression or protective GK activation.

h–i, Quantification of urea and NO levels (**h**) and viability (**i**) in human islets expressing PC or ARG2 and cultured as in (f); n = 3 donors in duplicates for urea and NO, and n=5 donors for viability assays. N and GT denote normal and glucotoxic conditions, respectively. Data are means \pm s.e.m. from independent experiments with statistical analyses using one-way (a-e) and two-way (h-i) ANOVA with Tukey adjustment for multiple comparisons.

Table 1 |

Statistical comparisons of blood glucose levels in diabetic mice transplanted with human donor islets shown in Fig. 6

Comparison	P-value
Vehicle vs. RO0281675	0.7507
Vehicle vs. BAD SAHB _A SD	0.0014
Vehicle vs. BAD SAHB _A AAA	0.5685
Vehicle vs. BAD SAHB _A SD + PAA	0.9411
BAD SAHB _A SD vs. RO0281675	0.0100
$\operatorname{BAD}\operatorname{SAHB}_{\operatorname{A}}\operatorname{SD}\operatorname{vs.}\operatorname{BAD}\operatorname{SAHB}_{\operatorname{A}}\operatorname{AAA}$	0.0085
$BAD SAHB_A SD vs. BAD SAHB_A SD + PAA$	0.0114
BAD SAHB _A AAA vs. RO0281675	0.9997
BAD SAHB _A SD + PAA vs. BAD SAHB _A AAA	0.0933
BAD SAHB _A SD + PAA vs. RO0281675	0.0828

One-way ANOVA with multiple comparisons test comparing means of each conditions versus the other for every day that blood glucose as measured. Data are from n=6 (Veh, BAD SAHBA SD), n=3 (RO0281675, BAD SAHBA SD + PAA) and n=4 (BAD SAHBA AAA) mice. *P*-values were calculated using one-way ANOVA with Tukey adjustment for multiple comparisons. Additional details for statistical analyses are provided in Supplementary Tables 2 and 3 online.