MNK2 deficiency potentiates β-cell regeneration via translational regulation

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Regenerating pancreatic β-cells is a potential curative approach for diabetes. We previously identified the small molecule CID661578 as a potent inducer of β-cell regeneration, but its target and mechanism of action have remained unknown. We now screened 257 million yeast clones and determined that CID661578 targets MAP kinase-interacting serine/threonine kinase 2 (MNK2), an interaction we genetically validated in vivo. CID661578 increased β-cell neogenesis from ductal cells in zebrafish, neonatal pig islet aggregates and human pancreatic ductal organoids. Mechanistically, we found that CID661578 boosts protein synthesis and regeneration by blocking MNK2 from binding eIF4G in the translation initiation complex at the mRNA cap. Unexpectedly, this blocking activity augmented eIF4E phosphorylation depending on MNK1 and bolstered the interaction between eIF4E and eIF4G, which is necessary for both hypertranslational and β-cell regeneration. Taken together, our findings demonstrate a targetable role of MNK2-controlled translation in β-cell regeneration, a role that warrants further investigation in diabetes.

Both type 1 and type 2 diabetes manifest with elevated circulating glucose levels caused by the deregulation of insulin signaling and/or the loss of functional insulin-producing β-cells. Although daily insulin injection, lifestyle interventions and various drug treatments can manage the disease, there is currently no available cure. Therefore, stimulating endogenous β-cell regeneration is an attractive curative approach for diabetes; however, current efforts have failed to translate into a clinically approved drug.

Drug screening in vivo for chemicals stimulating β-cell regeneration has the potential to accelerate the drug discovery process as it is performed in a physiological, whole-organism setting. The zebrafish model has emerged as a powerful tool for performing unbiased, large-scale chemical/genetic screens that are directly coupled to phenotypic analyses. These types of screens have identified molecules that have already entered clinical development11, showcasing the translational potential of the approach. Metabolism, and in particular diabetes research, is an area in which chemical screens using zebrafish have grown in popularity. Such screens have already identified compounds that can stimulate β-cell proliferation14,15 and neogenesis from duct-residing pancreatic progenitors16,17, two of the main known mechanisms of endogenous β-cell regeneration. Moreover, zebrafish chemical screens for regulators of glucose metabolism have identified compounds with potential use as antidiabetic treatments16–18. Collectively, these studies have demonstrated the power of the zebrafish model for performing chemical screens to identify compounds that could be repurposed as antidiabetic drugs. However, many hits from phenotypic screens have unknown targets and mechanisms of action that create a bottleneck for further development but, if the targets are defined, can open up new research areas.

In this work, we aimed to identify the molecular mechanism of action of CID661578 (1), the most striking hit from a zebrafish chemical screen for stimulators of β-cell regeneration1. By performing a modified yeast two-hybrid screen suited for drug target deconvolution, we identified MAP kinase-interacting serine/threonine kinase 2 (MNK2) as the molecular target of CID661578 and validated this interaction in vivo. MNK2 participates in initiation of mRNA translation and has been postulated to modulate the process in a transcript-selective fashion19. Here, we show that kinase-independent blocking of MNK2 leads to bolstered protein synthesis in the pancreatic duct and that the effects are conserved across zebrafish, pig and organoid cultures of human pancreatic ducts. Overall, our results demonstrate a conserved pathway to stimulate β-cell neogenesis by boosting protein synthesis through targeting MNK2.

Results

Yeast chemical hybrid screen identifies MNK2 as the molecular target of CID661578. In a previous large-scale chemical screen, we identified five small molecules that potently drove β-cell regeneration in zebrafish larvae4. Four of the five hit compounds converged on the adenosine pathway and stimulated β-cell proliferation, while the fifth hit compound, named CID661578, had no known molecular target or cellular mechanism of action (Fig. 1a). To identify the molecular target of CID661578, we used yeast chemical hybrid (YChemH) screening technology, which takes advantage of the...
Fig. 1 | YChemH screen identifies MNK2 as the molecular target of CID661578. a, Schema for the screening of compounds increasing β-cell regeneration using a transgenic zebrafish model for β-cell ablation and approximately 10,000 compounds. The hits included four compounds affecting adenosine signaling and CID661578 with an unknown mechanism. b, Schematic showing the structures of CID661578 and the analog CID661578.6 along with the screening strategy (YChemH). The red circles highlight the structures that were altered in CID661578. Survival of yeast on selective histidine-free medium was the output of the screen for clones expressing interactors of the CID661578.6 bait; TMP, trimethoprim; AD, activation domain. c, Table summarizing the top hits of the YChemH screen from the two cDNA libraries. The A-classified hits (drl and acin1b from the zebrafish embryo library and MNK2 from the human islet library) have a higher probability of being true targets of CID661578.6 than B- and C-classified hits. d, Validation of the MNK2–CID661578.6 interaction with different concentrations of CID661578.6 bait and an MNK2-expressing yeast clone. DMSO demonstrates the sensitivity to the selective medium, and yeast clones did not survive in the selective histidine-free medium. The interaction between MNK2 and CID661578.6 promoted yeast survival, as illustrated by the multiple colonies at the four spots of inoculation (decreasing levels of inoculation from the top to the bottom). Each condition was tested in two replicates. e, Validation of the zebrafish Mnk2b–CID661578.6 interaction with different concentrations of CID661578.6 bait and two different DHFR hook vectors. Experiments using the original hook vector, N-LexA–DHFR-C, are listed as 1, 2, and 3. Experiments using the modified vector with the reverse order, N-DHFR–LexA-C, are listed as 4, 5, and 6. Both full-length zebrafish Mnk2b (3 and 6) and a fragment (2 and 4) corresponding to the original fragment of the human MNK2 identified in the screen were used. Human MNK2 was used as a positive control (1 and 4), and zebrafish Mnk2b only mediated binding when expressed by the hook vector with the reverse order (5 and 6) to the one used in the original screen (explaining why zebrafish Mnk2b did not show up as a hit in the original screen).
classical yeast two-hybrid system for protein–protein interactions but enables screening for protein targets of small molecules. First, we generated a series of structural analogs of CID661578 to simplify the structure and identified one (termed CID661578.6 (2)) that exerted similar effects on β-cell regeneration as CID661578 (Extended Data Fig. 1). We used yeast clones with a construct that expressed a DNA-binding domain (LexA) coupled to the enzyme dihydrofolate reductase (DHFR) along with a GAL4 activation domain fused to each cDNA in the libraries. We screened two different cDNA libraries, one derived from human islets and another derived from zebrafish embryos, with a total of 135 million and 122 million clones, respectively. The rationale of this approach is that when yeast are incubated with the CID661578.6-derived bait (3), the attached trimethoprim will interact with DHFR on the DNA-binding site of LexA. Interaction between CID661578.6 and a protein fragment (that is, the prey) generated from the cDNA library will lead to yeast survival on the selective histidine-free medium (Fig. 1b). The most likely targets of CID661578 were classified as A hits, and less likely targets were classified as B hits, C hits and so forth, depending on the confidence in the interaction (Fig. 1c).

Next, we followed up on hits that had previously been shown to affect metabolism and could validate the interaction in an MNK2-expressing yeast clone with different concentrations of the CID661578.6-derived bait in the YChemH system (Fig. 1d). Subsequently, we assessed whether the zebrafish homolog of MNK2 (Mnk2b) could also bind CID661578.6 because it did not appear as a hit in the zebrafish cDNA library. To this end, we cloned the cDNA sequence of mknk2b (the genes encoding Mnk kinases are called mnk) in frame with the GAL4 activation domain and used two different DHFR hook vectors of the YChemH system. The YChemH assay results revealed that zebrafish Mnk2b only bound to CID661578.6 when the new, modified DHFR hook vector was used, while human MNK2 bound to CID661578.6 regardless of the vector conformation (Fig. 1e), explaining why Mnk2b did not show up as a hit in the original screen of the zebrafish cDNA library. In summary, through a series of in vitro experiments, we identified MNK2 as the likely molecular target of CID661578.

CID661578.6 promotes β-cell regeneration of ductal origin. To identify the cellular source of the newly formed β-cells, we first assessed their proliferation status. We used the β-cell ablation zebrafish model, Tg(ins:flag-NTR), in which the enzyme nitroreductase (NTR) is expressed under the control of the β-cell-specific insulin promoter. When the prodrug metronidazole (MTZ) is administered, NTR converts it to a toxic byproduct, resulting in the specific ablation of β-cells.14-15 We treated the zebrafish β-cell ablation model with CID661578 in the presence of EdU to label dividing cells, and quantification of EdU phase-contrast images showed no alteration in proliferation status between the control and CID661578-treated groups (Extended Data Fig. 2a–c). Next, we performed lineage tracing of the pancreatic ductal cell population using the inducible Tg(tp1:creER2) line (tp1 is a Notch-responsive element characterizing the duct population) and the responder Tg(abwish:split) line. We induced Cre-mediated recombination (5–6 days post-fertilization (d.p.f.)) and let the larvae grow to the juvenile stage, followed by β-cell ablation and treatment with CID661578.6 (Fig. 2a). Quantification of β-cells along the tail of the pancreas (often referred to as secondary islets) revealed that treatment with CID661578.6 increased the number of regenerating β-cells derived from the ductal cell population (Fig. 2b–c). Moreover, we used a complementary lineage tracing approach to validate these findings, where double transgenic Tg(tp1:H2BmCherry);Tg(ins:GFP) zebrafish in which tp1 drives the expression of the stable fluorescent protein H2BmCherry as a ductal cell tracer were used, and we confirmed ductal-derived β-cell regeneration (Extended Data Fig. 3a–e). Furthermore, we excluded the possibility that the chemical treatment altered the number and proliferation of the Notch-responsive ductal cells by assaying for EdU incorporation into Tg(tp1:GFP) zebrafish (Extended Data Fig. 2d–g). Thus, by using two different lineage tracing strategies, we identified ductal cells as the cellular source of the newly formed β-cells.

We subsequently assessed glucose levels after ablation of β-cells and treatment of adult fish with CID661578.6 for 3 d. As expected, β-cell ablation caused an increase in blood glucose at 3 d after ablation in the control fish, whereas fish treated with CID661578.6 had significantly lower blood glucose than controls (Fig. 2f). Glucose levels were also reduced in zebrafish larvae following CID661578 or cercosporamide (4) (a previously described selective inhibitor of MNK2 (ref. 16)) treatment (Extended Data Fig. 2h). The newly formed β-cells also appeared mature in terms of mnx1 expression (Extended Data Fig. 4a–d) as well as being devoid of glucagon and insulin coexpression (Extended Data Fig. 4e–g). Taken together, these results show that compounds interfering with Mnk2 lower glucose levels in both larval and older zebrafish.

Next, we explored mknk2b expression in the adult zebrafish pancreas using a recently published single-cell RNA-sequencing (RNA-seq) dataset17 that contains all the major pancreatic cell types (Fig. 2g and Extended Data Fig. 5). The expression of mknk2b was not ductal specific, yet its highest expression was in a cluster of ductal cells (Fig. 2h). Taken together, these results showed that CID661578.6 increased the regeneration of functional β-cells in both larval and older zebrafish by promoting β-cell neogenesis from a ductal origin.

CID661578 targets Mnk2b in vivo to drive β-cell regeneration. Subsequently, we wanted to determine whether the in vivo engagement of Mnk2b is responsible for the observed phenotypes of CID661578 treatment. We treated zebrafish larvae with CID661578, cercosporamide or their combination from 4 to 6 d.p.f. during β-cell regeneration. We observed a dramatic increase in regenerating β-cells in the primary islets of zebrafish larvae following treatment with either chemical, but no additive effect was observed when they were combined (Fig. 3a–e). This result suggested that cercosporamide and CID661578 have similar effects on β-cell regeneration, and the absence of additive/synergistic effects indicated that they converge on a common molecular target/pathway. Further, we found that neither CID661578 nor cercosporamide treatment affected the development of any of the endocrine cell populations in zebrafish larvae, suggesting that the effects of this pathway are restricted to the regenerative state (Extended Data Fig. 6a–i).

We also tested if the effects of the chemicals targeting Mnk2 could be reproduced using genetic approaches. The mknk2 gene is duplicated in the zebrafish genome, and an investigation of the expression of the two paralogs mknk2a and mknk2b in published RNA-seq data revealed that mknk2b is the predominantly expressed paralog in ductal cells. Therefore, we generated a mknk2b full-body knockout zebrafish using CRISPR–Cas9 mutagenesis to target the N-terminal part of the protein. First, we observed physiological β-cell development in the homozygous mutants (Fig. 3f–i). However, β-cell regeneration was enhanced in the homozygous mknk2b mutants following β-cell ablation (Fig. 3j–m), suggesting that the absence/inhibition of Mnk2b is responsible for this phenotype. We reproduced the mknk2b-knockout phenotype using a splice-blocking morpholino to knockdown mknk2b (validated by quantitative PCR with reverse transcription (RT–qPCR) (Extended Data Fig. 7k)). Knockdown of mknk2b increased the number of β-cells, while no additive effect was observed from simultaneous mknk2b knockdown and CID661578.6 treatment (Extended Data Fig. 7a–e). In addition, morpholino knockdown of the other two A hits from the zebrafish library, acin1b and drl, did not increase β-cell regeneration, suggesting that these two genes are not responsible for...
Fig. 2 | CID661578.6 increases β-cell regeneration from a pancreatic ductal origin and lowers glucose levels. a, Schematic of the lineage tracing experiment. Briefly, larvae were treated with 4-hydroxytamoxifen (4-OHT) for 24 h (5–6 d.p.f.) to induce recombination of the reporter. At 28 d.p.f., the fish were treated with MTZ for 24 h to ablate the β-cells, followed by 48 h of treatment with DMSO or CID661578.6. b–e, Representative images of Tg(ubi:switch); Tg(tp1:creERT2); Tg(ins:flag-NTR) fish treated with DMSO (b) or 2 μM CID661578.6 (c) and immunostained for insulin at 31 d.p.f.; scale bars, 20 μm. Quantifications of the number of β-cells in the secondary islets along the tail of the pancreas (d) as well as the number of β-cells derived from Notch-responsive cells (e) are shown; n = 21 (control) and n = 18 (CID661578.6) for d–e. An unpaired two-tailed Student’s t-test was used to assess significance for d (⁎P = 0.0393), and a two-tailed Mann–Whitney test was used for e (⁎⁎P = 0.0087). Data are presented as mean values ± s.e.m. The experiment shown in b and c was repeated twice with similar results. f, Blood glucose was measured 3 d post-β-cell ablation (d.p.a.) in 4-month-old fish treated with DMSO or CID661578.6. Blood glucose levels in zebrafish without β-cell ablation were included as a basal-state reference; n = 7 (control), n = 10 (control, 3 d.p.a.), n = 10 (CID661578.6, 3 d.p.a.). A one-way ANOVA followed by Šidák’s multiple comparisons test was used to assess significance for f (⁎⁎P = 0.0078). Data are presented as mean values ± s.e.m. g,h, UMAP plots showing the different cell types present in the adult zebrafish pancreas after reanalysis of published single-cell RNA-seq data (g) and expression of mknk2b (h) at various levels in the different clusters.
the observed phenotypes (Extended Data Fig. 7f–m). These results further supported that β-cell regeneration increased in the absence of mknk2b.

Finally, we reasoned that because the knockout and knockdown of mknk2b had similar effects as CID661578.6 treatment, overexpression of the protein would sequester CID661578.6 and reduce β-cell regeneration. To this end, we cloned zebrafish mknk2b and human MKNK2 and overexpressed them under the control of the tp1 promoter in the β-cell ablation model, which was subsequently treated with CID661578.6. Overexpression of either mknk2b or MKNK2 significantly decreased the effect of CID661578.6 on β-cell regeneration (Fig. 3n–r). These experiments using mosaic overexpression were also confirmed in a stable line overexpressing mknk2b, which showed an even stronger reversal of the CID661578.6 effect on β-cell regeneration (Extended Data Fig. 7n–r). Collectively, these data support Mnk2b as the molecular target of CID661578.6 in vivo and that Mnk2b can restrict β-cell neogenesis from a ductal origin.

CID661578 boosts translation to increase β-cell regeneration. To better understand the molecular mechanism induced by CID661578, we treated zebrafish larvae with CID661578 for 24 h before global metabolomics characterization(15). After creating a metabolite profile of CID661578-treated zebrafish, we identified differentially regulated metabolites (Fig. 4a). An interesting observation was that the levels of many amino acids were altered following CID661578 treatment. MNK2 interacts with a complex of eukaryotic translation initiation factors and thereby plays a role in protein synthesis(16). Thus, the metabolomics data indicated possible changes in protein synthesis as a key effect of CID661578, consistent with the known role of Mnk2. Moreover, α-d-glucose was significantly downregulated, an observation that we replicated using glucose measurements with an in vitro assay for both CID661578 and cercosporamide (Fig. 4a and Extended Data Fig. 6b). Metabolite set enrichment analysis of zebrafish-specific pathways showed that downregulated metabolites following CID661578 treatment were related to non-essential amino acid metabolism (Fig. 4b), while the pathways related to upregulated metabolites were less impacted and limited to changes in pyrimidine metabolism (Extended Data Fig. 8a). Further, enrichment analysis of the single-cell RNA-seq data used to examine the expression of mknk2b showed that genes enriched in the ducetal cells have a role in mRNA translation (Extended Data Fig. 8b). Taken together, these results further strengthen the hypothesis that CID661578 affects global changes in protein synthesis and glucose metabolism, in agreement with protein synthesis being a highly energy-consuming process that should reduce nutrient levels, including glucose.

To further investigate the changes in protein synthesis in vivo, we measured the incorporation of O-propargyl-puromycin (OPP), a modified amino acid that is incorporated in proteins during translation and can be visualized with a Click-IT reaction. Following β-cell ablation, larvae were incubated with OPP for 20 h concomitant with treatments of CID661578.6, 4EGI-1 or their combination. 4EGI-1 inhibits the interaction between the translation initiation factors eIF4E and eIF4G, which together with Mnk2 function within the translation initiation complex(17). The rationale of this assay was to assess the outcome of altering the translation initiation complex composition during CID661578.6 treatment. OPP incorporation was predominantly observed in the ducetal cells of the pancreas, indicating a high protein synthesis rate in this population. CID661578.6 treatment drastically increased the incorporation of OPP in the ducetal cells (and the intestine), an effect abolished following cotreatment with the inhibitor 4EGI-1 (Fig. 4c–h). These data suggest that CID661578.6 boosts protein synthesis and that its effect is dependent on the translation initiation complex.

Encouraged by the observation that 4EGI-1 inhibited CID661578.6-induced protein synthesis, we examined whether 4EGI-1 could also inhibit the induced β-cell regeneration. To this end, zebrafish larvae were treated during the regenerative period with CID661578.6, 4EGI-1 or their combination. Interestingly, 4EGI-1 treatment was also sufficient to inhibit CID661578.6-induced β-cell regeneration (Fig. 4l–m). Thus, through a combination of metabolomics and protein synthesis measurements, we demonstrated that CID661578.6 induces protein synthesis in vivo and that the effect on both protein synthesis and β-cell regeneration is blocked by targeting the translation initiation complex that Mnk2 is a part of.

CID661578.6 modulates the translation initiation complex. We sought to identify how interference with Mnk2 affects translation initiation complex composition. First, we performed an in vitro characterization of the kinase activity of Mnk2 in the presence of CID661578, CID661578.6 and the known Mnk2 inhibitor cercosporamide. We decided to assess the inhibitory effect of the chemicals on the kinase activities of Mnk2, Mnk1 and Jak3, which all have been shown to be inhibited by cercosporamide(18). We observed that the kinase activity of all three kinases remained unchanged after treatment with either CID661578 or CID661578.6, whereas cercosporamide potently inhibited Mnk2 and partially inhibited Mnk1 and Jak3 (Fig. 5a–c). Subsequently, we performed a kinome screen in which we assessed the kinase activity of 140 human kinases.
after treatment with CID661578.6 or cercosporamide. We did not observe any drastic changes in kinase activity with CID661578.6 treatment, whereas cercosporamide decreased the kinase activity of numerous kinases (Extended Data Fig. 9a,b). These results suggest that CID661578.6 does not affect protein synthesis by inhibiting the kinase activity of MNK2.

We then reasoned that CID661578.6 binding to MNK2 could alter the composition of the translation initiation complex. Given that inhibiting the interaction between translation initiation factors eIF4E and eIF4G was sufficient to block the effects of CID661578.6 in vivo, we hypothesized that CID661578.6 binding to MNK2 may enhance the interaction between eIF4E and eIF4G at the mRNA cap.
To test our hypothesis, we treated the COLO 320HSR cell line with CID661578.6, 4EGI-1 or their combination. Subsequently, we pulled down the cap-binding protein eIF4E from cell lysates using beads with an immobilized m’GTP structure of the mRNA cap. Treatment with CID661578.6 stabilized the eIF4E–eIF4G interaction, an effect that could be reversed following treatment with the inhibitor 4EGI-1 (Fig. 5d). Subsequently, we performed the same m’GTP pulldown assay in vitro using rabbit reticulocytes. The advantage of using this in vitro system is that rabbits do not have an ortholog of MNK2. CID661578.6 did not increase the eIF4E–eIF4G interaction in rabbit reticulocytes, indicating that the increase in the eIF4E–eIF4G interaction after CID661578.6 treatment is dependent on MNK2 (Fig. 5e). Lastly, we asked how the interaction between CID661578 and MNK2 could affect the recruitment of MNK2 to the translation initiation complex. For this experiment, we used the human pancreatic cancer cell line Panc-1, as it was more efficiently transfected than the COLO 320HSR cell line. We began by validating that both CID661578 and its analog CID661578.6 increased the eIF4G–eIF4E interaction in Panc-1 cells by using the m’GTP pulldown assay (Fig. 5f). Next, we transfected Panc-1 cells with a FLAG–MNK2 plasmid, added DMSO/CID661578 and performed immunoprecipitation of MNK2 using anti-FLAG. Immunoblotting against eIF4G (which is the protein that directly interacts with MNK2 in the complex) revealed that CID661578 treatment largely abolished the interaction between MNK2 and eIF4G (Fig. 5g). Finally, we assessed whether phosphorylation of eIF4E was affected by CID661578-induced changes in the translation initiation complex composition. Unexpectedly, we observed that phosphorylation of eIF4E was increased after treatment with CID661578 and csecospomamide (that is, selective interference of MNK2) but was nearly abolished with two broader inhibitors blocking both MNK1 and MNK2 (that is, CGP57380 and eFT508; Fig. 5h). To address whether increased eIF4E phosphorylation is important for β-cell regeneration, we cotreated fish with CID661578 and eFT508 and observed that eFT508 could inhibit the increase in β-cell numbers induced by CID661578 (Fig. 5i). Taken together, these data suggest that CID661578 binds to MNK2, preventing it from interacting with eIF4G, which can increase MNK1-dependent phosphorylation of eIF4E and bolsters the interaction between eIF4E, eIF4G and the mRNA, resulting in increased protein synthesis.

As alterations in both eIF4E phosphorylation and increased eIF4F–complex (that is eIF4E, eIF4G and eIF4A) formation affects mRNA translation in a transcript-selective fashion21,22, we sought to identify translationally regulated mRNAs following both CID661578 and csecospomamide treatments. To this end we performed poly-some profiling of Panc-1 cells after chemical treatments and used RNA sequencing to quantitate total mRNA and mRNA associated with more than three ribosomes (Fig. 5j). We then identified mRNA whose translational efficiency was modulated along with mRNAs with changed abundance and translationally buffered (Fig. 5k,l and Methods). We focused our analysis on transcripts with altered translational efficiencies predicted to affect protein levels and found that there was a highly significant overlap of hypo- and hypertranslated mRNAs between CID661578 and csecospomamide treatments (Extended Data Fig. 10a–c). Overall, we identified a total of 270 hypertranslated and 99 hypotranslated mRNAs that were shared between treatments (Supplementary Data 1–3). This further highlights that csecospomamide and CID661578 target overlapping molecular pathways in Panc-1 cells. Gene ontology (GO) analysis identified several GO terms enriched among proteins encoded by mRNAs that were hypotranslated in response to both compounds, with the most striking being mitochondrial-related processes (Extended Data Fig. 10d). By contrast, there were no significantly enriched pathways among shared hypertranslated mRNAs. Lastly, we examined the 5’ untranslated region (5’ UTR) sequences in search of features of translationally regulated mRNAs. Our analysis demonstrated differences in GC content among translationally regulated mRNAs following chemical treatments (Extended Data Fig. 10e,f), a feature underlying 5’ UTR structures. Therefore, both CID661578 and csecospomamide modulate mRNA translation in a selective fashion where the hypertranslated mRNAs had 5’ UTRs with low GC content and the hypotranslated mRNAs had 5’ UTRs with high GC content, consistent with previous studies on eIF4E-regulated translation21,22.

**CID661578-induced β-cell neogenesis translates to mammals.** To examine whether our findings were translatable to mammals, we took advantage of an in vitro culture system of neonatal pig islet aggregates. Pancreata from 3-d-old pigs were digested, and the islet aggregates were generated and cultured in vitro for 3 d before a 5-d treatment with CID661578 or csecospomamide. These islet aggregate preparations are highly enriched in intrasislet ductal cells, making them an ideal model to study the effect of the assayed chemicals. MNK2 is expressed in the duct as well as in islets of juvenile and adult pigs (Extended Data Fig. 9c–e). Treatment with either CID661578 or csecospomamide increased the number of insulin+ cells in the islet aggregates (Fig. 6a–d). The number of CK7+ ductal cells decreased after treatment with CID661578, while the number of cells coexpressing insulin and CK7 increased following either treatment (Fig. 6e,f). These results indicated that the new β-cells also have a ductal origin in neonatal pig islets and showed that the increase of β-cells in the zebrafish could be translated to a mammalian model.

Lastly, we stained human pancreatic sections for MNK2 to assess its expression. We observed MNK2 expression in islets without prominent expression in β-cells (Fig. 6g). Interestingly, we observed a distinct strong expression of MNK2 in a sparse...
**Pathway impact**

- **log (P)**

**D-Glutamine and D-glutamate metabolism**

| Compounds | Impact |
|-----------|--------|
| Alanine, aspartate and glutamate metabolism | 6.4 |
| OPP; tp1:GFP; Hoechst | 4.0 |

**OPP fluorescence intensity of tp1:GFP+ cells (AU)**

- **Control**
- **CID661578.6**
- **4EGI-1**
- **CID661578.6 + 4EGI-1**

**β-Galactosidase activity (mean)**

- **Control**
- **CID661578.6**
- **4EGI-1**
- **CID661578.6 + 4EGI-1**

**DMSO**

- **DMSO1**
- **DMSO2**
- **DMSO6**
- **DMSO5**
- **DMSO4**
- **DMSO3**

**CID1**

- **CID661578.64**
- **EGI-1**
- **CID661578.6**
- **6**
- **4**
- **EGI-1**

**Downregulated after treatment with CID661578**

- Upregulated after treatment with CID661578
population of cells along the pancreatic ducts. The MNK2-expressing cells that were located just outside the luminal ductal lining most often did not coexpress MNK2 and CK19. However, we also observed MNK2+CK19+ ductal cells at other positions in the same ducts (Fig. 6h,i). To address whether CID661578 or cercosporamide can stimulate differentiation of human ductal cells toward β-cells, we generated ductal organoid cultures from healthy human donors (Fig. 6j). Encouragingly, treatment of the human organoid cultures with either CID661578 or cercosporamide increased the expression of INS mRNA compared to the DMSO-treated controls (Fig. 6k). In sum, taking into account the potent effect of MNK2-interfering drugs on β-cell differentiation in neonatal pig islets and human ductal organoids as well as the intriguing expression pattern of MNK2 in humans, further investigation of the translational potential of this class of drugs is warranted.

Discussion

In the current study, we identified the molecular target of CID661578 as MNK2. The MNK2–CID661578 interaction potently induced β-cell regeneration from a pancreatic ductal cell origin and was sufficient to improve glucose control in both larval and adult diabetic zebrafish models. CID661578 was the only hit from our previous chemical screen that did not have a known target. An important step in the drug discovery process is the identification of protein interactors and target engagement in vivo. Here, we used the YChemH system and identified MNK2 as one of the molecular targets of CID661578. Additionally, we used a combination of polysome profiling and in vitro biochemical experiments to shed light on the molecular mechanism of action of CID661578 in pancreatic ductal cells.

The neogenesis of β-cells from pancreatic ductal cells has been previously observed in the zebrafish model and has now been accepted as an endogenous path for β-cell regeneration. However, whether a similar pancreatic progenitor population resides within the ductal cell compartment in adult mammalian models remains unclear. Different lineage tracing methods in mouse models have yielded various results regarding the contribution of ductal cells to β-cell neogenesis. This controversy highlights the need to study ductal cells and their progenitor potential in the zebrafish model to identify new effectors and markers that could be translated to mammalian systems. Interestingly, CID661578 treatment also increased the formation of new β-cells from intraislet ductal cells in cultures of neonatal pig islet aggregates and stimulated INS expression in ductal-derived human organoids. These results demonstrated that the pathway could also be exploited to increase the differentiation of β-cells from a ductal source in mammalian systems.

We also observed that the Mnk2b–CID661578 interaction potently lowered glucose levels in zebrafish, suggested to be due to a combination of increased β-cell regeneration and protein synthesis (one of the most energy-consuming processes in the cell). This was supported by the fact that glucose lowering also occurred during homeostasis following chemical treatment, suggesting that increased glucose consumption is possibly the fuel for the increased protein synthesis. Interestingly, a recent report has linked analogs of cercosporamide to a glucose-lowering effect in mice. Furthermore, Mnk1- and Mnk2-knockout mice exhibit beneficial metabolic outcomes when challenged with a high-fat diet. These data are consistent with our observations and open new avenues for exploiting MNK2 in metabolic diseases.

During the course of our study, we performed metabolomics to assess the effects of CID661578 treatment in vivo, which highlighted global changes related to glucose metabolism and protein synthesis. We observed that the chemical treatment increased protein synthesis in vivo, an effect that was most profound in the ductal cell population. Hypertranslation as a mechanism that governs stem cell differentiation has recently been demonstrated for cell types belonging to a few different organs. Our study expands this concept to the pancreas and to a regenerative setting in vivo, suggesting that targeting initiation of translation could represent a conserved process stimulating differentiation and regeneration in multiple systems.

MNKs belong to the MAPK interacting protein kinase family and were identified in screens for interactors of the ERK and p38 MAP kinases. MNKs primarily phosphorylate eIF4E and thereby have a context-dependent role in protein synthesis. However, although our data indicated that CID661578 treatment increased protein synthesis, it did not affect the kinase activity of MNK2 in vitro. Instead, we observed that CID661578 increased the interaction between mRNAs, the cap-binding protein eIF4E and the scaffold protein eIF4G, which also binds to MNK1/MNK2 (ref. ), together with an increase in phospho-eIF4E levels. Notably, previous studies in, for example, cancer and immune cells using cercosporamide observed reduced phosphorylation of eIF4E. This discrepancy could be attributed to differential activities and upstream regulation of the MNKs. MNK2 contributes to a basal level of phospho-eIF4E, while the activity of MNK1 can be potentiated by upstream signaling to increase phospho-eIF4E in the absence of MNK2 (ref. ). Consistently, reduced expression of MNK2 was previously reported to be associated with increased levels of MNK1 (ref. ). Nevertheless, our studies using pan-MNK inhibitors support that the increased phospho-eIF4E following CID661578 depends on hyperactive MNK1. Our interpretation is that β-cell regeneration is increased when MNK2 is inhibited but is unaffected when both MNK1 and MNK2 are inhibited. Therefore, deleting/interfering with the less efficient kinase might open up for the more efficient kinase, leading to a net increase in phospho-eIF4E. Additionally, we observed that

![Fig. 5](CID661578 increases the interaction between eIF4G and eIF4E and leads to translational changes, without affecting the kinase activity of MNK2. a–c. Dose-response of CID661578, CID661578.6 or cercosporamide on MNK2 (a), MNK1 (b) and JAK3 (c) kinase activity in vitro; n=2 for each concentration tested. Data are presented as mean values ±s.e.m. d, Immunoblotting against eIF4G and eIF4E after an m7GTP pulldown assay in lysates of COLO 320HSR cells after 6-h treatment with DMSO, CID661578.6, 4EGI-1 or CID661578.6 together with 4EGI-1. For a loading control, 5% of the input was used. e, Immunoblotting against eIF4G and eIF4E after an m7GTP pulldown assay in rabbit reticulocytes treated with the indicated concentrations of CID661578.6. f, Immunoblotting against eIF4G and eIF4E after an m7GTP pulldown assay in lysates of PANC-1 cells treated with DMSO, CID661578 or CID661578.6 for 6h. For a loading control, 1% of the input was used. g, Immunoblotting against eIF4G and eIF4E after an immunoprecipitation (IP) assay with anti-FLAG in lysates of PANC-1 cells that were treated for 6 h with DMSO or CID661578. For a loading control, 1% of the input was used; I8, immunoblot. h, Immunoblotting against phospho-eIF4E (Ser209; p-eIF4E), total eIF4E and actin in lysates of PANC-1 cells after 6-h treatment with DMSO, CID661578, cercosporamide, CGP57380 or eFT508. i, Quantification of the number of β-cells in 6 d.p.f. zebrafish larvae following β-cell ablation and treatment for 48 h with DMSO, CID661578, eFT508 or a combination of CID661578 and eFT508; n=15 (control), n=14 (CID661578), n=17 (eFT508) and n=15 (CID661578 + eFT508). A one-way ANOVA followed by Dunnett’s multiple comparisons test was used to assess significance for i (**P = 0.0014 (control versus CID661578) and *P = 0.0283 (CID661578 versus CID661578 + eFT508)). Data are presented as mean values ±s.e.m. Experiments in d–h were repeated at least two times. j, Representative polysome tracings from optimized sucrose gradients of PANC-1 cells treated with DMSO, CID661578 or cercosporamide. k, Scatter plots showing log, fold changes for total mRNA (x axis) and polysome-associated mRNA (y axis) for the comparisons of CID661578 (k) and cercosporamide (l) to DMSO. Color codes indicate significantly affected mRNAs identified by anota2seq analysis.)
CID661578 treatment drastically decreases the binding of MNK2 to eIF4G, resulting in increased interaction of eIF4E and eIF4G at the mRNA cap. Further, our polysome profiling analysis uncovered a total of 369 common translationally regulated mRNAs following treatment with CID661578 and cercosporamide. Coupled with the observed increase in phospho-eIF4E, this is one of the most extensive signatures of the effect of the phospho-eIF4E modification on the translatome described to date.

The zebrafish model has emerged as a powerful system for coupling large-scale screens with desired phenotypic outcomes in vivo. The ability to screen in vivo rather than using in vitro culture systems offers the advantage of performing chemical screening in a setting where all of the tissues are present and can interact. Here, we report the identification of Mnk2 as the molecular target of CID661578, the most striking hit from a chemical screen for drivers of β-cell regeneration. Our results identified a previously unknown role for Mnk2 in β-cell neogenesis from cells residing within the ductal cell compartment of the pancreas, thereby paving the way for an alternative path to stimulate β-cell neogenesis, and hence regeneration, for the management of diabetes.
**Fig. 6 | CID661578/cercosporamide treatment increases β-cell differentiation in ductal cells from neonatal pigs and human organoids.**

**a-f.** Images of neonatal pig islets treated with DMSO (a), CID661578 (b) or cercosporamide (c) and stained for insulin (red) and the ductal cell marker CK7 (green). Quantification results showed that treatment with either CID661578 or cercosporamide increased the number of insulin+ β-cells (d), decreased the number of CK7+ duct cells (e) and increased the number of double-positive (insulin+CK7+) cells (f); n=6; **P = 0.0041 and *P = 0.0116 (d); *P = 0.0401 and P = 0.1671 (NS, not significant) (e); **P = 0.0034 and *P = 0.0137 (f). A Kruskal–Wallis test followed by Dunn’s multiple comparisons test was used to assess significance for d-f. Data are presented as mean values ± s.e.m.

**g-i.** Images of human pancreatic sections from different donors stained for MNK2, with insulin used as a marker of β-cells and CK19 used to mark the pancreatic duct. Similar results have been reproduced in stainings from pancreatic sections of multiple human donors.

**j.** Schema showing the procedure for generating and treating human ductal-derived organoids (j). Brightfield images of representative examples of human ductal-derived organoids before differentiation and after treatment with cercosporamide are shown; scale bar, 200 µm. INS mRNA expression is shown in **k** for three different organoid preparations (that is, from three different donors) for cercosporamide and two for CID661578. The experiment was reproducible in at least two different organoid preparations.
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Tg(tp1:H2BmCherry)s939
Tg(ins:flag-NTR)s950
approved Swedish ethical permit for the usage of animals. Previously generated mknk2b, reverse primer 5′ ATTGTTGCT-3′. To eef1a1l1 sequences were cloned into the middle entry vector pDONR221 of the Gateway-AT forward primer 5′-CGACTCACTCAAAACACAGAATC-3′. The AB strain was used for experiments with wild-type zebrafish. The morpholinos targeted against mknk2b, d1 or acinb1 were synthesized by Gene Tools and had the following sequences: M:5′-ATTCACAGACGCATCCTTGC-3′; d1: forward primer 5′-AGCAGTCATCAAAACACAAGATC-3′, reverse primer 5′-ATGTGACTGAAGTGTCGAT-3′; the products of the PCR were amplified by DreamTaq PCR master mix (Thermo Fischer Scientific); the products of the PCR were confirmed in pooled injected embryos after DNA extraction from a pool of 10 embryos using a Quick-RNA Microprep MO (5µL), following sequences: mknk2b′ was annealed with the universal Alt-R′′activating crRNA (tracrRNA) sequence (IDT) by incubating the solution for 5′ to E3 medium to a final concentration of 100µM for 24 h. For m7GTP pulldown experiments, 24 h before treatment, cells were plated with fetal bovine serum. PANC-1 cells were obtained from ATCC and cultured with RPMI 1640

### Chemicals

**Chemical names**: Thermo Fisher Scientific; Bio-Rad; C10-spacer beads purchased from Acros Organics; DMSO, 40µM mercaptopropyl-m7GTP (C10-spacer) beads were purchased from MYBioSource, MBS448092. For experiments in juveniles, the insulin:GFP-β′′area was measured on a flattened projection (average intensity). For the OPP intensity measurements, all larvae were imaged using the same parameters on a confocal microscope. Quantification was performed using the Fiji parameter (mean gray value). The mean gray value was measured from eight tGF P 1+ cells transfected for each line as a reference for each cell, and the average of the mean gray value of the eight cells was calculated. All images were acquired with LAS X (v3.5.5.19976) software. The contrast was adjusted for visualization purposes in some experiments. In the case where the same adjustments were made for all displayed images from the same experiment. The original unmodified pictures were used for analysis.

**Chemical synthesis of C1661578 analogs.** A detailed report that includes the steps used to synthesize all the analogs described in this study is provided in Supplementary Note 2. For the zebrafish Mknk2b binding experiments, full-length or truncated mknk2b (corresponding to the fragment of the human MKNK2 identified in the original screen) fused into the pET28a vector. Two different map pools were used to validate the binding of zebrafish Mknk2b to the 1N-Lexa-eDHFR-C (original hook vector) and 2-N-eDHFR-LexA-C (reverse order hook vector).

**Metabolomics.** Metabolites were extracted from pools of 10 zebrafish larvae at 5 d.p.f. using a methanol-based extraction, and metabolite analysis was performed using liquid chromatography coupled to tandem mass spectrometry. Two different map pools were used to validate the binding of zebrafish Mknk2b to the 1N-Lexa-eDHFR-C (original hook vector) and 2-N-eDHFR-LexA-C (reverse order hook vector).

**Cell culture, immunofluorescence and mGTP pulldown experiments.**

**Immunofluorescence and confocal analysis.** Immunofluorescence staining and confocal analysis of larvae and 1-month-old fish were performed as previously described. Primary antibodies were used against green fluorescent protein (GFP; to amplify the GFP signal, 1:500, Aves Labs, GFP-1020), glucagon (1:200, Sigma, GCL), insulin (1:100; custom antibody produced in Cambridge Research and tdTomato (1:500; MYBioSource, MBS448092). For experiments in juveniles, the insulin:GFP-β′′area was measured on a flattened projection (average intensity). For the OPP intensity measurements, all larvae were imaged using the same parameters on a confocal microscope. Quantification was performed using the Fiji parameter (mean gray value). The mean gray value was measured from eight GFP+ cells transfected for each line as a reference for each cell, and the average of the mean gray value of the eight cells was calculated. All images were acquired with LAS X (v3.5.5.19976) software. The contrast was adjusted for visualization purposes in some experiments. In the case where the same adjustments were made for all displayed images from the same experiment. The original unmodified pictures were used for analysis.

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For the immunoprecipitation experiments, PANC-1 cells were plated in 10-cm dishes and transfected using the DharmaFECT Duo transfection reagent (Active Motif) with 2 μg of the MNK2–FLAG plasmid, obtained from Origene (NM_198570.4). Cells were cultured in serum-free medium (87788), and 1 μg of protein was incubated for the immunoprecipitation with anti-FLAG (Sigma-Aldrich, 6 μg, F3165) overnight. As a negative control for the immunoprecipitation assay, we used the lysine buffer alone incubated overnight with anti-FLAG. The following day, the samples were incubated for 1 h with 30 μl of a 1:100 (in PBS) of the thermo Fisher Scientific, 88803) and washed four times with NET-2 buffer (50 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1 mM MgCl2, 0.5% Nonidet P-40 and one tablet of protease inhibitors), and proteins were eluted from the beads with loading buffer (4X Laemmli sample buffer, Bio-Rad).

For western blotting, after separating proteins on 4–15% gradient gels (Bio-Rad), proteins were transferred to membranes that were then blocked with 5% milk in TBST. Antibodies against the following proteins were used: eIF4E (1:200; Cell Signaling Technologies, 2498), eIF4F (1:2.5; Cell Signaling Technologies, 2948), MNK2 (1:10, Sigma–Aldrich, SAB2101483), β-actin (1:5,000; Sigma-Aldrich, A5441, clone AC-15), phospho-eIF4E (Ser 209; 1:1,000; Cell Signaling Technologies, 9741), anti-rabbit coupled to horseradish peroxidase (1:5,000; Thermo Fischer Scientific, 32040), anti-MNK2 coupled to horseradish peroxidase (1:5,000; Jackson Immunoresearch, AB_2307347).

For the reticulocyte mGTP pulldown assay, untreated rabbit reticulocyte lysates (Promega) were incubated with the specified concentrations of CID661578.6 for 1 h at 30°C followed by mGTP pulldown and western blotting, as described above.

Polysome profiling. To isolate fractions of efficiently translated mRNAs, polysome profiling, using an optimized sucrose gradient, was performed using a recently described optimized sucrose gradient35. Briefly, 4X 10^6 PANC-1 cells were seeded in 15-cm plates 24 h before treatment. Cells were treated with cercosporamide (5 μM), CID661578 (40 μM) or DMSO for 24 h, and polysomes were collected in TRIsol reagent (Thermo Fisher Scientific) and pooled, allowing for isolation of efficiently translated polysome-associated mRNA.

RNA extraction was performed using the TRIzol reagent protocol (Thermo Fisher Scientific) followed by additional purification using an mRNAeasy MinElute Cleanup kit (Qiagen). RNA quality was assessed using a Bioanalyzer 2100 with an RNA 6000 Nano kit (Agilent). Smartseq2 sequencing libraries were prepared as previously described36 using 10 μg of mRNA as input. Libraries were prepared for total cytoplasmic and polysome-associated fractions from four biological replicates of cells treated with cercosporamide, CID661578 or DMSO. Libraries were pooled and sequenced on an Illumina NovaSeq 6000 platform using a 50-base pair paired-end setup.

RNA sequencing read quality was evaluated using MultiQC (1.7). Adapters and reads mapping to ribosomal RNA were removed using BBduk (36.59) from the BBTools suite (http://gigi.doc.nl/data-and-tools/bb-tools/) before alignment to hg38 using HISAT2 (2.1.0) with default settings. Reads were summarized and reads mapping to ribosomal RNA were removed using BBDuk (36.59) from BBTools. Genes were distinguished from each sample for isolation of total cytoplasmic RNA. The remaining lysates were layered onto optimized sucrose density gradients (5% 34% 55% wt/vol) and were centrifuged at 4°C for 35,000 rpm for 2 h followed by fractionation. Fractions containing mRNAs bound to polysomes were collected in TRIsol reagent (Thermo Fisher Scientific) and pooled, allowing for isolation of efficiently translated polysome-associated mRNAs.

Human pancreatic ductal organoid culture. Human pancreatic exocrine tissue, obtained after an organ donation, was minced and organoids were generated using the Ricordi method from cadaveric organ donors with informed written consent, was processed to isolate ductal fragments and generate organoid cultures. Ethical approval for processing pancreatic samples from deceased organ donors was granted by the Clinical Research Ethics Committee of Hospital de Bellvitge (PR030/22). Ductal fragments were embedded in GFR Matrigel and cultured in human organoid expansion medium60. After three to four passages, organoid expansion medium was replaced by a basic medium containing Advanced DMEM/F12, ITS-X (1X), heparin (0.1 mg ml–1), N-acetylcysteine (0.25 mM), FGF10 (0.1 μM) or 1 μM CID661578 (Sigma-Aldrich) or 1 μM cercosporamide (Tocris). The organoids were treated for 3d, and the medium was replaced with identical fresh medium every 48 h. Samples were collected from each condition, and immunohistochemical staining was performed as described previously37. Antibodies against the following proteins were used to stain the pig sections: insulin (1:5, Dako, MR002), CK7 (1:10, Dako, clone OV-TL 12/30) and MNK2 (1:200, Sigma-Aldrich, SAB101483). Human pancreatic sections were stained using the same protocol as described for the porcine sections. All images were acquired with NIS-Elements (version 4.30) software. Human tissues were kindly provided by the Alberta Diabetes Institutes Islet Core, and ethical approval for the use of human samples was obtained from the University of Alberta Human Research Ethics Board, protocol PRO00001416. Informed written consent was provided at the institutions where the organs were collected.

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Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability
The raw metabolomics data were uploaded to Metabolomics Workbench with the study ID ST002119 (https://doi.org/10.21228/M80D9F). The raw polysome profiling data were uploaded to GEO with the accession number GSE200477. Expression of mknk2b in zebrafish was assessed using data downloaded from GEO under the accession number GSE106121 and sample number GSM3032164. The rest of the data are included in the current manuscript. The datasets generated during the current study are available from the corresponding author on reasonable request. Source data are provided with this paper.

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Author contributions
C.K., C.L.M., J.M., and O.A. performed the zebrafish and translation experiments. X.L., C.K. and J.W.L. performed the metabolomics analysis. H.R. and G.S.K. performed the experiments with pig and human samples. K.W and O.L. designed, performed and analyzed the polysome profiling experiments. A.F.R. and M.R. performed the human organoid experiments. C.K. and O.A. designed experiments and wrote the manuscript with help from all authors.

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Competing interests
The authors declare no competing interests.

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Extended Data Fig. 1 | See next page for caption.
Extended Data Fig. 1 | Effect of different analogues of CID661578 on β-cell regeneration. **a**, Chemical structures of the different analogues of CID661578 synthesized for the screen. **b**, Quantification of the number of regenerating β-cells in Tg(ins:kaede);Tg(ins:CFP-NTR) treated with the indicated analogue, tested in three different concentrations from 4-6 dpf after β-cell ablation. The absence of bars denote that the chemical treatment was toxic to the larvae. $n$ = 32 (DMSO), $n$ = 1 (CID661578.8-100 μM), $n$ = 13 (CID661578.8-30 μM), $n$ = 15 (CID661578.8-10 μM), $n$ = 0 (CID661578.8-100 μM), $n$ = 13 (CID661578-100 μM), $n$ = 16 (CID661578-30 μM), $n$ = 16 (CID661578-10 μM), $n$ = 0 (CID661578.1-100 μM), $n$ = 16 (CID661578.1-30 μM), $n$ = 16 (CID661578.1-10 μM), $n$ = 0 (CID661578.2-100 μM), $n$ = 13 (CID661578.2-30 μM), $n$ = 16 (CID661578.2-10 μM), $n$ = 16 (CID661578.3-30 μM), $n$ = 16 (CID661578.3-10 μM), $n$ = 0 (CID661578.4-100 μM), $n$ = 13 (CID661578.4-30 μM), $n$ = 16 (CID661578.4-10 μM), $n$ = 16 (CID661578.5-100 μM), $n$ = 14 (CID661578.5-30 μM), $n$ = 16 (CID661578.5-10 μM), $n$ = 1 (CID661578.6-100 μM), $n$ = 16 (CID661578.6-30 μM), $n$ = 16 (CID661578.6-10 μM), $n$ = 0 (CID661578.7-100 μM), $n$ = 16 (CID661578.7-30 μM), $n$ = 16 (CID661578.7-10 μM), $n$ = 14 (CID661578.8-100 μM), $n$ = 14 (CID661578.8-30 μM), $n$ = 15 (CID661578.8-10 μM), $n$ = 8 (CID661578.9-100 μM), $n$ = 12 (CID661578.9-30 μM) and $n$ = 16 (CID661578.9-10 μM). Data are presented as mean values ±SEM.
Extended Data Fig. 2 | See next page for caption.
Extended Data Fig. 2 | CID661578 does not induce β- or ductal-cell proliferation. a–c, Representative confocal images of Tg(ins:H2BGFP);Tg(ins:flag-NTR) larvae at 6 dpf after β-cell ablation (3-4 dpf) and incubation with DMSO (a) or 5µM CID661578 (b) during the regenerative period (4-6 dpf). β-cells that proliferated are marked by EdU and displayed as yellow overlap (arrowheads). The number of β-cells that proliferated was not altered by CID661578 (c). Scale bar, 10µm. n=13 (Control) and n=15 (CID661578). Unpaired two-tailed Student’s t test was used to assess significance for (c) ns=not significant (P=0.8340). Data are presented as mean values ±SEM. The experiment shown in (a–b) has been repeated two times with similar results. d–g, CID661578 treatment does not alter the number or the proliferation of notch-responsive ductal cells. Confocal images of DMSO- (d) or CID661578-treated (e) Tg(tp1:GFP);Tg(ins:flag-NTR) larvae at 6 dpf after β-cell ablation and incubation with EdU. White dashed lines outline the pancreata of the larvae. Scale bar, 10µm. Quantification of the number of notch-responsive cells (f) and the number of proliferating, EdU+ notch-responsive ductal cells (g) per larva is shown. n=11 (Control) and n=12 (CID661578). Data are presented as mean values ±SEM. h, Glucose levels of Tg(ins:flag-NTR) larvae treated with DMSO, cercosporamide or CID661578 during the regenerative period (4-6 dpf). Four larvae were pooled for each replicate. n=4 (all treatments – 3 dpf), n=5 (all treatments – 4 dpf), n=5 (all treatments – 5 dpf), n=4 (Control and Cercosporamide – 6 dpf) and n=4 (CID661578 – 6 dpf). 2way ANOVA test followed by Tukey’s multiple comparisons test was used to assess significance for *P=0.0430 (5 dpf Control vs Cercosporamide), ****P<0.0001 (5 dpf Control vs CID661578), ****P<0.0001 (6 dpf Control vs CID661578) and ****P<0.0001 (6 dpf Control vs Cercosporamide). Data are presented as mean values ±SEM.
Extended Data Fig. 3 | Validation of the notch-responsive cells as the source of the newly formed β-cells in juvenile fish. a-e, CID661578.6 promotes β-cell neogenesis in juvenile fish. Single-plane confocal images of Tg(ins:GFP);Tg(tp1:H2BmCherry);Tg(ins:flag-NTR) pancreata from 1-month-old zebrafish treated with DMSO (a) or CID661578.6 (b). Briefly, 1-month-old zebrafish were incubated with MTZ (1 mM) for 24 hours to ablate β-cells followed by chemical treatment for 2 days. Pancreata are outlined with white dashed lines. Quantification of β-cells in the secondary islets in the tail of the pancreas (c) shows an increase in β-cell regeneration, and quantification of the overlap between the notch-responsive cell tracer and β-cell marker demonstrated an increase in the number of β-cells derived from notch-responsive cells (d). CID661578.6 treatment also doubled the ins^+ area normalized to the body length of the fish (μm²/mm) (e). Scale bar, 50 μm. (c and d), n = 11 (Control) and n = 13 (CID661578.6); (e), n = 10 (Control) and n = 13 (CID661578.6). Unpaired two-tailed Student’s t test was used to assess significance for (c) *P = 0.0229; two-tailed Mann-Whitney test was used for (d) ****P < 0.0001; two-tailed Mann-Whitney test was used for (e) **P = 0.0099. Data are presented as mean values ± SEM.
Extended Data Fig. 4 | Newly-derived β-cells express mnx1, another marker for β-cell identity/maturation, and are devoid of glucagon expression. a-d. Single-plane confocal images of 6 dpf Tg(mnx1:GFP);Tg(ins:flag-NTR) larvae treated with DMSO (a) or CID661578 (b) (4-6 dpf) after β-cell ablation (3-4 dpf) and staining for insulin. Quantification results showed that CID661578 treatment increased the number of β-cells (c), but there was no difference in the percentage of mnx1::GFP⁺, insulin⁺ β-cells (d). Scale bar, 10 μm. For (c) and (d) n = 10 (Control) and n = 12 (CID661578). Unpaired two-tailed Student’s t test was used to assess significance for (c) *P = 0.0497. Data are presented as mean values ±SEM. e-g. Representative confocal images of 6 dpf Tg(ins:H2BGFP); Tg(tp1:H2BmCherry);Tg(ins:flag-NTR) larvae treated with DMSO (e) or CID661578.6 (f) (4-6 dpf) after β-cell ablation (3-4 dpf) and stained for glucagon. White arrowheads point to bihormonal cells (glucagon⁺, ins:H2BGFP⁺) derived from notch-responsive cells. Quantification of bihormonal cells derived from notch-responsive cells (g) showed no difference between treatments. Scale bar, 10 μm. n = 9 (Control) and n = 7 (CID661578.6). Data are presented as mean values ±SEM.
Extended Data Fig. 5 | Gene expression of the primary pancreatic lineage genes in the single-cell RNA-Seq dataset. **a–f**, UMAPs showing the expression of amy2a (acinar cells) (**a**), cftr (ductal cells) (**b**), ins (β-cells) (**c**), gcga (α-cells) (**d**), sst2 (δ-cells) (**e**) and sst1.1 (**f**) in the dataset used to analyze mknk2b expression in adult zebrafish pancreata.
Extended Data Fig. 6 | CID661578-treated zebrafish in the basal state exhibit no changes in endocrine cell numbers. a–c, Maximum projections confocal images of primary islets in Tg(ins:H2BGFP) larvae at 6 dpf after treatment with (a) DMSO, (b) CID661578 or (c) cercosporamide from 3–6 dpf. Quantification (g) of the total number of β-cells revealed no changes resulting from the treatments. Scale bar, 10 µm. n = 12 (Control), n = 11 (CID661578) and n = 11 (Cercosporamide). Data are presented as mean values ± SEM. d–f, Single-plane confocal images of primary islets in Tg(gcga:GFP);Tg(sst2:dsRed2) larvae at 6 dpf after treatment with (d) DMSO, (e) CID661578 or (f) cercosporamide from 3–6 dpf. Quantification of the total number of α-cells (h) and δ-cells (i) revealed no changes resulting from the treatments. Scale bar, 10 µm. For both (h and i), n = 14 (Control), n = 9 (CID661578) and n = 9 (Cercosporamide). Data are presented as mean values ± SEM. j, Glucose measurements in zebrafish larvae in the basal state treated with DMSO, CID661578 or cercosporamide for 1 day (3–4 dpf). Four larvae were pooled for each replicate. n = 5 (Control), n = 4 (CID661578) and n = 5 (Cercosporamide). One-way ANOVA test followed by Dunnett’s multiple comparisons test was used to assess significance for (j) ****P < 0.0001 (Control vs CID661578); *P = 0.0182 (Control vs Cercosporamide).
Extended Data Fig. 7 | See next page for caption.
Extended Data Fig. 7 | Morpholino-mediated knockdown of the A-hits from the yeast screen show that mknk2b, but not acin1b and drl, affects β-cell regeneration. a–e, Tg(ins:H2BGFP);Tg(ins:flag-NTR) embryos were injected at the 1-cell stage with a control morpholino (a,c) or a morpholino against mknk2b (b,d), β-cells ablated between 2-3 dpf, and treated with DMSO (a,b) or 2μM CID661578.6 (c,d) for 2 days. Quantification of regenerated β-cells (e) revealed an increase upon mknk2b knockdown, similar to the chemical inhibition of Mnk2. No additive effect was observed with the combined mknk2b MO and CID661578.6 treatment. Scale bar, 10μm. n = 24 (Control), n = 26 (mknk2b MO), n = 24 (CID661578.6) and n = 23 (mknk2b MO + CID661578.6). *P = 0.0210 (control vs mknk2b MO) and 0.0459 (mknk2b MO vs CID661578.6), ****P < 0.0001. Data in this graph are pooled from two independent experiments and are presented as mean values ±SEM.

f–j, Confocal images of primary islets of Tg(ins:H2BGFP);Tg(ins:flag-NTR) at 5 dpf, after injection at the one-cell stage with control (f), acin1b (g), drl (h) or mknk2b (i) morpholinos. Following the ablation of β-cells, only mknk2b knockdown significantly increased β-cell regeneration (j). Scale bar, 10μm. n = 12 (Control MO), n = 14 (acin1b MO), n = 11 (drl MO) and n = 9 (mknk2b MO). (j) *P = 0.0410. Data are presented as mean values ±SEM.

k–m, Agarose gel images of RT-qPCR validating the knockdown of mknk2b (k), drl (l) and acin1b (m) at two stages (2 dpf and 5 dpf). Amplification of eef1a1l1 was used as control. Arrows point to the band of the expected size. The experiment shown in (k–m) has been repeated twice with similar results. n–r, Single-plane confocal images of Tg(ins:H2BGFP);Tg(ins:flag-NTR) islets from control (n and o) or larvae overexpressing mknk2b in notch-responsive cells (p and q), treated with DMSO (n and p) or CID661578.6 (o and q). Quantification results (r) showed that the stable Tg(tp1:mknk2b) overexpression blocked the effect of CID661578.6 on β-cell regeneration, similar to that observed in the transient overexpression model (see Fig. 3). Scale bar, 10μm. n = 13 (Control), n = 12 (CID661578.6), n = 14 (Tg(tp1:mknk2b)) and n = 11 (Tg(tp1:mknk2b)+CID661578.6). (r) ***P = 0.0001 and *P = 0.0172. Data are presented as mean values ±SEM.
Extended Data Fig. 8 | Enrichment analysis for the upregulated metabolites and differentially expressed genes in the single-cell RNA-Seq data.

a, Pathway analysis, assessing 81 characterized metabolic pathways in zebrafish, using the significantly upregulated metabolites. Only pyrimidine metabolism was significantly enriched in this dataset (FDR < 0.05).

b, Enrichment analysis for KEGG pathways in differentially expressed genes, that is, genes upregulated in ductal cells (‘low-notch’ cluster, see Fig. 2g) compared to all endocrine cells.
Extended Data Fig. 9 | CID661578.6 does not potently affect the activity of any kinase in an in vitro screen; and MNK2 expression in the pig pancreas.

**a, b.** An in vitro screen to assess the specificity of 1 µM CID661578.6 and 1 µM cercosporamide against a panel of 140 kinases. CID661578.6 did not alter the activity of any kinase by more than 50% (**a**), while cercosporamide inhibited the activity of several kinases, including MNK1 and MNK2 (**b**). Red arrows point to MNK1 and MNK2.

**c-e.** Images of neonatal (**c, d**) and adult (**e**) pig pancreatic sections immunostained for MNK2, with insulin used as a marker of β-cells and CK7 as a marker of ductal cells. Scale bar, 50 µm. Similar staining results were obtained in sections from multiple pig pancreata.
Extended Data Fig. 10 | See next page for caption.
Extended Data Fig. 10 | Polysome profiling expands on the effects and similarities between CID661578 and cercosporamide treatments on PANC-1 cells. a, Scatterplot indicating that genes translationally regulated by CID661578 are similarly regulated by treatment with cercosporamide. b,c, Empirical cumulative distributions of log2 fold changes for polysome-associated (translated) and total mRNA when comparing cercosporamide and DMSO treatments. Transcripts whose translation was suppressed (dark red) or activated (light red) by CID661578 differed from background (grey) and showed a similar directionality following cercosporamide treatment. P-values (Wilcoxon test) comparing each set to background mRNAs and shifts (log2) relative to background transcripts at multiple quartiles are also shown. d, Significantly enriched GO terms among proteins encoded by mRNAs that were hypotranslated upon both CID661578 and cercosporamide treatments relative to DMSO. e,f, Violin plots showing the 5'UTR GC percentage among transcripts translationally activated or suppressed following CID661578 (e) or cercosporamide (f) treatment. The background (grey; that is mRNAs not in regulated subsets) is also shown together with P-values from two-sided Wilcoxon Rank Sum test for the indicated comparisons. For (e): (background) n = 9853, minima:16.7; maxima:100; centre:63.6; upper bound:95.4; lower bound:32.25th per:55.7; 75th per:71.6; (translation up) n = 319, minima:31.1; maxima:85.9; centre:56.5; upper bound:85.9; lower bound:31.1; 25th per:47.7; 75th per:66.8; (translation down) n = 127, minima:32.5; maxima:88.4; centre:66.7; upper bound:88.4; lower bound:33.9; 25th per:59.8; 75th per:77. For (f): (background) n = 9853, minima:16.7; maxima:100; centre:63.6; upper bound:95.4; lower bound:32.25th per:55.7; 75th per:71.6; (translation up) n = 1850, minima:25; maxima:86.2; centre:58.6; upper bound:86.3; lower bound:25.3; 25th per:50; 75th per:66.5; (translation down) n = 1468, minima:35.2; maxima:93.7; centre:67.6; upper bound:93.7; lower bound:35.8; 25th per:59.6; 75th per:75.4.
Reporting Summary

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Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

☐ n/a Confirmed
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☐ A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
☐ The statistical test(s) used AND whether they are one- or two-sided
☐ Only common tests should be described solely by name; describe more complex techniques in the Methods section.
☐ A description of all covariates tested
☐ A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
☐ A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) and variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
☐ For null hypothesis testing, the test statistic (e.g. t, F, r) with confidence intervals, effect sizes, degrees of freedom and P value noted
☐ Give P values as exact values whenever suitable.
☐ For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
☐ For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
☐ Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

LAS X v3.5.5.19976 (for confocal microscopy), NIS-Elements 4.30 (for qPCR data collection), QuantStudio Software V1.2.4 (for qPCR data collection and analysis), Abbott glucometer (for adult zebrafish glucose measurements).

Data analysis

Fiji/ImageJ (for image analysis - version 2.0.0-r-c165), Excel (version 16.16.27), GraphPad PRISM 8.0b9.0, RStudio 1.4.1717, R package Seurat 3.5.1, R clusterProfiler package (3.10.1), QuantStudio Software V1.2.4 (for qPCR data analysis), MultiQC (version 1.7 - polsencode quality check), BBDuk from the BBTools suite (version 36.59 - remove ribosomal RNA reads of the polsencode), HISAT2 (version 2.1 - genome alignment of polsencode data), RSSubset (version 2.6.4 - Summarize reads of the polsencode data), R package anota2seq (version 1.14.0 - analysis of polsencode data), ClueGO plug-in (version 2.5.8) within CytoScape (version 3.8.2 - gene ontology analysis), Siene 2.2 was used for chromatographic alignment and peak integration. Morpheus tool from Broad Institute and Metaboanalyst 4.0 was used to analyze the metabolomics data.

For manuscripts utilizing custom algorithms or software that are not central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.
Data

Policy information about availability of data
All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

The raw data of the metabolomics study is uploaded to Metabolomics Workbench with the study ID ST002119 [http://dx.doi.org/10.21228/M80D9F]. The raw data of the polysome profiling is uploaded to GEO with the accession number GSE200047. Expression of mink2b in zebrafish was assessed using data downloaded from GEO under the accession number GSE106121 and sample number GSM303164. The rest of the data are included in the current manuscript. The datasets generated during the current study are available from the corresponding author on reasonable request. Source data are provided with this paper.

Field-specific reporting

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size
No predetermination of sample size was made. We considered a sufficient final sample size when statistical value for group comparison was reached.
All in vivo biological experiments in the present manuscript have been repeated at least two times. Each biological replicate included more than seven, embryo or adult, zebrafish. Experiments with pig islets were performed at least two times.
All in vitro experiments containing cell lines were repeated at least two times. The knockdown was performed once. During the course of the study the YCHE14 experiments were repeated multiple times. The polysome-seq data were performed with biological replicates, the human ductal endodermoids experiments were replicated with tissue from 2 human donors (CID661578 treatment) and 3 human donors (cyclosporamide treatment).

Data exclusions
No data were excluded from the final analysis.

Replication
Experiments have been repeated at least two times, with the majority of them having more than three biological replicates. All experiments are reproducible.

Randomization
Zebrafish embryos, pig islet preparations and human duct-derived organoids were randomly assigned to the respective treatment groups, prior to treatment. Adult zebrafish used for the experiments were randomly assigned to treatment groups.
Regarding cell culture experiments, cells were randomly assigned to treatment groups after plating.

Blinding
Investigators were blinded during the identification of the activity of the analogue CID661578.6 (Supplementary figure 1) and for all the experiments with the mink2b mutant zebrafish line.
In all other experiments investigators were not blinded. When the experimental design required allocation to specific groups or treatments no blinding was required. When different treatment groups were involved the data collection was unbiased and the different groups were blinded for the analysis of these datasets.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|
| ☑   | Antibodies            |
| ☑   | Eukaryotic cell lines |
| ☑   | Palaeontology and archaeology |
| ☑   | Animals and other organisms |
| ☑   | Human research participants |
| ☑   | Clinical data |
| ☑   | Dual use research of concern |

Methods

| n/a | Involved in the study |
|-----|-----------------------|
| ☑   | ChIP-seq              |
| ☑   | Flow cytometry        |
| ☑   | MRI-based neuroimaging |
**Antibodies**

**Antibodies used**

For the zebrafish work: #1 - chicken anti-GFP (Aves Labs - GFP-1020, 1:500), #2 - mouse anti-glucagon (1:200; Sigma G62554), #3 - rabbit anti-insulin (1:100; custom made by Cambridge Research Biochemicals), #4 goat anti-tdT (1:500; MyBioSource MBS448092).

For pig islet and human sections stainings: #1 - guinea-pig anti-insulin (1:5; DAKO, code# R002), #2 - mouse anti-CX7 (3:100; DAKO, clone OV7L 12/30), #3 - rabbit anti-MNK2 (1:200, Sigma-Aldrich, SAB2101483), #4 - mouse anti-CX19 (1:50, M08880-1-2, Clone RCK108, Agilent).

For the cell lines and in vitro reticulocyte western blot experiments: #1 - rabbit anti-EIF4G (1:2000; Cell Signaling Technologies Z498), #2 rabbit anti-EF4E (1:2000; Cell Signaling Technologies 9742), #3 rabbit anti-MNK2 (Sigma-Aldrich SAB2101483, 1:1000), #4 mouse anti-b-ACTIN (1:5000; Sigma-Aldrich A5441 clone AC-15), #5 anti-rabbit coupled to HRP (1:5000; Thermo Fischer Scientific – 31460), #6 anti-mouse coupled to HRP (1:5000; Jackson ImmunoResearch AB_2307347) #7 mouse anti-FLAG antibody (1:2000, Sigma-Aldrich, F3165), #8 phospho-eIF4E (SER209) (1:1000; Cell Signaling Technologies 9741).

**Validation**

For the zebrafish stainings: antibody #1 has been cited 1055 times in the literature (e.g. https://doi.org/10.15252/embb.2019192903), antibody #2 antibody has been cited 264 times in the literature (e.g. zebrafish example: doi:10.2337/db16-1587) #3 has been validated using transgenic lines fluorescently labelling the mature beta-cells in zebrafish (e.g. see here for validation:DOI: https://doi.org/10.7554/eLife.65758) and antibody #4 has been (e.g. DOI: https://doi.org/10.7554/eLife.65758).

For antibodies used in the pig aggregates and human section stainings, the antibodies were used in previous studies [Hassouna et al; Transplantation; 2018].

All the antibodies used for the western blot experiments have been widely used and cited in numerous previous studies. #1EIF4G antibody has been validated for western blot with genetic knockdown in mammalian cells (e.g. see here as an example https://doi.org/10.3390/jms20071580). #2 EIF4E antibody has been validated numerous times for western blot with genetic knockdowns in human cell lines in published studies (e.g. see here doi: 10.1158/1535-7163.MCT-17-2357). #4 actin antibody has been cited more than 8000 times for western blot applications in mammalian systems (e.g. recently here DOI: 10.7554/eLife.74650). #7 anti-FLAG antibody has been cited numerous times for immunoprecipitation experiments (e.g. https://doi.org/10.1038/s41467-017-02861-2) and we also validated in Figure 5g where we can see enrichment of the flag tagged protein after immunoprecipitation compared to the input control. #8 p-EF4E antibody for western blot has been validated by genetic manipulation of MNK kinase (e.g. see here: DOI: 10.1172/jcrg.202466). The #3 MNK2 antibody was validated by overexpression of MKN2 protein in COLO320 followed by western blot as part of this study.

**Eukaryotic cell lines**

**Policy information about: cell lines**

**Cell line source[s]**

COLO 320HSR and PANC-1 cells were purchased from ATCC.

**Authentication**

Cells were verified by ATCC using STR profiling and the morphology of the cells was monitored for any changes during experiments.

**Mycoplasma contamination**

Cell cultures were checked for mycoplasma contamination frequently and no mycoplasma infection was identified during the experiments.

**Commonly misidentified lines (See [ICCLAC register])**

None of the cell lines used are listed in the ICCLAC database.

**Animals and other organisms**

**Policy information about: studies involving animals / ARRIVE guidelines**

**Recommended for reporting animal research**

**Laboratory animals**

For zebrafish experiments either the AB or the TL background strain was used. The transgenic lines used were: Tg(insl:flag-NTR), Tg(insl:GFP), Tg(tpl:H2B::mCherry), Tg(tpl:GFP), Tg(slp:GFP), Tg(insl:CFP::NTR), Tg(insl:Kaede), Tg(insl:2BFP), Tg(slp:Cre:ERT2), Tg(3:5ubibloxp::FGFP-loxp::mCherry), Tg(tpl:19sa MKN2), Tg(tpl:mnkn2b) and mnkn2b-4bp deletion mutant line.

Three ages of zebrafish were used for the experiments: Larvae (up to 6 dpf) - juvenile (1 month old) – adult (4 months old) Sex is not possible to be determined at the larvae and juvenile stage. For the adult experiments the majority of the zebrafish were males. This was done due to availability of males/females in these clutches and not due to a consciously taken decision to exclude more males.

Wild type 3 days old neonatal pigs of either sex were sacrificed to obtain the islet cultures for experiments.

**Wild animals**

The study did not involve wild animals.

**Field-collected samples**

The study did not involve samples collected from the field.

**Ethics oversight**

Work with zebrafish was approved and performed following the guidelines of Stockholms djurförsöksnämnd under an approved ethical permit.

Work with neonatal pig islets was performed under the guidelines from Canadian Council on Animal Care.
Human research participants

Policy information about studies involving human research participants

Population characteristics
General information of the organ donors used for the immunostainings in Fig. g-i: Donor 1: Female, Age: 33, Height: 1.61[m], Weight: 70[kg], BMI: 27.3, Cold Ischemia Time (h): 12. Donor 2: Female, Age: 60, Height: 1.7[m], Weight: 75.1[kg], BMI: 25.9, Cold Ischemia Time (h): 13.5. Donor 3: Male, Age: 47, Height: 1.61[m], Weight: 106.9[kg], BMI: 41.2, Cold Ischemia Time (h): 15.3. No information about chronic medication was available for these donors.

For the human donors whose pancreas was used to generate the ductal organoids in Fig 6 j-k: Donor 1: Male, Age: 45, BMI: 27.70, Cause of death: Stroke. Donor 2: Male, Age: 48, BMI: 24.77, Cause of death: Anoxia after cardiac arrest. Donor 3: Male, Age: 58, BMI: 26.23, Cause of death: Anoxia after cardiac arrest. No information on chronic medication was available for donors 1 & 2. Donor 3 had asthma and was medicated with a bronchodilator when needed, but no other chronic medication was reported.

Recruitment
Human tissues were kindly provided by the organ donors to the Alberta Diabetes Institutes Islet Core and the Hospital de Bellvitge respectively. No preselection of donor/tissue material was done and the selection was based on availability of the human cadaveric material at the time of both studies.

Ethics oversight
1. Ethical approval for the use of human samples for staining was obtained from the University of Alberta’s Human Research Ethics Board protocol PRO00001416.

2. Ethical approval for processing pancreatic samples from unidentified organ donors was granted by the Clinical Research Ethics Committee of Hospital de Bellvitge (PR030/22) for the generation of the human ductal-derived organoids. Informed consent was obtained for use of the cadaveric material for research purposes.

Note that full information on the approval of the study protocol must also be provided in the manuscript.