Glycolysis–Wnt signaling axis tunes developmental timing of embryo segmentation

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Abstract

The question of how metabolism impacts development is seeing a renaissance [1, 2]. How metabolism exerts instructive signaling functions is one of the central issues that need to be resolved. We tackled this question in the context of mouse embryonic axis segmentation. Previous studies have shown that changes in central carbon metabolism impact Wnt signaling [3-6] and the period of the segmentation clock [7], which controls the timing of axis segmentation. Here, we reveal that glycolysis tunes the segmentation clock period in an anti-correlated manner: higher glycolytic flux slows down the clock, and vice versa. Transcriptome and gene regulatory network analyses identified Wnt signaling and specifically the transcription factor Tcf7l2, previously associated with increased risk for diabetes [8, 9], as potential mechanisms underlying flux-dependent control of the clock period. Critically, we show that deletion of the Wnt antagonist Dkk1 rescued the slow segmentation clock phenotype caused by increased glycolysis, demonstrating that glycolysis instructs Wnt signaling to control the clock period. In addition, we demonstrate metabolic entrainment of the segmentation clock: periodic changes in the levels of glucose or glycolytic sentinel metabolite fructose 1,6-bisphosphate (FBP) synchronize signaling oscillations. Notably, periodic FBP pulses first entrained Wnt signaling oscillations and subsequently Notch signaling oscillations. We hence conclude that metabolic entrainment has an immediate, specific effect on Wnt signaling. Combined, our work identifies a glycolysis-FBP-Wnt signaling axis that tunes developmental timing, highlighting the instructive signaling role of metabolism in embryonic development.

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1 Introduction

Central carbon metabolism impacts gene expres-2 sion and signal transduction via modulating epigenetic and protein post-translational modifications, while exerting its bioenergetic function by producing energy, reducing equivalents, and cellular building blocks to fuel biological processes [1, 2, 10–14]. While such a widespread role of metabolism is well-known, how metabolism acts as an instructive rather than a permissive signal 10 to control phenotypic outcomes remains a central 11 question. In the definition we use, an instructive 12 signal is information-rich, hence having the capa-13 bility of tuning a phenotypic outcome, as opposed 14 15 to a permissive signal leading to a binary effect [15, 16].16

To reveal tunability, it is crucial to be able 17 to tune metabolism dynamically and to monitor 18 its impact, for instance at the level of signaling, 19 in real time and in a quantitative manner. Such 20 an approach is applicable to the study of verte-21 brate embryo mesoderm segmentation. Presomitic 22 mesoderm (PSM) is segmented into somites, the 23 precursors for vertebrae and skeletal muscles, in 24 a periodic fashion [17]. The timing of this pro-25 cess is tightly regulated by a molecular oscilla-26 tor known as the segmentation clock, which is 27 best characterised by oscillatory activity of the 28 Notch signaling pathway [18]. Temporal period-29 icity of Notch signaling oscillations is translated 30 into spatial periodicity of somites by integrating 31 additional information encoded by graded signal-32 33 ing pathways such as Wnt, FGF, and retinoic acid [17, 19–21]. In the mouse PSM, FGF and 34 Wnt signaling pathways are also the components 35 of the segmentation clock, exhibiting oscillatory 36 activities coupled to Notch signaling oscillations 37 [19, 22, 23]. Importantly, this highly complex net-38 work of interconnected signaling pathways can 39 be dynamically perturbed and functionally stud-40 ied by using a combination of quantitative live 41 imaging and a dynamical systems approach. For 42 instance, using microfluidics-based entrainment, 43 we previously showed that the segmentation clock 44 network can be efficiently controlled via external 45 periodic pulses of Notch and Wnt signaling cues, 46 achieving synchronization and tuning of signaling 47 oscillation period [19, 24]. 48

Here, we build on the quantitative live imaging, genetics, and entrainment approach that provide a powerful experimental framework to tackle the central question of how metabolism plays an instructive role. In the PSM, changes in central carbon metabolism impact Wnt signaling [3–6] and the period of the segmentation clock [7]. In particular, glycolysis has been shown to establish an activity gradient from the posterior to anterior PSM [3, 4], being functionally linked to graded signaling activity within the mouse PSM [4-6]. Furthermore, it has been shown that active glycolysis is required for maintaining the segmentation clock oscillation [3]. In this work we addressed whether and how glycolysis plays an instructive role in regulation of developmental timing of mammalian embryo segmentation.

2 Results

2.1 Glycolytic flux tunes the period of the segmentation clock

We first asked whether changes in glycolytic flux would have any effect on the segmentation clock period. To manipulate glycolytic flux using genetics, we utilised a conditional cytoPFKFB3 (hereafter termed as TG) transgenic mouse line that we previously generated [6]. In this TG line, a cytoplasmic, dominant active form of the glycolytic enzyme PFKFB3 [25] is expressed from the Rosa26 locus upon CRE-recombination, leading to a glucose-dose dependent increase of glycolytic flux in PSM explants [6]. To quantify the segmentation clock period using real-time imaging, we used a fluorescent reporter mouse line, which reflects the oscillatory gene activity of Notchtarget gene Lfng [26].

Using this experimental strategy, we found that in TG explants cultured in 2.0 mM glucose, the segmentation clock slowed down by about 20% compared to control explants, without arrest of segmentation clock or morphological segmentation defects (Fig. 1A, 1B, Supplementary Video 1). The slowing down of segmentation clock oscillations was also evident when using a Wnt reporter line, i.e., *Axin2-Achilles* knock-in reporter [24] (Extended Data Fig. 1, Supplementary Video 2). To test whether the observed effect on the segmentation clock oscillations is indeed due to an increased glycolytic flux, and not merely the effect

of the overexpression of cytoPFKFB3 protein per 145 97 se, we cultured TG explants in reduced glucose 146 98 concentrations, in order to reduce glycolytic flux 147 99 (Extended Data Fig. 2A). Indeed, lowering glucose 148 100 concentration rescued the clock period phenotype 149 101 in TG explants (Fig. 1B), indicating that the seg- 150 102 mentation clock period responds to glycolytic flux 151 103 rather than cytoPFKFB3 protein per se. 152 104

To further probe whether glycolytic flux 153 105 instructs the segmentation clock period, we inves-154 106 tigated the impact of tuning (i.e., increasing and 155 107 decreasing) glycolytic flux in wild-type explants. 156 108 Importantly, we found that the segmentation clock 157 109 period was tunable in wild-type explants by mod- 158 110 ulating glycolytic flux. Increasing glucose led to 159 111 a slower segmentation clock (Fig. 1C), which 160 112 we also observed when fructose 1.6-bisphosphate 161 113 (FBP), a glycolytic sentinel metabolite [6, 27], 162 114 was supplemented to the medium (Fig. 1D). On 163 115 the other hand, replacing glucose with galactose, 164 116 which leads to minimum glycolytic flux (Extended 165 117 Data Fig. 2B) [28], resulted in the acceleration 166 118 of the segmentation clock (Fig. 1D). Therefore, 167 119 minimizing glycolytic flux speeds up the segmen- 168 120 tation clock, while increasing glycolytic flux has 169 121 the opposite effect. 170 122

Taken together, our data shows that glycolytic 171 flux tunes the segmentation clock period in an 172 anti-correlated manner. 173

2.2 Characterizing glycolytic flux-induced transcriptional responses in PSM cells

To gain insight into the mechanism underlying the glycolytic flux-dependent control of the segmentation clock period, we next looked into flux-induced transcriptional responses and their potential mechanisms operating in the PSM. 183

we built PSM-specific enhancer- 184 134 First. mediated gene-regulatory network (eGRN) using 185 135 the GRaNIE (Gene Regulatory Network Infer- 186 136 ence including Enhancers) method [29], which 187 137 constructs eGRN based on co-variation of chro- 188 138 matin [i.e., transcription factor (TF) binding site] 189 139 accessibility, TF expression and corresponding 190 140 target gene expression across samples. We gener- 191 141 ated paired transcriptome [i.e., RNA sequencing 192 142 (RNA-seq)] and chromatin accessibility [i.e., 193 143 assay for transposase-accessible chromatin with 194 144

sequence (ATAC-seq)] data from wild-type, noncultured PSM tissues. The PSM tissues were microdissected into tailbud, posterior PSM, anterior PSM, and somite regions, so that a resulting eGRN is linked to gene expression changes following PSM cell differentiation along the embryonic axis, which also mirrors metabolic state changes [3, 4] (Extended Data Fig. 3A).

The resulting eGRN includes 2522 genes out of 28629 (= 9%) genes expressed in the PSM and consists of 69 regulons, where each regulon represents a set of target genes regulated by a TF through their accessible enhancer regions (Extended Data Fig. 3A). These regulons include those associated with TFs that regulate PSM cell differentiation, such as Cdx2 [30] and T [31], providing evidence for the validity of the PSM-specific eGRN inferred with the GRaNIE method.

For the identification of glycolytic fluxresponsive genes, we performed transcriptome analysis using explants from control and TG explants cultured in different glucose concentrations for three hours. We limited our analysis to the tailbud region, where the clock period phenotype is most apparent. Combined with the dataset from our previous study [6], this analysis revealed 617 flux-responsive differentially expressed genes (DEGs) that were either upregulated (Cluster (C) 1 and C3) or downregulated (C2, C4, and C5) by increasing glycolytic flux (Fig. 2A, Supplementary Table 1).

By matching the flux-responsive DEGs to the PSM-specific eGRN, we revealed that 132 DEGs are part of the regulons. Intriguingly, the vast majority of those (90 out of 132 DEGs) are part of the Tcf7l2 regulon (Fig. 2B, Extended Data Fig. 3B). Gene expression of the Tcf7l2 regulon is downregulated with both increased glycolytic flux and FBP supplementation (Fig. 2C, Extended Data Fig. 3C), conditions that we found to cause slowing down of the segmentation clock (Fig. 1).

Tcf7l2 is tightly linked to Wnt signaling [32, 33], and identified as a repressor in our eGRN analysis (Extended Data Fig. 3A). Therefore, these results reveal a glycolysis-Wnt-signaling axis where increased glycolytic flux activates the Tcf7l2 regulon, providing the mechanistic basis for the anti-correlation between glycolytic flux and Wnt signaling target gene expression. Functionally, the glycolysis-Wnt-signaling axis could hence

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Fig. 1 Glycolytic flux tunes the segmentation clock period in an anti-correlated manner. (A) Kymographs showing the dynamics of the Notch signaling reporter (= LuVeLu [26]) in control (Ctrl) and cytoPFKFB3 (TG) PSM explants in 2.0 mM glucose condition. (B-D) Quantification of the segmentation clock period in various metabolic conditions. The clock periods were determined as a mean of LuVeLu periods between 400-600 min of the imaging. Since the clock period is highly sensitive to temperature, the comparisons are always made within each experiment. (B) The clock period in VI explants cultured in 0.5 mM or 2.0 mM glucose. (C) Effects of glucose titration on the clock period in wild-type explants [0.5 mM (0.5G) vs. 2.0 mM (2.0G) vs. 10 mM (10G) glucose]. (D) Effects of fructose 1,6-bisphosphate (FBP) or galactose (GALA) on the clock period in wild-type explants [CTRL, culture medium with 2.0 mM glucose; FBP, culture medium with 2.0 mM glucose and 10 mM FBP; GALA, culture medium with 2.0 mM glactose (without glucose)]. One-way ANOVA with Tukey's post hoc test (*p <0.05, **p <0.01, ***p <0.001). Mean \pm standard deviation (SD) are shown in the graph, and individual data points represent biological replicates.

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underlie the observed tuning of segmentation clock 204
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¹⁹⁷ 2.3 Glycolysis-Wnt signaling axis ¹⁹⁸ controls the segmentation clock ¹⁹⁹ period

To functionally test whether the glycolysis-Wnt- ²¹¹ signaling axis underlies the flux-dependent tuning ²¹² of the segmentation clock period, we performed ²¹³ a genetic rescue experiment using a mutant for ²¹⁴

Dickkopf-1 (Dkk1) [34, 35], a developmentally critical Wnt signaling inhibitor that acts at the level of ligand-receptor interaction. We asked whether partial deletion of Dkk1 could rescue the clock period phenotype observed in TG embryos, where elevated glycolytic flux correlated with Wnt signaling downregulation. Excitingly, we indeed found that in TG embryos in which one allele of Dkk1 was deleted, the segmentation clock period was rescued in most of the samples (Fig. 3A). Critically, we found that lactate secretion was not



Fig. 2 Tcf7l2 regulon responds to glycolytic flux changes within PSM cells. (A) A heatmap showing glycolytic flux-responsive differentially expressed genes (DEGs) between wild-type (WT) and cytoPFKFB3 (TG) PSM explants cultured for three-hour in various (i.e., 0.5 mM, 2.0 mM, and 10 mM) glucose conditions (adjusted *p*-value <0.01, WT vs. TG for each glucose condition). Normalized counts by variance stabilizing transformation (VST) were used to calculate the z-scores. The datasets were integrated with the datasets from Miyazawa *et al.* (2022) [6]. DEGs that are parts of the PSM-specific eGRN are marked by green. (B) A table showing the number of the flux-responsive DEGs that are included in each PSM-specific regulon. (C) A box plot showing fold changes in gene expression of the flux-responsive Tcf7l2 targets between different metabolic conditions.

affected by Dkk1 heterozygosity (Fig. 3B). TG 228 215 explants maintained high glycolytic flux even in a 229 216 Dkk1 heterozygous background, despite showing 230 217 a rescued clock phenotype. These findings indi- 231 218 cate that the proximate cause of the observed 232 219 clock phenotype in TG embryos are changes in 233 220 signaling, rather than cellular metabolic state. 234 221 To further probe the mechanism underlying 235

To further probe the mechanism underlying ²³⁵ the clock period phenotype, we also examined ²³⁶ whether there is a correlation between cellular ²³⁷ redox state and the segmentation clock period, ²³⁸ as recently suggested in an embryonic stem cell ²³⁹ (ESC)-based model for the segmentation clock [7]. ²⁴⁰ To this end, we quantified the NAD⁺/NADH ratio in control and TG explants under different culture conditions. As expected, the NAD⁺/NADH ratio changed in response to alterations in glycolytic flux (Extended Data Fig. 4). Importantly however, the NAD⁺/NADH ratio was comparable between control explants cultured in 10 mM glucose and TG explants cultured in 2.0 mM glucose (Extended Data Fig. 4), which showed a significant difference in the segmentation clock period (Fig. 1).

Taken together, these data provide strong evidence that the tuning effect of glycolytic flux on 292

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the segmentation clock period is not mediated via 289
changes in cellular bioenergetic state but rather, 290
via modulation of Wnt signaling. 291

244 2.4 Metabolic entrainment of the 245 segmentation clock

To further investigate how glycolytic flux is linked 246 to oscillatory signaling and the segmentation 247 clock, we used a dynamical systems approach 248 208 based on entrainment. Entrainment offers a quan-249 200 titative and non-disruptive approach to reveal 250 300 functional dependencies within a dynamical sys-251 301 tem. We had previously established microfluidics-252 302 based entrainment of the mouse embryo segmen-253 tation clock, using periodic pulses of signaling 304 254 pathway modulators, such as a Notch signaling 305 255 inhibitor and a Wnt signaling activator [19, 24]. 306 256 Based on our finding of a functional glycolysis-257 Wnt-signaling axis, we wondered whether the seg-258 mentation clock network could also be entrained 259 300 by periodic changes in glycolytic flux. 260

As glycolytic flux in PSM cells can be con-₃₁₁ 261 trolled via the concentration of glucose in the $_{312}$ 262 culture medium (Extended Data Fig. 2), we 313 263 used microfluidics to implement periodic changes 314 264 in glucose concentration during the culture of 315 265 PSM explants and monitored segmentation clock ₃₁₆ 266 dynamics using real-time imaging of a Notch 317 267 signaling reporter. Strikingly, we found that peri- $_{318}$ 268 odic alternations of glycolytic flux are indeed 319 269 sufficient to entrain Notch signaling oscillations 320 270 underlying the segmentation clock (Fig. 4A, 4A', 321 271 4B, 4B', Supplementary Video 3). We quanti-₃₂₂ 272 fied entrainment based on phase-locking (Fig. 4A', 323 273 4B', Extended Data Fig. 5B) and also using the 324 274 first Kuramoto order parameter (Fig. 4A, 4B), 325 275 which effectively measures how synchronous dif-276 ferent samples are oscillating. 277

In addition to periodic changes in glucose, we 278 also tested whether periodic pulses of the sentinel 279 metabolite FBP would be sufficient to entrain the 280 segmentation clock. Indeed, our results revealed $^{\scriptscriptstyle 328}$ 281 evidence for Notch signaling entrainment by peri- 329 282 odic application of FBP (Fig. 4C, 4C', Extended 330 283 Data Fig. 5B, Supplementary Video 4). In con- 331 284 trast, periodic application of pyruvate, the end 332 285 product of glycolysis, was not sufficient to entrain 333 286 the segmentation clock (Extended Data Fig. 5A, 334 287 5A', 5B, Supplementary Video 5). These results 335 288

show that transient, periodic perturbations of glycolysis, specifically at the level of the sentinel metabolite FBP, can entrain the segmentation clock. This provides additional, independent support for glycolytic flux-signaling closely linked to developmental timing.

Importantly, we used metabolic entrainment to further disentangle the functional dependencies between glycolysis, Wnt and Notch signaling pathways. Do periodic FBP pulses entrain Wnt signaling directly or indirectly through Notch signaling entrainment? We previously had shown that Wnt and Notch signaling oscillations are coupled within the segmentation clock network [19]. This means that entrainment of Notch signaling oscillations eventually leads to entrainment of Wnt signalling oscillations with a time delay, and vice versa. Thus, we next quantified the timing of metabolic entrainment in regard to both Notch and Wnt signaling oscillations, in order to distinguish direct from more indirect dependencies between glycolysis and Wnt signaling. Notably, we found that periodic FBP pulses first entrained Wnt signaling oscillations, while entrainment of Notch signaling oscillations followed with considerable delay (Fig. 4C, 4D, Supplementary Video 6). Hence, this dynamic entrainment analysis provides strong evidence that glycolysis/FBP has a direct effect on Wnt signaling within the segmentation clock network.

Combined, we show for the first time metabolic entrainment of the segmentation clock, which further establishes a signaling function of glycolysis. Moreover, our analysis of entrainment dynamics supports a specific, direct functional connection of glycolytic flux-signaling to the Wnt signaling pathway.

3 Discussion

3.1 Glycolysis-FBP-Wnt signaling axis within the PSM.

In this study, we show that glycolytic flux tunes the timing of axis segmentation through its instructive function on Wnt signaling. This is supported by our finding that in conditions of increased glycolytic flux, the partial deletion of Dkk1 rescued the segmentation clock period (Fig. 3). Previously, several mechanisms have been proposed regarding how glucose metabolism impacts



Fig. 3 Genetic rescue of the slow segmentation clock phenotype in cytoPFKFB3 embryos without affecting glycolytic flux. (A) Quantification of the segmentation clock period in control (Ctrl) and cytoPFKFB3 (TG) explants with one allele of Dkk1 (HET), compared to samples with wild-type Dkk1 copy number (WT). The clock period under 2.0 mM glucose condition was determined as a mean of LuVeLu periods between 400-600 min of imaging. One-way ANOVA with Tukey's post hoc test (*p <0.05, **p <0.01, ***p <0.001). Mean \pm SD are shown in the graph, and individual data points represent biological replicates. (B) Lactate secretion was quantified as a proxy for glycolytic flux within PSM cells. After 12 h *ex vivo* culture in 2.0 mM glucose, the amount of lactate secreted from control (Ctrl) and cytoPFKFB3 (TG) PSM explants was quantified in wild type (WT) samples with normal Dkk1 copy number and in samples with one allele of Dkk1 (HET). Welch's unpaired t-test (n.s., not significant). Mean \pm SD are shown in the graph, and individual data points represent biological replicates.

Wnt signaling via post-translational modifications 362 337 [5, 36, 37]. Our results presented here reveal 363 338 a key signaling role for the glycolytic sentinel 364 339 metabolite FBP. We propose that the 'glycolysis- 365 340 FBP-Wnt signaling axis' is a module that connects 366 341 metabolism, signaling and developmental timing. 367 342 More specifically, combined with our previous 368 343 study [6], we provide in vivo evidence that glycol- 369 344 ysis controls Wnt signaling in a dose-dependent, 370 345 anti-correlated manner (Fig. 1-3). Hence, while 371 346 increasing glycolytic flux leads to a decrease in 372 347 Wnt-signaling target gene expression and a slow- 373 348 ing down of segmentation, we also see evidence 374 349 for the inverse: decreasing glycolytic flux within 375 350 a physiological range correlates with increased 376 351 Wnt target gene expression and accelerated seg- 377 352 mentation. Furthermore, we showed that periodic 378 353 application of FBP first synchronizes Wnt signal- 379 354 ing oscillations and subsequently Notch signaling 380 355 oscillations during metabolic entrainment (Fig. 4). 381 356 These findings indicate that glycolytic flux, or its 382 357 dynamics, tunes Wnt signaling activity to control 383 358 the timing of the segmentation clock. 359 384

These results appear to contrast with findings in studies using *in vitro* stem cell models for 386 mesoderm specification, in which glycolytic inhibition led to downregulation, not upregulation, of Wnt signaling [4, 5, 38-40]. One potential reason for this apparent discrepancy could be rooted in the strength of perturbation applied to glycolytic flux. In the studies mentioned above, glycolysis was either strongly impaired pharmacologically or bypassed altogether (i.e., no glucose condition), which caused downregulation of Wnt (and other signaling) activity. This indeed shows that ongoing glycolysis is required, permissively, for signaling. In contrast, we show that tuning glycolytic flux within the physiological range, both lowering and increasing flux, leads to an anticorrelated response at the level of Wnt signaling targets and segmentation clock period. Combined, the available evidence hence suggest the existence of multiple functional dependencies between glycolvsis and signaling. First, a permissive glycolvtic function for signaling is evident, i.e., some glycolytic activity is *per se* required. In addition, we show here that an instructive, tunable glycolysis-FBP-Wnt signaling axis exists, controlling the period of the segmentation clock in vivo. Future mechanistic studies will further resolve both the permissive and instructive glycolytic function in different contexts.

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Fig. 4 Metabolic entrainment of the segmentation clock. (A-D) Detrended (via sinc-filter detrending, cut-off period = 240 min) time-series of LuVeLu (A-C) and Axin2-Achilles (D) intensity oscillations in wild-type PSM explants during metabolic entrainment (dashed lines: individual samples, bold black line: median values, grey shades: the first to third quartile range). Changes in the first Kuramoto order parameter are shown in magenta. Samples were incubated either in a constant (i.e., 2.0 mM) glucose condition with periodic mock pulses (gray) (A) or alternating culture conditions (B-D) with a period of 140-min and a pulse length of 30-min [alternating between: (B) 2.0 mM (white) to 0.5 mM (yellow) glucose conditions; (C, D) the medium with (cyan) or without (white) 20 mM FBP on top of 2.0 mM glucose]. To keep molarity of the medium at constant during experiments, non-metabolizable glucose (i.e., 3-O-methyl-glucose) was added to the medium when necessary. (A'-D') Stroboscopic maps showing step-wise changes in the phase of LuVeLu (A'-C') and Axin2-Achilles oscillations (D') during metabolic entrainment. At each pulse of metabolic perturbations with glucose (A', B') or FBP (C', D'), the phase of the oscillator (i.e., new phase) is plotted against its phase at the previous pulse (i.e., old phase). Darker dots represent later time points. Stroboscopic maps of a single representative sample are shown on the right (the numbers in the plots indicate the number of the pulses).

Intriguingly, during metabolic entrainment, we
noticed that periodic changes in glycolytic flux
and FBP levels induce periodic changes in tissue
shape (Supplementary Video 3-6). This suggests a
potential additional link between glycolysis, Wnt
signaling, and tissue shape changes. Importantly,

however, periodic pulse of pyruvate also caused a similar shape change phenotype but did not result in segmentation clock entrainment. While we therefore conclude that tissue shape changes are not sufficient for entrainment, their link to

400 metabolic signaling needs to be a focus of future 449
401 studies. 450

To reveal the detailed mechanisms of glycolytic 451 402 flux signaling, it will be crucial to identify FBP 452 403 sensor molecules that mechanistically link intra- 453 404 cellular FBP levels and Wnt signaling. Probing 454 405 FBP-protein interactions is one exciting direction 455 406 that in different contexts have already indicated 456 407 the widespread regulatory potential of FBP [41–457 408 44]. In addition, our transcriptome and eGRN 458 409 analyses identified genes within the Tcf7l2 regulon 459 410 as particularly glycolytic flux-sensitive (Fig. 2). 460 411 This raises the possibility that Tcf7l2 is a part of 461 412 the FBP sensing mechanisms and hence FBP-Wnt 462 413 signaling axis. Notably, Tcf7l2 has been strongly 463 414 associated with type 2 diabetes and is involved in 464 415 gluocse homeostatis and insulin secretion in pan-465 416 creatic β -cells [8, 9]. An exciting possibility that 466 417 requires further investigation is that FBP directly $_{\rm 467}$ 418 impacts Tcf7l2 activity in an allosteric manner 468 419 within the segmentation clock network but poten- 469 420 tially also in other biological contexts including 470 421 pancreatic β -cells. 422

3.2 Glycolytic flux control of the segmentation clock period.

The primary function of the glycolysis-FBP-Wnt 474 425 signaling axis that we revealed in this study is 475 426 the control of segmentation clock period in mouse 476 427 embryos. Previously, Wnt signaling had been func- 477 428 tionally linked to the regulation of the segmen- 478 429 tation clock period [45], although the underly- 479 430 ing mechanisms were not addressed. Our work 480 431 reveals the direct impact of metabolic state on 481 432 Wnt-signaling and clock period and hence empha- 482 433 sises the need for future studies to identify how 483 434 Wnt signaling impacts the period of segmenta-484 435 tion clock oscillations. Recently, a series of studies 485 436 have reported on potential mechanisms of how, 486 437 in general, the oscillation period can be tuned. 487 438 Accordingly, a study using in vitro stem cell sys-439 tem reported that the segmentation clock period 489 440 is controlled by mitochondrial respiration, cellu- 490 441 lar redox state, and ultimately protein translation ⁴⁹¹ 442 rate [7]. Additionally, several in vitro studies 492 443 emphasized that differences in protein turnover 493 444 rates underlie species-specific developmental tim- 494 445 ing [7, 46–49]. How our *in vivo* findings on the 495 446 link of glycolysis, Wnt signaling and develop-496 447 mental timing relate to these in vitro studies is 497 448

not resolved yet. In principle, our finding that increased glycolytic flux leads to a slowing down of the segmentation clock is compatible with a role of mitochondrial respiration, since glycolysis and respiration are considered to be inversely correlated (i.e., Crabtree effect). However, we found that glycolytic flux-signaling shows specificity at the level of FBP, as periodic pulses of pyruvate are not sufficient to entrain the segmentation clock, which could argue against an involvement of mitochondrial respiration. In addition, our findings revealed that glycolysis functions via Wnt signaling (Fig. 3), and not via cellular redox state (Extended Data Fig. 4). We also found clear evidence for a direct immediate effect of FBP on Wnt signaling using metabolic entrainment (Fig. 4). Combined, our findings hence argue against a widespread, bioenergetic mechanism. Instead we identifies a non-bioenergetic metabolic signaling role and reveals the glycolysis-FBP-Wnt signaling axis as a regulator of the segmentation clock period.

3.3 Future direction

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In conclusion, our study provides evidence that glycolysis is instructive in regulation of Wnt signaling. This regulatory function is crucial for controlling developmental timing and potentially embryonic patterning. The association between glycolysis and Wnt signaling in many biological contexts, ranging from development to disease states [9, 50–52], underscores the critical need to now explore how ubiquitously the glycolysis-FBP-Wnt signaling axis functions in living systems.

These findings also raise the more general question about the significance of the functional link between metabolic activity and developmental timing. We discuss here two, potentially interconnected, hypotheses regarding the broader implications of this relationship.

One appealing hypothesis is that the intrinsic temporal organization of metabolism, which manifests as metabolic rhythms and cycles at various temporal and spatial scales [53], serves as the core template for biological timing and oscillations [54]. In this study, we provide the first demonstration that if present, metabolic cycles (in our case experimentally generated via entrainment) can potently entrain the segmentation clock and developmental timing. Thus, as a next logical step, efforts

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need to be intensified to elucidate the presence of 542 498 metabolic rhythms and cycles in living systems. 543 499 In addition, the link between metabolism, 544 500 developmental signaling and timing could serve 545 501 the integration of environmental cues, such as 546 502 changes in nutritional resources. Interestingly, we 547 503 show that access to higher glucose concentrations 548 504 slows down the pace of embryonic segmentation. 549 505 In order to understand the significance of this 550 506 functional dependence between metabolism, sig- 551 507 naling and timing, it will be critical to study the 552 508 dynamic interplay of organisms with their natural 553 509 environment, considering the entire life cycle. 554 510

511 4 Methods

512 4.1 Animal work

All animals were housed in the EMBL animal 558 513 facility under veterinarians' supervision and were 559 514 treated following the guidelines of the European 560 515 Commission, revised directive 2010/63/EU and 561 516 AVMA guidelines 2007. All the animal experi-517 ments were approved by the EMBL Institutional 563 518 Animal Care and Use Committee (project code: 564 519 21–001_HD_AA). The detection of a vaginal plug $_{565}$ 520 was designated as embryonic day (E) 0.5, and all $_{566}$ 521 experiments were conducted with E10.5 embryos. 567 522

523 4.2 Mouse lines

570 The following mice used in this study were 524 571 described previously and were genotyped 525 572 using primers described in these references: 526 Axin2-Achilles [24], Hprt^{Cre} [55], LuVeLu [26], 527 Rosa26^{loxP-stop-loxP-cytoPFKFB3}[6], Dkk1 mutant 573 528 [35]. While the Dkk1 mutant line was maintained 574 529 on C57BL/6j genetic background, the other 530 575 mouse lines were maintained on CD1 genetic 531 576 background. For the genetic rescue experiments, 532 the following primers were used to detect the 533 578 mutant allele of Dkk1 [56]: forward, 5'-GCT 534 579 CTA ATG CTC TAG TGC TCT AGT GAC- 3'. 535 580 Reverse, 5'-GTA GAA TTG ACC TGC AGG 536 581 GGC CCT CGA-3'. 537 582

⁵³⁸ 4.3 *Ex vivo* culture of PSM explants

⁵³⁹ Dissection and *ex vivo* culture of PSM explants ⁵⁸⁶ ⁵⁴⁰ were performed as described before [6]. In brief, ⁵⁸⁶ ⁵⁴¹ E10.5 embryos were collected in DMEM/F12 ⁵⁸⁷ (without glucose, pyruvate, glutamine, and phenol red; Cell Culture Technologies) supplemented with 2.0 mM glucose (Sigma-Aldrich, G8769), 2.0 mM glutamine (Sigma-Aldrich, G7513), 1.0% (w/v) BSA (Cohn fraction V; Equitech-Bio, BAC62), and 10 mM HEPES (Gibco, 15360–106). PSM explants were isolated using a micro scalpel (Feather Safety Razor, No. 715, 02.003.00.715) and were cultured in DMEM/F12 supplemented with 0.5–2.0 mM glucose, 2.0 mM glutamine, and 1.0% (w/v) BSA (Cohn fraction V; Equitech-Bio, BAC62) at 37°C, under 5% CO₂, 60% O₂ condition.

4.4 Live imaging of Notch and Wnt signaling reporter lines

To monitor Notch and Wnt signaling activity using real-time imaging, LuVeLu [26] and Axin2-Achilles knock-in [24] reporter lines were utilized, respectively. Following dissection, PSM explants were washed once with culture medium and were transferred into agar wells (600 nm-width, 3% low Tm agarose, Biozyme, 840101) in 4-well slides (Lab-Tek, #155383). Imaging was performed with a LSM780 laser-scanning microscope (Zeiss), at $37^{\circ}C$, under 5% CO₂, 65% O₂ condition. Samples were excited by a 514 nm-wavelength argon laser through 20×Plan-Apochromat objective (numerical aperture 0.8). Image processing was done using the Fiji software [57]. For extracting period and phase of signaling oscillations, wavelet analysis was performed using pyBOAT [58].

4.5 NAD⁺/NADH and lactate measurements

PSM explants without somites were cultured for one hour in DMEM/F12 supplemented with varying amounts of glucose or galactose (Sigma, G0750). The explants were flash frozen by liquid N₂ following one hour *ex vivo* culture and were stored at -80° C until use. NAD⁺/NADH measurements were performed according to the manufacturer's instructions (Promega, G9071). In brief, eight explants were lysed in 90 µl of 0.1N NaOH with 0.5% DTAB and were split into two tubes (40 µl per tube). Samples were then incubated at 60°C for 15 min with or without adding 20 µl of 0.4N HCl for NAD⁺ and NADH measurements, respectively. After neutralisation either by

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0.5M Trizma base solution (for NAD⁺ samples) 638
or Trizma-HCl solution (for NADH samples), the 639
lysates were used for NAD⁺/NADH measure- 640
ments. Lactate measurements were performed as 641
described before [6]. 642

4.6 ATAC- and RNA-sequencing analysis

PSM explants of E10.5 wild-type embryos (CD1 596 647 genetic background) were microdissected into 648 597 tail bud, posterior PSM, anterior PSM, and $_{649}$ 598 somite regions by micro scalpel in cold PBS. $_{650}$ 590 Each tissue region was transferred into a micro 651 600 well (ibidi, #80486) and mechanically dissoci-₆₅₂ 601 ated to a cell suspension in 4.2 µl cold PBS. 653 602 Finally, 0.7 μ l and 3.3 μ l cell suspensions were 654 603 used for RNA-sequencing (RNA-seq) and ATAC-604 sequencing (ATAC-seq), respectively. For the com- $_{656}$ 605 parison between control and cytoPFKFB3 PSM 657 606 explants, explants were cultured for three-hour ex 658 607 vivo before collecting tail buds for RNA-seq anal-608 ysis. 609 660

ATAC-seq. We followed the Omni-ATAC proto-610 col [59] with some modifications. For transposi-611 662 tion reactions, 3.3 µl cell suspensions were mixed 612 663 with 5.0 µl 2x TD buffer (20 mM Tris-HCl pH 664 613 7.6, 10 mM MgCl₂, 20% dimethyl formamide), $_{665}$ 1.0 μl TDE1 (Illumina, #15027865), 0.1 μl 1% 666 615 digitonin (Promega,#G9441), 0.1 µl 10% Tween- $_{\rm _{667}}$ 616 20 (Sigma, #11332465001), 0.1 µl 10% NP-40 668 617 (sigma, #11332473001), and 0.4 µl nuclease-free ₆₆₉ 618 water. After 30 min incubation at 37°C on a 670 619 thermomixer set at 600 rpm, the samples were $_{671}$ 620 purified by a DNA Clean and Concentrator-5 672 621 (Zymo Research, D4014) and DNA concentrations 673 622 were determined by Qubit Fluorometer (dsDNA 674 623 High Sensitivity Kit, ThermoFisher, Q32851). The 675 624 samples were diluted to 20 ng/µl and used as 676 625 templates for library preparations by PCR. PCR 677 626 reactions were performed using primers from Nex-627 tera XT Index Kit (Illumina, FC-131-1001) and 679 628 NEBNext High Fidelity 2X PCR Master Mix 680 (NEB, M0541). After purification with Qiagen 681 630 MinElute PCR Purification Kit (Qiagen, 28004), 682 631 individual libraries were size selected (100–800 $_{683}$ 632 bp) with Ampure XP beads (Beckman Coul-633 ter, #A63881). Libraries were quantified using 634 the Qubit Fluorometer (dsDNA High Sensitiv-635 ity Kit) and average fragment length distribu-636 tion was determined by the Bioanalyzer (Agilent, 637

High Sensitivity DNA kit, 5067-4626). Prepared libraries were multiplexed in pools of equimolar concentrations and sequenced on the NextSeq 500 (Illumina) platform with 75-bp paired-end readings. After demultiplexing and barcode trimming (Trimmomatic Galaxy Version 0.36.6), sequencing reads were quality checked (FastQC Galaxy Version 0.73) and aligned to Mus Musculus genome (GRCm38) with the Bowtie2 aligner (Galaxy Version 2.3.4.2, options -I 0 -X 2000 –dovetail –sensitive). Multi-mapping and duplicate reads were removed; finally only reads mapping to major chromosomes were kept [60].

RNA-seq. We followed the Smart-seq2 protocol [61] with some modifications. In brief, dissociated cells were lysed with three times volume of cell lysis buffer (0.02% Triton-X with RNasin), snap frozen by liquid N_2 , and stored at $-80^{\circ}C$ until cDNA synthesis. cDNAs were synthesized using SuperScript IV Reverse Transcriptase (Thermo Fisher Scientific) and amplified by PCR with HiFi Kapa Hot start ReadyMix (Kapa Biosystems, KK2601). After clean-up with SPRI beads, concentrations of cDNA (50-9000 bp) samples were determined by the Bioanalyzer (Agilent, High Sensitivity DNA kit). 250 pg cDNAs were then used for tagmentation-based library preparation. Libraries were quantified using the Qubit Fluorometer (dsDNA High Sensitivity Kit) and average fragment length distribution was determined by the Bioanalyzer (Agilent, High Sensitivity DNA kit, 5067-4626). Prepared libraries were multiplexed in pools of equimolar concentrations and sequenced on the NextSeq 500 (Illumina) with 75-bp paired-end (for the wild-type, non-cultured PSM explants) or single-end (for the comparison between control and cytoPFKFB3 explants) readings. After demultiplexing and barcode trimming (TrimGalore Galaxy Version 0.4.3.1), sequencing reads were quality checked (FastQC Galaxy Version 0.69) and aligned to Mus Musculus genome (GRCm38) with the with the STAR aligner (version 2.5.2b, default options) [60]. Multi-mapping reads were removed and RNA-seq quality assessed with Picard CollectRnaSeqMetrics (Galaxy version 2.7.1.1)

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4.7 GRaNIE analysis

Enhancer-mediated gene regulatory network 730 685 (eGRN) was constructed from the matched RNA-₇₃₁ 686 seq and ATAC-seq data (24 samples for each) of 732 687 the PSM explants from E10.5 wild-type embryos $_{\scriptscriptstyle 733}$ 688 using the developer's version of the now pub-689 734 lished GRaNIE package (https://bioconductor. 735 690 org/packages/release/bioc/html/GRaNIE.html) 691 736 [29]. Raw gene counts from RNA-seq data 737 692 were produced with a summarizeOverlaps func-738 693 tion from the GenomicAlignments R package 739 694 (https://bioconductor.org/packages/release/ 695 bioc/html/GenomicAlignments.html) [62], cor-741 696 rected for different experimental batches using 742 69 Combat-seq function from the R package sva 743 698 [63]and log2 normalised. ATAC-seq peak 744 699 counts were generated using DiffBind R package 700 745 (https://bioconductor.org/packages/DiffBind/), 701 746 and peak positions were identified using MACS2 702 software (https://genomebiology.biomedcentral. 748 703 com/articles/10.1186/gb-2008-9-9-r137) [64]. The 749 704 details of the GRaNIE approach are described 750 705 here [29]. Briefly, in the first step the expression $_{751}$ 706 of each TF was correlated with accessibility of $_{\scriptscriptstyle 752}$ 707 each of the accessible regions (=ATAC-seq peak) 753 708 with and without a known binding site of the 754 709 TF (foreground and background, respectively). 755 710 Known binding sites were defined using the 756 711 HOCOMOCO database v.10 [65]. Significantly 757 712 correlated TF-peak links were identified using 758 713 empirical FDR of 30% (calculated separately for 759 714 each TF) and an absolute correlation Pearson's 760 715 coefficient of >0.4. In the second step chromatin ₇₆₁ 716 accessibility at the ATAC-seq peaks was corre-762 717 lated with the expression of all genes less than $_{763}$ 718 250kb away from the peak and peak-gene links $_{\scriptscriptstyle 764}$ 719 were retained if they were positively and signif- $_{\scriptscriptstyle 765}$ 720 icantly (P <0.05) correlated (our assumption is $_{766}$ 721 that accessibility at the regulatory region pos-722 itively correlates with expression of the linked $_{768}$ 723 gene), and if their Pearson's correlation coefficient 724 was >0.4. This resulted in the eGRN consisting $_{770}$ 725 of 69 TFs, 5154 TF-peak-gene connections of $_{771}$ 726 2522 unique genes. TF regulons were defined as 727 all TF-gene links of each TF within the network. 728 772

4.8 Microfluidics-based segmentation clock entrainment

PDMS chips and PTFE tubing (inner diameter: 0.6 mm, APT AWG24T) for microfluidicsbased entrainment experiments were prepared as described before [19, 24]. Culture media were prepared on the day of experiments by adding a metabolite of interest [either glucose, FBP, pyruvate (Sigma, P4562), or 3-OMG (Sigma, M4879)] to DMEM/F12 supplemented with 2.0 mM glutamine (Sigma-Aldrich, G7513), 0.01% (w/v) BSA (Cohn fraction V; Equitech-Bio, BAC62), and 1% penicillin-streptomycin (Gibco, 15140122). The PDMS chip (soaked in PBS) and the culture medium (filled in 10 mL syringes; BD Biosciences, 300912, diameter 14.5 mm) were degassed before use for at least one hour in a vacuum desiccator chamber.

Following dissection, PSM explants with two intact somites were transferred to the PDMS chip and sample inlets were plugged with a PDMSfilled PTFE tubing. The tubings connected to the syringes with medium were then connected to the medium inlets and the samples were placed in the incubator ($37^{\circ}C$, 5% CO₂, 65% O₂) installed on a LSM780 laser-scanning microscope (Zeiss) for preculture. Pumping was started for both the control and treatment medium at the flow rate of 20 µl/hr. A half hour later, only the control medium was pumped into the chip for another 30 min at the flow rate 60 µl/hr. After the pre-culture, imaging was started under constant or alternating culture conditions.

For data analysis, moving ROIs (30-pixel in diameter) were placed in the posterior PSM to obtain intensity profiles of LuVeLu or Axin2-Achilles reporters over time. To extract the period and phase of LuVeLu and Axin2-Achilles oscillations, the intensity profiles were analysed using a wavelet analysis workflow [58]. Entrainment of Notch and Wnt signaling oscillations was analysed using stroboscopic maps and the first Kuramoto order parameter as described before [24].

4.9 Data availability

The ATAC-seq and RNA-seq data generated inthis study were deposited in the BioStudies under

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the accession number E-MTAB-13692, E-MTAB13693, and E-MTAB-13694. For identifying glycolytic flux-responsive genes, the RNA-seq data ⁸²⁰
from our previous study [available in the European ⁸²¹

779 Nucleotide Archive (ENA) under the accession

number PRJEB55095] were also used [6].

781 5 Acknowledgments

824 We thank Irene Miguel-Aliaga, Kristina Staporn-782 825 wongkul, and Vikas Trivedi for their feedback 783 826 on the manuscript, Vladimir Benes and Laura 784 Villacorta for technical advice and support for 827 785 RNA-seq analysis and Jonathan Landry for help-786 ing RNAseq data analysis. This work is sup-787 ported by the EMBL Advanced Light Microscopy 788 Facility (ALMF), Genomics Core Facility, and 789 all the member of Laboratory Animal Resource 790 831 (LAR). H.M. was supported by the EMBL 791 Interdisciplinary Postdoc (EI3POD) programme 792 833 under H2020 Marie Skłodowska-Curie Actions 793 COFUND (grant number 664726) and the Japan 794 834 Society for the Promotion of Science (JSPS). 795 E.E. was supported by the Human Frontier Sci-796 836 ence Program (HFSP) fellowship. This work was 797 supported by the European Molecular Biology⁸³⁷ 798 Laboratory and received funding from the Euro-⁸³⁸ 799 pean Research Council under an ERC consolida-800 839 tor grant agreement n.866537 to A.A. and the 801 German Research Foundation/DFG (project SFB⁸⁴⁰ 802

⁸⁰³ 1324 – project number 331351713) to A.A.

⁸⁰⁴ 6 Author contributions

- $_{\tt 805}$ H.M.: Conceptualization, Methodology, Formal
- analysis, Investigation, Writing Original Draft,
 Visualization, Supervision
- ⁸⁰⁸ J.R.: Conceptualization, Methodology, Formal
- analysis, Investigation, Writing Original Draft,
 Visualization
- 811 P.G.L.S.: Methodology, Software, Investigation
- 812 E.E.: Methodology, Formal analysis, Investigation
- 813 D.B.: Software, Formal analysis, Investigation
- ⁸¹⁴ C.G.: Software, Formal analysis
- ⁸¹⁵ J.Z.: Supervision, Funding acquisition
- 816 A.A.:Conceptualization, Methodology, Writing –
- 817 Original draft preparation, Supervision, Project
- 818 administration, Funding acquisition

Declarations

The authors declare that they have no conflict of interests.

Appendix A Extended Data

- A.1 Extended Data Fig. 1 Increasing glycolytic flux slows down Wnt signaling oscillations.
- A.2 Extended Data Fig.2 Glycolytic flux shows glucose-dose dependency in PSM cells.
- A.3 Extended Data Fig.3 Building a PSM-specific eGRN using the GRaNIE method.
- A.4 Extended Data Fig.4 Response of cellular redox state to alterations in glycolytic flux within PSM cells.
- A.5 Extended Data Fig.5 Segmentation clock entrainment by periodic, transient glycolytic cues.



Extended Data Fig. 1 Increasing glycolytic flux slows down Wnt signaling oscillations. (A) Kymographs showing the dynamics of the Axin2-Achilles knock-in reporter in control (Ctrl) and cytoPFKFB3 (TG) PSM explants in 2.0 mM glucose condition. (B) Quantification of the Wnt signaling oscillation periods in Ctrl and TG explants cultured in 2.0 mM glucose. The periods were determined as a mean of Axin2-Achilles periods between 400-600 min of the imaging. Welch's unpaired t-test, ***p <0.001. Mean \pm SD are shown in the graph, and individual data points represent biological replicates.



Extended Data Fig. 2 Glycolytic flux shows glucose-dose dependency in PSM cells. (A, B) Lactate secretion was quantified as a proxy for glycolytic flux within PSM cells. The amount of lactate secreted from PSM explants during 12 h ex vivo culture was quantified. (A) Comparison of lactate secretion between control (Ctrl) and cytoPFKFB3 (TG) explants cultured in 0.5 mM or 2.0 mM glucose (the data for 2.0 mM glucose condition is adapted from Miyazawa et al. 2022 [6]). (B) The effect of replacing glucose with galactose on lactate secretion from wild-type explants. Welch's unpaired t-test, *p <0.05, **p <0.01 vs. Ctrl. Mean \pm SD are shown in the graph, and individual data points represent biological replicates.



Extended Data Fig. 3 Building a PSM-specific eGRN using the GRaNIE method. (A) A heatmap showing gene expressions of each PSM-specific regulon (i.e., means of all the targets) identified by the GRaNIE method. Normalized counts by variance stabilizing transformation (VST) were used to calculate the z-scores. (B) A network showing TFs (colored squares) and their glycolytic flux-responsive target genes (colored circles). (C) Box plots showing fold changes in gene expressions of flux-sensitive DEGs that constitutes each PSM-specific regulon. The fold changes were calculated between different metabolic conditions.



Extended Data Fig. 4 Response of cellular redox state to alterations in glycolytic flux within PSM cells. Quantification of NAD⁺/NADH ratio following one-hour ex vivo culture of control (Ctrl) and cytoPFKFB3 (TG) PSM explants under various culture conditions. For the galactose (GALA) condition, 2.0 mM galactose was supplemented to the culture medium instead of glucose. Mean \pm SD are shown in the graph, and individual data points represent biological replicates. Welch's unpaired t-test, *p <0.05.



Extended Data Fig. 5 Segmentation clock entrainment by periodic, transient glycolytic cues. (A) Detrended (via sinc-filter detrending, cut-off period = 240 min) time-series of LuVeLu intensity oscillations in wild-type PSM explants exposed to periodic pulses of 20 mM pyruvate (dashed lines: individual samples, bold black line: median values, grey shades: the first to third quartile range). Changes in the first Kuramoto order parameter are shown in magenta. To keep molarity of the medium at constant during experiments, 20 mM non-metabolizable glucose (i.e. 3-O-methyl-glucose) was added to the basal medium containing 2.0 mM glucose. (A') Stroboscopic maps showing step-wise changes in the phase of LuVeLu oscillations in response to periodic pyruvate pulses. Darker dots represent later time points (the numbers in the plots indicate the number of the pulses). (B) Stroboscopic maps showing the phase of Notch (i.e., LuVeLu) and Wnt (i.e., Axin2-Achilles) oscillations at the last pulse of metabolite. Filled circles represent entrained samples, while open circles represent non-entrained samples. Samples are considered to be entrained when a phase difference between the last and second last pulses is less then $\pi/8$. CON-Gluc, constant (2.0 mM) glucose condition; ALT-Gluc, alternating (from 2.0 mM to 0.5 mM) glucose condition.

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