1	Identification of a small molecule that stimulates human $\beta$ -cell proliferation
2	and insulin secretion, and protects against cytotoxic stress in rat insulinoma
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#### 35 Abstract

36 A key event in the development of both major forms of diabetes is the loss of functional 37 pancreatic islet  $\beta$ -cell mass. Strategies aimed at enhancing  $\beta$ -cell regeneration have long been pursued, but methods for reliably inducing human  $\beta$ -cell proliferation with full retention 38 39 of key functions such as glucose-stimulated insulin secretion (GSIS) are still very limited. 40 We have previously reported that overexpression of the homeobox transcription factor 41 Nkx6.1 stimulates β-cell proliferation, while also enhancing GSIS and providing protection 42 against  $\beta$ -cell cytotoxicity through induction of the VGF prohormone. We developed an Nkx6.1 pathway screen by stably transfecting 832/13 rat insulinoma cells with a VGF 43 44 promoter-luciferase reporter construct, using the resultant cell line to screen a 630,000 compound chemical library. We isolated three compounds with consistent effects to 45 46 stimulate human islet cell proliferation. Further studies of the most potent of these 47 compounds, GNF-9228, revealed that it selectively activates human  $\beta$ -cell relative to  $\alpha$ -cell 48 proliferation and has no effect on  $\delta$ -cell replication. In addition, pre-treatment, but not short 49 term exposure of human islets to GNF-9228 enhances GSIS. GNF-9228 also protects 50 832/13 insulinoma cells against ER stress- and inflammatory cytokine-induced cytotoxicity. 51 In contrast to recently emergent Dyrk1a inhibitors that stimulate human islet cell 52 proliferation, GNF-9228 does not activate NFAT translocation. These studies have led to 53 identification of a small molecule with pleiotropic positive effects on islet biology, including 54 stimulation of human  $\beta$ -cell proliferation and insulin secretion, and protection against 55 multiple agents of cytotoxic stress.

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#### 57 Introduction

58 Both major forms of diabetes involve loss of functional  $\beta$ -cell mass, but methods for 59 reliably inducing human  $\beta$ -cell proliferation with full retention of key functions such as glucose-60 stimulated insulin secretion (GSIS) are still very limited. Early approaches focused on 61 manipulation of oncogenes or core elements of the cell replication machinery such as cyclin 62 D1, cdk6, p16 or p57 [1-3], but such strategies either led to functional impairment or were not 63 therapeutically tractable due to the common use of these factors for cell cycle control in all 64 tissues in the body. More recently, several groups reported that inhibition of the tyrosine kinase Dyrk1A with harmine or other small molecules stimulates human  $\beta$ -cell replication via 65 activation of the NFAT/calcineurin pathway [4-6]. However, the wide tissue distribution of 66 67 Dyrk1a suggests that inhibitors could trigger promiscuous cellular proliferation, and indeed,

systemic administration of harmine activates islet  $\alpha$ -,  $\delta$ -, and ductal cell proliferation in addition to its effects on  $\beta$ -cells [5]. Therefore, additional strategies for expansion of functional human  $\beta$ -cell mass are needed, including mechanisms that are orthogonal to the Dyrk1A/NFAT pathway.

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73 Our laboratory has performed a series of studies on overexpression of the homeobox transcription factor Nkx6.1 in pancreatic islets [7-12]. 74 Key findings include: 1) 75 Overexpression of Nkx6.1 in rat islets results in increased  $\beta$ -cell replication accompanied by improved GSIS [7,8]; 2) Nkx6.1 induces expression of a prohormone, VGF, in both rat and 76 77 human islets, which is processed to yield multiple peptides, including TLQP-21 [9]. TLQP-21 78 protects  $\beta$ -cells from apoptotic cell death, and VGF plays an important role in insulin vesicle 79 trafficking [9, 12]. However, VGF or its peptides do not activate  $\beta$ -cell proliferation [9]; 3) 80 Nkx6.1 activates  $\beta$ -cell replication by a pathway that is additive to and distinct from the 81 pathway activated by overexpression of Pdx-1 [10,11]. Importantly, overexpression of Nkx6.1 82 under control of the constitutive CMV promoter (causing expression in all islet cells) 83 selectively stimulates proliferation of  $\beta$ -cells [8, 11], whereas overexpression of Pdx-1 via the 84 same vector stimulates proliferation of both  $\alpha$ - and  $\beta$ -cells via a secreted factor or factors [11, 85 13]. These results support the idea that Nkx6.1 activates a replication pathway with intrinsic  $\beta$ -86 cell specificity.

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Based on these findings, we stably transfected the INS-1-derived 832/13 rat insulinoma cell line [14] with a plasmid containing the VGF promoter driving expression of luciferase, and used the resultant cell line to screen a 630,000 compound small molecule library. Given the positive impact of Nkx6.1 and VGF in islet cells [7-12], the screen was designed with a VGFluc construct to capture small molecules that activate Nkx6.1 expression, stabilize or activate

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93 Nkx6.1 function, or serve as direct inducers of VGF. We ultimately isolated three compounds 94 with robust and consistent effects to stimulate human islet cell proliferation. Further studies of 95 the most potent of these compounds, GNF-9228, revealed that it selectively activates human 96  $\beta$ -cell relative to  $\alpha$ - or  $\delta$ -cell proliferation. Moreover, pre-treatment of human islets with GNF-97 9228 enhances insulin secretion. GNF-9228 also protects a rat  $\beta$ -cell line, INS-1 832/13, 98 against cytotoxicity. Finally, GNF-9228 does not activate NFAT translocation, suggesting a 99 mechanism of action distinct from Dyrk1a inhibitors.

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#### 102 Materials and methods

103 Cell culture and reagents. INS-1-derived 832/13 rat insulinoma cells were cultured as 104 previously described [14]. Human islets were obtained from the Integrated Islet Distribution 105 Program (IIDP) or from the University of Alberta Human Islet Core. Viability of human islets 106 was tested by staining with propidium iodide as described in the IIDP SOP (http://idp.coh.org), 107 and were used only if viability was >75%. Human islets were cultured in RPMI-1640 108 (Invitrogen 11879) supplemented with 5.5 mmol/L glucose, 10% FCS (Sigma), 100 U/ml 109 penicillin, 100 µg/ml streptomycin and 250 ng/ml amphotericin B (Gibco #15240). An 110 adenovirus vector containing the cDNA for NFATc1 was obtained from Welgen, and stock 111 solutions were diluted 1:8000 when added to Ins1 cells in 384 well plates.

Generation of the 832/13-VGF-Luc Cell Line. Primers GAA GAT CTA TGG TCG AGGGCT GGC G and GGG GTA CCT GAC CCC CCT TCT CAG C corresponding to +77 to -2022 of the rat *VGF* promoter were used to amplify rat genomic DNA. The amplified PCR product was digested with BgIII and KpnI for ligation upstream of a luciferase reporter gene that is contained in vector pGL4.21 (Promega). 832/13 cells were transfected with the resultant VGF promoter-luciferase reporter plasmid by electroporation and selected with puromycin [14] to yield stably transfected 832/13-VGF-Luc cells.

**Gene expression analyses**. RT-PCR was used to measure the following transcripts, using the indicated Taqman probes: Rat Nkx6.1 : Rn01450076\_m1, human NKX6.1: Hs00232355\_m1, human VGF: Hs00705044\_s1, human MYC: Hs00153408\_m1. Gene expression was normalized against expression of cyclophilin A: human PPIA: Hs04194521 s1, rat Ppia: Rn00690933 m1.

Use of 832/13-VGF-Luc cell line for small molecule high-throughput screen. High
 throughput screening was performed using 832/13-VGF-Luc cells. Briefly, 2,000 cells per well
 were plated in 5 μL RPMI in 1536-well plates. Individual molecules from a 630,000 compound

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127 chemical library assembled at the Genomics Institute of the Novartis Foundation (GNF) were 128 added to single wells at a final concentration of 10 µM and incubated for 6 h. 2 µL of Bright 129 Glo (Promega) was added to each well and luciferase-generated luminescence was 130 measured with a plate reader (GNF Systems). Data were normalized to assay plate median 131 and hits were called based on a robust deviation of 3. Hits in the primary screen were 132 counterscreened against Ins1 cells stably transfected with a cyclic AMP response element 133 (CRE)-luciferase reporter or HEK293T cells containing a PLTP-DR4B luciferase construct 134 (pGL4.24 luciferase vector) [15]. The three compounds with human islet proliferative activity, 135 GNF-9228, GNF-4088, GNF-1346 and the Dyrk1a inhibitor GNF-4877 [4] were dissolved in DMSO. 136

Islet cell proliferation measured by EdU incorporation. For EdU labeling, a 1:1,000 dilution of EdU-labeling reagent (Invitrogen) was added to islet culture medium during the last 18h (Figure 3) or 72h (Figure 7) of cell culture. Islets were harvested and processed for immunofluorescence analyses as described previously [13]. Images were captured and analyzed with the Cellomics CX5 High Content (HC) cell based imaging system (Thermo) as described [13].

Glucose-Stimulated Insulin Secretion. Human islets were pretreated with 10 µM GNF-9228 or DMSO for 72h. Insulin secretion was measured by static incubation of 30 islets in 4 replicate groups as previously described [9]. Secreted insulin was measured by ELISA (80-INSMR-CH10, ALPCO) and normalized to insulin content [9]. To measure the acute effects of GNF-9228, some batches of human islets were treated with GNF-9228 during the secretion assay only (no pre-treatment).

Viability and cell toxicity assays. 832/13 cells were seeded in 96-well tissue culture dishes.
To induce cell toxicity, cells were treated with 200-500 nM thapsigargin or with a mixture of 1

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151 ng/ml rat IL-1 $\beta$  (Sigma) + 100 U/ml rat IFN- $\gamma$  (Thermo). Cell viability was measured using the 152 CellTiter96 assay (Promega), or for apoptosis by measurement of caspase-3 cleavage [16].

153 **Pharmacokinetics of GNF-9228 in mice.** The following studies were approved by the Duke 154 University Institutional Animal Care and Use Committee (IACUC). We delivered 30 mg/kg of 155 GNF-9228, GNF-1346, or GNF-4088 suspended in DMSO by intraperitoneal injection into 156 normal C57BL6 mice and measured levels of the compounds in the blood by mass 157 spectrometry over the following 24 hours. Six C57BL6 mice also received daily injections of 30 158 mg/kg GNF-9228 for one week, while 6 control mice received injections of DMSO. For all 159 mice. BrdU was added to the drinking water at a concentration of 0.8 mg/ml. 24 h after the 160 final GNF-9228 or DMSO injection, the mice were euthanized and pancreata were dissected, 161 fixed in neutral-buffered formalin, and paraffin embedded. Slides were incubated overnight 162 with guinea pig anti-insulin (Dako) and mouse anti-BrdU (Dako) antibodies, followed by 163 detection with an AlexaFluor 488 conjugated goat anti-guinea pig and AlexaFluor 555 164 conjugated goat anti-mouse secondary antibody (Invitrogen), and counterstaining with DAPI. 165 Images were captured and analyzed using OpenLab software. A minimum of 4 slides per 166 pancreas spaced 75-100 µm apart were analyzed, comprising a total of approximately 10,000 167 cells per condition.

168 Statistical Analyses.

Data are presented as means + standard errors of mean (SEM) and graphed using GraphPad
Prism 8.1 software. The differences between groups were compared using a Student's t-test
and values of P < 0.05 were considered significant.</li>

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#### 176 **Results**

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178 Cell-based screen for inducers of Nkx6.1 and VGF. We engineered a rat insulinoma (INS-1)-derived cell line, 832/13 [14], by stable transfection with a rat VGF promoter-luc reporter to 179 180 generate a new cell line, 832/13-VGF-luc. 832/13-VGF-luc cells were screened with the 181 630,000 compound small molecule library assembled at the Genomics Institute of the Novartis 182 Research Foundation (GNF). A total of 4437 compounds caused significant (standard 183 deviation of three from the mean) stimulation of luciferase expression in the screen. These 184 hits were counterscreened in HEK-293 cells stably transfected with an unrelated promoter-185 luciferase construct (PLTP-luc), or Ins1 cells transfected with a CREB-luciferase construct. 186 The counterscreening protocol yielded 41 compounds with activity profiles similar to those shown for compounds GNF-9228 and GNF-7169, featuring robust, dose-dependent activation 187 188 of the VGF-luc construct, with no activation of PLTP-luc (Fig 1). Since the VGF promoter 189 contains a consensus cyclic AMP response element-binding protein (CREB) binding 190 sequence [17], compounds were also counterscreened against a CREB-luc reporter. 191 Compound GNF-7169 had no activity against the CREB-luc construct, whereas GNF-9228 192 had a weak activating effect (Fig 1).

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Fig 1. Representative "Hits" from small molecule screen. A high throughput screen was performed with 832/13 cells stably expressing a VGF promoter-luciferase reporter construct (832/13-VGF-luc cells). Hits were called based on a robust deviation of 3 from the mean. Hits were then confirmed in 8 point dose response studies in 832/13-VGF-Luc cells (VGF) and counter-screened against HEK-293T PLTP-Luc (PLTP) and Ins1 CRE-Luc (CRE) cell lines. Dose response curves for two representative hits, GNF-7169 and GNF-9228, are shown.

201 We studied the effects of the 41 lead compounds on islet cell proliferation, GSIS, and 202 protection against apoptotic cell stress. Through this, we isolated three compounds (GNF-

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203 9228, GNF-1346, and GNF-4088) with robust and consistent effects to stimulate human islet cell proliferation. GNF-9228 and GNF-4088 are variants of a common chemotype, whereas 204 GNF-1346 has a distinct structure (Fig 2A). Consistent with the design of the screen, both 205 206 GNF-9228 and GNF-1346 caused potent induction of Nkx6.1 mRNA expression in 832/13 207 cells (Fig 2B). However, when tested on human islets, the two compounds caused a nonsignificant trend to increase Nkx6.1 mRNA, and had no effect on expression of VGF or the c-208 209 myc oncogene, a known activator of islet cell replication (Figs 2C, 2D, 2E) [18]. GNF-9228 210 was found to be the most potent of the three compounds for stimulation of human islet cell proliferation, and studies that follow therefore focus on the activities of this molecule. 211

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213 Fig 2. Structures of compounds GNF-9228, GNF-4088 and GNF-1346 and their effects 214 on possible target genes. **A.** Structures of three compounds with human islet cell 215 proliferative activity isolated from our 832/13-VGF-luc screen are shown. GNF-9228 and 216 GNF-4088 have very similar structures, whereas GNF-1346 is distinct. Β. RT-PCR measurement of Nkx6.1 levels in 832/13 cells following exposure to 10 µM GNF-1346, 10 µM 217 218 GNF-9228 or vehicle (DMSO) for 6 hours. Five human islet samples from different donors 219 were treated with 10  $\mu$ M GNF-9228, 10  $\mu$ M GNF-1346 or DMSO for 6 hours, followed by measurement of the following mRNAs by RT-PCR: C. Nkx6.1; D. VGF; E. c-myc. Data in 220 221 panels B-E are expressed as fold-increase in 832/13 cells or islets treated with GNF-1346 or 222 GNF-9228 relative to DMSO-treated, with n = 6 for 832/13 cells and n = 5 for human islet 223 experiments (\*\* p< 0.0001 comparing GNF-9228 or GNF-1346-treated cells to DMSO-treated 224 cells in panel B).

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**GNF-9228 selectively stimulates proliferation of human islet**  $\beta$ -cells. Human islet cell replication was measured by EdU incorporation, and specific islet cell types were identified by insulin, glucagon, or somatostatin co-staining. A total of 7 human islet aliquots, each from a different donor, were tested for the effect of 10 µM GNF9228 on islet  $\beta$ -cell and  $\alpha$ -cell replication. The dose of GNF-9228 was chosen as the maximally active dose based on

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231 titration studies in rat islets (S1 Fig). The individual human islet preps had substantial variation 232 in the percent of  $\beta$ - and  $\alpha$ -cells that were positive for EdU staining under control (DMSO-233 treated) conditions, including zero values for two of the  $\alpha$ -cell experiments. The data are 234 therefore presented as fold-response to normalize the differences in basal values. All 7 235 human islet preps exhibited an increase in  $\beta$ -cell and total cell EdU incorporation in response 236 to GNF-9228 relative to DMSO treatment, with statistically significant average responses of 6-237 fold in total islet cells, and 7.3-fold in  $\beta$ -cells (**Fig 3**). The 5 human islet samples with 238 detectable  $\alpha$ -cell EdU staining at baseline all increased EdU incorporation into  $\alpha$ -cells in 239 response to GNF-9228, but here the average was 4.3-fold, and of marginal statistical 240 significance (p=0.06). Additional information about the number of cells assayed and the 241 percentages of EdU-positive islet cells,  $\beta$ -cells, and  $\alpha$ -cells, is provided in (S1 Table). In 242 addition, in a subset of human islet preparations with detectable co-staining of EdU and 243 somatostatin under basal conditions, GNF-9228 caused no increase in EdU incorporation into 244  $\delta$ -cells (**S2 Table**). Taken together, these data suggest that under the conditions used in 245 these experiments (EdU exposure of 18 hours) GNF-9228 activates EdU incorporation into 246 human islet cells in a  $\beta$ -cell selective fashion, with a marginal effect on  $\alpha$ -cells and no detectable effect on  $\delta$ -cells. 247

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249 Fig 3. GNF-9228 selectively stimulates human Islet β-cell replication. Intact human islets 250 were cultured for 72 h in the presence of 10 µM GNF-9228 or DMSO (vehicle control). EdU 251 was added for the last 18 h of culture. Islets were dispersed, stained for EdU incorporation 252 and treated with antibodies against EdU, insulin or glucagon, and immunofluorescent signals 253 were detected and quantified with a Thermo Scientific Cellomics CX5- High Content (HC) cell 254 imaging system. A. Low magnification image showing increase in islet cell EdU incorporation 255 (yellow) in GNF-9228 versus DMSO-treated cells, with cells visualized by DAPI staining 256 B. High magnification image showing two Edu (orange nuclei), insulin (green (blue). 257 cytoplasm) co-positive cells and one Edu (orange nucleus), glucagon (blue cytoplasm) co-

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258 positive cell. **C.** Bar graph summary of data expressed as the fold-increase in EdU positive 259 cells across all islet cells assayed (left), in  $\beta$ -cells (EdU and insulin co-positive cells (center), 260 and  $\alpha$ -cells (EdU and glucagon positive cells (right), representing experiments performed on 7 261 independent human islet aliquots (total cells and  $\beta$ -cells) and 5 human islet samples ( $\alpha$ -cells), 262 expressed as mean +/- S.E.M.(\* p < 0.05). See S1 Table for further details.

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264 GNF-9228 stimulates insulin secretion in rat and human islets. We next tested the 265 impact of GNF-9228 on insulin secretion. Acute (less than 24 hour exposure) of islets to GNF-9228 had no impact on GSIS (S1 Fig). However, pre-exposure of human islets to 5 or 10 266 267 µM GNF-9228 for 72 hours caused a 75-140% increase in insulin secretion in the presence of 268 stimulatory (16.7 mM) glucose compared to human islets treated with DMSO (Fig 4). Insulin 269 secretion was normalized to insulin content in these experiments, which showed a non-270 significant trend to decrease in response to GNF-9228 treatment (human islets treated for 72 271 h with 5  $\mu$ M or 10  $\mu$ M GNF-9228 had 86 ± 9.1% and 83 ± 12.9% of the insulin content 272 measured in DMSO-treated islets, respectively). This non-significant change in insulin content 273 does not explain the enhanced insulin secretion elicited by GNF-9228 treatment. We also note 274 that insulin secretion at basal (2.8 mM) glucose levels increased significantly at 5  $\mu$ M, and 275 showed a non-significant trend to increase at 10 µM GNF-9228 (Fig 4). This increase in basal 276 insulin secretion caused the stimulation index (insulin secreted at 16.7 mM relative to 2.5 mM 277 glucose) to be essentially identical in the GNF-9228 and DMSO-treated islets. Nevertheless, pretreatment with, but not acute exposure to GNF-9228 enhances insulin secretion at 278 279 stimulatory glucose in human islets.

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Fig 4. Pre-treatment of human islets with GNF-9228 enhances insulin secretion. Intact human islets were pre-treated for 72 h with 5 or 10  $\mu$ M GNF-9228 or DMSO (vehicle control). Glucose-stimulated insulin secretion was then measured by static incubation in secretion buffer containing 2.5 mM glucose followed by 16.7 mM glucose for 1 h each. Secreted insulin is reported as a percentage of insulin content. Data represent the mean +/- S.E.M. from 4

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independent human islet preparations, each assayed in quadruplicate. (# p < 0.005 compared to DMSO treated cells at 16.7 mM glucose; p < 0.01 compared to DMSO treated cells at 2.5 mM glucose).

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290 GNF-9228 protects 832/13 cells from ER stress and cytokine-induced cytotoxicity. To 291 test GNF-9228 for its potential pro-survival effects, 832/13 cells were exposed to 500 nM 292 thapsigargin (TG) for 6 hours in the presence and absence of 10 µM GNF-9228. Co-treatment 293 of 832/13 cells with TG + GNF-9228 resulted in a clear reduction in cleaved caspase-3 compared to control cells incubated with TG + DMSO (Figs 5A and 5B). In another set of 294 295 experiments, 832/13 cells were pretreated with GNF-9228 or vehicle for 24 h followed by 24 h 296 exposure to 200 or 250 nM thapsigargin (TG) or a mixture of the cytotoxic cytokines IL-1 $\beta$  and 297  $\gamma$ -IFN in the presence or absence of GNF-9228 (Fig 5C). In these experiments, cell viability 298 was measured with the mitochondrial activity dye MTS. Exposure to 250 nM TG + DMSO 299 decreased cell viability by 67%, whereas cells treated with TG + GNF-9228 suffered only a 300 31% reduction in viability. Similarly, exposure of vehicle-treated 832/13 cells to the 301 inflammatory cytokines IL-1 $\beta$  +  $\gamma$ -IFN caused a 67% reduction in viability, whereas GNF-9228 limited the effect to a 40% reduction. Thus, GNF-9228 exhibits anti-apototic and pro-cell 302 303 survival effects in the INS-1-derived 832/13 cell line. Note that we chose to perform our 304 experiments in these cells based on our finding that they are far more sensitive to activation of 305 apoptosis than primary islet cells [16], suggesting that the cell line provides the most stringent 306 model for demonstrating cytoprotective effects of GNF-9228 at this stage of the work.

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Figure 5. GNF-9228 protects 832/13 cells from ER stress and cytokine-induced cytotoxicity. A. 832/13 cells were treated with 500 nM thapsigargin (TG) for 6h in the presence and absence of 10  $\mu$ M GNF-9228. A representative immunoblot of cleaved caspase-3 is shown. B. Bar graph summary of 4 independent densitometric measurements of cleaved caspase-3 in 832/13 cells exposed to 500 nM TG ± 10  $\mu$ M GNF-9228 or DMSO for

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6 hours. **C.** 832/13 cells were pre-treated with 10 μM GNF-9228 or DMSO for 24 h followed by 24 h incubation with two doses of thapsigargin (TG) or a mixture of IL-1β and γ-IFN. Cell viability was measured by MTS assay. ( $^{\text{k}}$ p< 0.001;  $^{*}$ p < 0.05;  $^{\text{s}}$ p < 0.01, comparing GNF-9228 to DMSO- treated cells).

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318 Comparison of the effects of GNF-9228 and the Dyrk1A inhibitor GNF-4877. To test 319 whether GNF-9228 activates islet cell proliferation by a mechanism similar to that reported for Dyrk1a inhibitors [4-6], we compared the effects of GNF-9228 and GNF-1346 to those of 320 321 GNF-4877 [4] and harmine [5] on NFAT nuclear translocation. Treatment with increasing doses of these agents demonstrated clear activation of NFAT nuclear translocation by GNF-322 4877 and harmine, but not GNF-9228 or GNF-1346 (Figs 6A and 6B). Consistent with these 323 findings, the robust proliferative effect of GNF-9228 in rat islets was not affected by co-324 325 treatment with cyclosporin A, a potent NFAT pathway inhibitor (S3 Fig). Also indicative of a unique mechanism of action, GNF-9228 caused a significant 3.9-fold increase in VGF-luc 326 327 activity relative to DMSO treatment (p < 0.005), whereas GNF-4877 failed to cause significant 328 luc activation in this assay (Fig 6C).

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Figure 6. GNF-9228 and GNF-1346 signal by distinct mechanisms relative to Dyrk1a 330 331 inhibitors. A. Ins1 cells treated with an NFATc1-GFP adenovirus were treated with a range 332 of doses of the Dyrk1a inhibitors GNF-4877 or harmine, GNF-9228, or GNF-1346 or DMSO for 2h. Following treatment, cells were fixed and imaged for GFP nuclear localization. Data is 333 quantified as percent of cells with nuclear localization of GFP for n = 3 experiments. B. 334 335 Representative images showing subcellular localization of NFATc in the presence of 10 µM 336 GNF-4877, harmine, GNF-9228, GNF-1346, or DMSO; C. 832/13 cells were transiently 337 transfected with the VGF-luc plasmid used to generate 832/13-VGF-luc cells and treated with 10 µM GNF-9228, 5 µM GNF-4877, or 5 µM forskolin for 24 h followed by measurement of 338 339 luc-generated luminescence. Data are expressed as fold-change relative to DMSO-treated cells in four independent experiments ( $^{\#}$  p<0.005). 340

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We also compared the proliferative effects of 10  $\mu$ M GNF-9228 to those of 5  $\mu$ M GNF-4877, a 342 343 concentration reported as maximally active for islet cell proliferation [4]. In these experiments 344 (Figure 7) human islets were exposed to EdU for 72 hours, in contrast to the studies in Figure 345 3 where EdU was provided for 18 hours. All 6 human islet preps responded to GNF-9228 or 346 GNF-9228 + GNF-4877 with an increase in EdU incorporation into total islet cells and  $\beta$ -cells, with statistically significant average responses of approximately 10-fold to both treatment 347 348 conditions (**Fig 7 and S3 Table**). GNF-4877 also increased EdU incorporation into  $\beta$ -cells in 349 all 6 human islet preps, with an average increase of 4-fold (p = 0.06). (Fig 7). Combining the 350 two compounds had no additive effect on total cell or  $\beta$ -cell proliferation. GNF-9228 or GNF-351 9228 + GNF-4877 also caused an average, statistically significant 5-fold increase in  $\alpha$ -cell 352 EdU incorporation in 5 separate human islet aliquots (Fig 7 and S3 Table). GNF-4877 alone 353 also increased  $\alpha$ -cell EdU incorporation in all 5 preps surveyed, with an average 3-fold 354 increase (p = 0.06). We also measured EdU incorporation into  $\delta$ -cells in human islet preps 355 treated with EdU for 72 hours and again found no consistent effect of GNF-9228 (S1 Table). 356 Thus, with longer duration of EdU exposure, we find that GNF-9228 increases both  $\beta$ -cell and 357  $\alpha$ -cell but not  $\delta$ -cell replication, with a preferential effect on  $\beta$ -cells.

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359 Fig 7. Comparison of effects of GNF-9228 and the Dyrk1a inhibitor GNF-4877 on human islet cell proliferation. Intact human islets were cultured for 72h in the presence of 10 µM 360 361 GNF-9228, 5 µM GNF-4877, a combination of both, or DMSO (vehicle control). EdU was 362 added for the last 72 h of culture. Islets were dispersed, stained for EdU incorporation and treated with antibodies against insulin or glucagon, and immunofluorescent signals assayed 363 with a Thermo Scientific Cellomics CX5- High Content (HC) cell imaging system. Data are 364 365 expressed as the fold-increase in total EdU positive cells (left),  $\beta$ -cells (EdU and insulin co-366 positive cells) (center), and  $\alpha$ -cells (EdU and glucagon positive cells) (right), representing

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experiments performed on 6 independent human islet aliquots (total cells and β-cells) and 5 human islet samples ( $\alpha$ -cells), expressed as mean +/- S.E.M. (\* p< 0.05, # p < 0.005 relative to DMSO control). See S3 Table for more details.

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#### 371 Compounds GNF-9228, GNF-1346, and GNF-4088 have poor *in vivo* pharmacodynamic

372 **properties.** To test the effects of our compounds in the *in vivo* setting, we first injected 30 373 mg/kg of GNF-9228 suspended in DMSO into normal C57BL6 mice and measured its blood 374 levels over the following 24 hours. Although high levels of GNF-9228 were detected at the 375 earliest time points (Cmax = 8493 nM), the compound was cleared rapidly, with levels falling 376 to 853 nM at 1 hour post-injection (half life = 8 minutes; S4 Fig). The pharmacokinetic 377 properties of GNF-1346 and GNF-4088 were even less encouraging. Nevertheless, we 378 performed a study in which GNF-9228 was injected into C57BL6 mice daily at a dose of 30 379 mg/kg for one week, with mice receiving BrdU in the drinking water at a concentration of 0.8 380 mg/ml throughout the study. GNF-9228 treatment caused no incease in BrdU incorporation 381 into mouse islet cells compared to DMSO-treated mice, likely due to its rapid clearance.

#### 383 Discussion

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385 Both major forms of diabetes involve loss of functional  $\beta$ -cell mass. In type 1 diabetes (T1D), islet  $\beta$ -cells are destroyed by an autoimmune mechanism, whereas in type 2 diabetes (T2D), 386 387 the decline in  $\beta$ -cell mass and function is mediated by metabolic fuel overload and 388 inflammatory pathways [19]. Thus, for both T1D and T2D, strategies for regeneration of 389 functional  $\beta$ -cell mass should ideally address all of the contributory elements of  $\beta$ -cell failure. 390 including susceptibility to cytotoxic agents, loss of proliferative capacity, and impairment of 391 insulin secretion. In addition, the strategy chosen should target regeneration of islet  $\beta$ -cells 392 specifically, or at least selectively, with minimal activity against other islet or peripheral cells. 393 In light of these challenges, it is not surprising that no  $\beta$ -cell regenerative drugs currently exist 394 for humans with diabetes.

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396 The current study was built upon work showing that overexpression of the homeobox 397 transcription factor Nkx6.1 stimulates islet  $\beta$ -cell replication in a selective fashion, while also 398 enhancing GSIS [7-12]. VGF is a gene that is robustly induced by Nkx6.1 overexpression. 399 and subsequent studies demonstrated several salutary effects of the VGF prohormone and its 400 encoded peptides such as TLQP-21 on  $\beta$ -cell survival and function [9,12]. Based on these 401 findings, we designed a cell-based screening strategy involving stable transfection of 832/13 402 cells with a VGF promoter-luciferase reporter gene. The screen yielded three compounds 403 representing two chemotypes (Fig 2) with a robust capacity to stimulate human islet cell 404 replication.

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406 The most potent of these compounds, GNF-9228, has a remarkable combination of 407 properties. First, it selectively activates human  $\beta$ -cell EdU incorporation, with a lesser effect 408 on  $\alpha$ -cell replication and no effect on  $\delta$ -cells. Second, pre-exposure of human islets to GNF-409 9228 enhances insulin secretion at stimulatory glucose levels (16.7 mM). Third, pre-treatment of 832/13 rat insulinoma cells with GNF-9228 is cytoprotective against the ER stress-inducing 410 411 agent thapsigargin or a mixture of the cytotoxic cytokines IL-1 $\beta$  +  $\gamma$ -IFN. We recognize that the compound is not entirely  $\beta$ -cell specific, with a lesser, but significant effect to stimulate  $\alpha$ -412 413 cell replication (Figure 7). We also acknowledge that the compound tends to increase insulin 414 secretion at basal glucose levels, an effect that if mimicked in the *in vivo* setting could result in 415 increased risk for hypoglycemia. Further evaluation of these potential shortcomings will 416 require development of GNF-9228 analogs with improved in vivo pharmacodynamics.

417

418 In addition to our work on Nkx6.1 and Pdx-1 as upstream inducers of islet cell proliferation [7-419 13], other signaling pathways that have been identified with potential for activating human  $\beta$ -420 cell proliferation include the PDGF pathway [21], signaling by TGF- $\beta$  family members 421 [13.20.22], and glucose-regulated activation of proliferation mediated by ChREBP and c-myc 422 [17,23]. Pathways that regulate translocation of the NFAT transcription factor have also 423 received attention stemming from early studies showing that conditional ablation of NFAT 424 caused a reduction of  $\beta$ -cell mass in mouse models [24-26]. The tyrosine kinase Dyrk1A is a 425 potent regulator of NFAT translocation [24]. Recently, Dyrk1A inhibitors with islet cell 426 proliferative effects have emerged including harmine, identified in a small molecule screen for c-myc inducers [5], GNF-4788, synthesized as a derivative of an aminopyrazine scaffold 427 428 previously shown to have proliferative activity [4, 27], and 5-IT, an adenosine kinase inhibitor 429 that cross-reacts with Dyrk1a [6]. All of these agents cause substantial increases in human  $\beta$ -

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cell proliferation, with either no impairment of function [4,5], or an actual increase in GSIS with
prolonged (12 days) treatment [6].

432

433 While these results provide encouragement for further investigation of Dyrk1a inhibitors for 434 diabetes therapy, concerns about their ultimate practical utility include: 1) When added to 435 human islets, harmine caused a significant increase in proliferation of non- $\beta$ -cells, including  $\alpha$ -436 cells,  $\delta$ -cells and ductal cells [5]. This coupled with the broad expression of Dyrk1a suggests 437 that unwanted proliferative responses might occur in peripheral tissues in response to 438 systemic administration of Dyrk1a inhibitors; 2) Harmine activates c-myc, explaining at least a 439 portion of its replicative activity [5], GNF-4788 inhibits the Dyrk1a-related kinase GSK- $\beta$  [4], 440 and 5-IT was originally isolated as an adenosine kinase inhibitor [6], suggesting possible off-441 target effects of these agents; 3) Overexpression of c-myc has independent effects on human 442 islet cell proliferation [17], but also appears to drive islet de-differentiation and impairment of 443 GSIS [28]; 4) In contrast to our work with GNF-9228, none of the studies on Dyrk1a inhibitors 444 report a cytoprotective effect.

445

Importantly, our studies suggest that GNF-9228 activates signaling pathways distinct from those used by Dyrk1a inhibitors. Unlike GNF-4788 and harmine, GNF-9228 does not induce nuclear translocation of an NFAT-GFP fusion gene. Moreover, the proliferative effects of GNF-9228 are not blocked by NFAT translocation inhibitors such as cyclosporin A, and GNF-9228 but not GNF-4788 activates the VGF-luc reporter in 832/13-VGF-luc cells. However, GNF-9228 and GNF-4788 have no additive effects on human islet cell proliferation, possibly suggesting some mechanistic commonality of these agents that remains to be defined.

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Although we have identified differences in GNF-9228 and Dyrk1a inhibitor cell signaling, we have not yet defined the mechanism(s) of action of GNF-9228. In human islets, a dose of GNF-9228 (10  $\mu$ M) that activates  $\beta$ -cell proliferation and GSIS causes a non-significant trend for increase of Nkx6.1 mRNA and has no effect on VGF or c-myc mRNA. This suggests that Nkx6.1 does not mediate the replicative effect of GNF-9228, and VGF is unlikely to explain its effects on insulin secretion and cell survival. Further studies will be required to discern the mechanism(s) of action of GNF-9228.

461

In sum, we report the identification of a small molecule that stimulates human  $\beta$ -cell proliferation, enhances insulin secretion, and protects against cytotoxic agents. An obvious next step is to test the impact of GNF-9228 on islet cell proliferation, function, and survival *in vivo*, but unfortunately all three of the compounds that emerged from our screen have poor pharmacodynamic properties. Our current focus is on development of chemically modified versions of GNF-9228 with enhanced bioavailability to allow *in vivo* testing. Further studies are also required to understand its mechanism(s) of action.

469

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472

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479

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483

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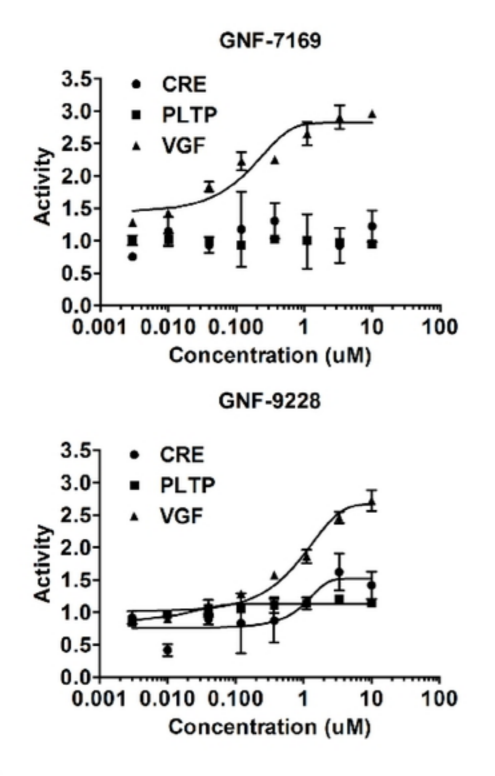
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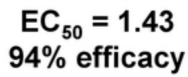
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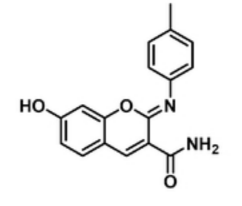


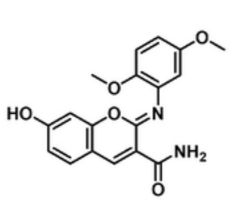
EC<sub>50</sub> = 0.11 98% efficacy

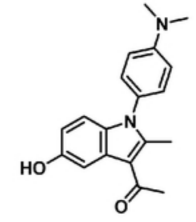




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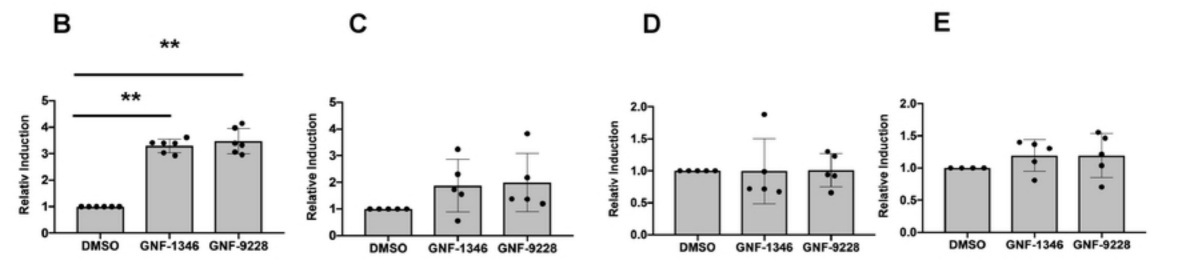




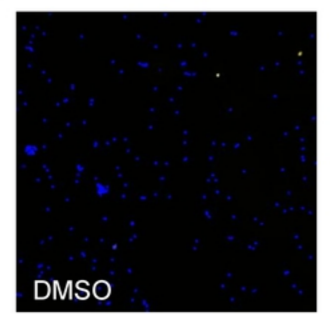
GNF-9228

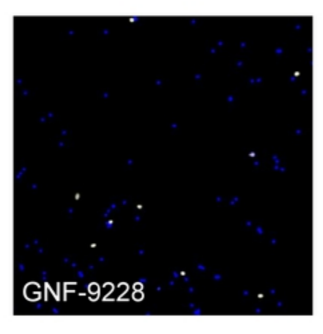
GNF-4088

GNF-1346

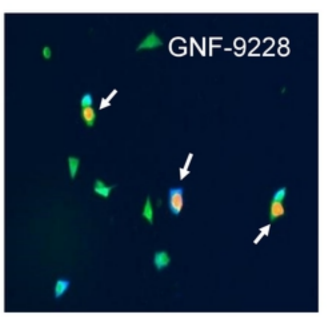


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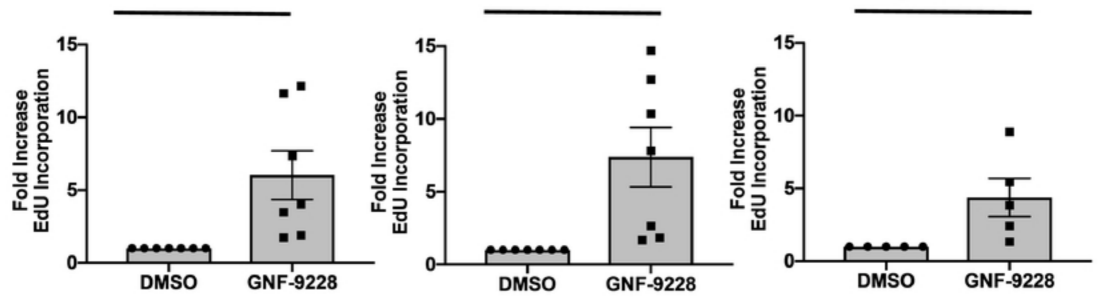
C Islet Cell proliferation

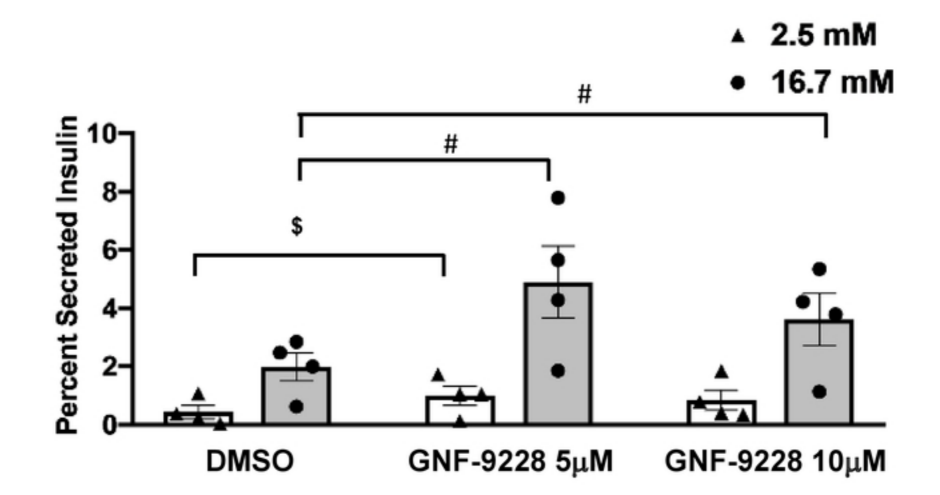
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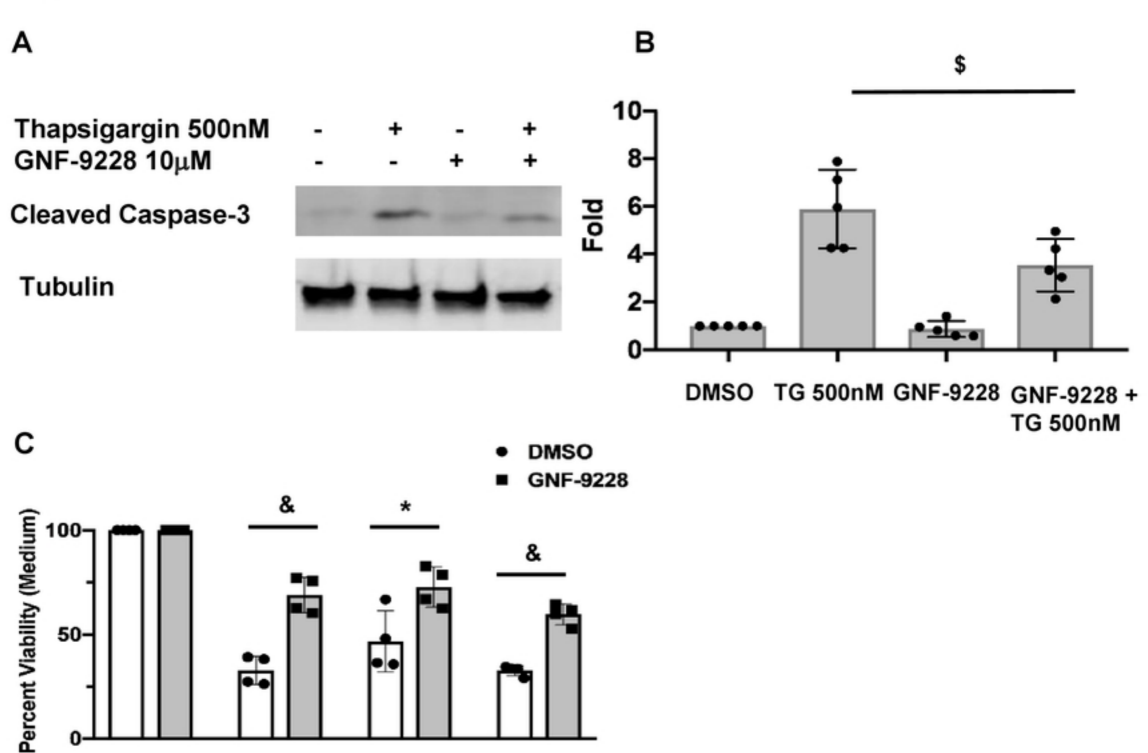
### β-Cell proliferation

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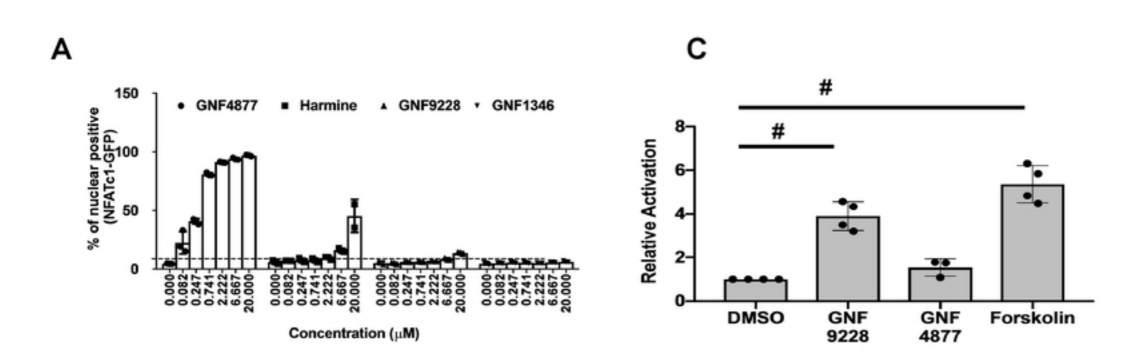
## α-Cell proliferation p=0.06







Medium TG 250nM TG 200nM IL-1β+IFN-γ



В

