

Slow oscillations persist in pancreatic beta cells lacking phosphofructokinase M

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1 **Slow oscillations persist in pancreatic beta cells lacking** 2 **phosphofructokinase M**

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6

7 **Abstract**

8 Pulsatile insulin secretion by pancreatic beta cells is necessary for tight glucose control in
9 the body. Glycolytic oscillations have been proposed as the mechanism for generating the
10 electrical oscillations underlying pulsatile insulin secretion. The glycolytic enzyme 6-
11 phosphofructokinase-1 (PFK) synthesizes fructose-1,6-bisphosphate (FBP) from fructose-
12 6-phosphate (F6P). It has been proposed that the slow electrical and calcium oscillations
13 (periods of 3-5 min) observed in islets result from allosteric feedback activation of PFKM
14 by FBP. Pancreatic beta-cells express three PFK isozymes, PFKL, PFKM and PFKP. A
15 prior study of mice that were engineered to lack PFKM using a gene trap strategy to delete
16 *Pfkm* produced a mosaic reduction in global *Pfkm* expression but the islets isolated from
17 the mice still exhibited slow Ca^{2+} oscillations. However, these islets still expressed residual
18 PFKM protein. Thus, to more fully test the hypothesis that beta cell PFKM is responsible
19 for slow islet oscillations, we made a beta-cell-specific knockout mouse that completely
20 lacked PFKM. While PFKM deletion resulted in subtle metabolic changes *in vivo*, islets
21 that were isolated from these mice continued to exhibit slow oscillations in electrical
22 activity, beta-cell Ca^{2+} concentration and glycolysis, as measured using PKAR, a FBP
23 reporter/biosensor. Furthermore, simulations obtained with a mathematical model of beta-
24 cell activity shows that slow oscillations can persist despite PFKM loss provided that one
25 of the other PFK isoforms, such as PFKP, is present, even if its level of expression is
26 unchanged. Thus, while we believe that PFKM may be the main regulator of slow

oscillations in wild type islets, PFKP can provide functional redundancy. Our model also suggests that PFKM likely dominates, *in vivo*, because it outcompetes PFKP by its higher FBP affinity and lower ATP affinity. We thus propose that isoform redundancy may rescue key physiological processes of the beta-cell in the absence of certain critical genes.

Significance

Pancreatic β -cells secrete insulin in pulses, reflecting bursting electrical activity and subsequent oscillations in the intracellular Ca^{2+} . It has been hypothesized that these events are generated by intrinsic glycolytic oscillations. We investigated the role of the muscle isoform of the glycolytic enzyme phosphofructokinase (PFKM) in the oscillatory activity of β -cells. PFKM is subjected to allosteric regulation conducive to glycolytic oscillations and has been previously shown to dominate other PFK isoforms in total enzymatic activity *in situ*. We show that β -cell specific depletion of *Pfkm* in mice does not affect the slow electrical bursting, Ca^{2+} oscillations, or glycolytic oscillations of islets *ex vivo*. Mathematical modelling provides an explanation for these results based on compensation by existing PFK isoforms of β -cells.

Introduction

In both human and mouse, pancreatic islets secrete insulin in a pulsatile fashion, and this pulsatility is lost in type 2 diabetes, thus reflecting its importance in regulation of blood glucose [1]. Pulsatile insulin release is necessary for the efficacious action of insulin without provoking insulin resistance in its target tissues, which include liver, fat and skeletal muscle [2].

In isolation, mouse islets are capable of robust oscillations over a range of periods from less than 1 min (e.g. 15 seconds) to 3 – 5 min [3]. While the faster oscillations are likely mediated by interactions between ion channels and sustained by increases in ATP/ADP triggered by glucose metabolism [4], the origin of the slower oscillations has been more elusive. An interesting and compelling early hypothesis for their generation is that they are triggered by slow oscillations in ATP/ADP due to intrinsic glycolytic oscillations in β -cells. Tornheim and associates proposed that the glycolytic enzyme phosphofructokinase, specifically its muscle isoform (PFKM), was responsible. This conjecture was based on the fact that autocatalytic activity of this enzyme wherein its product, fructose-1,6-bisphosphate (FBP), provides positive feedback to PFKM in an allosteric manner, and that the accelerated depletion of its substrate, fructose-6-phosphate (F6P), provides negative feedback [5, 6]. They also demonstrated that PFKM activity dominates that of the other isoforms in β -cells [5], even though later studies (including studies using RNAseq) reported similar levels of expression of M- and P-type isoforms [7-9] or even more P than M [10]; most all found that L is the least expressed isoform. Mathematical modelling of this process [11] was incorporated into two previous models from our group, the Dual Oscillator Model [4] and more recently, the Integrated Oscillator Model [12, 13]. While both models are capable of accounting well for the oscillations observed experimentally in mouse islets, several issues remained unclear, prompting the current investigation.

To test the hypothesis that slow oscillations are driven by glycolytic oscillations mediated by PFKM, Richard et al [8] made a whole-body mouse model using a gene trap to knock down PFKM. They found that reducing PFKM did not abolish or even significantly change the oscillatory properties of Ca^{2+} or insulin oscillations. However, PFKM expression was

not completely abolished by this approach, raising the possibility that residual PFKM protein was sufficient to support oscillatory activity. In addition, this knockdown approach was not β -cell specific, meaning that the contributions of other tissues could not be ruled out.

To critically revisit this issue, we constructed a mouse using contemporary genetic approaches to delete PFKM specifically and completely in β -cells, which we refer to as β -PFKM-KO mice. To further improve on the earlier study, we not only measured free Ca^{2+} oscillations with fura-2 and electrical oscillations using patch clamp, but directly tested for changes in glycolytic oscillations using a FRET probe of glycolytic activity of our own design, pyruvate kinase activity reporter (PKAR), which monitors the FBP level in the β -cell [14]. We report here that the electrical activity, slow oscillations in islet free Ca^{2+} , and the FBP level remained largely undisturbed in the islets of the knockout mice in which PFKM was completely knocked out in β -cells, and consequently the metabolic phenotype of the animals was also largely unaffected, although some small changes were observed. These results indicate that other PFK isoforms (PFKP or PFKL) provide the FBP needed for glucose metabolism in β -cells. Indeed, it is well established that mouse islets contain multiple PFK isoforms [5]. The question then becomes, can a non-M-type PFK isoform take over the role played by PFKM in generating oscillations in β -cell activity? These other isoforms have a lower affinity for the allosteric activator FBP, and a higher affinity for the inhibitor ATP [5]. Could a PFK enzyme having these properties sustain oscillations driven by oscillations in ATP, as has been proposed for PFKM?

In the second part of our study, we use the IOM mathematical model to show that when PFKM is absent, a non-PFKM isoform (which we refer to for specificity as PFKP, though PFKL could work as well) can assume the role of PFKM quite successfully. That is, with a model that incorporates both PFKM and PFKP, the activity of the former dominates under normal conditions by virtue of the increased activity, not by increased expression of PFKM. If PFKM is removed, however, PFKP takes over the role of PFKM, and provides the FBP necessary for sustaining metabolic oscillations. These results thus support the hypothesis that while PFKM likely dominates the other isoforms in ATP production necessary for

slow oscillations in wild type animals, the other isoforms are able to supply an alternative pathway when PFKM is disabled.

Material and Methods

Mathematical Model

We used a modified version of the Integrated Oscillator Model (IOM) to investigate the contribution of different PFK isoforms to the generation of β -cell oscillatory activity. The model is described in detail in Supporting Material. The differential equations were integrated numerically using MATLAB version 2020b (MathWorks Inc., Natick, MA) and the computer code can be downloaded from <https://www.math.fsu.edu/~bertram/software/islet/>.

The basic model for PFK activity that we employed was developed by Smolen [11]. We modified this model, which includes only one PFK isoform, to account for a second PFK isoform. This isoform, which could be either the liver-type (PFKL) or the platelet type (PFKP), has a lower affinity for FBP [15] and a higher affinity for ATP [16, 17] compared to PFKM. Due to the similar affinities of PFKL and PFKP, we included only one of the non-PFKM enzymes, PFKP.

In our model, the total flux through the PFK reaction, J_{PFK} , is the sum of the contributions of each of the two isoforms (PFKM and PFKP)

$$J_{\text{PFK}} = J_{\text{PFK-M}} + J_{\text{PFK-P}} , \quad (1)$$

where $J_{\text{PFK-M}}$ and $J_{\text{PFK-P}}$ are portions of flux mediated by PFKM and PFKP, respectively. Each component $J_{\text{PFK-I}}$, where I can be M or P, is described by

$$J_{\text{PFK-I}} = v_{\text{PFK-I}} \frac{w_{1110\text{-I}} + k_{\text{PFK}} \sum_{i,j,l} w_{ij1l\text{-I}}}{\sum_{i,j,k,l} w_{ijkl\text{-I}}} . \quad (2)$$

where the indices i, j, k , and l take on values of 0 and 1, and the maximum rate $v_{\text{PFK-I}}$ and weights w_{ijkl-I} are isoform-specific. The weights are given by

$$w_{ijkl-I} = \frac{\left(\text{AMP}/K_1\right)^i \left(\text{FBP}/K_{2-I}\right)^j \left(\text{F6P}^2/K_3\right)^k \left(\text{ATP}^2/K_{4-I}\right)^l}{f_{13}^{ik} f_{23}^{jk} f_{41}^{il} f_{42}^{jl} f_{43}^{kl}} . \quad (3)$$

The weights of the two isoforms differ only in their affinities for FBP and ATP. In (3), these affinities are represented by the parameters K_{2-I} and K_{4-I} , which identify FBP and ATP dissociation constants, respectively. We therefore set $K_{2-P} > K_{2-M}$ to reflect the lower affinity of PFKP for FBP than PFKM, and $K_{4-P} < K_{4-M}$ to account for the higher affinity for ATP of PFKP than PFKM. The parameter values are shown in Table 1.

The knockout of PFKM was simulated by decreasing the maximum rate through the PFK reaction mediated by the M-type isoform, $v_{\text{PFK-M}}$, from $0.01 \mu\text{M ms}^{-1}$ (baseline value) to $0 \mu\text{M ms}^{-1}$, while $v_{\text{PFK-P}}$ was unchanged.

Parameter	Value	Parameter	Value	Parameter	Value
$v_{\text{PFK-P}}$	$0.01 \mu\text{M ms}^{-1}$	K_{2-P}	$2 \mu\text{M}$	f_{23}	0.2
$v_{\text{PFK-M}}$	$0.01 \mu\text{M ms}^{-1}$	K_3	$5 \times 10^4 \mu\text{M}^2$	f_{41}	20
k_{PFK}	0.06	K_{4-M}	$1000 \mu\text{M}^2$	f_{42}	20
K_1	$30 \mu\text{M}$	K_{4-P}	$100 \mu\text{M}^2$	f_{43}	20
K_{2-M}	$1 \mu\text{M}$	f_{13}	0.02		

Table 1. Parameters used for the PFK subsystem of the model.

Molecular biology

Construction of β -cell specific PFKM null mice

CRISPR/Cas9 was used to identify a suitable target for Cas9 endonuclease by submitting genomic *Pfkm* DNA sequence to an algorithm (<http://www.crispor.tefor.net>; [18]). sgRNAs for Cas9 targets were obtained from Synthego.com [19] and recombinant Cas9

endonuclease was from MilliporeSigma [20]. Mouse zygotes microinjected with Cas9/sgRNA ribonucleoprotein complexes (RNP) identified a sgRNA to cleave exon 3 of *Pfkm*. DNA from blastocysts was subjected to PCR and DNA sequencing to identify small insertions/deletions at Cas9/sgRNA cut sites [21] and a single stranded DNA donor replaced the critical exon with a floxed exon [22, 23]. Premature termination codons introduced using this approach have been shown to block protein production [24, 25]. RNP and ssDNA donor were microinjected into mouse zygotes (50 ng/μl Cas9 protein, 30 ng/μl sgRNA, 10 ng/μl ssDNA donor) as described [26]. Surviving zygotes were then transferred to pseudopregnant females. Potential G0 founder pups were screened for floxed *Pfkm* using PCR [21, 27-29]. G0 founders carrying floxed *Pfkm* were mated to wild type mice and resulting G1 pups screened for floxed *Pfkm* transmission. The sequences of G1 pups were determined by cloning of genomic DNA flanking the insertion site [29]. A floxed *Pfkm* mouse line derived from independent founders was crossed with RIP2-cre (expressing rat insulin promoter 2) mice to generate β -cell specific *Pfkm* nulls. Islets isolated from these mice had selective loss of PFKM mRNA and protein compared to Cre-positive wild-type PFK mice, by RT-PCR and western blotting, respectively.

Gene expression analysis

Total RNA was extracted from isolated islets using miRNeasy micro kit and treated with DNase I according to the manufacturer's instructions (Qiagen). cDNA was synthesized using High-Capacity cDNA Reverse Transcription Kit (Thermo Fisher). Quantitative real time RT-PCR (RT-qPCR) was performed on a StepOnePlus™ System (Applied biosystem) using TaqMan Universal PCR Master Mix (Thermo Fisher). Each sample was run in duplicates, and the gene expression was calculate using the change in threshold ($\Delta\Delta CT$) method with TATA box binding protein (TBP) as internal control. The TaqMan gene expression probes used in the study are summarized in Table S1 in the Supporting Material. Genotyping was done on a regular basis by sending tail samples to a commercial lab (Transnetyx, Cordova, TN).

Islet preparation

Islets were isolated from 3-4-month-old mice using collagenase injection using an established protocol [30]. The animal protocol used was approved by the University of Michigan Committee on the Use and Care of Animals (UCUCA). Islets were hand-picked into saline and then transferred into culture media consisting of RPMI1640 supplemented with FBS (10%), glutamine, and pen/strep. Islets were kept in culture overnight in an incubator at 37° C.

Live cell imaging methods

Adenoviruses were used to express the PKAR FRET biosensor in pancreatic islet β -cells under control of the rat insulin promoter as in [30]. Islets were placed in a glass-bottomed chamber (54 mL volume) (Warner Instruments, Hamden, CT) on a model No. IX71 inverted microscope (Olympus, Melville, NY) equipped with a 20x/0.75 NA objective (Nikon Instruments, Melville, NY). The chamber was perfused at 0.3 mL/min and temperature was maintained at 33° C using inline solution and chamber heaters (Warner Instruments). Excitation was provided by a TILL Polychrome V monochromator set to 10% output. Excitation (x) or emission (m) filters (ET type; Chroma Technology, Bellows Falls, VT) were used in combination with an FF444/521/608-Di01 dichroic (Semrock, Lake Forest, IL) as follows: 430/24x, 470/24m and 535/30m (430x – R535m/470m). Fluorescence emission was collected with a QuantEM:512SC camera (PhotoMetrics, Tucson, AZ) or an ORCA-Flash4.0 V2 Digital CMOS camera (Hamamatsu, Skokie, IL) at 0.125–0.2 Hz. A single region of interest was used to quantify the average response of individual β -cells using MetaMorph (Molecular Devices, LLC, San Jose, CA.) software.

Calcium measurements

Islets were loaded with fura-2/AM (2.5 μ M) for 45 min in medium containing 5 mM glucose prior to imaging. Islets were then transferred to a 1 mL perfusion chamber containing 5 mM glucose imaging buffer for 6 min, followed by 10 to 30 min perfusion with this solution at approximately 1 mL/min. Imaging buffer contained (in mM): 140 NaCl, 3CaCl₂, 5 KCl, 2 MgCl₂, 10 HEPES and 5 glucose. Ratiometric fura-2 imaging was

carried out using 340/380 nm excitation and collecting 502 nm emission, as previously described [30]. The fluorescence data were acquired using Metafluor, with a single region of interest used to quantify the average response of individual islets.

Electrophysiology

Patch pipettes were pulled from filament-containing borosilicate glass capillaries (WPI Instruments) using a Sutter P-97 puller (Sutter Instruments, Novato, CA) and had resistances of 4-6 M Ω when filled with solution containing an internal buffer containing (in mM): 28.4 K₂SO₄, 63.7 KCl, 11.8 NaCl, 1 MgCl₂, 20.8 HEPES, and 0.5 EGTA at pH7.2. Electrodes were then backfilled with the same solution but containing amphotericin B at 0.36 mg/ml to allow membrane perforation. Islets were transferred from culture dishes into a 0.5 ml recording chamber. Solutions held at 32–34°C were driven through the bath by a gravity system at a rate of 1 ml/min. Islets were visualized using an inverted microscope (Olympus IX50). Pipette seals obtained were >2 G Ω . Perforation was judged to be successful when the series resistance decreased to a steady-state level and membrane capacitance increased. Recordings were made using an extracellular solution containing (in mM): 140 NaCl, 3 CaCl₂, 5 KCl, 2 MgCl₂, 10 HEPES, and 11.1 or 2.8 glucose. β -cells were identified by their lack of activity in 2.8 mM glucose and by the appearance of regular electrical bursting in external solution containing 11.1 mM glucose. Drugs were dissolved directly into saline solution daily using DMSO stocks; the final concentration of DMSO used was always <0.1%.

One β -cell in each intact islet was typically patched. After the perforated patch configuration in voltage clamp mode was established, membrane potential was recorded in the current clamp mode.

RT-PCR

Total RNA was extracted from islets using the RNeasy Mini Kit (Qiagen, Ann Arbor, MI) according to the manufacturer's instructions. 0.4 μ g of total islet RNA was reverse-transcribed using Superscript RT II. Real-time experiments were carried out using an SYBR green PCR master mix (Applied Biosystems) with the primers shown in Table S1

in the Supporting Material. Raw threshold-cycle (CT) values were obtained using Step One software, and mean CT values were calculated from triplicate PCR reactions for each sample. Data were presented as RQ values (2-DD CT) with expression presented relative to an endogenous control, HPRT1.

Western blotting

Islets were handpicked in ice-cold PBS immediately after isolation and gently spun to remove the supernatant. Next, islets were lysed in RIPA lysis buffer supplemented with total protease inhibitor cocktail and stored at -80°C. Next, islet lysate was prepared by passing through 30-gauge needle. Protein concentration was determined using BCA reagent kit. Next, 25 µg protein lysate was mixed with loading buffer and boiled at 70°C for 10min. and separated by electrophoresis on NuPAGE 4-12%, Bis-Tris mini gels. Next, proteins from the gels were transferred on nitrocellulose membrane (iBlot2 transfer stacks) using iBlot 2 Gel Transfer Device with a preset 7min transfer protocol. Next, the membrane was incubated in 5%BSA in TBS tween 20 (TBST) 0.1% for the blocking of non-specific epitopes. Next, PFKM (1:1000 in TBST 0.1%) isoform and GAPDH (Rb 1:10,000 in 5%BSA in TBST 0.1%) primary antibodies were used overnight at 4°C. The primary antibodies were probed with secondary rabbit HRP (1:5000 5% BSA TBST 0.1%) by incubation at room temperature for 1hr. HRP signals was detected by incubating membranes in SuperSignal™ West Femto Chemiluminescent Substrate for 5 min at room temperature and membranes were subsequently imaged on ChemiDoc System (BioRad). The protein ladder was used to locating protein on membrane. Islets were collected from 10 control and five $\beta^{Pfk\text{m-/-}}$ animals.

Chemicals and reagents

Gels (ThermoFisher Scientific, cat#NP0321PK2), RIPA (Boston Bioproduct, cat#BP-115), iBlot transfer stack nitrocellulose membrane (Invitrogen, cat#IB23002), PFKM (Invitrogen; #PA5-29336), Gapdh (CST; G9545), Protease inhibitor (CST, cat#5872), Rb HRP (Invitrogen; cat#7074P2), Protein ladder (Biorad, cat#1610377), SuperSignal™ West

250 Femto Chemiluminescent Substrate (ThermoFisher, cat#62237), iBlot2 transfer stacks;
251 ThermoFisher#IB23002; BCA protein assay kit (ThermoScientific; cat#23225).

252 *Metabolic measurements and data analysis*

253 Whole animal measurements were made by the University of Michigan Animal
254 Phenotyping Core. For IPGTT, mice were fasted for 5h, given 1.25g/kg glucose IP and tail
255 vein blood were sampled before and then 0, 5, 15, 30, 60 or 120 min after injection. Blood
256 glucose was measured using a glucometer (Accucheck, Roche) and plasma insulin using
257 ELISA (Millipore). Tail vein blood was sampled before and then 0, 15, 30, 45, 60, 75 and
258 90 min after injection of glucose.

259 *Analysis of membrane potential and live cell imaging data*

260 Quantitative analysis of membrane potential, islet free calcium, and PKAR oscillations was
261 performed using MATLAB to measure oscillation period and plateau fraction. For
262 membrane potential and free calcium, traces were linearly detrended within time intervals
263 corresponding to a fixed glucose level, then lightly low-pass filtered (Savitzky-Golay filter,
264 cutoff period of ~45-60 s). Oscillation periods were detected as repeated crossings of a
265 threshold value of 55% of the trace amplitude. Plateau fraction was recorded as the time
266 between the beginning of a period and the subsequent time when the trace dropped below
267 45% of trace amplitude. The mean period and plateau fraction of all oscillations at each
268 applied glucose level were then computed.

269 For PKAR recordings, traces corresponding to individual β -cells were first normalized to
270 deviations from the trace mean value, $[x - \text{mean}(x)]/\text{mean}(x)$. Traces were then
271 detrended using a low-pass filtered trend line (Savitzky-Golay filter, cutoff period of ~15
272 min), and traces from β -cells corresponding to the same islet were averaged. The islet-
273 averaged traces were then low-pass filtered (Savitzky-Golay filter, cutoff period of ~2
274 min), and oscillation properties detected as described above.

275 Statistics were done with R (version 4.0.3) and MATLAB. Repeated measures ANOVA
276 was done with R command `aov`, followed by t-test. Linear mixed effect models were fit

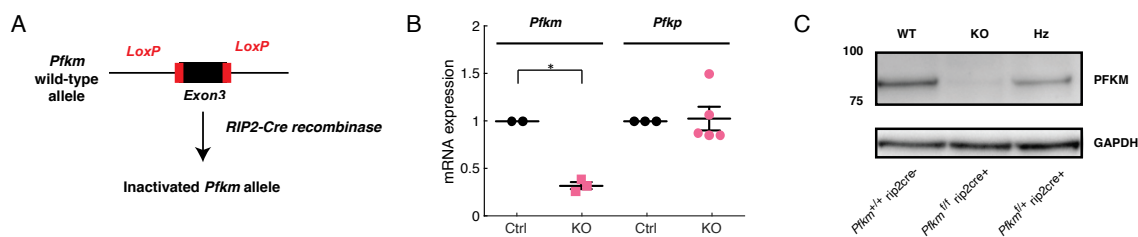


Fig. 1. β -PFKM-KO mice were generated using CRISPR/Cas9 to flox exon 3 of the *Pfk* gene (panel A). Employing this method, we were able to selectively delete *Pfk* mRNA transcript in KO mouse islets, while no statistically significant difference in *Pfkp* mRNA was observed between wild type controls and the KO islets (panel B). Western blot analysis done using a PFKM specific antibody confirmed the loss of PFKM protein in the KO islets, while reduced PFKM protein was evident in islets from heterozygous mice (panel C).

with the R commands `lm` and `lmer` to assess the dependence of oscillation period and plateau fraction on PFKM knockout status, sex, and glucose concentration. Details of those results are in Supporting Material.

Results

Oscillations persist in islets from β -PFKM-KO mice

Mouse islets have been previously shown to express PFKP, L, and M isoforms [5, 8]. As shown diagrammatically in Fig. 1A, exon 3 of *Pfk* was floxed using CRISPR/Cas9 and the resulting progeny were crossed with *RIP2-cre* mice to generate β -cell-specific *Pfk* null mice. Islets from *Pfk* null mice (knockout) shows depletion of *Pfk* transcripts compared to islets from litter-mate controls. Conversely, we found no statistically significant difference in *Pfkp* mRNA between islets from controls or *Pfk* null mice (Fig. 1B). Western blot analysis done with PFKM specific antibody (Fig. 1C), confirmed a loss of PFKM protein in the KO, whereas reduced protein was present in islets from heterozygotes. Taken together, our β -PFKM null mice exhibit β -cells specific depletion of

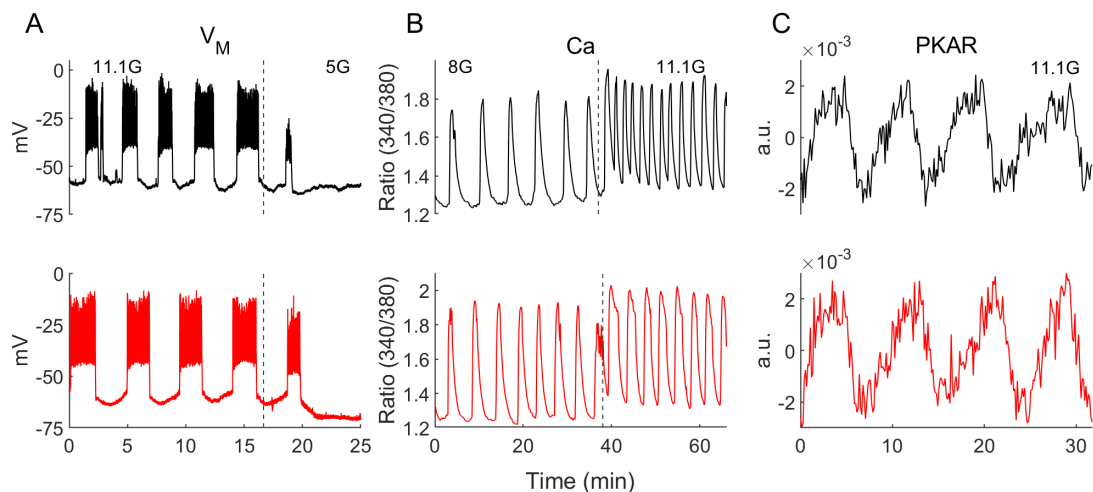


Fig. 2. β -PFKM-KO islets exhibit typical oscillations. Representative examples of oscillations at several glucose levels in control (black) and β -PFKM-KO islets for membrane potential (V_M , panel A), intracellular Ca^{2+} concentration (Ca, panel B), and normalized, detrended PKAR FRET ratio (panel C). Traces are representative of 21 V_M recordings, 10 Ca recordings, and 23 PKAR recordings. Control traces shown were from Cre⁺ control mice.

PFKM and appropriate model to investigate the role of PFKM in generating glycolytic oscillation [8].

To test whether the loss of PFKM in β -cells altered the oscillatory properties of the islet, as has been previously predicted [6], we used perforated patch-clamp to record oscillations of islet membrane potential, fura-2 to monitor islet Ca^{2+} , and PKAR to dynamically monitor FBP in both knockout and wild-type mouse islets. As shown in Fig. 2, there were no obvious changes in the oscillations of electrical activity (Fig. 2A; black traces correspond to WT, red to KO), or Ca^{2+} (Fig. 2B), which are typically observed in response to glucose concentrations $> \sim 7$ mM. Furthermore, reducing glucose concentration from 11.1 mM to 5 mM promptly shut off the oscillations as is typically seen in normal islets, and increasing glucose from 8 to 11.1 mM led to similar changes in the oscillations.

Finding that loss of PFKM did not strongly affect islet oscillations suggested that there may be redundant mechanisms regulating glycolytic oscillations and concomitant insulin release that are revealed by the absence of PFKM. To address this as well as further

improve on the earlier study, we next examined the glycolytic oscillations of WT and knockout islet β -cells using Pyruvate Kinase Activity Reporter (PKAR), a probe of our own design that was described previously [14]. Quantitative analysis of the periods and plateau fractions of islet membrane potential recordings from a total of 21 islets from 14 mice, Ca^{2+} recordings of 112 islets from 6 mice (10 recordings), and PKAR recordings of 52 islets from 14 mice (23 recordings) was carried out (as described in Table S7 of the Supporting Material). Fig. 3 summarizes this analysis, showing that β -PFKM-KO islets (red) had oscillations with similar periods and plateau fractions as compared to control islets (black) for membrane potential (Fig. 3A), Ca^{2+} (Fig. 3B), and PKAR oscillations (Fig. 3C) at different glucose levels. To statistically analyse islet oscillations across different groups, we needed to account for the hierarchical nature of islet measurements at three levels: islets were exposed to one or more glucose levels (repeated measures), batches of one or more islets from an animal were recorded in each recording, and one or more recordings were made from each animal. We addressed this using linear mixed effects modelling of oscillation properties as a function of glucose stimulus, PFKM status, and sex, with random effects for islet, recording, and mouse (see Tables S4, S5, and S6 and Fig. S1 in Supporting Material for details). This analysis indicated there was a significant dependence on glucose concentration, as expected. However, only a slightly smaller period was detected in the β -PFKM-KO islets, which was close to, but did not achieve, statistical significance at the $p=0.05$ level (V_M : -1.12 min, $p=0.067$; Ca : -0.71 min, $p=0.15$; PKAR: -1.44 min, $p=0.053$). Plateau fraction effect sizes were very small and not different statistically between β -PFKM-KO and control (V_M : -0.02, $p=0.58$; Ca^{2+} : 0.0, $p=0.98$; PKAR: -0.02, $p=0.40$). These results are consistent across all three recording types. We did note that the islets of male mice tended to have slightly higher oscillation period (~ 0.63 - 0.72 min higher than control), but this was not significant in any assay. Plateau fraction was slightly higher in islets from male animals for V_M and Ca recordings (V_M : +0.13, $p=0.046$; Ca : +0.13, $p=0.088$), but was slightly lower in the PKAR recordings (-0.06, $p=0.029$). These results show that PFKM is clearly not needed to produce slow oscillations. It is to be noted that the differences in oscillation periods reported here, especially between V_M or calcium oscillations and those of PKAR reflect the differing experimental conditions used, as viral induction of PKAR required adenoviral transduction followed by three days

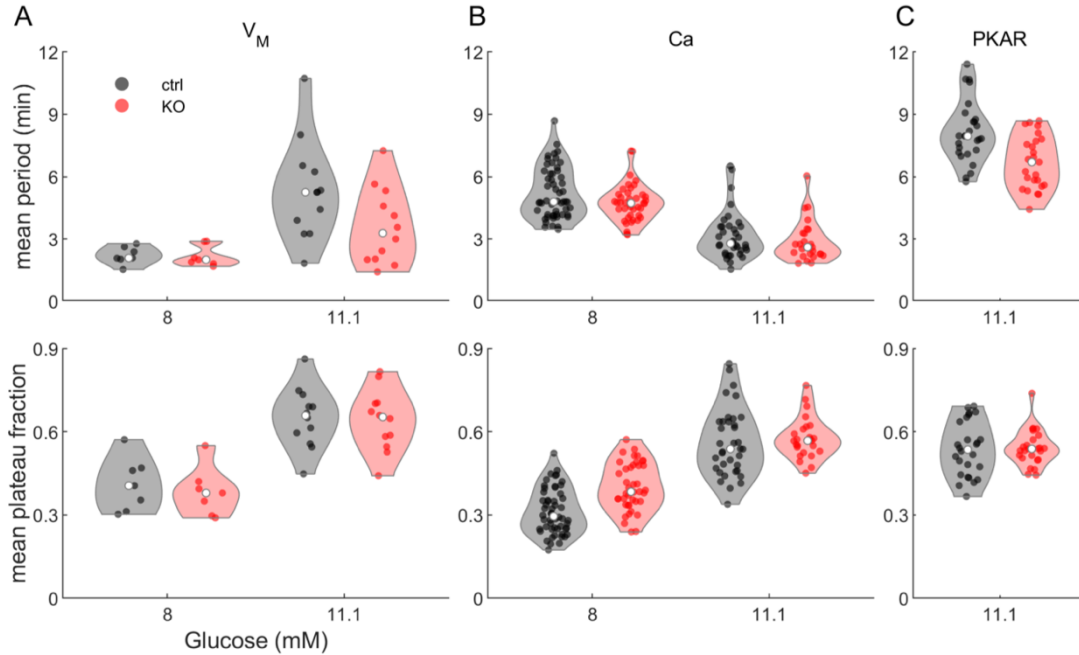


Fig. 3. Comparison of oscillation period and plateau fraction between β -PFKM-KO and control islets. Violin plots showing mean oscillation period (top panels) and plateau fraction (bottom panels) for islets exposed to specific glucose levels: 8 mM and 11.1mM glucose for membrane potential (panel A) and Ca^{2+} concentration (panel B), or at 11.1 mM glucose for PKAR (panel C). White dots indicate the median across all islets. All β -PFKM-KO islets oscillated at 8 mM and 11.1 mM glucose. Linear mixed effects modeling (see Material and Methods and Supporting Material) found that the modest reduction in period in β -PFKM-KO compared to control oscillations, was close to but did not achieve statistical significance at the $p=0.05$ level. The differences in plateau fraction were not significant. Control islets consisted of mostly $+/+$ Cre + islets (59%), with the remaining from WT or fl/fl Cre – mice. As no differences were noted among the controls, the results were pooled.

338 in culture [14]. In contrast to the results reported here, when simultaneous recordings of
 339 V_M and PKAR were made, the periods measured were strikingly similar [30].

340 To test whether the very small differences we observed in the properties of isolated islets
 341 were reflected in the *in vivo* metabolic profile of WT ($n = 21$; 11 F, 10 M) and null mice (n
 342 $= 25$; 15 F, 10 M), we conducted intraperitoneal glucose tolerance tests of the mice

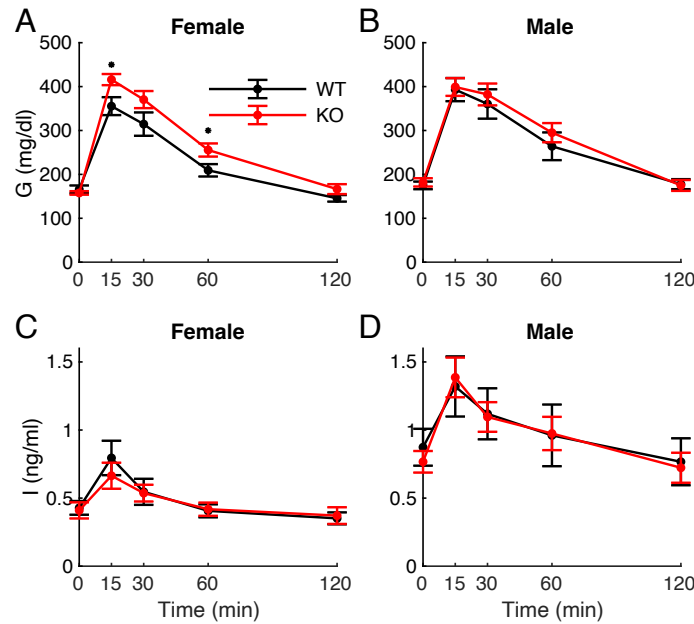


Fig. 4. In vivo metabolic measurements using intra-peritoneal glucose tolerance tests. Panels A and B, female mice; panels C and D male; panels A and C, glucose, panels B and D, insulin. The only significant difference is glucose for the females ($p = 0.04$) by repeated measures ANOVA. The values at $t = 15$ min and $t = 60$ min are different by unpaired t-test ($p < 0.05$).

(IPGTT). Female mice with the β -cell specific PFKM deletion showed a slight increase in glucose but no difference in their insulin (Fig. 4), suggesting a mild impairment in insulin secretion. We tested this using HOMA-Beta [31] and the insulinogenic index (IGI) (increment of insulin/increment of glucose during the first 30 minutes of the IPGTT) [32], and, though both measures of beta-cell function were numerically smaller in KO mice, neither was statistically significant (HOMA-Beta: 2.35 ± 1.7 control vs 2.00 ± 1.2 KO, $p = 0.4$; IGI: 0.0024 ± 0.004 control vs 0.0020 ± 0.001 KO, $p = 0.5$).

The overall increase in glucose was significant ($p = 0.04$) as assessed by repeated measures ANOVA, and the glucose values taken at the 15- and 60-minute time points were significantly different by t-test ($p < 0.05$). There was no significant difference in the insulin, however, by repeated measures ANOVA. Male mice showed no effect of PFKM deletion in either their glucose or insulin (repeated measures ANOVA was not significant).

The lack of effect of deleting PFKM in β -cells that are known to express other PFK isoforms suggests the possibility that another PFK isoform could be more functionally important in β -cells. While this is a tempting conclusion, we instead asked a subtler question: could these other isoforms take over the role of PFKM in driving metabolic oscillations, despite their differences in biochemical properties and despite the fact that PFKP expression, at least in terms of mRNA transcript levels, was unchanged in the knockouts (Fig. 1B)? To answer this, we next sought to take advantage of what is known biochemically about the differences between the isoforms and their regulation and examine their ability to generate oscillations in a mathematical model.

Slow oscillations may persist because PFKP takes over from PFKM.

In the Integrated Oscillator Model (IOM), modified to include equations for PFKP as well as PFKM (see Material and Methods), slow bursting electrical activity and associated slow Ca^{2+} oscillations occur under wild-type conditions (Fig. 5A, black trace). This activity is accompanied by slow oscillations in the ATP level (Fig. 5B). The rapid changes in V_M and contrasting slow changes in ATP have been universally seen in experiments, such as [30, 33], and also replicated in previous models, such as [34]. This is a consequence of the much slower kinetics of ATP compared to Ca^{2+} . During the active phase of each burst, the cytosolic Ca^{2+} concentration is elevated, so ATP is utilized to power Ca^{2+} pumps in the plasma membrane and endoplasmic reticulum membrane [35], resulting in a decline in the ATP concentration due to ATP consumption. Between bursts (i.e., silent phase), the Ca^{2+} concentration is low, reducing the ATP utilization and resulting in a rise in the ATP concentration. The fast jumps of Ca^{2+} in Fig.5 occur when the slow changes in ATP push the spiking dynamics back and forth across its thresholds for activity. The Ca^{2+} oscillations also induce oscillations in the FBP level through Ca^{2+} activation of pyruvate dehydrogenase [36](Fig. 5C). We refer to this mechanism as a passive metabolic oscillator (PMO), in which metabolic oscillations passively result from the Ca^{2+} oscillations. Although both PFKP and PFKM are present under wild-type conditions, the majority of the metabolic flux is through the M-type isoform (Fig. 5D, E, F), reflecting the fact that PFKM is the most active isoform [37].

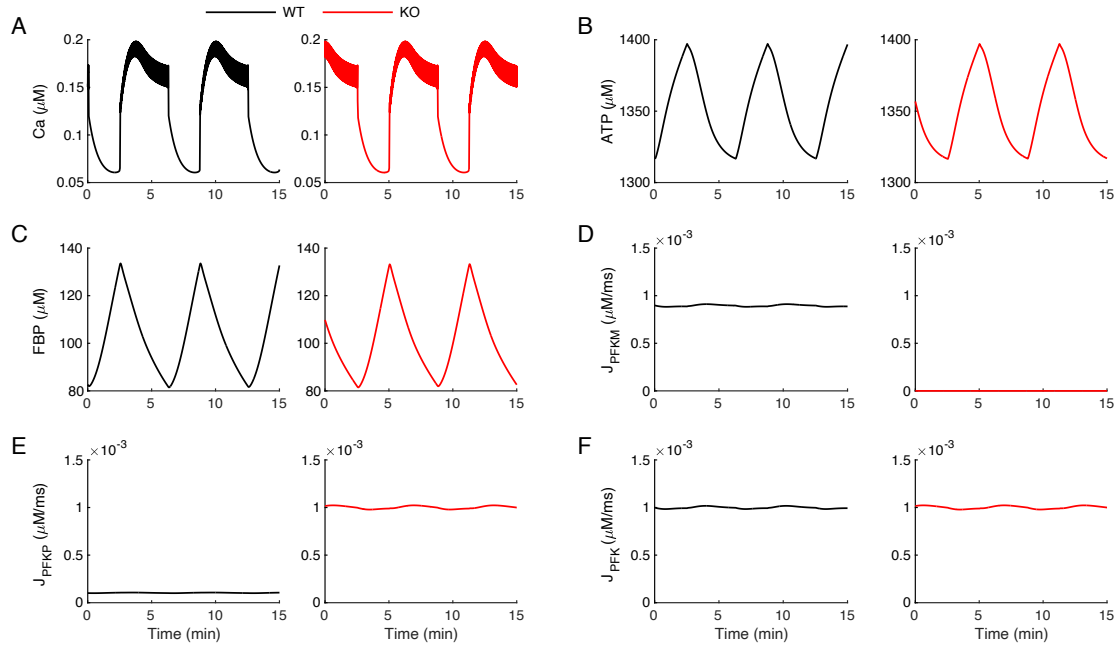


Fig. 5. Simulations showing the mechanism for persistence of slow oscillations in β -PFKM-KO islets. In the IOM, slow oscillations persist when PFKM is knocked out, due to increased activity of PFKP. The black traces are for wild-type conditions (with PFKM present), the red traces are after PFKM knockout. (A) The free cytosolic Ca^{2+} concentration exhibits slow oscillations for both wild-type and knockout conditions. The ATP concentration (B) and FBP concentration (C) exhibits slow oscillations before and after the knockout. (D) Metabolic flux through the PFKM enzymatic reaction is eliminated once the PFKM enzyme is knocked out. (E) The metabolic flux through PFKP is very small in the wild-type case, but in the knockout is comparable to the wild-type PFKM flux. (F) The total PFK metabolic flux is the same before and after the removal of PFKM. The time courses were generated with $g_{\text{K}(\text{Ca})} = 150 \text{ pS}$ and $v_{\text{PDH}} = 0.4 \text{ } \mu\text{M ms}^{-1}$. For the wild-type simulation (black), $v_{\text{PFK-M}} = 0.01 \text{ } \mu\text{M ms}^{-1}$ and $v_{\text{PFK-C}} = 0.01 \text{ } \mu\text{M ms}^{-1}$. PFKM is knocked out (red) by setting $v_{\text{PFK-M}} = 0 \text{ } \mu\text{M ms}^{-1}$.

384 When PFKM knockout is simulated, by setting the maximum activity rate of PFKM to 0
 385 (see Material and Methods), the oscillations in intracellular Ca^{2+} concentration, ATP
 386 concentration, and FBP concentration persist with no significant changes (Fig. 5A, B, C,
 387 red traces). This surprising result can be explained with an analysis of the distribution of

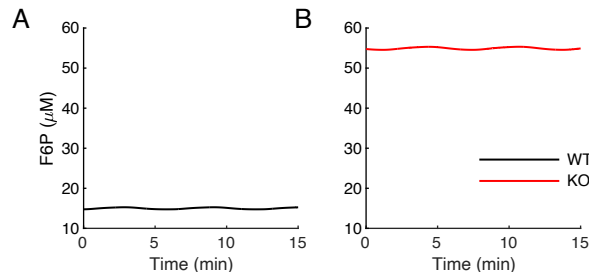


Fig. 6. The mathematical model predicts an increase in the F6P concentration in the β -PFKM-KO during slow bursting. The mean value of the F6P concentration is lower in the wild-type model islet (A) than in the model β -PFKM-KO islet (B).

PFK flux through the two isoforms. The total flux through PFK is the same after the knockout of PFKM as before its knockout (compare the black and red traces in Fig. 5F). However, the scenario at the single isoform level changes significantly once the PFKM is knocked out. In wild-type conditions, the flux through PFKP (Fig. 5E, black trace) is negligible compared with the PFKM flux (Fig. 5D). However, after the PFKM knockout, the PFKP flux reaches a level equal to that of the PFKM flux prior to its removal (compare the red trace in Fig. 5E to the black trace in Fig. 5D), while the PFKM flux is now zero.

One may expect the flux through PFK to be lower after removal of PFKM. How was it possible for PFKP to completely compensate for the loss of PFKM? This occurs because when PFKM is knocked out, the level of the PFK substrate F6P increases dramatically (Fig. 6). This higher substrate level compensates for the less-favourable allosteric affinities of PFKP, allowing it to produce the same metabolic flux as did PFKM. We note that none of the PFKP parameters were altered in this simulation. In particular, there is no upregulation of the PFKP enzyme. It is simply more active following the knockout of PFKM because the substrate level is much higher than before the knockout. This is consistent with mRNA expression data which shows that PFKP expression is not affected by the PFKM knockout (Fig. 1B).

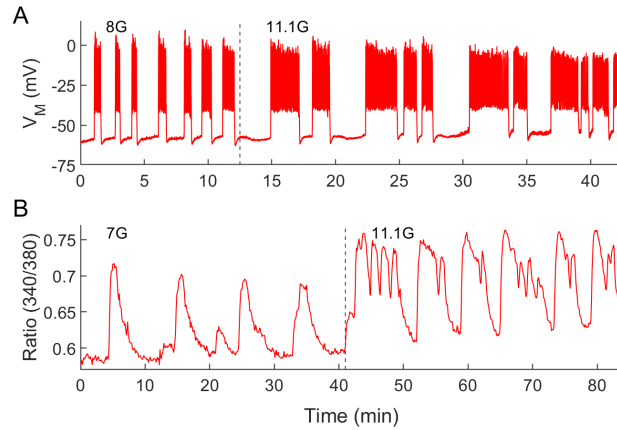


Fig. 7. Compound bursting oscillations are observed in β -PFKM-KO islets. Examples of compound bursting oscillations in membrane potential (A) and Ca^{2+} concentration (B), from two independent experiments.

407 **Compound oscillations also occur in β -PFKM-KO islets.**

408 In addition to the PMO mechanism presented in the previous section, the model can exhibit
 409 active metabolic oscillations (AMO). In this case, the intrinsic, active metabolic
 410 oscillations modulate the bursting activity. We have proposed that the AMO mode is
 411 required to generate compound bursting, which consists of fast bursts of electrical activity
 412 grouped into episodes of typically 3-5 min [38, 39]. Indeed, we have hypothesised that the
 413 slow wave that groups the fast bursts into episodes is due to intrinsic glycolytic oscillations,
 414 driven by the positive feedback of FBP onto PFKM and the subsequent depletion of
 415 substrate F6P [40]. Can this still work if PFKM is knocked out? Figure 7 shows that
 416 compound bursting does occur in β -PFKM-KO islets. Panel A shows a patch-clamp
 417 recording of electrical activity of a β -cell within an islet from a β -PFKM-KO mouse with
 418 several episodes of fast bursts. The number of bursts per episode is highly variable, as is
 419 typical for compound bursting oscillations. Panel B shows the same phenomenon in an
 420 independent recording of Ca^{2+} oscillations. To test whether the IOM with two PFK
 421 isoforms can replicate this finding, we first set model parameters so that the model cell was
 422 in a compound bursting mode prior to simulated β -PFKM knockout (Fig. 8, black trace).
 423 In the wild-type, there are FBP pulses (Fig. 8C) due to pulsatile activity of PFKM (Fig.

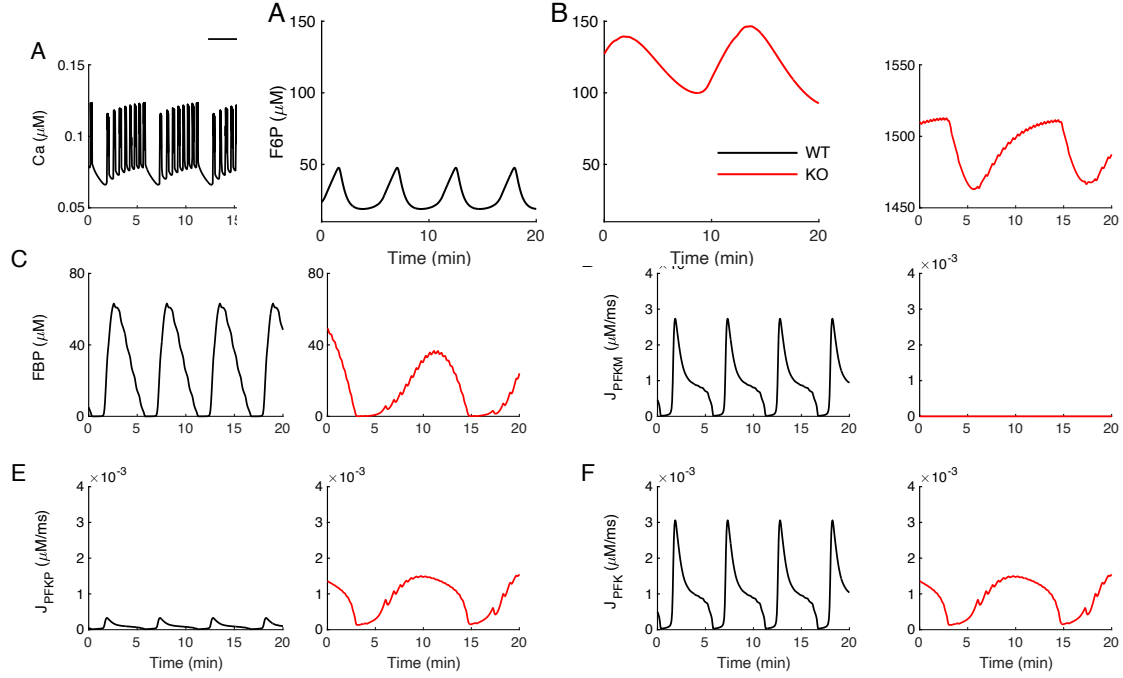


Fig. 8. Simulations showing the mechanism for persistence of compound oscillations in β -PFKM-KO islets. In the model, compound oscillations persist when PFKM is knocked out, due to increased activity of PFKP. (A) The free cytosolic Ca^{2+} concentration exhibits compound oscillations for both wild-type and knockout conditions. The ATP concentration (B) and FBP concentration (C) exhibits slow oscillations before and after the knockout. (D) Metabolic flux through the PFKM enzymatic reaction is eliminated once the PFKM enzyme is knocked out. (E) The metabolic flux through PFKP is very small in the wild-type case, but in the knockout is comparable to the wild-type PFKM flux. (F) Unlike the case of slow bursting (Fig. 5), the total PFK metabolic flux is different in the model wild-type and β -PFKM-KO islet, however, values are comparable. The time courses were generated using $g_{K(Ca)} = 650$ pS and $v_{PDH} = 2 \mu M ms^{-1}$. For the wild-type simulation (black), $v_{PFK-M} = 0.01 \mu M ms^{-1}$ and $v_{PFK-C} = 0.01 \mu M ms^{-1}$. PFKM is knocked out (red) by setting $v_{PFK-M} = 0 \mu M ms^{-1}$. In cases of compound bursting such as shown here, active phases of bursting are short relative to that of the compound oscillation and during the fast phase of activity, fast declines in ATP level due to Ca^{2+} influx drive very brief transient fluctuations, as described in [4].

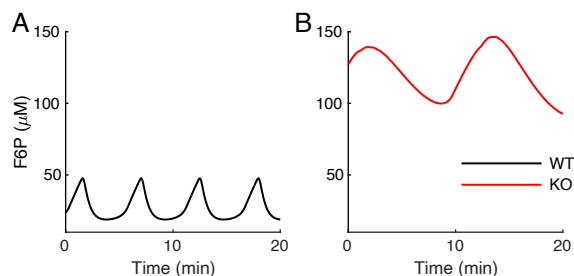


Fig. 9. The mathematical model predicts an increase in the F6P concentration in the PKFM-KO during compound bursting. Both the mean and the amplitude of oscillations in the F6P concentration are lower in the wild-type model islet (A) than in the model β -PKFM-KO islet (B).

8D), and the ATP level oscillates both due to the pulsatile production as well as Ca^{2+} -dependent consumption (Fig. 8B). The oscillation amplitude of PFKP flux is very small in the wild-type condition, but it increases dramatically when PFKM is knocked out (red trace) and PFKP is the only isoform present (Fig. 8E). In this scenario, compound oscillations in Ca^{2+} (Fig. 8A, red trace) and FBP pulses (Fig. 8C) persist. These pulses are now driven entirely by PFKP (Fig. 8E). As in the wild-type, the FBP pulses give rise to oscillations in ATP (Fig. 8B, red trace), which drive the episodes of electrical activity in compound bursting. As with slow bursting (Fig. 5), the P-type isoform of PFK provides the total PFK metabolic flux necessary to drive oscillations (Fig. 8F, red trace).

The PFKP isoform is again capable of rescuing oscillations in spite of its lower affinity for FBP allosteric feedback because the F6P substrate level rises to a much higher level when PFKM is removed (Fig. 9). While this prediction could in theory be measurable experimentally, we are not aware of any experimental approaches that have the requisite sensitivity and dynamics needed to perform this type of experiment at present.

There are profound oscillations in the F6P concentration both before and after removal of PFKM. These reflect the active glycolytic oscillation that drives the slow episodes of compound oscillations. These active glycolytic oscillations produce substrate oscillations

of much greater amplitude than in the case of passive glycolytic oscillations (Fig. 6). We once again note that the only parameter change made in the simulation of the knockout was to set the maximum PFKM flux rate, $v_{\text{PFKM-M}}$, to 0. In particular, no upregulation of PFKP protein is necessary to rescue the compound oscillations.

Discussion

A previous study found that slow oscillations in mouse islets persisted when PFKM levels were reduced using a gene trapping technique [8]. Our study used a gene knockout approach to determine whether oscillations persisted after the complete removal of PFKM (Fig. 1). We showed that oscillations in V_M , Ca^{2+} , and FBP are indeed present in β -PFKM-KO islets (Figs. 2, 3), and therefore do not rely on the M-type isoform. In addition, IPGTT tests performed on β -PFKM-KO mice revealed little or no difference in glucose or insulin responses when compared to wild-type mice (Fig. 4).

Previously, we predicted that oscillations could persist in PFKM knockout islets if the knockout of this isoform was compensated by increased protein expression of a different PFK isoform [13, 34, 41]. Since we showed here that the PFKP expression appeared to be the same in wild-type and knockout islets (Fig. 1), it is clear that a different compensation strategy must be involved.

Our explanation for how PFKP takes over from PFKM once the latter is knocked out is illustrated in the “competition scheme” shown in Fig. 10. Both PFK isoforms compete for the substrate F6P (Fig. 10A), and are subject to allosteric activation by FBP (green arrows) and inhibition by ATP (red arrows). However, because of the different affinities of PFKM and PFKP for FBP and ATP, the effects of activation and inhibition are different for the two isoforms [15, 16]. The M-type isoform has a high affinity for FBP (thick green arrow) and a low affinity for ATP (thin red arrow), while the P-type has low affinity for FBP (thin green arrow) and a high affinity for ATP (thick red arrow). When both isoforms are present (panel A), PFKP loses the competition for the substrate F6P because of the differences in

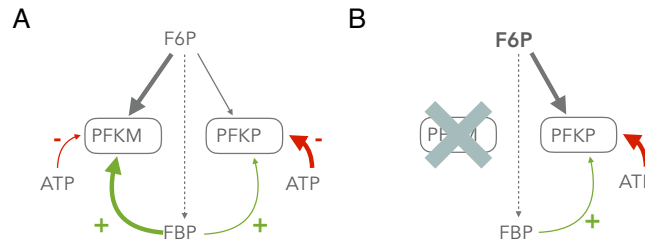


Fig. 10. Proposed mechanism for compensation to β -PFKM knockout. Grey arrows represent substrate flux (thicker arrows indicate greater flux). Other arrows represent positive (green) or negative (red) allosteric regulation (thicker arrows indicate greater affinity). (A) Wild-type. (B) β -PFKM knockout, leading to an increase in the F6P level.

these affinities, and the majority of PFK activity in this case is mediated by PFKM. Conversely, when PFKM is absent (panel B), PFKP is the only competitor for the substrate and thus wins the competition by default. However, since PFKP has weaker allosteric activation and stronger allosteric inhibition than PFKM did, the substrate level must now reach a much higher level (bold font in F6P) to produce the same metabolic flux through the enzyme.

For simplicity we model PFKP here as being modified allosterically by FBP in the same manner as PFKM but using a different parameter for FBP (and ATP) affinity. However, a possible alternative mechanism is that FBP increases PFKP activity instead by stabilizing PFKP in the active tetrameric form. FBP has in fact been shown to stabilize all three human PFK isoforms [42-44]. Having demonstrated the proof of concept in the present paper that PFKP can act as a reserve enzyme that takes over when PFKM is absent, we can explore this alternate mechanism in future versions of the model.

Our model predicts that the metabolic oscillations that occur during the slow bursting oscillations shown in Fig. 2 reflect a PMO in which oscillations in FBP and downstream ATP oscillations reflect the effects of Ca^{2+} on pyruvate dehydrogenase and the Ca^{2+} removal process that uses ATP hydrolysis to power Ca^{2+} pumps. The oscillations in Fig. 5 are generated in this way. This oscillation mechanism is robust to changes in the PFK parameters, so replacing the PFKM isoform with the PFKP isoform has little effect on the

V_M , Ca^{2+} , and FBP oscillations. It does, however, result in an increase in the level of F6P substrate (Fig. 6), and indeed this is an important model prediction whose validation awaits new methods having sufficient sensitivity for measuring the F6P levels of islets.

We hypothesize that compound oscillations, those where both fast and slow oscillations coexist simultaneously are fundamentally different from the slow oscillations, and their generation requires intrinsic glycolytic oscillations. That is, there is an active glycolytic oscillator that packages fast bursts into episodes through the action of ATP on K(ATP) channels. We showed, for the first time, that compound oscillations are produced even when PFKM is removed by genetic knockout (Fig. 7). These data therefore suggest that another PFK isoform is capable of producing the oscillations previously attributed to PFKM alone [6]. Here, we used mathematical modelling to demonstrate that the AMO can indeed be driven by PFKP (Figs. 8, 9). This AMO is, however, more sensitive to the parameter values of PFK than are PMOs. For example, for larger values of the maximum pyruvate dehydrogenase reaction rate, v_{PDH} , and/or lower values of the PFKP reaction rate, v_{PFK-P} , the compound oscillations present in simulated wild-type islets may be lost if PFKM is knocked out (results not shown). If the FBP production is too low (low v_{PFK-P}) or its consumption too high (high v_{PDH}), PFKP cannot sustain the amount of PFK activity necessary to generate intrinsic glycolytic oscillations and compound bursting is replaced by fast bursting. If compound oscillations persist, however, then the model predicts that F6P levels will be elevated in β -PFKM-KO islets relative to wild-type islets. This key prediction is therefore independent of whether oscillations are driven by PMOs or AMOs.

Conclusion

The redundancy of the PFK isoforms expressed in β -cells allows glycolytic and calcium oscillations to persist even if one of the isoforms is not expressed or otherwise active. Thus, despite PFKM being the physiologically most responsive isoform because of its higher affinities for FBP and ATP [5], it is dispensable for the production of oscillations. This molecular redundancy may reflect the importance of maintaining pulsatility in the actions of insulin in the maintenance of glucose homeostasis.

516

517 **Author Contributions**

518 All of the authors contributed to the experimental design, as well as writing and editing the
519 manuscript. VSP, BT, JR, and XT performed the experiments. IM, AS, RB, and PF
520 generated the mathematical models. PF, JH, and AS performed statistical analysis.

521

522 **Declaration of Interests**

523 The authors declare no competing interests

524

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Figure Legend

Fig. 1. β -PFKM-KO mice were generated using CRISPR/Cas9 to flox exon 3 of the *Pfkm* gene (panel A). Employing this method, we were able to selectively delete *Pfkm* mRNA transcript in KO mouse islets, while no statistically significant difference in *Pfklp* mRNA was observed between wild type controls and the KO islets (panel B). Western blot analysis done using a PFKM specific antibody confirmed the loss of PFKM protein in the KO islets, while reduced PFKM protein was evident in islets from heterozygous mice (panel C).

Fig. 2. β -PFKM-KO islets exhibit typical oscillations. Representative examples of oscillations at several glucose levels in control (black) and β -PFKM-KO islets for membrane potential (V_M , panel A), intracellular Ca^{2+} concentration (Ca, panel B), and normalized, detrended PKAR FRET ratio (panel C). Traces are representative of 21 V_M recordings, 10 Ca recordings, and 23 PKAR recordings. Control traces shown were from Cre+ control mice.

Fig. 3. Comparison of oscillation period and plateau fraction between β -PFKM-KO and control islets. Violin plots showing mean oscillation period (top panels) and plateau fraction (bottom panels) for islets exposed to specific glucose levels: 8 mM and 11.1mM glucose for membrane potential (panel A) and Ca^{2+} concentration (panel B), or at 11.1 mM glucose for PKAR (panel C). White dots indicate the median across all islets. All β -PFKM-KO islets oscillated at 8 mM and 11.1 mM glucose. Linear mixed effects modeling (see Material and Methods and Supporting Material) found that the modest reduction in period in β -PFKM-KO compared to control oscillations, was close to but did not achieve statistical significance at the $p=0.05$ level. The differences in plateau fraction were not significant. Control islets consisted of mostly +/+ Cre + islets (59%), with the remaining from WT or fl/fl Cre – mice. As no differences were noted among the controls, the results were pooled.

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673 Fig. 4. In vivo metabolic measurements using intra-peritoneal glucose tolerance tests.
674 Panels A and B, female mice; panels C and D male; panels A and C, glucose, panels B and
675 D, insulin. The only significant difference is glucose for the females ($p = 0.04$) by repeated
676 measures ANOVA. The values at $t = 15$ min and $t = 60$ min are different by unpaired t-
677 test ($p < 0.05$).

678

679 Fig. 5. Simulations showing the mechanism for persistence of slow oscillations in β -
680 PFKM-KO islets. In the IOM, slow oscillations persist when PFKM is knocked out, due to
681 increased activity of PFKP. The black traces are for wild-type conditions (with PFKM
682 present), the red traces are after PFKM knockout. (A) The free cytosolic Ca^{2+} concentration
683 exhibits slow oscillations for both wild-type and knockout conditions. The ATP
684 concentration (B) and FBP concentration (C) exhibits slow oscillations before and after the
685 knockout. (D) Metabolic flux through the PFKM enzymatic reaction is eliminated once the
686 PFKM enzyme is knocked out. (E) The metabolic flux through PFKP is very small in the
687 wild-type case, but in the knockout is comparable to the wild-type PFKM flux. (F) The
688 total PFK metabolic flux is the same before and after the removal of PFKM. The time
689 courses were generated with $g_{\text{K(Ca)}} = 150$ pS and $v_{\text{PDH}} = 0.4 \mu\text{M ms}^{-1}$. For the wild-type
690 simulation (black), $v_{\text{PFK-M}} = 0.01 \mu\text{M ms}^{-1}$ and $v_{\text{PFK-C}} = 0.01 \mu\text{M ms}^{-1}$. PFKM is
691 knocked out (red) by setting $v_{\text{PFK-M}} = 0 \mu\text{M ms}^{-1}$.

692

693 Fig. 6. The mathematical model predicts an increase in the F6P concentration in the β -
694 PKFM-KO during slow bursting. The mean value of the F6P concentration is lower in the
695 wild-type model islet (A) than in the model β -PFKM-KO islet (B).

696

Fig. 7. Compound bursting oscillations are observed in β -PFKM-KO islets. Examples of compound bursting oscillations in membrane potential (A) and Ca^{2+} concentration (B), from two independent experiments.

Fig. 8. Simulations showing the mechanism for persistence of compound oscillations in β -PFKM-KO islets. In the model, compound oscillations persist when PFKM is knocked out, due to increased activity of PFKP. (A) The free cytosolic Ca^{2+} concentration exhibits compound oscillations for both wild-type and knockout conditions. The ATP concentration (B) and FBP concentration (C) exhibits slow oscillations before and after the knockout. (D) Metabolic flux through the PFKM enzymatic reaction is eliminated once the PFKM enzyme is knocked out. (E) The metabolic flux through PFKP is very small in the wild-type case, but in the knockout is comparable to the wild-type PFKM flux. (F) Unlike the case of slow bursting (Fig. 5), the total PFK metabolic flux is different in the model wild-type and β -PFKM-KO islet, however, values are comparable. The time courses were generated using $g_{\text{K}(\text{Ca})} = 650 \text{ pS}$ and $v_{\text{PDH}} = 2 \text{ } \mu\text{M ms}^{-1}$. For the wild-type simulation (black), $v_{\text{PFK-M}} = 0.01 \text{ } \mu\text{M ms}^{-1}$ and $v_{\text{PFK-C}} = 0.01 \text{ } \mu\text{M ms}^{-1}$. PFKM is knocked out (red) by setting $v_{\text{PFK-M}} = 0 \text{ } \mu\text{M ms}^{-1}$. In cases of compound bursting such as shown here, active phases of bursting are short relative to that of the compound oscillation and during the fast phase of activity, fast declines in ATP level due to Ca^{2+} influx drive very brief transient fluctuations, as described in [4].

Fig. 9. The mathematical model predicts an increase in the F6P concentration in the PKFM-KO during compound bursting. Both the mean and the amplitude of oscillations in the F6P concentration are lower in the wild-type model islet (A) than in the model β -PFKM-KO islet (B).

723 Fig. 10. Proposed mechanism for compensation to β -PFKM knockout. Grey arrows
724 represent substrate flux (thicker arrows indicate greater flux). Other arrows represent
725 positive (green) or negative (red) allosteric regulation (thicker arrows indicate greater
726 affinity). (A) Wild-type. (B) β -PFKM knockout, leading to an increase in the F6P level.