Circadian transcriptional pathway atlas highlights a proteasome switch in intermittent fasting

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SUMMARY

While intermittent fasting is a safe strategy to benefit health, it remains unclear whether a “timer” exists in vivo to record fasting duration and trigger a transcriptional switch. Here, we map a circadian transcriptional pathway atlas from 600 samples across four metabolic tissues of mice under five feeding regimens. Results show that 95.6% of detected canonical pathways are rhythmic in a tissue-specific and feeding-regimen-specific manner, while only less than 25% of them induce changes in transcriptional function. Fasting for 16 h initiates a circadian resonance of 43 pathways in the liver, and the resonance punctually switches following refeeding. The hepatic proteasome coordinates the resonance, and most genes encoding proteasome subunits display a 16-h fasting-dependent transcriptional switch. These findings indicate that the hepatic proteasome may serve as a fasting timer and a coordinator of pathway transcriptional resonance, which provide a target for revealing the underlying mechanism of intermittent fasting.

Graphical Abstract

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AUTHOR CONTRIBUTIONS
G.L. designed the experiments. G.L., F.W., L.G., S.L., Y.Z., L.L., and Z.D. conceived experiments. G.L. performed the bioinformatics analysis. G.L., F.W., and R.X. contributed to the interpretation of results. G.L., F.W., and F.J.G. wrote the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.celrep.2022.111547.
In brief

While intermittent fasting benefits health, the optimal duration of each fasting remains an open question. Wei et al. map an atlas of canonical pathways in intermittent fasting, find that fasting for 16 h initiates circadian resonance of pathways in the liver, and identify the proteasome as a liver-specific fasting “timer”.

INTRODUCTION

Intermittent fasting has become a safe strategy to benefit health (de Cabo and Mattson, 2019; Stekovic et al., 2019). Growing evidence consistently shows that intermittent fasting can efficiently prevent a wide range of diseases, including diabetes, obesity, cancers, and neurodegenerative disorders (Di Francesco et al., 2018; Li et al., 2017; Nencioni et al., 2018). It even functions to increase lifespan or retard the aging process in different animal models (Anton et al., 2018; Catterson et al., 2018; Honjoh et al., 2009; Longo et al., 2021; Ulgherait et al., 2021; Xie et al., 2017). However, there are currently many intermittent fasting regimens (Anton et al., 2018; de Cabo and Mattson, 2019; Longo et al., 2021), while the strong evidence of exclusivity for the predominance of each regimen is limited. The temporal functional signatures of fasting remain to be elucidated.

It is assumed that flipping the metabolic switch from glucose to fatty-acid-derived ketones is a key mechanism responsible for the intermittent-fasting-induced health benefits (Anton et al., 2018; de Cabo and Mattson, 2019; Mattson et al., 2018), and plasma ketone levels
can serve as a biomarker of the metabolic switch (Anton et al., 2018). While there is a wide consensus that prolonged fasting can trigger the metabolic switch, and multifaceted evidence supports a crucial role of the metabolic switch in optimizing metabolism and improving metabolic diseases, the metabolic switch that occurs between 12 and 36 h after food deprivation varies greatly between individuals (Anton et al., 2018). In this context, it is difficult for the metabolic switch to accurately indicate functional switches at a given time point, and two open questions that arise are whether a “timer” exists in vivo to record the fasting duration, and whether and when does fasting spark transcriptional switches?

Theoretically, these temporal signatures should be associated with circadian clocks. Intriguingly, multiple lines of evidence have supported the interaction between circadian rhythms and intermittent fasting. Circadian clocks optimize the benefits of intermittent fasting through temporally orchestrating signaling pathways, metabolic activity, and other cellular functions (Manoogian and Panda, 2017; Mohawk et al., 2012). In turn, feeding behavior modifications in energy components and time can markedly alter rhythmic transcription (Greenwell et al., 2019; Kohsaka et al., 2007; Villanueva et al., 2019; Vollmers et al., 2009). Feeding-regulated hormones and metabolic oscillators can reset circadian rhythms in vivo (Bass, 2012; Crosby et al., 2019; Peek et al., 2013). Time-restricted feeding prevents metabolic diseases even in mice lacking a circadian clock (Chaix et al., 2019), while mistimed eating has deleterious effects on metabolic health (Challet, 2019). However, data from current transcriptomics studies have not revealed a transcriptional switch or fasting timer. In this case, we suspect that the temporal signatures may be tissue specific and present at the pathway level rather than the gene level.

Therefore, in the current study, with the use of gene set variation analysis (GSVA) (Hanzelmann et al., 2013) and canonical pathways gene sets (Mootha et al., 2003; Subramanian et al., 2005), we mapped a circadian transcriptional pathway atlas based on 600 samples across four metabolic tissues sampled every 4 h from mice placed on ad libitum (AL), acute fasting (AF), refeeding after acute fasting (AR), every-other-day fasting (EODF), and every-other-day refeeding (EODR) regimens. The evidence suggests that intermittent fasting significantly triggers liver-specific transcriptional switches of 43 canonical pathways after 16-h fasting or refeeding, and the liver proteasome may serve as a potential fasting timer and coordinator of pathway switches.

RESULTS

Canonical pathway profiling of mouse tissues reveals tissue-specific oscillating signatures in intermittent fasting

To determine the adaptive metabolic changes induced by intermittent fasting, the effect of EODF on body weight was first analyzed. Compared with the AL group, EODF did reduce the body mass gain (Figure S1). To obtain a comprehensive metabolic perspective of the temporal signature of intermittent fasting, we measured the transcriptome of four metabolic tissues (liver, skeletal muscle [biceps femoris], interscapular brown adipose tissue [BAT], and inguinal white adipose tissue [WAT]) isolated every 4 h from mice placed on AL, AF, AR, EODF, or EODR regimens (Figure 1A). RNA sequencing identified the expression of 17,848 genes at least in one group (Table S1), and the transcriptome measurements were
highly accurate and reproducible, with Spearman correlation coefficients more than 0.95 in biological replicates (Figures S2A and S2B) and more than 0.80 in samples across groups (Figures S2A and S2C).

Three-dimensional (3D) principal-component analysis (PCA) (Venables and Ripley, 2002) of transcriptomic data did not classify each group according to different feeding regimens (Figure S2D), indicating that it might be difficult to find the temporal signature of intermittent fasting at the gene level. Therefore, we further calculated pathway GSVA scores (Hanzelmann et al., 2013) for these genes based on 2,922 gene sets of canonical pathways that were downloaded from the MSigDB (c2.cp.v7.4) (Mootha et al., 2003; Subramanian et al., 2005). We identified 2,175 pathways at least in one group (Table S2), accounting for 74.4% of total pathways. Most (2,007 in 2,175) were commonly enriched in all four tissues (Table S2). Further 3D-PCA analysis with the GSVA scores improved the classification of samples across feeding regimens, but some samples still overlapped (Figure S2E). Given that more than half of the nuclear receptors and the majority of protein-coding genes exhibit rhythmic expression (Mure et al., 2018; Yang et al., 2006; Zhang et al., 2014), we suspected that pathways composed of multiple genes might also display rhythmic features and the effects of feeding regimens on pathways might be masked by the features.

As expected, we observed an obvious tissue-specific and feeding-regimen-specific circadian oscillation in canonical pathways (Figure 1B). In skeletal muscle of AL mice, the majority of pathways were upregulated in the dark cycle but downregulated in the light cycle (Figures 1B and 1C), which is synchronized with the characteristics that this nocturnal species is most active at night. The ratio of the dark cycle to the light cycle of upregulated pathways illustrated that skeletal muscle and WAT were more sensitive to AF than the other two tissues (Figure 1C). Further comparative analysis of circadian oscillating pathways revealed the effects of different feeding regimens on these pathways were tissue specific, and the total number of oscillating pathways varied among tissues (Figure 2). In AL mice, the percentage of oscillating pathways in muscle was just 8%, while that in other tissues was more than 25% (Figure 2). These data are consistent with previous evidence that 15% of hepatic transcripts and 4% of muscle transcripts were cyclic (Kinouchi et al., 2018). Compared with AL mice, all other feeding regimens tended to decrease the percentage of oscillating pathways in the liver and WAT, but, in muscle and BAT, the decrease only occurred in a specific feeding regimen (Figure 2). The correlations, amplitude, and phase distribution of oscillating pathways among different feeding regimens also showed tissue-specific signatures (Figure 2). It was reported that fasting can cease the oscillation of more than 60% of rhythmic transcripts in the liver and muscle (Kinouchi et al., 2018). Accordingly, we noticed that the oscillating pathway correlations of AL with AF in the liver and muscle were very low (Figure 2). If we considered the phase, there were few overlapping pathways between AL and AF (Figure 2; Table S3). Due to the low overlap of cyclic pathways among feeding regimens and tissues, a total of 2,079 rhythmic pathways were detected at least in a specific feeding regimen of one tissue (Table S3), which accounts for 95.6% of the 2,175 identified pathways (Table S2). Considering the high rhythmicity of genomic transcription (Mure et al., 2018) and the potential influence of feeding behavior on rhythmic transcription (Vollmers et al., 2009), the high oscillation of these pathways is logical. Notably, the phases in AF mice seemed to be chronological among different tissues: WAT peaked at 8–10 h after
fasting, then BAT at 12 h, liver at 20 h, and muscle at 22 h (Figure 2), suggesting that the
length of fasting time to initiate a response is also tissue specific. All these data indicate
that canonical pathways are highly rhythmic, and intermittent fasting can spin oscillating
signatures of these pathways in a tissue-specific manner.

Canonical pathway profiling all day suggests an adaptation to “acute fasting-refeeding” in
intermittent fasting

Owing to the high rhythmicity of genomic transcription and canonical pathways, to precisely
evaluate the effects of different feeding regimens on these pathways, the overlapping effects
of circadian oscillation on the data should first be eliminated. Thus, the mean expression
values (EDi) of each gene for the whole day were calculated from the expression values (Eij)
of six time points in a 24-h day within each feeding regimen and tissue (Figure 3A). The
means were then used for GSVA, 3D-PCA, and other analyses.

Based on these means, we did observe that different feeding regimens caused mild and clear
alterations in the transcripts (Figure 3B) and canonical pathways (Figure 3C), respectively,
as indicated by the generalized 3D-PCA distances. Surprisingly, different tissues appeared to
have coordinated responses to intermittent fasting, as the Spearman correlation coefficients
of canonical pathways exhibited similar patterns of change across tissues (Figure 3D);
that is, the coefficients within each feeding regimen were apparently positive, while those
between AF and EODR or between AR and EODF were obviously negative (Figure 3D).
These data challenged our conventional view that AF and AR or EODF and EODR are two
pairs of inverse effects. Especially, the coefficients between AF and AR in muscle, and those
between EODF and EODR in muscle, WAT, and BAT were positive (Figure 3D). These
results can be compared with the classic concept of ischemic-reperfusion injury. Although
the restoration of blood flow (reperfusion) is a routine means of handling tissue ischemia,
some functional and structural changes become apparent during reperfusion after a period
of ischemia (Soares et al., 2019). Therefore, these data indicate that acute fasting-refeeding
might be a pair of linked nutritional challenges, and that intermittent fasting, by adapting to
the challenges, may yield beneficial effects on an organism.

The functional rhythmic pathways exhibit a manifest switch after 16 h fasting or refeeding
in the liver

Since the pathway GSVA scores refer to genes in pathways that are concordantly over-
expressed or under-expressed and are approximately normal (Hanzelmann et al., 2013),
we performed differential expression analysis on the whole-day GSVA scores using limma
(Smyth, 2004). With the cutoff of $\log_2$(fold change) > $\log_2(1.5)$ and adjustedp < 0.05, we
detected 1,343 differentially expressed pathways (DEPs) at least in one compared pair of
a tissue (Table S4). The overlapping analysis within each tissue illustrated that only 16.6%–
24.9% of rhythmic pathways overlapped with DEPs (Figures 4A–4D), indicating that most
rhythmic pathways did not cause changes in transcriptional function throughout the whole
day. These pathways showing both oscillation and differential expression will be referred to
as functional rhythmic pathways (FRPs) in the following descriptions.
To explore the detailed oscillating characteristics of these FRPs, we first drew their heatmaps with GSV A scores calculated from the original expression data (Table S2). Intriguingly, the heatmap in the liver demonstrated an obvious expression switch at the 16th hour of fasting or refeeding (Figure 4E, black arrow), while that did not occur in other tissues (Figures 4F–4H) or the liver of AL mice (Figure 4E, black arrow). These results indicate that there is a liver-specific timer in vivo to record the duration of fasting and refeeding.

Therefore, we further analyzed the feeding-specificity and phase distribution of FRPs within each tissue. Results demonstrated that the major FRPs were clustered into AL and acute fasting-refeeding (Figures 5 [top bar charts] and S3A). Since an oscillating pathway that only occurs in AL mice actually represents a pathway that loses rhythmicity during AF and cannot restore rhythmicity in other feeding regimens, it also reflects the effect of AF. In this context, these data suggest that most of the overlapping pathways might directly stem from the AF-induced circadian fluctuations or indirectly derive from the adaptation to the AF-induced circadian fluctuations.

At each time point, the fraction of feeding-regimen-specific DEPs in FRPs fluctuated (Figure 5 [side bar charts]). Compared with the top bar charts, the proportion of EODF and EODR increased in the side bar charts, while that of AF decreased (Figure 5), indicating intermittent-fasting-induced changes in transcriptional function (as indicated by DEPs) originated more from the adaptation to AF-induced circadian fluctuation than from itself. Curiously, the peak at the 16th hour observed above (Figure 4E) did not appear, while the peak of FRPs varied across feeding regimens (Figures 5 [last column] and S3B). However, consistent with the above results (Figure 3D), we noticed several pairs of typical inverse effects between AR and EODF in DEPs of different tissues at the same oscillating time point (Figure 5 [green arrow]), and most of these pairs occurred in the rhythmic pathways of AF mice. The phase distribution of FRPs in AF mice highlighted the temporal effects of fasting in the liver, as only the liver sharply peaked at a single time point, the 20th hour of fasting (Figures S3C–S3F). Therefore, the pathways displaying inverse effects between AR and EODF at the 20th hour of fasting in the liver were chosen for further analysis.

Intriguingly, of the 45 pathways upregulated in AR mice and 57 pathways downregulated in EODF mice (Figure 5 [green arrow]), 43 pathways were shared by both, which accounts for 18.8% of the total 229 FRPs in the liver (Figure 4A). The oscillating curves of the 43 pathways displayed a typical circadian resonance (Figure 6A). Although the fitted curve in AF mice peaked at the 20th hour of fasting, almost all pathways were clearly switched at the 16th hour of fasting (Figure 6A), which might explain the discrepancy between the peak of FRPs (Figure 5) and the switching time of the heatmap (Figure 4E). Consistent with the above data (Figure 4E), these 43 pathways only oscillated in fasting or refeeding mice, but not in AL mice (Figure 6A). Moreover, across feeding regimens, the switching time was fixed at the 16th hour of fasting or refeeding (Figure 6A). These data therefore confirm our assumption that intermittent fasting may trigger transcriptional switches at the pathway level.
The circadian resonance of 43 pathways in the liver highlights a proteasome switch in intermittent fasting

While the circadian resonance of 43 pathways is interesting, the question that arises is why does this happen? To answer this question, we first classified these pathways according to the Reactome Pathway Database (Jassal et al., 2020), as most of the 43 pathways were Reactome pathways. The results revealed that, although 42% belonged to signal transduction, their functions were distributed over several categories (Figure 6B). Obviously, they cannot be classified into one functional class.

Then, the genes that constituted these pathways were extracted for overlap analysis. Of the total of 837 genes, 14 were shared by all 43 pathways (Figure 6C). Surprisingly, they all are genes encoding proteasome subunits (Figure 6C), implying that the proteasome may coordinate the circadian resonance of these pathways. If it is the case, the transcription of the genes encoding proteasome subunits would be expected to be switched at the time after fasting or refeeding for 16 h. To verify this assumption, the transcription data of all genes encoding proteasome subunits were extracted for oscillation analysis. As expected, the expression of these genes in the liver did switch at the time point in fasting or refeeding (Figure 7A), which did not occur in other tissues (Figures 7B–7D). Consistent with the concept that the amount of proteasomes in Mammalia is regulated at the transcriptional level (Meiners et al., 2003), the expression of three proteins encoded by Psma4, Psmb3, and Psmb6 in the liver that were randomly selected out from 14 shared genes in Figure 6C were also oscillated in a similar manner with the transcription of proteasome subunits (Figures 7E and S4A–S4E). These data suggest that the liver proteasome may be a fasting timer that coordinates the resonance of the 43 pathways. Since the proteasome plays a prominent role in the control of multiple basic cellular activities (Tanaka, 2009), this may explain why the 43 pathways belong to several categories.

One question that remains is whether the time point of a transcriptional switch is unique to fasting (refeeding) duration or the combined effect of dark/light switch and fasting (refeeding) duration. Fortunately, the data from a paper published in Cell Reports may be helpful to answer this question. In contrast to our fasting protocol that used a fixed start time point, that study fasted six groups of mice at six start time points with a 4-h interval between each (Kinouchi et al., 2018). Following this protocol, mice that were killed every 4 h over a day were all fasted for 24 h. Therefore, if fasting duration solely determines the transcriptional switch, the switching phenotype will be lost in these mice. As expected, the genes encoding proteasome subunits in these mice did not show obvious transcriptional switch at zeitgeber time 4 h or other times in the liver (Figure S5A), although some genes displayed transcriptional changes after fasting. Therefore, it might be logical to consider the transcriptional switch in these genes as a fasting-specific effect. Nevertheless, the fasting-induced transcriptional changes of some genes in muscle exhibited a seeming oscillatory effect (Figure S5B), suggesting a tissue-specific role for fasting. Together, these data indicate that it is the 16-h fasting that initiates the transcriptional switch of genes encoding proteasome subunits in the liver.

There is a wide consensus that prolonged fasting can promote the production of ketone bodies (Balasse and Fery, 1989). Recent evidence suggests that ketones serve as an
The important signal in hepatic response to fasting (Geisler et al., 2019) and a key mechanism responsible for the intermittent-fasting-induced health benefits (Anton et al., 2018; de Cabo and Mattson, 2019; Mattson et al., 2018). To explore the potential role of ketones in intermittent-fasting-induced transcriptional switches, the oscillations in serum ketone bodies across the different groups were analyzed. The results showed that fasting increased serum acetoacetate and ω-hydroxybutyrate levels (Figures S6A and S6B; Table S5), but they did not display obvious oscillations during fasting (Figures S6A and S6B; Table S5). These data suggest that ketone signaling should not orchestrate the transcriptional switches, and, therefore, our current evidence indicates that it is proteasome rather than ketone signaling that serves as a fasting timer.

**DISCUSSION**

In this study, we mapped the circadian transcriptional pathway atlas in four metabolic tissues of mice under five feeding regimens (AL and four components of intermittent fasting), which serves as an important resource for revealing the regulatory mechanism of intermittent fasting. Current evidence highlighting the importance of circadian rhythm in the beneficial effects of intermittent fasting (specifically time-restricted feeding) on health has been repeatedly reported (Chaix et al., 2019; Crosby et al., 2019; Greenwell et al., 2019; Manoogian and Panda, 2017; Ulgherait et al., 2021; Villanueva et al., 2019; Vollmers et al., 2009). Although the circadian transcriptome atlases in mice and primates have been mapped (Mure et al., 2018; Zhang et al., 2014) and rhythmic feeding has been shown to trigger the oscillations in most peripheral tissues (Damiola et al., 2000; Stokkan et al., 2001), the pathway atlas across major metabolic tissues of mice under intermittent fasting is still lacking. Here, we created high-resolution multi-organ transcriptional over-expression or under-expression of canonical pathways (Mootha et al., 2003; Subramanian et al., 2005) based on the GSVA analysis (Hanzelmann et al., 2013), and then characterized these pathways by oscillating signature and differential expression throughout the whole day. The data revealed that the transcriptional pathways were highly rhythmic, and the oscillating signature of these pathways was tissue specific and varied dramatically across feeding regimens. Although the oscillating pathways in muscle and WAT were more sensitive to AF than the other two tissues, BAT accounted for the most DEPs. These data imply that the changes in circadian oscillation of pathways cannot directly deduce the corresponding changes in transcriptional function. In fact, we found that only less than 25% of the oscillatory pathways lead to changes in transcriptional function all day.

A striking finding from this study is that fasting for 16 h can awake a liver-specific cellular quality controller (the proteasome). Although intermittent fasting has been proved to benefit health from many aspects, fasting programs vary across laboratories (Chaix et al., 2019; Li et al., 2017; Mihaylova et al., 2018; Ulgherait et al., 2021) and even within each laboratory (Chaix et al., 2019; Hatori et al., 2012; Villanueva et al., 2019). The main reason is that the exact time point in vivo at which organisms exhibit an apparent response to fasting has not been found. Here, we revealed that fasting for 16 h triggered an evident switching effect of many canonical pathways, and that, once this effect is initiated, the trough and peak would punctually occur in the following refeeding and fasting. Intriguingly, all the shared genes of these pathways were genes encoding proteasome subunits, indicating...
the proteasome coordinated the pathway switches. Moreover, almost all genes encoding proteasome subunits in the liver demonstrated a fasting-dependent transcriptional switch similar to these pathways, indicating that the proteasome might be a liver-specific fasting timer. Although these data cannot directly imply that the fasting-induced switching effects in canonical pathways stem from those in the proteasome, the high synchronization of the transcriptional switch of these pathways and genes implies they are closely associated with each other. Since the proteasome is a critical component of cellular quality control systems (Pohl and Dikic, 2019; Tanaka, 2009), and previous evidence suggests a crucial role for fasting in increasing the activity of the proteasome and promoting selective degradation of specific proteins (VerPlank et al., 2019), the regulation of proteasome-coordinated pathways by fasting and refeeding implies that intermittent fasting may benefit health from the improved function of the cellular quality control systems. Although an open question remains for the mechanism of the pulse-like upregulation and downregulation of genes and pathways at a single time point in response to fasting and refeeding, respectively, this finding highlights the essential position of 16-h fasting in the beneficial role of intermittent fasting. Consistent with this notion, recent evidence indicates that fasting for more than 16 h is necessary for extending the lifespan of *Drosophila* (Ulgherait et al., 2021), and the underlying mechanism for the lifespan extension is circadian-regulated autophagy (Ulgherait et al., 2021), which is another component of the cellular quality control systems (Pohl and Dikic, 2019; Shaid et al., 2013). In this context, the regulation of the hepatic proteasome by 16-h fasting may modulate the function of the cellular quality control systems, which contributes to the health benefits of intermittent fasting. This finding might therefore be helpful to reach a consensus on intermittent fasting programs.

An intriguing finding of the present study is that intermittent fasting initiates the transcriptional circadian resonance of many canonical pathways in the liver. It is assumed that some environmental or behavioral perturbations may trigger transcriptional resonance (Bass, 2012; Bass and Takahashi, 2010). Currently, no typical evidence exists for this concept in animals, but it does occur in *Arabidopsis* (Harmer et al., 2000). Theoretically, intermittent fasting, synchronized daily cycles of fasting and refeeding through the removal and supply of food at a fixed time point, may provoke the transcriptional resonance. However, although it was reported that time-restricted feeding can restore transcriptional rhythmicity in the liver of oscillator-deficient mice (Vollmers et al., 2009), the resonant expression of genes has not hitherto been revealed. One possibility for this is that the resonance in animals may happen on pathways rather than on genes. Here, we did find distinct transcriptional resonances of many canonical pathways in the liver. While the re-establishment of circadian transcription by intermittent fasting has been reported to prevent obesity and metabolic syndrome in mice lacking a circadian clock (Chaix et al., 2019), the importance of the resonance in health still needs to be explored.

Another important finding of the current study is that, analogous to ischemic reperfusion, acute fasting-refeeding may pose a synergistic challenge in mice. Simply from a nutritional perspective, AR is the inverse process of AF. However, it cannot be taken for granted that all the effects of AF can be reversed by AR. The Spearman correlation coefficients of canonical pathways in this study clearly indicated that it was EODR rather than AR that exhibited an obviously negative correlation with AF. From the temporal oscillation curve of resonant
pathways, the effects of AR even looked like a spillover effect of AF. By contrast, we noticed an obvious reverse effect between AR and EODF or AF and EODR. If we consider acute fasting-refeeding as a synergistic challenge, all these phenotypes are logical, and the reverse effects between AR and EODF or AF and EODR reflect the adaptation to acute fasting-refeeding after several daily cycles of fasting and refeeding. The health benefits of intermittent fasting may originate from the adaptation.

In summary, the present work establishes a temporal transcriptional atlas of canonical pathways under five feeding regimens. Fasting for 16 h can awaken a liver-specific cellular quality controller (the proteasome), which coordinates the transcriptional resonance of canonical pathways and may contribute to the health benefits of intermittent fasting. This resource can serve as a basis for targeting the proteasome to reveal the underlying mechanism of intermittent fasting, reaching a consensus on intermittent fasting programs, exploring the transcriptional mechanism of intermittent fasting, and developing intermittent fasting mimics.

**Limitations of the study**

While this temporal transcriptional atlas of canonical pathways may be helpful to reveal the underlying mechanisms for the health benefits of intermittent fasting, the findings from this study are primarily based on RNA sequencing datasets. Therefore, more experiments, especially genetic and pharmacological loss-of-function experiments targeting the hepatic proteasome, may be needed to further scrutinize the findings. Moreover, additional studies are required to answer several open questions that arise from this study, including what the underlying mechanisms are for the 16-h fasting-dependent and liver-specific proteasome provocation, which factors trigger the pathway resonance, and what the interaction is among proteasome activation, pathway resonance, and health benefits in intermittent fasting.

**STAR METHODS**

Detailed methods are provided in the online version of this paper and include the following:

**RESOURCE AVAILABILITY**

**Lead contact**—Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Guolin Li (hnsdlg1@hunnu.edu.cn).

**Materials availability**—This study did not generate new unique reagents.

**Data and code availability**

- The accession number for the sequences reported in this paper is GEO: GSE154797 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE154797). The DOI for the raw data of blots reported in this paper is 10.17632/x6s7p7cb7.1 (Mendeley Data: https://doi.org/10.17632/x6s7p7cb7.1). These data will be publicly available as of the date of publication.

- This paper does not report original code.
• Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animals—All mouse studies were approved by the Institutional Review Board of the Hunan Normal University and performed according to the Laboratory Animal Resources guidelines. Six-week-old male C57BL/6N mice were purchased from Beijing Vital River Laboratory Animal Technology Co., Ltd. Mice were housed in a temperature-controlled vivarium (25 ± 2°C) on a 12-h light-dark cycle with free access to water and standard rodent chow food (18.9% protein, 5.5% fat, and 61.3% carbohydrate). All experiments were started with 8-week-old mice.

METHOD DETAILS

Intermittent fasting treatment—Mice were randomly grouped to ad libitum (AL) group, acute fasting (AF) group, refeeding after acute fasting (AR) group, every other day fasting (EODF) group, and every other day refeeding (EODR) group. All mice were singly housed for two weeks before study initiation to allow for acclimation to the animal facility. The AL group mice were allowed unrestricted access to food. The mice of AF groups were allowed free access to food except for the last day of acute fasting for 4 h, 8 h, 12 h, 16 h, 20 h, and 24 h, respectively, before being killed. The mice in the AR groups were allowed free access to food except the last two days followed acute fasting for 24 h, and then refeeding for 4 h, 8 h, 12 h, 16 h, 20 h, and 24 h, respectively, before being killed. The mice in the EODF and EODR groups were fed with 15 cycles of alternating 24-h periods of free access to food followed by 24-h fasting, then EODF mice were killed after fasting for 4 h, 8 h, 12 h, 16 h, 20 h, and 24 h, respectively, while EODR mice were killed after refeeding for 4 h, 8 h, 12 h, 16 h, 20 h, and 24 h, respectively. Considering the nocturnal eating characteristics of mice, all fasting and refeeding switches were started at zeitgeber time 12 h.

Western Blot analysis—Frozen liver tissues were lysed in RIPA buffer supplemented with Halt Protease and Phosphatase Inhibitor Cocktail (Servicebio, Wuhan, China) and 1 mM PMSF. Protein concentrations were determined using BCA Protein Assay Kit (Servicebio, Wuhan, China) by a microplate reader (PERL ONG, Beijing, China). Twenty micrograms of protein per lane were loaded onto an 8–12% polyacrylamide Gel and then transferred to PVDF membranes using an electrophoresis chamber (Millipore, Shanghai, China) for 1.5 h. Membranes were blocked in 5% non-fat milk Tris-buffered saline (TBS) followed by overnight incubation in primary antibody at 4°C. Primary antibodies used are as follows: anti-PSMB3 (#K009840P, Solarbio, Beijing, China), anti-PSMB6 (#K003745P, Solarbio, Beijing, China), anti-PSMA4 (#K003378P, Solarbio, Beijing, China) and anti-ATCB (#YT0099, Immunoway, TX, USA). Following primary antibody incubation, blots were washed four times for 10 min each time with TBS and 0.1% Tween 20, and incubated with an anti-rabbit IgG HRP-conjugated secondary antibody (#AS014, ABclonal, Wuhan, China). Blots were stripped with Enhanced Chemiluminescent Liquid (ECL, NCM, Suzhou, China) and re-probed with anti-ATCB for normalization. Blot imaging was performed on a chemiluminescence analyzer (Tanon-5200 Multi, Shanghai, China).
Ketone bodies assays—Serum acetoacetate and β-hydroxybutyrate across the different groups were assessed in a 96-well microplate using commercial ketone bodies assay kits (#BC5060, Solarbio, Beijing, China) and monitored at 340 nm with a microplate reader (Perlong, Beijing, China).

RNA extraction and processing—Total RNA was extracted from frozen tissue using TRIzol reagent (Invitrogen) following the manufacturer’s instructions. The purity and concentration of the total RNA were determined by a NanoDrop spectrophotometer (ND-1000, Thermo Fisher). The quality/integrity of total RNA was evaluated by a Bioanalyzer (2100, Agilent) with RIN number >7.0, and confirmed by gel electrophoresis.

Library generation and sequencing—Poly(A) RNA was selected from 1 μg total RNA in each sample using Dynabeads Oligo (dT)25 (Thermo Fisher) through two rounds of purification and fragmented into small pieces using Magnesium RNA Fragmentation Module (New England Biolabs) under 94°C for 5–7 min. The cleaved RNA fragments were firstly reverse transcribed using SuperScript™ II Reverse Transcriptase (Invitrogen). Then the second strand of cDNA was synthesized using E. coli DNA polymerase I (New England Biolabs), RNase H (New England Biolabs), and marked with dUTP Solution (Thermo Fisher). Subsequently, after end polishing, A-tailing, adapter ligation, and size selection (using AMPureXP beads), the dUTP-marked strand was selectively degraded by Uracil-DNA-Glycosylase (UDG). The remaining strand was amplified with PCR (95°C for 3 min, 15 cycles of “98°C for 15 s, 60°C for 15 s, and 72°C for 30 s”, and 72°C for 5 min) to generate a cDNA library with 300 ± 50 bp. Libraries were multiplexed in equimolar concentrations and sequenced using an Illumina Novaseq™ 6000 with 2 × 150 bp paired-end sequencing (LC-Bio Technology CO., Ltd.) following the vendor’s recommended protocol.

Data processing—Quality control was assessed using Cutadapt (Martin, 2011), then HISAT2 aligner (Kim et al., 2015) was used to map reads to the genome (Mus musculus.GRCm38 downloaded from Ensembl). The mapped reads of each sample were assembled using StringTie (Pertea et al., 2015) with default parameters. Finally, fragments per kilobase of exon model per million mapped fragments (FPKM) values were calculated by StringTie and ballgown (Frazee et al., 2015; Pertea et al., 2015). Only genes with the average of FPKM values of the 6 time-points in each feeding regimen >0.1 (Mele et al., 2015) or expressed in at least four samples in each sample group were included in the analysis. One sample from muscle in EODF at ZT20 was excluded from analysis due to RNA degradation.

Rhythmicty assay—Rhythmicty was evaluated with DiscoRhythm (Carlucci et al., 2020) using Cosinor assay (Cornelissen, 2014) and JTK Cycle assay (Hughes et al., 2010). Pathways were considered as rhythmic only when Q values of the JTK Cycle assay were less than 0.01.

Gene set variation analysis (GSVA)—The pathway GSV A scores (Hanzelmann et al., 2013) were calculated from the FPKM values based on 2922 gene sets of canonical pathways that were downloaded from the MSigDB (c2.cp.v7.4) (Mootha et al., 2003;
Subramanian et al., 2005). The differential expression analysis was performed on the whole day GSVA scores using limma (Smyth, 2004). The pathways with log2(fold change) > log2(1.5) and adjusted p < 0.05 was considered significant.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**Statistical analysis**—All results are expressed as means ± SEM. Computations assumed that all groups were samples from populations with the same scatter. The investigators involved in this study were not completely blinded during sample collection or data analysis in the animal experiments. Outliers were identified by outlier analysis using the ROUT method of Prism 7.0 software and excluded from statistical analyses. Significance was determined by a two-tailed unpaired t test using Prism 7.0 software. A p < 0.05 was considered significant.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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Highlights

- Canonical pathways are highly rhythmic and sensitive to feeding regimens
- Fasting for 16 h awakens a liver-specific quality controller
- Fasting for 16 h triggers circadian resonance of canonical pathways
- The hepatic proteasome is a fasting “timer” and coordinator of pathway resonance
Figure 1. Canonical pathway profiling of mouse tissues reveals tissue-specific oscillating signatures in intermittent fasting

(A) Experimental design. Four metabolic tissues (liver, skeletal muscle [biceps femoris], interscapular brown adipose tissue [BAT], and inguinal white adipose tissue [WAT]) were collected every 4 h across the light/dark cycle from a single cohort of C57BL/6N mice. Six time points were profiled for all these tissues. n = 5 mice/group.

(B) Heatmaps showing tissue-specific and feeding-regimen-specific oscillatory pathways in four metabolic tissues. Cells were shaded according to GSV A scores (blue, downregulated; red, upregulated). Six columns in each heatmap from left to right are zeitgeber time (ZT) 16, 12, 8, 4, 0, and 24.

(C) Graph showing the ratio of upregulated pathways (dark/light) in liver, muscle, BAT, and WAT tissues.

Cell Rep. Author manuscript; available in PMC 2022 November 17.
ZT20, ZT0, ZT4, ZT8, and ZT12, respectively. The dark/light cycle was indicated by black and white bars (black, dark; white, light). Feeding regimens were indicated on the top of each heatmap (each color from light to dark represented feeding or fasting duration from 4 to 24 h with a step of 4 h). n = 5 mice/group.

(C) Tissue-specific dark/light ratio of upregulated pathways under different feeding regimens. Data are presented as mean ± SEM (n = 5 mice/group). AF, acute fasting; AL, ad libitum; AR, refeeding after acute fasting; BAT, brown adipose tissue; EODF, every-other-day fasting; EODR, every-other-day refeeding; GSVA, gene set variation analysis; WAT, white adipose tissue (see also Figures S1 and S2 and Tables S1 and S2).
Figure 2. Comparative analysis of circadian oscillating pathways highlights the tissue-specific response to different feeding regimens
Circadian oscillating pathways ($Q < 0.01$, Jonckheere-Terpstra-Kendall [JTK] cycle) for each tissue were plotted from left to right according to (1) total number of (#) oscillating pathways, (2) relative percentage of (%) oscillating pathways to detected pathways, (3) graphical summary of the correlations of oscillating pathways across different feeding regimens (ribbon thickness represents the number of correlated pathways), (4) amplitude of oscillation, and (5) phase distribution (the numbers outside the circle are ZTs; the blue-colored circle from light to dark refer to the fasting or feeding time from 0 to 24 h with

Cell Rep. Author manuscript; available in PMC 2022 November 17.
the step of 2 h; gray indicates the dark cycle; the numbers inside radial plot are the number of cycling pathways; different colors of radial plots refer to different feeding regimens as indicated on the top). n = 5 mice/group (see also Tables S2 and S3).
Figure 3. Canonical pathway profiling all day suggests an adaptation to acute fasting-refeeding in intermittent fasting

(A) Data processing. The mean expression values of genes throughout the whole day were calculated from the expression values of six time points in a 24-h day within each feeding regimen and tissue. n = 5 mice/group.

(B and C) The 3D-PCA plots of classes of the whole day gene expression (B) and pathway GSV A score (C) derived from five feeding regimens. n = 5 mice/group.

(D) The pathway Spearman correlation heatmaps showing the evident negative correlation of AF with EODR or AR with EODF in all four metabolic tissues. Cells were shaded according to pathway Spearman correlation coefficients (blue, negative; red, positive) that were calculated from the whole-day GSV scores. Feeding regimens are indicated on the
left and/or the top of each heatmap (each color represents a feeding regimen, and each cell refers to a replicate). n = 5 mice/group.
Figure 4. FRPs exhibit a manifest switch after 16 h fasting or refeeding in the liver
(A–D) The overlap of oscillating pathways (green) and DEPs (brown) in the liver (A), muscle (B), BAT (C), and WAT (D). Numbers in the overlapping regions are the number of FRPs (that is, pathways with both oscillating and differentially expressed), and other numbers in the circle are the number of pathways showing only oscillating or differentially expressed, respectively. n = 5 mice/group.
(E–H) Heatmap exhibiting the changes of GSVA scores of FRPs across different times of five feeding regimens in the liver (E), muscle (F), BAT (G), and WAT (H). Black arrows referred to the 16th hour of treatments (fasting or feeding), which was equal to the ZT4. Top color bars, different feeding regimens; bottom filled bars, dark; hollow bars, light. n = 5 mice/group. DEPs, differentially expressed pathways; FRPs, functional rhythmic pathways (see also Tables S2 and S4).
**Figure 5. Intersection of oscillating and DEPs**

Detailed heatmap of alteration frequencies in numbers displaying the intersections of the oscillating pathways (rows) and DEPs (columns) in different feeding regimens and tissues. Each column shows the distribution of DEPs by feeding-regimen-specific oscillating phases, and each row shows the distribution of phase-specific oscillating pathways by feeding-regimen-specific DEPs. Red, upregulated events; blue, downregulated events (color intensity increases with the frequency of occurrence). Color top bar charts show the fraction of feeding-regimen-specific oscillating pathways in DEPs of a specific feeding regimen.
and tissue. Color side bar charts show the fraction of feeding-regimen-specific DEPs in each phase of oscillating pathways in a specific feeding regimen. n = 5 mice/group. dn, downregulated; ET, executing time duration for fasting or feeding; up, upregulated (see also Figure S3 and Tables S2 and S3).
Figure 6. Circadian resonance of 43 pathways in the liver highlights a proteasome switch in intermittent fasting

(A) The circadian resonance of 43 functional pathways in the liver of AF mouse. n = 5 mice/group.

(B) The categories of the 43 resonant functional pathways according to the Reactome Pathway Database.

(C) List of shared genes by the 43 resonant functional pathways (see also Figure S2 and Table S2).
Figure 7. Circadian transcriptional changes of proteasome subunits confirm the switching effect after 16 h of fasting or refeeding in the liver

(A–D) The circadian transcriptional changes of proteasome subunits under different feeding regimens in the liver (A), muscle (B), BAT (C), and WAT (D). Each dotted red line refers to a proteasome subunit, and the black line refers to the average trend of all proteasome subunits. Shadow, dark; blank, light. n = 5 mice/group.

(E) The circadian transcriptional changes of three proteasome proteins under different feeding regimens in the liver. Each dotted red line refers to normalized mean densitometry values of a proteasome protein from western blot in Figure S4, and the black line refers to
the average trend of all three proteasome proteins. Densitometry values were normalized to actin beta (ACTB) expression for each lane and presented as fold change of AL24 mice. Shadow, dark; blank, light. n = 2 mice/group (see also Figures S4, S5, and S6 and Tables S1 and S5).
## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Rabbit PSMB3 antibody | Beijing Solarbio Science & Technology Co., Ltd | Cat# K009840P |
| Rabbit PSMB6 antibody | Beijing Solarbio Science & Technology Co., Ltd | Cat# K003745P |
| Rabbit PSMA4 antibody | Beijing Solarbio Science & Technology Co., Ltd | Cat# K003788P |
| Rabbit Actin β Polyclonal antibody | ImmunoWay Biotechnology Co. | Cat# YT0099; RRID:AB_2885029 |
| HRP Goat Anti-Rabbit IgG (H + L) | Clinitrines | Cat# AS014; RRID:AB_2769854 |
| **Chemicals, peptides, and recombinant proteins** | | |
| Anhydrous ethanol | Sinopharm Chemical Reagent Beijing Co Ltd | Cat# c003860 |
| Alcohol isopropyl | Sinopharm Chemical Reagent Beijing Co Ltd | Cat#CFSR-40049962 |
| β-Mercaptoethanol | Sinopharm Chemical Reagent Beijing Co Ltd | Cat# 0482 |
| **Critical commercial assays** | | |
| AMPureXP beads | Beckman | Cat# A63881 |
| Dynabeads Oligo (dT)25-61005 | Thermo Fisher | Cat# 61005 |
| dUTP Solution | Thermo Fisher | Cat# R0133 |
| E. coli DNA polymerase I | New England Biolabs | Cat# M0209 |
| Magnesium RNA Fragmentation Module | New England Biolabs | Cat# E6150 |
| RNase H | New England Biolabs | Cat# M0297 |
| Uracil-DNA Glycosylase | New England Biolabs | Cat# M0280 |
| SuperScript™ II Reverse Transcriptase | Invitrogen | Cat# 1896649 |
| Ketone body Assay Kit | Beijing Solarbio Science & Technology Co., Ltd | Cat# BC5060 |
| Enhanced Chemiluminescent Kit | New Cell & Molecular Biotech Co., Ltd | Cat# P10300 |
| **Deposited data** | | |
| RNA sequencing data | This paper | GEO: GSE154797 |
| Western Blot data | This paper; Mendeley Data | Mendeley Data: https://doi.org/10.17632/x64s7p7cb7.1 |
| **Experimental models: Organisms/strains** | | |
| Mouse: C57BL/6N | Beijing Vital River Laboratory Animal Technology Co., Ltd. | Strain Code: 213 |
| **Software and algorithms** | | |
| ImageJ | Schneider et al., 2012 | https://imagej.nih.gov/ij/; RRID:SCR_003070 |
| cutadapt-1.9 | https://cutadapt.readthedocs.io/en/stable/ | https://doi.org/10.14806/ej.17.1.200; RRID: SCR_011841 |
| DESeq2 | http://www.bioconductor.org/packages/release/bioc/html/DESeq2.html | https://doi.org/10.18129/B9.bioc.DESeq2; RRID:SCR_015687 |
| DiscoRhythm 1.2.1 | https://mcarlucci.shinyapps.io/discoRhythm/ | N/A |
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| edgeR               | https://bioconductor.org/packages/release/bioc/html/edgeR.html | doi:10.18129/B9.bioc.edgeR |
| GraphPad Prism 7.0  | http://www.graphpad.com/ | RRID:SCR_002798 |
| gffcompare-0.9.8.Linux_x86_64 | http://ccb.jhu.edu/software/stringtie/gffcompare.shtml | |
| HISAT2-2.0.4        | https://daehwankimlab.github.io/hisat2/ | RRID:SCR_015530 |
| StringTie-1.3.4d.Linux_x86_64 | http://ccb.jhu.edu/software/stringtie/ | RRID:SCR_016323 |
| GSV A 1.42.0        | https://www.bioconductor.org/packages/release/bioc/html/GSV A.html | doi:10.18129/B9.bioc.GSV A; RRID:SCR_021058 |
| scatterplot3d 0.3.41 | https://cran.r-project.org/web/packages/scatterplot3d/index.html | doi:10.18637/jss.v008.i11 |
| STAT 0.1.0          | https://cran.r-project.org/web/packages/STAT/index.html | N/A |
| limma 3.50.0        | https://bioconductor.org/packages/release/bioc/html/limma.html | doi:10.18129/B9.bioc.limma; RRID:SCR_010943 |
| Circos online      | http://mkweb.bcgsc.ca/tableviewer/ | RRID:SCR_011798 |