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Accessibility
Adenosine kinase inhibition selectively promotes rodent and porcine islet β-cell replication

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Diabetes is a pathological condition characterized by relative insulin deficiency, persistent hyperglycemia, and, consequently, diffuse micro- and macrovascular disease. One therapeutic strategy is to amplify insulin-secretion capacity by increasing the number of the insulin-producing β cells without triggering a generalized proliferative response. Here, we present the development of a small-molecule screening platform for the identification of molecules that increase β-cell replication. Using this platform, we identify a class of compounds [adenosine kinase inhibitors (ADK-Is)] that promote replication of rodent cells in three species (mouse, rat, and pig). Furthermore, the replication effect of ADK-Is is cell type-selective: treatment of islet cell cultures with ADK-Is increases replication of β cells but not that of α cells, PP cells, or fibroblasts. Short-term in vivo treatment with an ADK-I also increases β-cell replication but not exocrine cell or hepatocyte replication. Therefore, we propose ADK inhibition as a strategy for the treatment of diabetes.

Type 2 diabetes mellitus (T2DM) is a progressive disorder of glucose homeostasis that results in diffuse vascular disease and end-organ dysfunction. Although multiple pharmacological therapies for T2DM exist, none of these, as single agents or in combination, prevents the progressive decline in β-cell function or the macrovascular disease complications associated with T2DM (1, 2). Over the next 25 years, the incidence of diabetes is expected to double and reach a prevalence of 25% in the United States, making evident the need for effective new therapies (3).

Insufficient β-cell mass is increasingly recognized as a primary defect in T2DM that contributes to an impaired insulin-secretion capacity (4, 5). The potential therapeutic role of increasing the insulin-secretion capacity of an individual is highlighted by the successful treatment of insulin-dependent type 1 diabetic patients with pancreatic islet transplantation (6–8). However, the ability to restore the endogenous β-cell mass of an individual has not been achieved. Indeed, β cells possess a significant proliferative capacity in rodents and, potentially, in humans (9–11). Studies by our laboratory and others have demonstrated that simple self-duplication is the primary source of new β cells in mice (12–15). Although sources of new β cells other than self-duplication have been indicated under specific circumstances such as pancreatic duct ligation and near-complete β-cell ablation, the potential to harness these sources for β-cell generation is uncertain (16–19). Although the majority of human β-cell proliferative capacity may be lost with age, and the origin of new adult human β cells remains controversial, limited levels of human β-cell replication are observed in association with metabolically demanding conditions and human-to-mouse islet transplantation studies (20–24). Hence, the identification of a method to enhance β-cell replication is of great interest.

Despite the therapeutic potential of a factor that can safely increase the β-cell mass of an individual, no such biological entity has been unambiguously established; initial optimism that glucagon-like peptide receptor (GLP-1R) agonists might be capable of restoring islet β-cell mass has not been sustained (25, 26). Previously, a chemical screen for inducers of β-cell replication was performed with the intent of finding compounds for the treatment of T2DM (27). Unfortunately, this study did not identify compounds with β-cell-selective replication-promoting activity. Therefore, it is likely that these molecular targets have an unacceptable risk profile for use in vivo. One explanation for why only nonspecific mitogen compounds were found is that the screen was performed on a reversibly transformed cell line that may no longer retain metabolic characteristics of primary β cells. Here, we present the development of a screen for compounds that specifically promote primary β-cell replication. Using this platform, we have identified a class of compounds that selectively increase β-cell division and reveal a metabolic governor of β-cell proliferation.

Results

Development and Performance of a High-Throughput Primary β-Cell Replication Assay. To identify compounds that increase β-cell replication, we developed a screening platform using freshly isolated rat islet cells (Fig. 1A). Although the use of primary cells limits the supply of β cells and might be expected to introduce variability between culture preparations, this approach maximizes retention of in vivo metabolic characteristics pertinent to the mitotic behavior of β cells. Our cultures contained ~75% β cells/8 cells (pancreatic and duodenal homeobox 1; PDX-1⁺), ~18% α cells (glucagon⁺), ~3% fibroblasts (vimentin⁺), and ~5% other cell types (Fig. 1B). Although the mixed cell type composition of our culture complicates analysis of β-cell replication, the presence of multiple cell populations within a single well is also advantageous as it allows one to determine whether compound treatment has a β-cell-selective effect: an important feature of a useful therapeutic.

We used the transcription factor PDX-1 to identify β cells (28). Although PDX-1 is expressed by both β and δ cells, the vast majority of PDX-1⁺ cells are insulin⁺ β cells (Fig. 1C). We used nuclear PDX-1 staining as our primary β-cell marker because islet cells grow in dense irregular clusters that cause a cytoplasmic stain, such as insulin, to be ambiguously associated with multiple nuclei. As a result of this ambiguity, ki-67⁺ nuclei from rapidly replicating cells such as fibroblasts have the potential to be incorrectly attributed to insulin⁺ cells. The basal in vitro β-cell replication rate showed moderate inter-experiment variability (4.3–3.5%) and was typically higher than the in vivo β-cell replication rate (0.8 ± 0.2%) of animals of a similar age, as determined by the percentage of PDX-1⁺ cells that coexpressed ki-67.

Adenosine Kinase Inhibitors Promote β-Cell Replication. We screened ~850 compounds from a carefully selected library of cell-permeable bioactive compounds for their ability to increase β-cell replication and identified a class of compounds [adenosine kinase inhibitors (ADK-Is)] that selectively promote primary β-cell replication. Using this platform, we have identified a class of compounds that selectively increase β-cell division and reveal a metabolic governor of β-cell proliferation.


The authors declare no conflict of interest.

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replication, as measured by the percentage of PDX-1\(^+\) cells that coexpressed ki-67 (Fig. S1). Although replication rates were determined using automated image acquisition and analysis, dividing PDX-1\(^+\) cells (yellow arrows) are easily distinguished from dividing PDX-1\(^-\) cells (white arrows) by the colocalization of PDX-1 and ki-67 with visual inspection (Fig. 2A). Two hit compounds were identified, both well-characterized adenosine kinase (ADK) inhibitors (ADK-Is). 5-Iodotubercidin (5-IT) (CAS 24386-93-4) and ABT-702 (CAS 214697-26-4) increased the percentage of dividing PDX cells two- to threefold above the background, independent of a variable baseline PDX cell replication rate, and have a significant effect on PDX cell number (see below). To confirm the results using PDX-1 to identify β cells, similar experiments were performed using PDX-1 and insulin co-staining to identify β cells (Fig. S2A). Indeed, insulin\(^+\) cell proliferation is enhanced by ADK-I treatment. To confirm that ADK is the likely target, we tested additional ADK-Is for their ability to promote β-cell replication. Two additional ADK-Is that demonstrate similar efficacies to the primary hit compounds are shown (Fig. S2 B and C) (29).

Next, we tested the replication effect of ABT-702 on murine and porcine islets to exclude the possibility that these findings are unique to rat β cells. For the porcine β-cell studies, 2- to 3-y-old pigs were selected to test whether enhanced replication would be seen in older animals. Compound treatment of islet cells from both of these species and rats caused a dose-dependent induction of PDX cell replication (Fig. 2B and Fig. S3 A and B). Although 5-IT is more potent than ABT-702, both compounds have a maximum induction of approximately two- to threefold. One notable observation is that the analysis of ABT-702 at concentrations above ~20 μM is limited by background fluorescence. In addition, the replicative effect of these compounds was confirmed using both the mitotic phase marker phosphohistone H3 (PH3) and bromodeoxyuridine (BrdU) incorporation (Fig. S3 C–E). The ability of the ADK-Is to cause a similar induction of β-cell replication across multiple species and induce the expression or incorporation of multiple replication markers (ki-67, PH3, BrdU) confirm that these compounds activate a conserved mitotic pathway in β cells. Furthermore, culturing islet cells in the presence of the 5-IT for 6 d significantly increased the number of PDX-1\(^+\) cells compared with DMSO-treated cultures (Fig. 2C). At day 6, the number of PDX cells in the 5-IT-treated wells had increased by 40% compared with a 20% increase compared to DMSO-treated wells and suggests a change from a basal replication rate of ~3% per day to ~6% per day in our cultures.

**ADK Is Expressed by β Cells and Negatively Regulates β-Cell Replication.** ADK is a member of the sugar kinase group of enzymes, composed of three families (hexokinases, ribokinases, and galactokinases) that play important roles in cellular metabolism (30). ADK is a ribokinase that regulates the intracellular and extracellular adenosine levels through its ability to catalyze the phosphorylation of adenosine to AMP using ATP as the phosphate donor (31). Although this enzyme is broadly expressed, it is highly expressed in the liver and pancreas (32). ADK has two known forms, a long
ADK immunostaining of islet cultures revealed nuclear expression of ADK in β cells. In contrast, ADK staining was in the cytoplasm, not the nucleus, in fibroblasts and α cells (Fig. 3 A–C). Although ADK localization in δ cells (somatostatin+ cells) was variable, ADK was generally present in the nucleus of these cells (Fig. 3D). To ensure that our ADK staining was specific, we validated our antibody by transient transfection of cells with a full-length ADK cDNA, which caused strong nuclear expression (Fig. S4A). ADK expression was also determined in mouse liver (primarily nuclear), adipose (mixed cytoplasmic and nuclear), and muscle (primarily cytoplasmic; we noted high and low ADK-expressing fiber bundles, which may reflect the different muscle fiber types) tissue sections (Fig. S4B). The presence of nuclear staining in β cells, but not α cells, indicates that the long form of ADK is expressed in β cells and not α cells.

We next tested whether the ADK in β cells acts as a negative regulator of replication. Lentiviral infection was used to direct the expression of GFP and either a nonspecific inhibitory RNA (RNAi) or one targeted to ADK. The ability of the ADK-directed RNAi to decrease ADK protein levels was confirmed by Western blot and immunostaining (Fig. S4 C and D). By infecting approximately half of the islet cell culture, as determined by GFP cell expression, we could separately analyze the PDX cell replication rate of infected and noninfected cells within the same well. If ADK acts as a cell-autonomous regulator of β-cell replication, then PDX cells that receive the negative control plasmid and PDX cells that remained uninfected would have the same replication rate, whereas PDX cells infected with the ADK-targeted siRNA virus would have an increased replication rate. Indeed, the results confirmed this prediction (Fig. 3E). Uninfected PDX cells (Fig. 3E, blue bars) and control infected β cells (Fig. 3E, Left, red bar) all had the basal proliferation rate of ~2%. In contrast, PDX cells that received the ADK-directed siRNA demonstrated a 2.5-fold increase in their replication rate (Fig. 3E, Right, red bar). This suggests that ADK is a cell-autonomous negative regulator of β-cell replication and is likely to be the molecular target of the ADK-Is.

ADK-Is Promote β-Cell Replication via mTOR Activation. Several signaling molecules have previously been shown to participate in the regulation of β-cell replication (36–39). To explore the mechanism of ADK-Is-dependent β-cell replication, we determined the β-cell replication rate was measured in islet cultures simultaneously treated with 5-IT plus replication-pathway inhibitor (Fig. 4). Whereas the p38 mitogen-activated protein kinase (MAPK) inhibitor SB203580 caused a small but statistically significant increase in β-cell replication, the phosphoinositide 3-kinase (PI3K) inhibitor wortmannin and the mammalian target of rapamycin (mTOR) inhibitor rapamycin both suppressed ADK-Is-dependent β-cell replication. Although the SB203580 increased β-cell replication in conjunction with 5-IT, we did not see this effect when this compound was tested alone. The inhibitory effects of wortmannin and rapamycin suggest that activity of the PI3K/mTOR signaling pathway might mediate the β-cell replication response to ADK-Is. To test this hypothesis, we treated the rat β-cell line INS1E cells with ADK-Is and determined the phosphorylation status of the ribosomal protein S6 (RPS6), a downstream target of the mTOR signaling cascade (Fig. 4B) (40). Indeed, treatment with either 5-IT or ABT-702 resulted in increased in RPS6 protein phosphorylation. Similar but less-pronounced results were obtained when intact primary rat islets, which comprise multiple cell lineages, were treated with ABT-702 (Fig. S5A). This suggests a molecular mechanism by which ADK-Is promote β-cell replication is likely via activation of the mTOR signaling pathway. It is interesting to note that mTOR is both a cytoplasmic and nuclear kinase, and this may be relevant to the presence of nuclear ADK within β cells (41).

ADK-Is and Glucose or GLP-1R Agonists Have an Additive Effect on β-Cell Replication. Hyperglycemia is considered to be a driver of β-cell proliferation despite relatively little in vitro evidence to support this widely accepted principle (42–44). We took advantage of our β-cell replication platform and demonstrated that glucose has a concentration-dependent effect on β-cell proliferation (Fig. 4C). However, the response kinetics appears to be different from that of ADK-Is. Whereas ADK-Is increase ki-67 staining by 24 h, an effect of glucose is not seen until later (48 h). A distinct mechanism of action for glucose and ADK-Is is supported by the additive effect of 5-IT at all of the tested glucose concentrations (Fig. 4C). The combination of high glucose and 5-IT caused a fivefold induction of the replication rate above the baseline rate. The results of this experiment indicate that the addition of ADK-Is to a hyperglycemic environment might significantly increase β-cell growth in the setting of diabetes. Similar to glucose, GLP-1R agonists are able to increase β-cell replication (45). In our experimental conditions, GLP-1R agonists only modestly increased PDX cell replication (~1.5-fold) compared with 5-IT (~2.5-fold); however, the addition of 5-IT to the GLP-1R agonists led to an
ADK-Is Selectively Promote β-Cell Replication. We assessed the replication rate of multiple cell types in our islet culture: pancreatic polypeptide (PP) cells, α cells, δ cells, and fibroblasts (Fig. 5A and B). We expected that δ cells might show increased replication in response to ADK-Is because, like β cells, δ cells express nuclear ADK (Fig. 3D), secrete their hormone in response to glucose, and share the expression of several key transcription factors including PDX-1 (46). Indeed, δ cells did demonstrate a significant increase in replication in response to ADK-I treatment, whereas fibroblasts, α cells, and PP cells did not (Fig. 5B). The replication rate of PP cells is not shown because their division is extraordinarily rare in culture. In addition to β cells, hepatocytes also express high levels of nuclear ADK and, therefore, might be expected to proliferate in response to ADK-Is (Fig. 5A) (35). However, hepatocyte replication is not increased in response to drug treatment (Fig. 5C). Therefore, the replication effect of ADK-Is is selective and not solely dependent upon nuclear ADK expression.

ADK-Is Promote β-Cell Replication in Vivo. Encouraged by the in vitro results, we tested whether ABT-702 could selectively promote β-cell replication in vivo. ABT-702 was chosen because of its longer half-life compared with 5-IT (47). Indeed, a single intraperitoneal (i.p.) injection of ABT-702 resulted in a twofold increase in BrdU incorporation by β cells in PDX-1-enriched islets (Fig. 5D). These results were confirmed in two separate cohorts of animals in which (i) β cells were identified by the presence of insulin rather than PDX-1 and (ii) replicating cells were identified by the presence of ki-67 rather than BrdU (Fig. S6). Notably, treatment with ABT-702 did not increase the replication rate of exocrine cells, again highlighting the selectivity of ADK-Is (Fig. 5F and Fig. S6D). In addition, BrdU incorporation by hepatocytes was examined in response to ABT-702 treatment (Fig. 5G), and these cells did not show an increased rate of cell division. Therefore, ABT-702 selectively promotes β-cell replication in vitro and in vivo.

Discussion

Historically, T2DM therapies have attempted to augment insulin secretion (e.g., sulfonylureas) or reduce insulin demand (e.g., biguanides, thiazolidinediones) by lowering peripheral resistance. However, T2DM patients appear to have a limited capacity for adaptive β-cell expansion, and these approaches do not address this deficiency. Here, we present a platform to identify pharmacological agents that promote increased β-cell division. In contrast to previously identified growth-promoting compounds, which lack cell type specificity (27, 48, 49), ABT-702, a well-identified ADK-Is as a class of agents that is capable of promoting β-cell replication in vitro and in vivo. Of critical importance is that these compounds preliminarily appear to have a selective proliferation effect on β cells and not α cells, hepatocytes, exocrine cells, or fibroblasts. The molecular mechanism for β-cell selectivity is not apparent. Although we believe that the presence of nuclear ADK may be relevant to the observed selectivity, other cell types, such as hepatocytes, also have nuclear ADK but do not replicate in response to ADK-Is. We hypothesize that intracellular (possibly nuclear) adenosine levels within β cells activates the mTOR pathway with unique consequences within β cells. A notable limitation of the current study is that we have tested the proliferative effect of ADK-Is on a restricted number of cell types. This issue and whether ADK inhibition can promote the replication of normal and diabetic human β cells will need to be addressed in future studies.

The primary physiological cue for β-cell replication is thought to be glucose (44). Because the diabetic condition is defined by insufficient insulin supply despite excess glucose, diabetes, in part, represents a failure of the compensatory β-cell response to excess glucose. Here, we have identified a pharmacological stimulus for β-cell replication that is glucose-independent and, therefore, has the potential to circumvent defects in the glucose-response pathway. The additive replication effect of ADK-Is and GLP-1R agonists raises the possibility for simultaneous use of these agents in the treatment of T2DM. An important question
is whether long-term in vivo treatment with ADK-Is will improve glucose tolerance by augmenting β-cell mass in rodents and humans. To answer this experimental question, it may be necessary to develop ADK-Is with improved potency (the identified compounds are active in the 1–10 μM range) and less central nervous system accessibility to avoid central nervous system effects that have been observed with ADK inhibition (50).

Although inhibition of ADK causes cellular efflux of adenosine, we do not believe that the growth-promoting effect of ADK-Is on PDX-1+ cells is mediated by paracrine/autocrine adenosine signaling (51). First, we observe a cell-autonomous increase in β-cell proliferation in response to ADK knockdown. If this effect were mediated by extracellular adenosine, a paracrine effect would be anticipated. Second, the addition of adenosine receptor agonists and antagonists that were included in our screening libraries had no effect on β-cell proliferation in our assay. Third, β-cell replication in response to ADK-Is requires mTOR activity. Interestingly, prior work has suggested that mTOR activity is directly influenced by cellular ATP levels and that mTOR can be found in the nucleus of normal cells (41, 52). Whether the nuclear localization of ADK in β cells and the primary function of ADK in maintaining nucleotide pools is relevant to the mechanism of mTOR activation and the induction of β-cell replication by ADK-Is remains to be determined.

The identification of ADK as a regulator of β-cell replication is an unexpected finding that highlights the value of using chemical screening to reveal new biology. The methodology established here uses primary islet β cells may be usefully applied toward the study of γ-cells (somatostatin), and values were obtained using two-tailed t test.

Methods

Islet Isolation and Primary Screen Protocol. Rat (250-g Sprague–Dawley; Charles River) and mouse (12-wk-old animals; C57BL/6; Jackson Laboratory) islets were isolated as described previously (53). The use of animals was approved and carried out in accordance with our institutional animal care and use committee. Porcine islets from retired breeders were provided by VitaCyte. Islets were incubated (37 °C; 5% CO2) overnight in islet media (99–786-CV (Mediatech); 10% (vol/vol) FBS serum (Valley Biomedical); BS3033; 8.3 mM glucose (Sigma; G7528); 1x penicillin/streptomycin (Invitrogen; 15070–063); 1x Gentamycin (Invitrogen; 35050–079)). The following morning, islets were trypsinized into cellular clusters of 1–3 cells, resuspended in islet media and plated into the wells of a 96-well plate (Sigma; CLS3904) that had been coated with 804G (a rat bladder carcinoma cell line) conditioned media. The cellular plating density was 70,000 cells/well, and >95% viability was confirmed at the time of plating. The islet cells were allowed 48 h to adhere; at that time, the media were changed (as above, except 2% (vol/vol) serum, 5 mM glucose), and the cells were compound-treated. For screening, compounds were tested at 1 and 10 μM concentrations in duplicate on a single occasion. After 24 h of compound treatment, cells were fixed with fresh 4% (wt/vol) paraformaldehyde. See SI Methods for additional details regarding immunohistochemical and automated replication analysis.

Quantification of in Vivo Replication. Twelve-week-old C57/86 female animals were injected with BrdU (Sigma; B5002; 10 μg/kg) and with either ABT-702 (21 mg/kg) or DMSO vehicle. Twenty-four hours posttreatment, the animals were killed, and the relevant organs were harvested. A similar experiment was performed in 6-wk-old C57/86 female animals without female treatment. In these animals, replication was assessed after 24 h by ki-67 staining. All experiments were performed with a minimum of four animals per treatment
group. Every fourth 12-μm section was used for analysis, and a minimum of 2,000 β cells, exocrine cells, and hepatocytes per organ per animal were counted. Analysis was performed by manual picture acquisition and cell counting. β Cells were identified by either PDX-1 staining or insulin staining, with similar results. Exocrine cells were approximated by counting all nuclei outside the islet structure. A minority of these cells were not exocrine cells. Hepatocytes were identified as DAPI+ albumin+ cells (Bethyl Laboratories; A90-234A). Dividing cells were BrdU+ (Amersham; RPN202).

### Statistics

Data are presented as the means of multiple replicates performed simultaneously. All of the experimental results presented were repeated more than twice. Error bars show the SD unless otherwise specified. Results were compared using the two-tailed t test. The EC₅₀ was calculated using nonlinear regression with the highest replication rate constrained.

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