

1 A role for gene expression and mRNA stability in nutritional compensation of the circadian clock

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35 **Abstract**

36 Compensation is a defining principle of a true circadian clock, where its approximately
37 24-hour period length is relatively unchanged across environmental conditions. Known
38 compensation effectors directly regulate core clock factors to buffer the oscillator's period length
39 from variables in the environment. Temperature Compensation mechanisms have been
40 experimentally addressed across circadian model systems, but much less is known about the
41 related process of Nutritional Compensation, where circadian period length is maintained across
42 physiologically relevant nutrient levels. Using the filamentous fungus *Neurospora crassa*, we
43 performed a genetic screen under glucose and amino acid starvation conditions to identify new
44 regulators of Nutritional Compensation. Our screen uncovered 16 novel mutants, and together
45 with 4 mutants characterized in prior work, a model emerges where Nutritional Compensation of
46 the fungal clock is achieved at the levels of transcription, chromatin regulation, and mRNA
47 stability. However, eukaryotic circadian Nutritional Compensation is completely unstudied
48 outside of *Neurospora*. To test for conservation in cultured mammalian cells, we selected the
49 top two hits from our fungal genetic screen, performed siRNA knockdown experiments of the
50 mammalian homologs, and characterized the cell lines with respect to compensation. We find
51 that the wild-type mammalian clock is also compensated across a large range of external
52 glucose concentrations, as observed in *Neurospora*, and that knocking down *CPSF6* or *SETD2*
53 in human cells also results in nutrient-dependent period length changes. We conclude that, like
54 Temperature Compensation, Nutritional Compensation is a conserved circadian process in
55 fungal and mammalian clocks and that it may share common molecular determinants.

56

57 **Introduction**

58 Circadian clocks exist at the cellular level to allow cell types, tissues, and organisms to
59 properly align physiology with time of day. True circadian clocks are sensitive to the external
60 environment in two distinct ways. Discrete pulses of bright light, temperature, nutrients,
61 hormones, or other chemicals reset circadian oscillators and re-orient the clock's phase to the
62 new environment (reviewed in: Johnson et al., 2003). This resetting feature of circadian clocks
63 is most commonly experienced when jetlagged humans travel across multiple time zones and
64 entrain to the destination's light/dark cycles. On the other hand, circadian clocks are also
65 shielded or buffered from changes in the ambient environment within the physiological range of
66 an organism (Hastings and Sweeney, 1957; Pittendrigh et al., 1959). In the filamentous fungus
67 *Neurospora crassa*, circadian period length is maintained at approximately 21.5 hours when
68 grown at constant temperatures ranging from 16°C to 32°C (Gardner and Feldman, 1981) or

69 under different nutrient conditions (Sargent and Kaltenborn, 1972). This circadian property is
70 known as compensation, and the molecular mechanisms underlying period length
71 compensation remain elusive.

72 At the molecular level, the circadian oscillator is a transcription-translation feedback loop
73 (TTFL) that is functionally conserved from fungi to animals. The positive arm of the clock is
74 composed of transcription factor activators (WC-1/WC-2 in fungi; BMAL1/CLOCK in mammals),
75 which form a heterodimeric complex, drive transcription of direct target genes, and recruit
76 chromatin modifiers (Crosthwaite et al., 1997; Koike et al., 2012; Menet et al., 2014). The
77 positive arm directly regulates the negative arm of the clock (FRQ in fungi; PERs/CRYs in
78 mammals) (Aronson et al., 1994; reviewed in: Dunlap, 1999; Philpott et al., 2021). Negative arm
79 clock components form a stable complex with Casein Kinase I (CKI) and other factors, leading
80 to feedback and posttranslational inhibition of the positive arm to close the circadian feedback
81 loop (Wang et al., 2019; Cao et al., 2021). The positive and negative arms are sufficient for
82 rhythmicity, although accessory feedback loops confer additional clock robustness (reviewed in:
83 Takahashi, 2017). Such individual cellular clocks are coordinated in a coupled network to align
84 organismal physiology in mammals (reviewed in: Finger et al., 2020).

85 Temperature Compensation was first proposed to be a circadian property in the
86 dinoflagellate *Gonyaulax polyedra* when increasing temperatures led first to period lengthening
87 (so-called “over-compensation”) and then to period shortening at even higher temperatures, a
88 result that plainly conflicted with models based on biochemical reaction rates strictly increasing
89 with temperature (Hastings and Sweeney, 1957). In genetic model systems like *Neurospora* and
90 *Drosophila* where organisms operate at ambient temperatures, and even in homeothermic
91 animals, cellular circadian clocks are temperature compensated (Zimmerman et al., 1968;
92 Gardner and Feldman, 1981; Barrett and Takahashi, 1995; Izumo et al., 2003; Tsuchiya et al.,
93 2003; Kidd et al., 2015). The forward and reverse genetics that have driven current models for
94 Temperature Compensation have led to casein kinases as central regulators across multiple
95 circadian model systems. In *Neurospora* and in plants, Casein Kinase II (CKII) is required for
96 Temperature Compensation (Mehra et al., 2009; Portolés and Más, 2010). *Neurospora* CKII
97 activity increases linearly with temperature and directly phosphorylates the negative arm of the
98 clock (Mehra et al., 2009). The *tau* mutant hamster (CKI ϵ^{R178C}) was the first characterized
99 mammalian Temperature Compensation defect (Ralph and Menaker, 1988; Tosini and
100 Menaker, 1998; Lowrey et al., 2000). Recent structural work has demonstrated that CKI *tau*
101 alters both priming and progressive phosphorylation events on the negative arm of the clock
102 (Philpott et al., 2020), which may account for its period shortening with temperature (so-called

103 “under-compensation”). Indeed, temperature-sensitive target binding has been shown for
104 mammalian CKI (Shinohara et al., 2017), and the interaction strength between CKI and FRQ in
105 *Neurospora* has been implicated in Temperature Compensation as well (Hu et al., 2021). Taken
106 together, CKI and CKII each have temperature sensitive aspects of enzyme activity, both can
107 directly phosphorylate core clock components, and casein kinases appear to play a conserved
108 role in Temperature Compensation of fungal, plant, and animal clocks.

109 In contrast to Temperature Compensation, mechanisms underlying Nutritional
110 Compensation (also known as Metabolic or Glucose Compensation) are poorly understood in
111 *Neurospora* and completely unstudied in other eukaryotic circadian systems. Period
112 compensation to a variety of ATP:ADP ratios has been well described in the prokaryotic
113 cyanobacteria *Synechococcus elongatus* (reviewed in: Johnson and Egli, 2014). A handful of
114 Nutritional Compensation defects have arisen sporadically in *Neurospora*, the most developed
115 involving the transcription factor repressor CSP-1 (Lambrechts et al., 2009). CSP-1 directly
116 regulates and is regulated by the clock’s positive arm White Collar Complex (WCC), forming an
117 accessory negative feedback loop. In a $\Delta csp-1$ mutant, period significantly shortens as a
118 function of glucose concentration (Sancar et al., 2012). In fact, direct overexpression of *wc-1*
119 also causes nutritional under-compensation (Dovzhenok et al., 2015). Nutritional Compensation
120 is defective in the absence of the general transcription repressor RCO-1 (Olivares-Yanez et al.,
121 2016), likely due its normal role in preventing WCC-independent *frq* transcription (Zhou et al.,
122 2013). Over-compensation was also found in loss-of-function mutants of an RNA helicase,
123 PRD-1, which normally localizes to the nucleus only under high glucose conditions (Emerson et
124 al., 2015). Nutrient sensing and signaling pathways should presumably also play a role in
125 Nutritional Compensation of the clock, and RAS2 and cAMP signaling have been implicated
126 (Gyöngyösi et al., 2017). Taken together, the current incomplete model for Nutritional
127 Compensation in *Neurospora* assembled from random hits implicates transcriptional and post-
128 transcriptional regulation of core clock factors by transcription factors, an RNA helicase, and
129 cAMP signaling.

130 Since the realization of circadian compensation, the field has speculated that
131 Temperature and Nutritional Compensation pathways may share common regulators
132 (Pittendrigh and Calderola, 1973; Roenneberg and Merrow, 1999; Johnson and Egli, 2014). We
133 directly test this model and find in *Neurospora* that previously reported compensation mutants
134 are specific to either Temperature or Nutritional Compensation. Given this separation of function
135 and relatively little mechanistic knowledge about Nutritional Compensation, we designed a
136 genetic screen to identify new compensation mutants in *Neurospora crassa*. We identify 16

137 gene knockouts with Nutritional Compensation phenotypes, greatly expanding the list of 4
138 previously characterized mutants. We also provide the first evidence that, like Temperature
139 Compensation, Nutritional Compensation is relevant in the mammalian circadian system, and
140 confirm that knockdowns of the human genes homologous to the top two hits from our genetic
141 screen, *CPSF6* and *SETD2*, impair the clock's ability to maintain its period length at different
142 glucose levels. These data establish a potentially conserved genetic basis for the phenomenon
143 of circadian Nutritional Compensation and anchor the phenomenon for further genetic and
144 molecular dissection.

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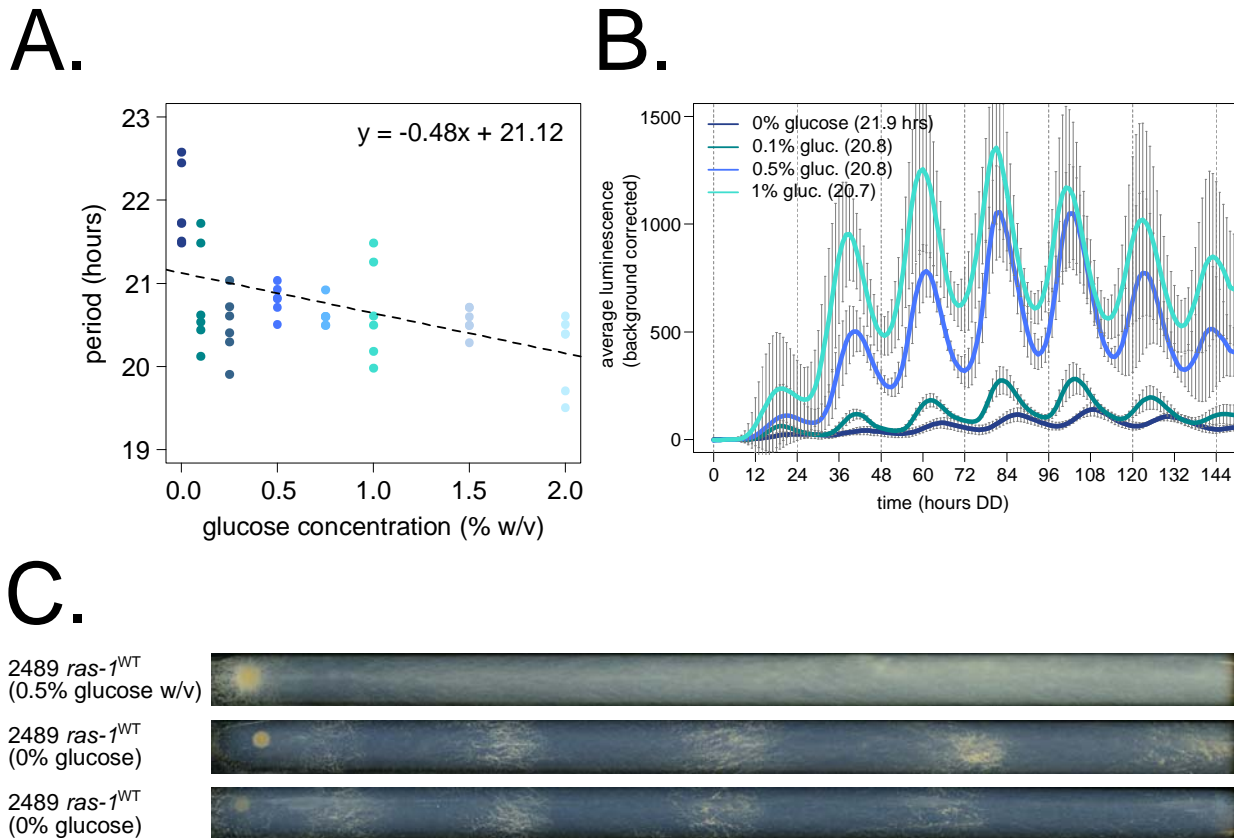
146 **Results**

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148 **Nutritional Compensation is distinct from Temperature Compensation in *Neurospora*.**

149 We first set out to characterize the properties of Nutritional Compensation in wild-type
150 *Neurospora*. Traditional circadian experiments have utilized the *ras-1^{bd}* mutant, which promotes
151 the formation of circadianly-regulated distinct bands of conidial spores in a race tube assay
152 (Sargent et al., 1966; Belden et al., 2007a). However, the *ras-1* gene (NCU08823) is implicated
153 in growth and regulation of reactive oxygen species. Thus to accurately profile normal
154 Nutritional Compensation, a wild-type *ras-1⁺* strain containing a *frq* clock box transcriptional
155 reporter was used to measure period length across glucose concentrations ranging from 0 –
156 111 mM (0 – 2% w/v). The *Neurospora* circadian period is slightly under-compensated to
157 nutrients (Figure 1A); under-compensation has also been observed with respect to temperature
158 (Mehra et al., 2009; Hu et al., 2021). Fungal biomass increases by orders of magnitude when
159 grown in the range of 0% – 0.75% w/v glucose (Supplementary Figure 1), and this increased
160 biomass accounts for the increased magnitude of luciferase rhythms observed at higher glucose
161 levels (Figure 1B) (Supplementary Movie 1).

162



164 **Figure 1. Nutritional Compensation properties of *Neurospora crassa*.** Circadian period
165 length was determined by bioluminescent recordings of a *ras-1*^{WT} strain cultured on race tubes,
166 where the fungal growth front encounters constant glucose concentrations (N = 6 race tubes per
167 concentration). No arginine was added to the medium. Period shortens slightly as a function of
168 glucose levels, which is indicated by the negative slope of the linear fit (g1m in R, Gaussian
169 family defaults, slope = -0.48 ± 0.12) (A). Averaged biological and technical replicates are
170 shown for 0 mM (0% w/v), 5.6 mM (0.1% w/v), 27.8 mM (0.5% w/v), and 55.5 mM (1% w/v)
171 glucose levels (standard deviation error bars). Period lengths are 21.9 ± 0.5 , 20.8 ± 0.6 , $20.8 \pm$
172 0.2 , and 20.7 ± 0.6 hours, respectively (average \pm SD) (B). Surprisingly, the *ras-1*^{WT} strain can
173 form distinct conidial bands when grown on 0% glucose starvation medium, contrasted with
174 constitutive conidiation seen at high glucose levels (C).

175
176 Having established that the *Neurospora* clock displays compensation for period length
177 across glucose concentrations, we asked whether Temperature Compensation mutants also
178 have Nutritional Compensation phenotypes, and vice versa, together in the same assay. CKII is
179 required for normal Temperature Compensation, and its catalytic subunit mutant *cka*^{prd-3} (Y43H)

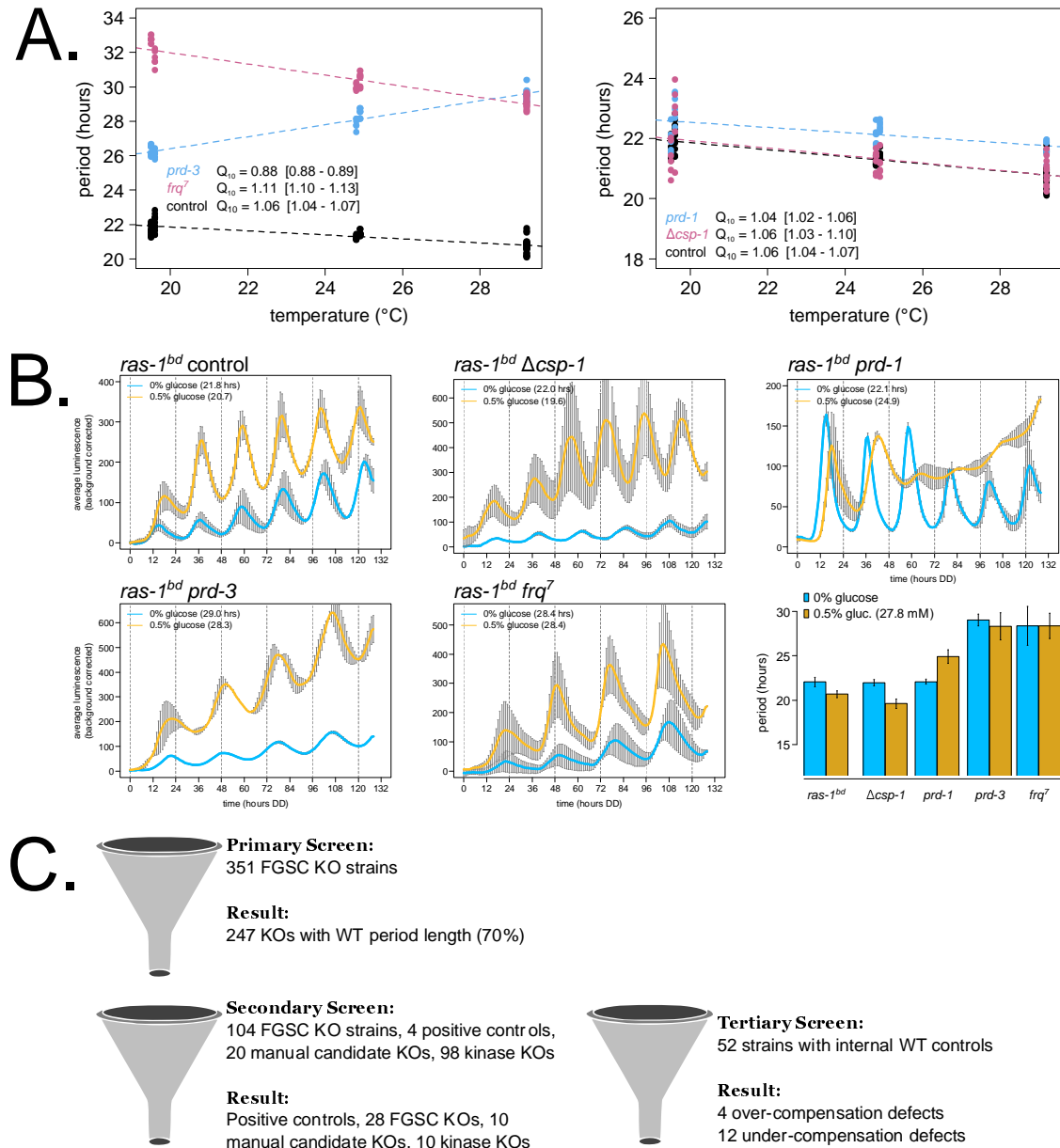
180 has an over-compensation phenotype by race tube assay (Mehra et al., 2009). The *frq*⁷
181 (G459D) point mutant is temperature under-compensated (Gardner and Feldman, 1981). The
182 *frq* clock box transcriptional reporter was integrated into the *cka*^{prd-3} and *frq*⁷ mutant
183 backgrounds, and temperature over- ($Q_{10} = 0.88$) and under-compensation ($Q_{10} = 1.11$)
184 phenotypes were confirmed by luciferase assay (Figure 2A). Known Nutritional Compensation
185 mutants *prd-1* and Δ *csp-1* were also tested, and each had normal Temperature Compensation
186 profiles ($Q_{10} = 1.04, 1.06$). The same 4 compensation mutant reporter strains were then grown
187 on race tubes containing zero or high glucose medium, and bioluminescence was recorded to
188 track period length and Nutritional Compensation phenotypes. Controls were slightly under-
189 compensated (Figure 1A, Figure 2B). Nutritional Compensation mutants *prd-1* and Δ *csp-1*
190 showed the over- and under-compensation phenotypes reported by previous studies (Sancar et
191 al., 2012; Emerson et al., 2015). Temperature Compensation mutants *cka*^{prd-3} and *frq*⁷ have
192 normal Nutritional Compensation (Figure 2B). We conclude that Temperature and Nutritional
193 Compensation are controlled by distinct pathways in *Neurospora*. These data suggest that
194 further examination of available mutants defective in Temperature Compensation will not inform
195 our understanding of Nutritional Compensation, and that a separate genetic screen is
196 warranted.

197 To achieve this, we leveraged the whole genome knockout collection in *Neurospora*
198 *crassa* (Colot et al., 2006) to initiate a screen for new Nutritional Compensation regulators. Two
199 major classes of gene knockouts were selected to search for compensation phenotypes.
200 Kinases are central to many aspects of cellular processes and regulation including responses to
201 the environment (pheromones, osmotic conditions, carbon/nitrogen regulation, etc.), and a
202 collection of ~100 different kinase knockout circadian reporter strains was available from
203 previous work (Dasgupta, 2015). In addition to posttranslational modifications by kinases,
204 posttranscriptional regulation is emerging as critically important for circadian output (Hurley et
205 al., 2018). Together with the dramatic nutritional over-compensation defect seen in *prd-1* RNA
206 helicase mutants (Emerson et al., 2015) (Figure 2B), we selected 351 putative RNA regulatory
207 protein knockouts to screen for compensation regulators. Our list of putative RNA-binding and
208 RNA regulatory proteins was derived from bioinformatic databases and from the literature (Ray
209 et al., 2013; Hogan et al., 2015; Zaveri et al., 2017; Basenko et al., 2018) to include proteins
210 with nucleotide-binding functional annotation but exclude known transcriptional regulators.
211 Multiple circadian period alterations were identified in a recent characterization of transcription
212 factor knockouts (Muñoz-Guzmán et al., 2021), and Nutritional Compensation defects among
213 transcription factors, in addition to CSP-1 and RCO-1, will be the subject of future study. Finally,

214 a handful of manually selected candidate genes were included in the compensation screen. 8
215 classical alleles with long or short circadian period (the series of *prd* mutants and *frq* point
216 mutants) were included, and 12 knockouts were manually selected based on reported roles in
217 nutrient sensing and signaling.

218 The genetic screen for Nutritional Compensation defects was divided into three phases
219 (Figure 2C). The 351 previously uncharacterized RNA regulatory protein knockout strains were
220 grown on glucose starvation race tubes (Figure 1C) to identify those with period length
221 differences ± 1 hour as compared to internal wild-type controls ($N = 2 - 8$ replicate race tubes
222 per KO strain). Approximately 70% of the putative RNA regulatory protein knockout strains
223 showed normal period length on starvation medium and were eliminated in the primary screen
224 (Figure 2C, Supplementary Table 1). During this phase of the screen, 7 knockout strains were
225 found with enhanced conidial banding patterns (and variable growth rates) relative to wild-type
226 controls, reminiscent of the *ras-1^{bd}* mutant phenotype (Supplementary Figure 2). In the
227 secondary screen, the *frq* clock box transcriptional reporter was integrated into all knockouts of
228 interest, along with the existing kinase deletion collection (Dasgupta, 2015) and manually
229 selected candidate strains. This collection of more than 200 strains was screened in a high
230 throughput 96-well plate format using glucose and amino acid starvation medium to identify
231 period length differences ± 1 hour from internal wild-type controls ($N = 6 - 12$ replicate wells per
232 KO strain). About 77% of the candidate strains were eliminated during the secondary screen
233 due to normal period length on starvation medium (Figure 2C, Supplementary Table 1). The ~ 50
234 remaining knockout strains and wild-type controls were then advanced to the lowest throughput
235 tertiary screen: bioluminescence race tube assays directly comparing zero versus high glucose
236 and amino acid medium (0.5% w/v glucose, 0.17% w/v arginine). It should be noted here that
237 the race tube assay is particularly well suited for a Nutritional Compensation mutant screen
238 because the growth front, which produces most of the bioluminescence, always encounters
239 fresh medium; thus there is no complicating effect of nutrient depletion over the course of a six-
240 day assay (Supplementary Movie 1). Circadian period lengths were quantified to assay
241 Nutritional Compensation phenotypes. 16 new mutants emerged from the genetic screen
242 showing large period changes between zero and high nutrient conditions (Figure 2C,
243 Supplementary Table 1).

244



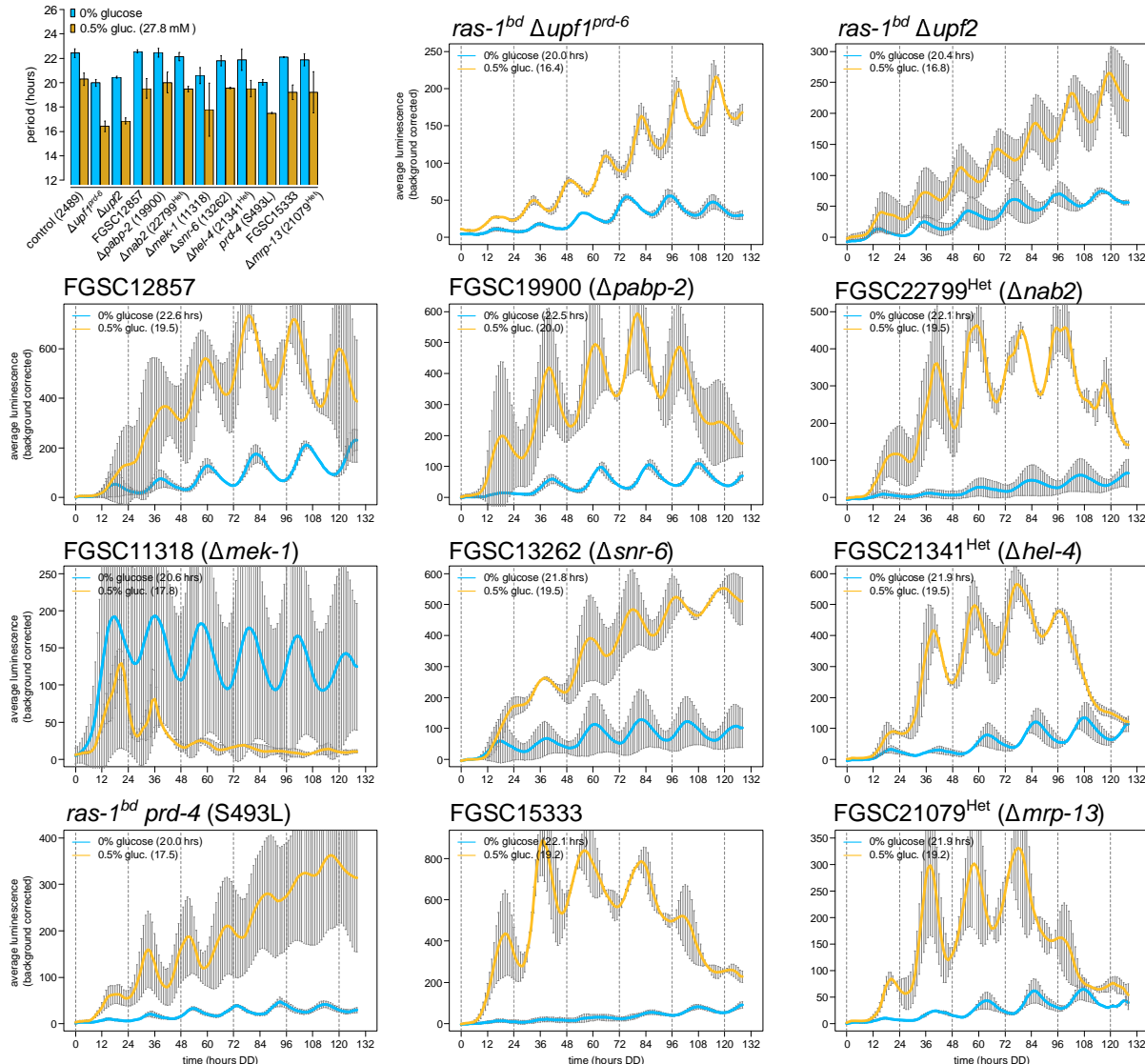
245

246 **Figure 2. Nutritional Compensation mutants have normal Temperature Compensation in**
 247 ***Neurospora*, catalyzing a genetic screen for new mutants.** 96-well plate luciferase assays
 248 were used to measure the circadian period length across temperatures in constant darkness (N
 249 = 12 replicates per strain, per temperature). A linear model was fit to period length data from
 250 each strain (`g1m` in R), and a Q_{10} temperature coefficient was calculated using the model-fitted
 251 period lengths at 20°C and 30°C. Shaded areas around the linear fit represent the 95%
 252 confidence intervals on the slope. Error ranges on Q_{10} values were computed from the 95% CIs
 253 (**A**). Circadian bioluminescence was recorded from race tube cultures of the indicated
 254 genotypes, as previously described (Larrondo et al., 2012) (Supplementary Movie 1). High

255 nutrient medium (yellow lines) contained 0.5% w/v glucose 0.17% w/v arginine, and zero
256 nutrient medium (blue lines) contained 0% glucose 0% arginine. Period lengths were computed
257 (N = 2 – 5 biological replicates per nutrient concentration) and summarized in a bar graph (**B**).
258 Cartoon depiction of the 3-phase genetic screen for Nutritional Compensation defects among
259 kinase, RNA regulatory proteins, and manual candidate knockout strains (**C**).
260

261 **mRNA stability and polyadenylation factors emerge as key regulators of circadian period
262 length across nutrients.**

263 We first examined the group of 12 under-compensation mutants, where period length
264 shortens as a function of nutrient levels (Figure 3, Supplementary Table 1). The two most
265 significant hits are subunits of the nonsense-mediated decay machinery in *Neurospora* and
266 showed more dramatic nutrient-responsive period shortening than $\Delta csp-1$ (Sancar et al., 2012).
267 Additionally, 3 of the 12 mutants share a common function in polyadenylation of nascent RNAs.
268 NCU02736 (FGSC12857) is designated as an uncharacterized gene in *Neurospora* but is
269 broadly conserved in fungi. Its *Saccharomyces cerevisiae* homolog is a component of the
270 mRNA cleavage and polyadenylation factor I complex (YGL044C, *RNA15*). PABP-2
271 (NCU03946, FGSC19900) binds in poly(A) tail regions and can broadly regulate mRNA stability.
272 NAB2 (NCU16397, FGSC22799^{Het}) shares a common domain with the yeast gene *NAB2*
273 (YGL122C), which plays a role in mRNA export and stability. Taken together, regulation of
274 polyadenylation and/or mRNA stability is a common axis of period maintenance with increasing
275 nutrients. The other 6 under-compensation mutants did not share an obvious functional
276 pathway, although some of these genes do show circadian rhythms at the mRNA or protein
277 level and/or are regulated in response to light (Supplementary Table 2). However, given the
278 slight under-compensation phenotype described for the wild-type clock in *Neurospora* (Figure
279 1A, Figure 2B, Figure 3), the remaining under-compensation mutants were not pursued further.
280



281

282 **Figure 3. Nutritional under-compensation mutants are enriched for regulators of mRNA**
283 **stability.** Circadian bioluminescence was recorded from race tube cultures of the indicated
284 deletion mutants. High nutrient medium (yellow lines) contained 0.5% w/v glucose 0.17% w/v
285 arginine, and zero nutrient medium (blue lines) contained 0% glucose 0% arginine. Period
286 lengths were computed ($N \geq 2$ biological replicate period estimates per nutrient concentration)
287 and summarized in a bar graph compared to controls. “Het” indicates heterokaryon strains
288 derived from the *Neurospora* whole genome deletion collection, which were maintained on
289 hygromycin selection medium prior to the bioluminescence race tube assays.

290

291 The nonsense-mediated decay (NMD) pathway is required for a normal circadian period
292 length due to its regulation of *casein kinase 1* mRNA levels (Kelliher et al., 2020a). We

293 hypothesized that NMD regulation of *ck-1a* also underlies its nutritional under-compensation
294 defect. NMD in *Neurospora* can be triggered by long 3' UTR sequences, and the *ck-1a* 3' UTR
295 is among the longest 1% of UTRs in *Neurospora* (Kelliher et al., 2020a). We generated a *ck-1a*
296 mutant strain lacking its entire 3' UTR, decoupling *ck-1a* transcripts from NMD targeting. The *ck-1a*
297 Δ 3'UTR strain does indeed have a short period length phenocopying that of NMD mutants;
298 however, the *ck-1a* Δ 3'UTR under-compensation phenotype is not as severe as observed in
299 NMD mutants (Supplementary Figure 3). This suggests that multiple NMD targets are required
300 for normal Nutritional Compensation. Interestingly, NMD mutants also show ~1.4-fold changes
301 in *wc-2* and *frh* gene expression levels (Kelliher et al., 2020a; Figure 5B; Wu et al., 2017). *wc-2*
302 is upregulated in NMD mutants, which could easily explain the under-compensation phenotype
303 because *wc-1* overexpression alone is sufficient to drive nutritional under-compensation
304 (Dovzhenok et al., 2015). Curiously, we find that *frh*, the obligate binding partner of the
305 disordered protein FRQ (Hurley et al., 2013), is 18-fold down-regulated in response to carbon
306 starvation in wild-type *Neurospora* (Wang et al., 2017) (Supplementary Figure 4). In fact, both
307 *prd-1* and *frh* are among the top 220 genes in the entire *Neurospora* transcriptome that
308 decrease in expression after glucose starvation. Future work will determine whether NMD
309 regulation of *ck-1a*, *wc-2*, *frh*, or all transcripts explains the nutritional under-compensation
310 defect.

311 Mutations in two genes, FGSC16956 and FGSC12033, revealed unique nutritional
312 phenotypes compared to the remaining set of 11 under-compensation mutants (Figure 3).
313 FGSC16956 (Δ NCU02152) had the shortest period phenotype observed in the entire primary
314 and secondary screens (Supplementary Table 1). NCU02152 was undescribed in the
315 *Neurospora* literature but contains protein domain homology to the mammalian Cleavage &
316 Polyadenylation Specificity Factor subunit 6 (CPSF6). CPSF6 is a member of the CFI_m
317 complex, which binds in the 3' end of nascent mRNAs and facilitates cleavage and poly(A) tail
318 placement. The CFI_m complex also contains a second essential component CPSF5. Using the
319 human CPSF5 protein sequence, NCU09014 was confidently identified as its *Neurospora*
320 homolog (reciprocal BLAST e-values = $1e^{-72}$ / $3e^{-74}$). The single mutant FGSC12033
321 (Δ NCU09014 / Δ cpsf5) had the same short period length as Δ cpsf6 (Supplementary Table 1),
322 which indicates an obligate multimeric complex as with the mammalian CFI_m complex. In the
323 tertiary screen, Δ cpsf5 and Δ cpsf6 mutants showed progressive period shortening after ~60
324 hours into the circadian free run, specifically when grown on high nutrient medium. Recalling the
325 *prd-1* region-specific Nutritional Compensation phenotype where period defects were seen at
326 the growth front encountering fresh medium, but not at the point of inoculation where nutrients

327 were depleted (Emerson et al., 2015), we checked period length from old/aging tissue
328 surrounding the point of fungal inoculation in $\Delta cpsf5$ and $\Delta cpsf6$ mutants. An ~18-hour short
329 period length was observed in the old tissue region of $\Delta cpsf5$ and $\Delta cpsf6$ mutants grown on high
330 arginine (Figure 4A), compared to a ~20-hour period length on zero nutrient medium
331 (Supplementary Table 1). This result indicates that $\Delta cpsf5$ and $\Delta cpsf6$ mutants must undergo a
332 transition from high-to-low amino acid levels in old tissue to reveal the 18-hour period defect
333 (Supplementary Movie 2). The $\Delta cpsf5$ and $\Delta cpsf6$ mutants differ from other under-compensation
334 mutants (Figure 3) because the short period defect was induced by amino acids, not by glucose,
335 and because the Nutritional Compensation phenotype is specific to old tissue (Figure 4A).
336

337 **The Alternative Polyadenylation (APA) landscape is altered in Nutritional Compensation
338 mutants.**

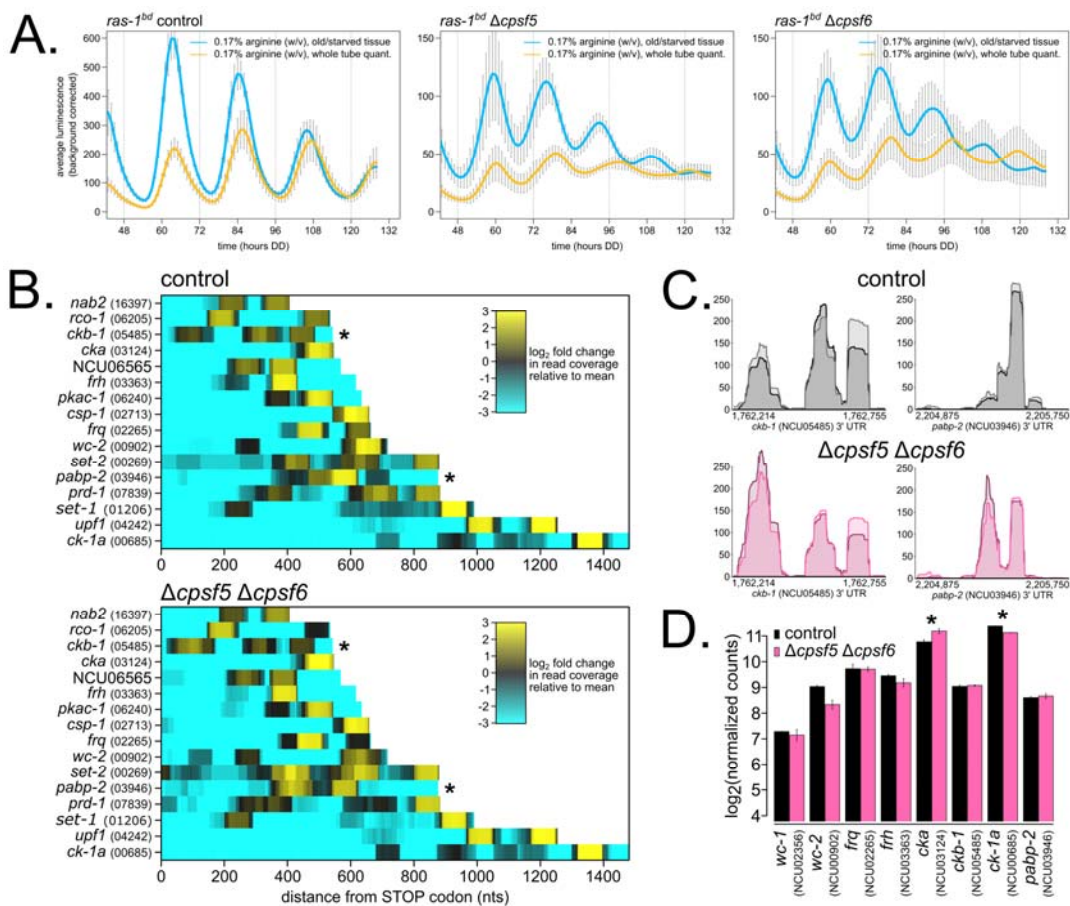
339 Given the large period defect observed in $\Delta cpsf5$ and $\Delta cpsf6$ mutants (Figure 4A) and
340 other under-compensation mutants related to polyadenylation (Figure 3), we hypothesized that
341 poly(A) tail maintenance and concurrently the stability of a core clock mRNA(s) is required for
342 Nutritional Compensation. To assay the biochemistry of $\Delta cpsf$ mutants, we required a solid
343 medium growth regime to carefully control nutrient levels and to confidently compare results to
344 our genetic screen for Nutritional Compensation phenotypes. *Neurospora* biochemistry is
345 traditionally accomplished using extracts from liquid-grown cultures (Nakashima, 1981; Kelliher
346 et al., 2020b), where nutrient consumption rates are less well defined and less relevant to the
347 ecological niche of the organism. A cellophane petri plate assay was developed to harvest
348 biomolecules from solid medium cultures of *Neurospora* (Materials and Methods). We confirmed
349 that the circadian clock is equally functional on cellophane plates and in liquid cultures
350 (Supplementary Figure 5). We next generated a $\Delta cpsf5 \Delta cpsf6$ double mutant strain (hereafter
351 referred to as ΔCFI) and confirmed that its period length and Nutritional Compensation
352 phenotype matched results from race tube assays (Supplementary Figure 5). Biological
353 duplicate wild-type control and ΔCFI mutant cellophane plates were grown at 25°C under
354 constant light for 3 days on high nutrient medium (containing 0.25% w/v glucose 0.17% w/v
355 arginine; high-to-low nutrient transition will occur in aged tissue before 72 hours growth), and
356 total RNA was extracted for 3' End Sequencing (Materials and Methods).

357 Nascent mRNA transcripts can contain multiple sites for polyadenylation to occur, which
358 is known as Alternative Polyadenylation (APA). APA events generate mRNA isoforms with
359 variable 3' UTR lengths and nucleotide sequences (reviewed in: Mayr, 2017). Along with other

360 factors involved in mRNA 3' end cleavage and polyadenylation, CFI m mutants have been
361 shown to alter APA patterns genome wide (Kubo et al., 2006; Martin et al., 2012; Zhu et al.,
362 2018). Changes in APA have been directly linked to RNA stability in yeast and in mammals
363 (Mayr and Bartel, 2009; Moqtaderi et al., 2022). We first sought to identify normal instances of
364 APA in wild-type *Neurospora* 3' UTRs by comparing our 3' End Sequencing dataset to an
365 existing 2P-Seq dataset (Zhou et al., 2018) (Materials and Methods). 843 consensus genes
366 (>10% of *Neurospora* 3' UTRs) contain multiple poly(A) sites after applying a strict intersection
367 cutoff for identification in all 4 wild-type datasets (Supplementary Table 3), which is a lower
368 genome-wide estimate than >60% APA in mammals. Next comparing Δ CFI m and controls, we
369 found 193 examples of APA events in controls collapsing to a single poly(A) peak in mutants
370 (21%), 123 examples of a single poly(A) peak in controls expanding to multiple APA events in
371 mutants (13%), and 155 examples of APA events in both control and mutant where the location
372 of the predominant poly(A) peak was significantly changed in mutants (16%) (Supplementary
373 Table 4). Taken together, ~50% of the APA landscape is altered in *Neurospora* Δ CFI m mutants.
374 Knockdown of the mammalian CFI m complex causes global 3' UTR shortening, as proximal
375 poly(A) sites become preferred over distal poly(A) sites (Kubo et al., 2006; Martin et al., 2012;
376 Zhu et al., 2018). In *Neurospora*, we find that a distal-to-proximal shift occurs in a majority (63%)
377 of the 155 altered APA events in the Δ CFI m mutant.

378 Nine core clock and compensation genes are among the consensus list of 3' UTRs with
379 APA events: *ck-1a* (NCU00685), *frq* (NCU02265), *upf1^{prd-6}* (NCU04242), *ckb-1* (NCU05485),
380 *rco-1* (NCU06205), *pkac-1* (NCU06240), NCU06565, *prd-1* (NCU07839), and *nab2* (NCU16397)
381 (Supplementary Table 3). Given the strict cutoff for consensus APA, we added 7 additional
382 genes after visual inspection: *set-2* (NCU00269), *wc-2* (NCU00902), *set-1* (NCU01206), *csp-1*
383 (NCU02713), *cka^{prd-3}* (NCU03124), *frh* (NCU03363), and *pabp-2* (NCU03946). 3' UTR regions
384 containing APA events were visualized in a heatmap (Figure 4B). 2 out of 16 genes of interest,
385 *ckb-1* and *pabp-2*, are among the list of genes with altered APA patterns in the Δ CFI m mutant
386 (Supplementary Table 4; Figure 4B asterisks *). poly(A) read pileups were visualized for 3'
387 UTRs of *ckb-1* and *pabp-2* to confirm significant re-organization of poly(A) tail locations in the
388 Δ CFI m mutant (Figure 4C). Following our original hypothesis, altered 3' UTR length in Δ CFI m
389 should lead to altered mRNA stability and changes in gene expression compared to control
390 samples. An extremely slight and statistically non-significant increase (t-test, $p = 0.5$) was
391 observed for both *ckb-1* and *pabp-2* gene expression levels (Figure 4D). *ckb-1* encodes the
392 regulatory subunit of *Neurospora* Casein Kinase II (CKII), and interestingly, the catalytic alpha
393 subunit of CKII (*cka^{prd-3}*) increased significantly in the Δ CFI m mutant, despite no visible changes

394 in 3' UTR poly(A) sequence coverage for *cka^{prd-3}* (Figure 4B). In addition, *ck-1a* gene expression
 395 levels are modestly (1.2 fold) but significantly decreased in Δ CFIm (Figure 4D), a result that is
 396 counterintuitive given the mutant's short period length phenotype (Kelliher et al., 2020a). Future
 397 work will determine whether overexpression of CKII underlies the Δ CFIm short period length
 398 and Nutritional Compensation phenotypes.
 399



400
 401 **Figure 4. The *Neurospora* CFIm complex is involved in Alternative Polyadenylation**
 402 **(APA), and a subset of core clock and compensation-relevant genes are altered in Δ CFIm**
 403 **mutants.** Circadian bioluminescence was recorded from race tube cultures of the indicated
 404 strains grown in high amino acid race tube medium (0.17% w/v arginine, 0% glucose).
 405 Luciferase signal was acquired from the entire race tube of fungal growth (yellow lines) or from
 406 an old tissue region of the race tube (blue lines) (see: Supplementary Movie 2). Circadian period
 407 lengths were computed for each region (Materials and Methods; N = 4 race tubes per genotype;
 408 standard deviation error bars): control: 21.7 ± 0.2 hrs (whole tube), 21.4 ± 0.3 hrs (old tissue);
 409 Δ cpsf5: 19.0 ± 0.5 hrs (whole tube), 17.2 ± 0.2 hrs (old tissue); Δ cpsf6: 19.4 ± 0.3 hrs (whole

410 tube), 17.3 ± 0.3 hrs (old tissue) (**A**). Wild-type control and $\Delta cpsf5 \Delta cpsf6$ double mutant
411 cellophane plate cultures were grown in constant light at 25°C for 72 hours on high nutrient
412 medium (0.25% w/v glucose, 0.17% w/v arginine). Total RNA was extracted, and 3' End
413 Sequencing was performed (N = 2 biological replicates per genotype). 16 core clock and
414 compensation genes of interest were selected, read pileup data from 3' UTR regions were
415 extracted, biological duplicate samples averaged together, and heatmaps were generated.
416 Read count pileups are depicted as \log_2 -fold changes, and both color scales are normalized to
417 mean counts in the wild-type dataset. Each point along the x-axis represents nucleotide
418 coordinates from the STOP codon for each mRNA (where data from any negative / Crick strand
419 genes are shown in reverse orientation), and genes are ordered along the y-axis by increasing
420 3' UTR lengths. Asterisks (*) indicate two genes, *ckb-1* and *pabp-2*, with significantly altered
421 APA patterns between control and mutant (**B**). Genomic tracks were generated using the Gviz
422 package in R to visualize poly(A) read pileups in the 3' UTR regions of *ckb-1* and *pabp-2*. In the
423 ΔCFI mutant, poly(A) tail locations are significantly changed for *ckb-1* and *pabp-2* (**C**). Gene
424 expression levels of core clock and compensation genes were measured by normalizing total
425 read counts for each gene (Materials and Methods). Log₂-transformed read counts are shown.
426 Asterisks indicate $p < 0.05$ (*) by student's t-test comparing mutant to control levels. The *cka*^{prd-3}
427 transcript is 1.34-fold upregulated, and the *ck-1a* transcript is 1.2-fold down-regulated in ΔCFI
428 (**D**).
429

430 **Chromatin modifiers emerge as key regulators of circadian period length across**
431 **nutrients.**

432 We next examined the 4 over-compensation mutants identified in our genetic screen,
433 where period lengthens as a function of nutrient levels (Figure 5A, Supplementary Table 1). The
434 wild-type *Neurospora* clock is slightly under-compensated (Figures 1 – 3), and therefore this
435 group of over-compensation mutants, together with *prd-1* (Figure 2B) (Emerson et al., 2015),
436 represent clear and bona fide Nutritional Compensation defects. Protein Kinase A (*pkac-1*)
437 shows extended compensation, or approximately the same period length at zero and high
438 nutrients (Figure 5A). The effect of loss of PKA on Nutritional Compensation is subtle and likely
439 due to its regulation of RCM-1 and WCC-independent *frq* transcription—RCM-1 normally acts
440 as a general transcription co-repressor with RCO-1 and prevents *frq* transcription in a $\Delta wc-1$ or
441 $\Delta wc-2$ background (Zhou et al., 2013; Liu et al., 2015); however, a $\Delta rcm-1$ mutant did have
442 normal Nutritional Compensation in our screen (Supplementary Table 1). FGSC16412
443 ($\Delta NCU02961$, $\Delta rbg-28$) is a broadly conserved ribosome biogenesis factor in fungi, currently

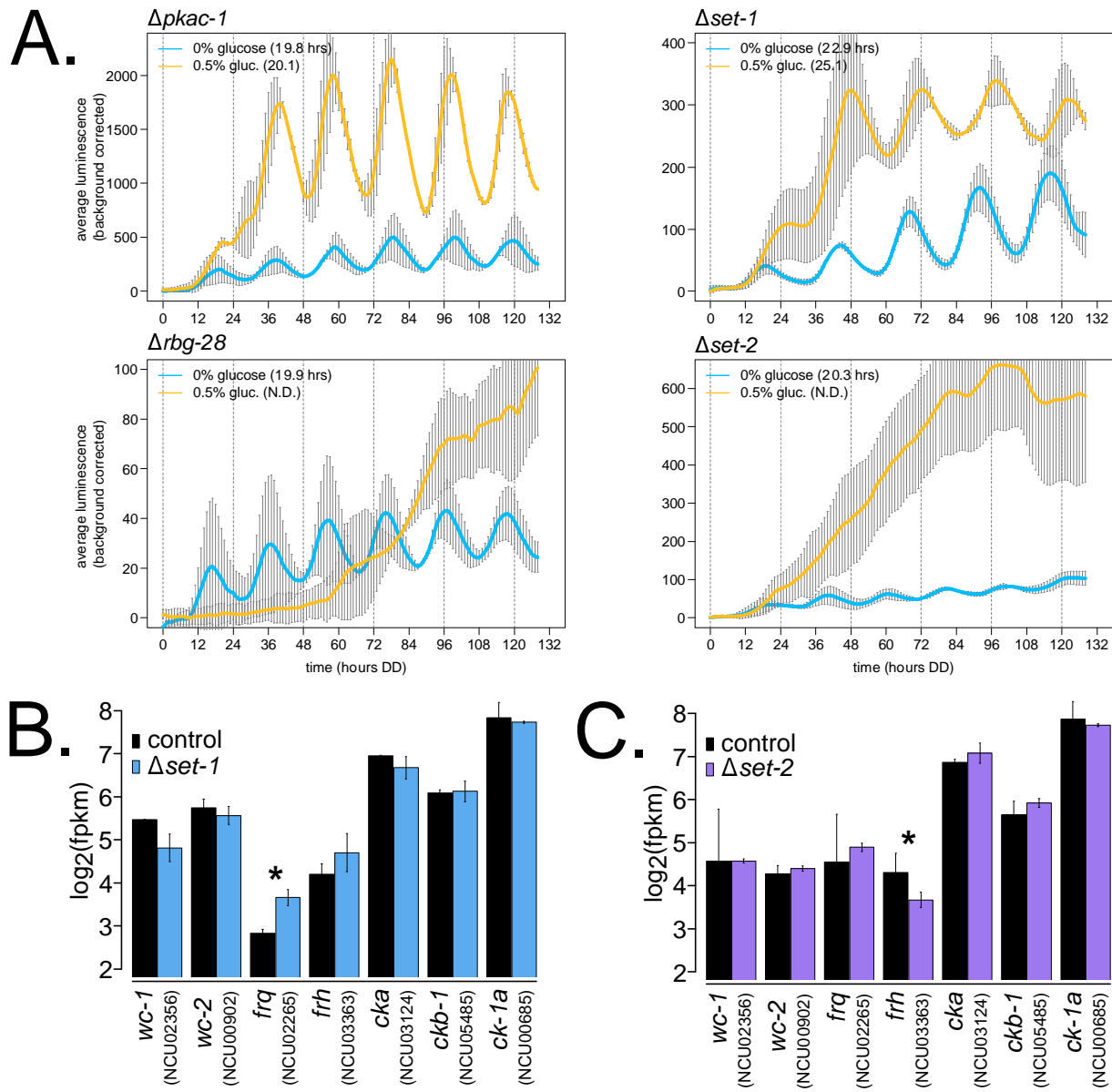
444 uncharacterized in the *Neurospora* literature but homologous to *RNH70* (YGR276C) in *S.*
445 *cerevisiae*. *Rnh70p* (Rex1p) is a 3'-to-5' exonuclease involved in 5S and 5.8S rRNA processing
446 (Van Hoof et al., 2000). The completely arrhythmic phenotype of $\Delta rbg-28$ on high nutrients
447 (Figure 5A) is intriguing and implicates translational machinery in Nutritional Compensation for
448 the first time (see Discussion).

449 Half of the novel over-compensation mutants are involved in chromatin regulation, and
450 both contain SET domains (reviewed in: Freitag, 2017). SET-1 (NCU01206) is a histone H3
451 lysine 4 (K4) methyltransferase and the catalytic subunit of the COMPASS complex in
452 *Neurospora*. $\Delta set-1$ was previously reported to be arrhythmic (Raduwan et al., 2013), but here
453 we find an increasing period length as a function of glucose levels (Figure 5A). SET-1 was
454 convincingly shown to regulate methylation levels and transcriptional repression at the *frq* locus
455 (Raduwan et al., 2013; Zhu et al., 2019), and upon re-analysis of the $\Delta set-1$ transcriptome, *frq* is
456 the only core clock gene significantly altered in the $\Delta set-1$ mutant due to loss of repression
457 (Figure 5B). The $\Delta set-1$ nutritional over-compensation phenotype is consistent with loss of
458 chromatin repression on *frq*—as glucose levels increase, more cellular energy is available for
459 transcription/translation, and circadian period is lengthened due to high *frq* levels and prolonged
460 negative feedback. This SET-1 nutritional mechanism is analogous to CKII's role in regulating
461 the larger pool of FRQ protein at higher temperatures in Temperature Compensation (Mehra et
462 al., 2009).

463 SET-2 (NCU00269) is a histone H3 K36 methyltransferase in *Neurospora*, which
464 deposits inhibitory chromatin marks in actively transcribed regions via a physical association
465 with RNA Polymerase II and prevents improper transcription initiation inside coding regions
466 (Adhvaryu et al., 2005; Bicocca et al., 2018). $\Delta set-2$ was also previously reported to be
467 arrhythmic (Zhou et al., 2013; Sun et al., 2016), but here we find that $\Delta set-2$ rhythms are
468 completely intact with a short period length on nutrient starvation medium (Figure 5A). SET-2 is
469 required to maintain H3K36me2 and H3K36me3 marks across the *frq* locus, and the $\Delta set-2$
470 mutant results in hyper-acetylation of *frq*, improper activation of WCC-independent *frq*
471 transcription, constitutively high *frq* expression levels, and, presumably, the arrhythmic clock
472 phenotype observed under high nutrients (Zhou et al., 2013; Sun et al., 2016). Our result
473 indicates for the first time that WCC-independent *frq* transcription is nutrient dependent and only
474 reaches levels sufficient for arrhythmicity at high nutrient levels. However, low WCC-
475 independent *frq* transcription does not explain the short period length of the $\Delta set-2$ mutant in
476 zero nutrient medium (Figure 5A), as low WCC expression results in lengthened periods (Cheng

477 et al., 2001). Curiously, upon mining existing data for the Δ set-2 transcriptome, *frq* levels were
478 not significantly increased, and instead *frh* levels were down-regulated (Figure 5C). *frh* is among
479 the top down-regulated genes during glucose starvation (Supplementary Figure 4), and perhaps
480 Δ set-2 further affects *frh* transcription, leading to a circadian period change. Future work using
481 cellophane plate assays (Supplementary Figure 5) will determine whether SET-2 regulation of
482 *frh* or another gene expression program explains its short period length on nutrient starvation
483 medium. Notably, two RNA helicases physically associated with the mammalian negative arm
484 complex, DDX5 and DHX9, show a short period length upon siRNA knockdown (Padmanabhan
485 et al., 2012), and so decreased levels of the *frh* helicase in Δ set-2 could potentially explain its
486 short period phenotype. Taken together, our genetic screen revealed mRNA stability,
487 polyadenylation, and chromatin modifier pathways converging on gene expression regulation to
488 enact circadian Nutritional Compensation in fungi.

489



490

491 **Figure 5. Nutritional over-compensation mutants are enriched for chromatin regulators.**

492 Circadian bioluminescence was recorded from race tube cultures of the indicated deletion
 493 mutants. High nutrient medium (yellow lines) contained 0.5% w/v glucose 0.17% w/v arginine,
 494 and zero nutrient medium (blue lines) contained 0% glucose 0% arginine. Period lengths are
 495 indicated as insets (N = 2 biological replicates per nutrient concentration) (A). RNA-Sequencing
 496 data were mined from a previous study (Zhu et al., 2019) (Materials and Methods), where 2%
 497 (high) glucose liquid cultures were harvested at circadian time point DD24. Log₂-transformed
 498 FPKM values are shown for core clock gene expression levels (N = 2 biological replicates per
 499 genotype). The asterisk indicates p = 0.05 (*) by student's t-test comparing mutant to control

500 levels. The *frq* transcript is 1.78-fold more abundant in Δ set-1 (**B**). RNA-Sequencing data were
501 mined from a previous study (Bicocca et al., 2018) (Materials and Methods), where 1.5% (high)
502 sucrose liquid cultures were sampled. Log₂-transformed FPKM values are shown for core clock
503 gene expression levels (N = 2 biological replicates for Δ set-2, and N = 6 biological replicates for
504 controls). The asterisk indicates p < 0.05 (*) by student's t-test comparing mutant to control
505 levels. The *frh* transcript is 1.61-fold less abundant in Δ set-2 (**C**).
506

507 **Nutritional Compensation also functions in a mammalian circadian system.**

508 Compensation is a defining principle of circadian oscillators. Temperature Compensation
509 mechanisms are active in mammalian tissue culture (Izumo et al., 2003; Tsuchiya et al., 2003),
510 despite long evolutionary timescales in homeothermic organisms. Thus, we hypothesized that
511 Nutritional Compensation is also a conserved feature between the fungal and mammalian
512 circadian clocks. Four previous studies hinted that Nutritional Compensation mechanisms may
513 be actively maintaining circadian period length across physiologically relevant nutrient
514 environments.

515 Inhibiting transcription using α -amanitin or actinomycin D against RNA Polymerase II
516 activity led to dose-dependent shortening of the NIH3T3 period length (Dibner et al., 2009). In
517 other words, the mammalian circadian oscillator is over-compensated with respect to
518 transcription rates, reminiscent of temperature over-compensation observed in *Gonyaulax*
519 (Hastings and Sweeney, 1957). Similarly, induction of autophagy and amino acid starvation
520 shortens period length in MEFs (Beesley et al., 2020), once again indicating an over-
521 compensation phenotype to amino acid levels in mammals. Varying levels of FBS do not alter
522 the circadian period length in NIH3T3 cells (Matsumura et al., 2014) despite substantial
523 changes in cell growth rate. Rhythms at the single-cell level of MEFs grown in microfluidic
524 devices show slight period shortening when cells were given fresh medium every hour
525 compared to a single medium supply at the beginning of the experiment, suggesting instead
526 nutritional under-compensation (Gagliano et al., 2021). Thus, preliminary evidence is consistent
527 with functional Nutritional Compensation in the mammalian circadian system.

528 We used a U2OS *Bmal1-dLuc* reporter cell line to compare rhythms in high versus low
529 glucose medium and found a slight under-compensation phenotype (Supplementary Figure 6),
530 similar to the single-cell rhythms report (Gagliano et al., 2021). Period length was 0.5 hours
531 shorter in 25 mM (high) glucose compared to 5.56 mM (low) glucose, trending in the same
532 direction as the ~1.5-hour period shortening seen for the wild-type *Neurospora* clock across
533 glucose concentrations (Figures 1 – 3). In this circadian assay, *Bmal1-dLuc* cells have reached

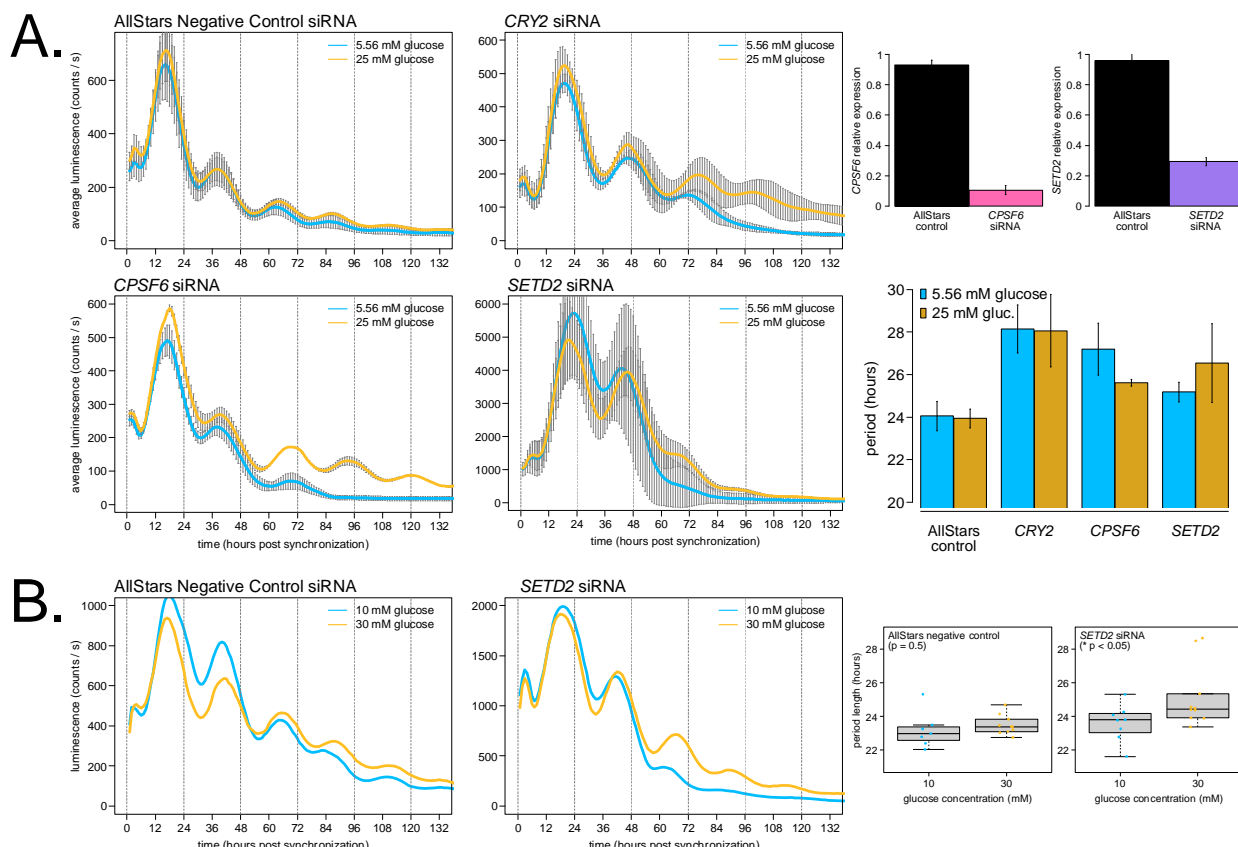
534 confluence and are no longer actively dividing, but metabolism and drug responses of cultured
535 cells are reported to be significantly different in high versus low glucose tissue culture medium
536 (reviewed in: Abbas et al., 2021).

537 Maintenance of circadian period length between high and low glucose medium and a
538 small amount of literature precedent does not, however, prove the relevance of Nutritional
539 Compensation to the mammalian clock. A mutant phenotype showing nutrient-dependent period
540 changes (i.e. defective Nutritional Compensation) would provide much stronger evidence for the
541 biological relevance of compensation. Therefore, we selected two of the most significant
542 compensation phenotypes from our fungal genetic screen— $\Delta cpsf6$ (NCU02152) and $\Delta set-2$
543 (NCU00269)—and identified the homologous human genes *CPSF6* and *SETD2* (reciprocal
544 BLAST e-values = $7e^{-6}$ / $7e^{-8}$ for *Cpsf6* and $4.4e^{-61}$ / $5.7e^{-62}$ for *Set-2*). If the circadian functions
545 of *CPSF6* or *SETD2* are indeed conserved with *Neurospora*, we expected to observe a period
546 length change as well as a Nutritional Compensation defect. *CPSF6* and *SETD2* were not
547 among the hits from a genome-wide screen for period length defects using U2OS cells (Zhang
548 et al., 2009) or tested in a kinase/phosphatase siRNA screen (Maier et al., 2009); however,
549 visual inspection of the genome-wide screen data confirmed that a subset of the pooled siRNAs
550 did show period effects after *CPSF6* or *SETD2* knockdown (source: BioGPS database,
551 “Circadian Genomics Screen” plugin).

552 *CPSF6* and *SETD2* were knocked down using siRNAs in *Bmal1-dLuc* cells, and period
553 length was measured from high and low glucose medium (Figure 6). AllStars Negative Control
554 siRNA, which does not target any known mammalian transcript, was used as an internal control
555 for each biological replicate assay, and *CRY2* siRNA knockdown was used as a positive control
556 for a known long period phenotype (Baggs et al., 2009; Lee et al., 2019). Control *Bmal1-dLuc*
557 cells had a slight under-compensation phenotype (Figure 6A), matching preliminary results
558 (Supplementary Figure 6). *CRY2* knockdown lengthened period by ~4 hours in both high and
559 low glucose conditions, indicating no effect on Nutritional Compensation (Figure 6A). *CPSF6*
560 knockdown lengthened period by ~1.5 hours in high glucose and further lengthened period by
561 ~3 hours in low glucose, indicating an under-compensation phenotype compared to control cells
562 (Figure 6A). The long period observed in *CPSF6* knockdowns was the opposite of the short
563 period phenotype in *Neurospora* (Figure 4A). Most interestingly, *SETD2* knockdown drastically
564 increased the amplitude of the *Bmal1-dLuc* transcriptional reporter compared to controls. As
565 with *Neurospora*, *SETD2* knockdown rhythms were less robust (Figures 5A & 6A). *SETD2*
566 knockdown lengthened period by ~2.5 hours in high glucose but only lengthened period by ~1
567 hour in low glucose (Figure 6A). To further validate nutritional over-compensation in *SETD2*

568 knockdowns, *Bmal1-dLuc* cells were compared across higher concentrations of glucose with
569 more robust rhythms (Figure 6B). Just like *Neurospora*, *SETD2* mutants show a nutritional over-
570 compensation phenotype. This genetic evidence strongly suggests that Nutritional
571 Compensation mechanisms also regulate the mammalian circadian clock in physiologically
572 relevant environments.

573



574

575 **Figure 6. CPSF6 and SETD2 are required for normal Nutritional Compensation of U2OS**
576 ***Bmal1-dLuc* cells cultured in high versus low glucose.** U2OS cells were transfected with the
577 indicated siRNAs (15 pmol total), incubated for 2 days, and *Bmal1-dLuc* rhythms were
578 measured for 5 – 6 days following dexamethasone synchronization. Knocked-down cells were
579 assayed in either MEM high glucose (yellow lines) or MEM low glucose (blue lines) medium.
580 Averaged luciferase traces are shown for each glucose concentration without detrending or
581 further data manipulation (N = 6 biological / technical replicates for AllStars negative controls; N
582 = 4 biological / technical replicates for other siRNA knockdowns per nutrient concentration;
583 standard deviation error bars). siRNA knockdown efficiency was measured using RT-qPCR
584 relative to non-transfected control samples (Materials and Methods). ~90% knockdown was
585 achieved for CPSF6, and ~70% knockdown for SETD2 under these experimental conditions (N

586 = 2 biological replicate samples each with N = 3 technical replicate RT-qPCR reactions). Period
587 lengths were calculated and averaged for all replicates (**A**). siRNA knocked-down cells were
588 assayed in either DMEM 30 mM glucose (yellow lines) or DMEM 10 mM glucose (blue lines)
589 medium. One representative luciferase trace is shown (N = 7 – 10 biological / technical
590 replicates per nutrient concentration). Period lengths were calculated and averaged for all
591 replicates and plotted as boxplots (**B**). Average period lengths were: 23.2 ± 1.1 hrs (low glucose
592 control) and 23.5 ± 0.6 hrs (high glucose control); 23.6 ± 1.1 hrs (low glucose *SETD2*
593 knockdown) and 25.2 ± 1.8 hrs (high glucose *SETD2* knockdown) (* p < 0.05, student's t-test).
594

595 **Discussion**

596 We present the largest genetic screen to date for mutants displaying altered circadian
597 Nutritional Compensation (Supplementary Tables 1 & 2, Figure 2C). Together with a recent
598 survey of 177 transcription factor knockouts (Muñoz-Guzmán et al., 2021), circadian functional
599 genomics is well underway utilizing the *Neurospora* deletion collection (Colot et al., 2006). In
600 this study, 16 new Nutritional Compensation mutants were identified and characterized along
601 with the 4 previously characterized mutants (*Δcsp-1*, *prd-1*, *Δrco-1*, and *Δras2*)—fungal
602 Nutritional Compensation occurs at the level of gene regulation and involves transcription
603 factors, RNA helicases, chromatin modifiers, and polyadenylation machinery. Nutritional
604 Compensation effectors are responsible for directly regulating the core oscillator to maintain the
605 circadian period length across different nutrient levels. In high nutrient environments, CSP-1
606 forms a negative feedback loop on WCC expression and activity by inhibiting *wc-1*
607 overexpression and preventing nutritional under-compensation (Sancar et al., 2012; Dovzhenok
608 et al., 2015). RCO-1, SET-2, and possibly PKA are required to block WCC-independent *frq*
609 transcription in high nutrient conditions (Figure 5) (Zhou et al., 2013; Olivares-Yanez et al.,
610 2016; Sun et al., 2016). More broadly, repressive chromatin marks at the *frq* locus, including
611 those regulated by SET-2, SET-1, and the COMPASS complex, are required to maintain
612 Nutritional Compensation, especially in high nutrients (Figure 5). The nonsense-mediated decay
613 machinery is required for circadian function through its regulation of *ck-1a* expression (Kelliher
614 et al., 2020a), and *ck-1a* transcript stability appears to undergo additional, nutrient-dependent
615 regulation because deletion of the *ck-1a* 3' UTR (its NMD-targeting sequence) causes nutritional
616 under-compensation (Figure 3, Supplementary Figure 3). The alternative polyadenylation (APA)
617 landscape in *Neurospora* is dynamic across nutrient conditions and requires CFIIm complex
618 activity (Figure 4).

619 This screen and associated candidates have opened up several avenues for further
620 work on Nutritional Compensation mechanisms in *Neurospora*. For instance, PRD-1 RNA
621 helicase target genes under high glucose conditions are completely unknown. In this study, we
622 provide some evidence that CKII activity is altered in Δ CFIm mutants (Figure 4), but the effect of
623 CKII overexpression on both circadian period length and Nutritional Compensation remains to
624 be tested. Genetic screens extending from this work can examine both transcription factor and
625 chromatin modifier knockouts for Nutritional Compensation phenotypes. Curiously, the subset of
626 chromatin readers and writers that have been screened to date (using standard medium
627 conditions) have an unusually high ~15% hit rate for circadian period defects (Belden et al.,
628 2007b; Wang et al., 2014). In this study, chromatin modifiers not only regulate chromatin state
629 at the *frq* locus, but dynamic expression of *frh* across nutrient levels may also be involved in
630 compensation, potentially mediated by SET-2 (Figure 5C). Intriguingly, quantitative modeling
631 has suggested that nutrient-dependent FRH sequestration away from the negative arm complex
632 is a plausible mechanism for Nutritional Compensation (Upadhyay et al., 2019). Mechanistic
633 work in prokaryotes has indicated that Nutritional Compensation can be derived from the same
634 core clock enzyme, KaiC, through its protein domains with equal and opposite balancing
635 reaction rates (Phong et al., 2013; Hong et al., 2020). In fungi and perhaps other eukaryotic
636 clocks, Nutritional Compensation appears to be maintained through regulation of multiple
637 different core clock factors. Nutritional Compensation mechanisms very likely extend to other
638 physiological environmental variables such as nitrogen sources and levels (Huberman et al.,
639 2021), vitamins, and soil pH (Ruoff et al., 2000).

640 The work establishes Nutritional Compensation in the mammalian circadian clock for the
641 first time, and its clinical relevance may be high. There is a direct link between pancreatic clock
642 function and risk for type 2 diabetes (Marcheva et al., 2010). Fasting glucose concentration in
643 human serum is approximately 5 mM in healthy controls, but glucose levels can increase by 3+
644 orders of magnitude in type 2 diabetic hyperglycemia (Shapiro et al., 1991; Radziuk and Pye,
645 2006). Thus, deleterious SNPs in Nutritional Compensation-relevant genes could exacerbate
646 disease outcomes when cellular clocks encounter nutrients outside of the physiological range.
647 In addition to Nutritional Compensation, there is a large body of quality literature describing the
648 set of metabolites and metabolic enzymes that can directly feed back and affect circadian
649 function, including adenosine monophosphate (AMP) and AMP kinase, nicotinamide adenine
650 dinucleotide (NAD⁺) and SIRT1 activity, acetylglucosamine (O-GlcNAc) (Liu et al., 2021), and
651 mTOR activity (Ramanathan et al., 2018) (reviewed in: Bass and Takahashi, 2010; Sancar and
652 Brunner, 2014; Asher and Sassone-Corsi, 2015; Dibner and Schibler, 2015). In *Neurospora*,

653 extensive metabolic rhythms are also present (Hurley et al., 2018; Baek et al., 2019), and
654 rhythmic metabolic reaction fluxes likely function similarly to the mammalian clock (Krishnaiah et
655 al., 2017; Thurley et al., 2017; Collins et al., 2021). For the mammalian clock, it will be critical to
656 define “physiologically relevant” nutrient levels, which could vary by organ or even cell type. A
657 high fat diet has been shown to lengthen circadian behavioral rhythms by ~0.5 hours (Kohsaka
658 et al., 2007), suggesting an over-compensation phenotype. In this study, we observed nutritional
659 under-compensation in U2OS osteosarcoma cells (Figure 6). Does a high fat diet push cells out
660 of the physiological range of compensation for the circadian oscillator, or are different
661 mammalian cell types differentially over- or under-compensated to nutrients? Notably, time-
662 restricted feeding has been shown to alleviate circadian disruption and some metabolic
663 consequences of a high fat diet (Hatori et al., 2012; Wehrens et al., 2017). Future work will
664 implement more metabolically relevant cell types, such as hepatocytes and adipocytes
665 (Ramanathan et al., 2014), to answer these questions about mammalian clock compensation.

666 Circadian control of polyadenylation is an emerging topic. Modern transcriptomic
667 approaches have identified hundreds of rhythmic Alternative Polyadenylation (APA) events in
668 mice and in plants (Liu et al., 2013; Gendreau et al., 2018; Greenwell et al., 2020; Yang et al.,
669 2020). We and others have begun to define the set of APA events in *Neurospora* (Zhou et al.,
670 2018) (Supplementary Table 3), which can be extended to circadian APA events. Two cleavage
671 and polyadenylation factors (CPSF1 and 7) were found to physically interact with the negative
672 arm complex in mice (Ju et al., 2020), and *CPSF6* oscillates at the transcriptional level in mouse
673 kidney and brain (Zhang et al., 2014). During preparation of this manuscript, circadian
674 colleagues reported the long period length of *CPSF6* knockdown (Schmal et al., 2021), which
675 our results further support (Figure 6A). Interestingly, *CPSF6* knockdown cells showed a
676 temperature under-compensation defect, and multi-omics identified *EIF2S1* as the key effector
677 gene upon *CPSF6* knockdown (Schmal et al., 2021). The *Neurospora* homolog of *EIF2S1* is
678 eIF2α / NCU08277 (reciprocal BLAST e-values = 6e⁻⁹¹ / 1e⁻⁸¹), which is a central hub of rhythmic
679 translation initiation peaking in the subjective evening (Karki et al., 2020). We identified a
680 ribosome biogenesis exonuclease, RBG-28, which is required for rhythmicity under high nutrient
681 conditions (Figure 5). Transcription, RNA processing reactions (such as nonsense-mediated
682 decay, splicing, and polyadenylation), and translation are tightly coupled processes. In fact,
683 rhythmic polyadenylation of rRNAs has been linked to translational rhythms in mouse liver
684 (Sinturel et al., 2017). Nutritional Compensation pathways likely occur at multiple steps in the
685 gene expression of multiple core clock components.

686 By suggesting *CPSF6* and *SETD2* as targets, *Neurospora* functional genetics has again
687 informed mammalian-relevant circadian mechanisms (reviewed in: Loros, 2020), and the over-
688 compensation defect of *SETD2* provides affirming evidence of eukaryotic Nutritional
689 Compensation outside of *Neurospora* (Figure 6). A handful of previous studies have implicated
690 histone methyltransferases in mammalian circadian function, including both activating (Katada
691 and Sassone-Corsi, 2010; Valekunja et al., 2013) and repressive chromatin marks (Etchegaray
692 et al., 2006). *SETD2* joins a growing list of circadianly-relevant histone methyltransferases and
693 chromatin modifiers. In fact, recent work has demonstrated a novel function for a key histone
694 methyltransferase in the circadian transcription-translation feedback loop (TRITHORAX in
695 insect, MLL1 in mammals), further highlighting the utility of circadian model systems for
696 understanding the mammalian clock (Zhang et al., 2022).

697

698 **Materials and Methods**

699

700 ***Neurospora* strains, growth conditions, and genetic screen.**

701 Strains used in this study were derived from the wild-type background (FGSC2489 *mat*
702 A), *ras-1^{bd}* background (87-3 *mat a* or 328-4 *mat A*), or the Fungal Genetics Stock Center
703 (FGSC) knockout collection as indicated (Supplementary Table 1). Strains were constructed by
704 transformation or by sexual crosses using standard *Neurospora* methods
705 (<http://www.fgsc.net/Neurospora/NeurosporaProtocolGuide.htm>). The *frq* clock box
706 transcriptional reporter was transformed and used as previously described (Kelliher et al.,
707 2020a). The fungal biomass reporter gene (Supplementary Figure 1) is composed of 430 bp of
708 the *gpd* promoter from *Cochliobolus heterostrophus* driving constitutive levels of codon
709 optimized *luciferase* and integrated at the *csr-1* (NCU00726) locus (Bartholomai, 2021).

710 All race tubes contain a base medium of 1X Vogel's Salts, 1.5% w/v Noble agar (Thermo
711 Fisher # J10907), and 50 ng/ml biotin. Noble agar was used instead of standard bacteriological
712 agar (Thermo Fisher # J10906) because impurities in bacteriological agar can be metabolized
713 by *Neurospora* and interfere with accurate quantification of Nutritional Compensation
714 phenotypes (Emerson et al., 2015). Glucose and arginine were supplemented into race tube
715 medium as indicated. High glucose was defined as 0.5% w/v (27.8 mM) based on literature
716 precedent (Sancar et al., 2012; Olivares-Yanez et al., 2016) and based on growth rate and
717 period length similarity to higher glucose concentrations (Figure 1A, Supplementary Figure 1G).
718 High arginine was defined as 0.17% w/v because concentrations higher than 0.5% w/v interfere
719 with circadian banding in race tube assays (Sargent and Kaltenborn, 1972). To optimally

720 visualize and quantify the circadian banding pattern in the primary genetic screen (Figure 2C),
721 screen medium contained: 1X Vogel's Salts, 1.5% Noble agar, 50 ng/ml biotin, and 0.17%
722 arginine. To quantify period lengths in carbon and nitrogen starvation conditions, the secondary
723 screen medium contained: 1X Vogel's Salts, 1.5% Noble agar, 50 ng/ml biotin, and 25 μ M
724 luciferin (GoldBio # 115144-35-9). The tertiary screen medium contained: 1X Vogel's Salts,
725 1.5% Noble agar, 50 ng/ml biotin, 25 μ M luciferin, and glucose/arginine levels as indicated
726 (Figures 3 – 5). For Temperature Compensation 96-well plate experiments (Figure 2A), the
727 standard medium recipe contained: 1X Vogel's Salts, 1.5% bacteriological agar, 50 ng/ml biotin,
728 25 μ M luciferin, 0.03% w/v glucose, and 0.05% w/v arginine.

729 Liquid medium cultures were grown from fungal plugs in Bird Medium + 1.8% w/v
730 glucose (Supplementary Figure 5) as previously described (Kelliher et al., 2020b). Solid medium
731 cultures were implemented to determine mRNA (or protein) levels from Nutritional
732 Compensation mutants of interest. Medium was poured into 100-mm petri plates and cooled to
733 solidify (~20 ml per plate; 1X Vogel's Salts, 1.5% Noble agar, 50 ng/ml biotin, 25 μ M luciferin,
734 and glucose/arginine levels as indicated). Cellophane (Idea Scientific # 1080) paper discs were
735 cut to the size of 100-mm plates, and autoclaved to sterilize. A sterile cellophane disc was then
736 placed on top of the solidified medium. Conidia from strains of interest were resuspended in 100
737 μ l of sterile water, vortexed to mix, pipetted on top of the cellophane disc, and spread with a
738 sterile plate spreader (in order to maintain approximately the same conidial age across the
739 cellophane plate). Inoculated plates were then covered with a Breathe-Easy strip for gas
740 exchange (USA Scientific # 9123-6100). After growing tissue on cellophane plates for the
741 indicated amount of time, mycelia and conidia were harvested from atop the cellophane layer by
742 scraping with a 1000 μ l pipette tip. Harvested fungal tissue (approximately the size of one US
743 quarter per each 100-mm cellophane plate) was rapidly hand dried using paper towels and an
744 Eppendorf tube rack, and flash frozen in liquid nitrogen for storage before biochemical
745 extraction.

746 Most strains were genotyped by growth on selective medium (5 μ g/ml cyclosporine A
747 and/or 200-300 μ g/ml Hygromycin). Key strains were genotyped by PCR as previously
748 described (Kelliher et al., 2020a) using genotyping primers:

749 *ck-1a^{LONG}-VHFΔ3'UTR::hyg^R* (NCU00685 Δ3'UTR): 5' GCTGCTGCTCGTAAGGAC 3'
750 and 5' CATCAGCTCATCGAGAGCCTG 3'

751 *Δcpsf5::hyg^R* (FGSC KO mutant): 5' CTCTGGTCGAGAACACTGCG 3' and 5'
752 CAGGCTCTCGATGAGCTGATG 3'

753 $\Delta c p s f 6::h y g^R$ (FGSC KO mutant): 5' CACCAACCCTAACCCGTGAT 3' and 5'
754 CAGGCTCTCGATGAGCTGATG 3'

755 $\Delta s e t - 2::h y g^R$ (FGSC KO mutant): 5' GACGTCATCGGTGTTGAGAC 3' and 5'
756 CAGGCTCTCGATGAGCTGATG 3'

757

758 ***Neurospora* luciferase reporter detection and data analysis.**

759 96-well plates were inoculated with conidial suspensions and entrained in 12 hour
760 light:dark cycles for 2 days in a Percival incubator at 25°C. Temperature inside the Percival
761 incubator was monitored using a HOBO logger device (Onset # MX2202) during entrainment
762 and free run. Race tubes were entrained in constant light at 25°C for 3 – 24 hours (mean
763 entrainment time for all experiments was 14 hours in LL / overnight). Entrained 96-well plates or
764 race tubes were then transferred into constant darkness to initiate the circadian free run.
765 Individual race tubes were separated by ~3 cm tall strips of 6-ply black railroad board paper to
766 prevent contamination of light signal between cultures. Luminescence was recorded using a
767 Pixis 1024B CCD camera (Princeton Instruments). Bioluminescent signal was acquired for 10 –
768 15 minutes every hour using LightField software (Princeton Instruments, 64-bit version 6.10.1).

769 The average bioluminescent intensity of each 96-well or race tube was determined using
770 a custom ImageJ Macro with background correction for each image (Larrondo et al., 2012,
771 2015). Most race tube period lengths reported in this study were derived from luciferase signal
772 measurements across the entire race tube (Figures 1, 3, and 5). However, the long period
773 defect in the *prd-1* mutant only occurs at the growth front (i.e. high nutrients) region of fungal
774 tissue (Emerson et al., 2015). For the *prd-1* strain, an ImageJ macro was modified to quantify
775 only the fungal growth front (Figure 2B). On the other hand, the $\Delta c p s f 5$ and $\Delta c p s f 6$ mutants
776 showed additional period shortening in aged tissue (Figure 4A) (Supplementary Movie 2). For
777 the *cpsf* mutants, an ImageJ macro was modified to quantify only the old tissue region. Custom
778 ImageJ Macros to quantify the growth front or old tissue regions of race tubes from Princeton
779 *.spe image files are available at: <https://github.com/cmk35>.

780 To calculate the circadian period length, background-corrected luminescence traces
781 were run through two different algorithms and averaged as previously described (Kelliher et al
782 2020 eLife). Race tubes period lengths were measured using Chron OSX 2.1 software. For
783 Temperature Compensation experiments, the Q_{10} temperature coefficient was calculated using
784 the formula: [(frequency of clock at 30°C) / (frequency of clock at 20°C)]^[10°C / (30°C - 20°C)], where
785 frequency = period length⁻¹.

786

787 **Neurospora RNA isolation and 3' End Sequencing analyses.**

788 Frozen *Neurospora* tissue was ground in liquid nitrogen with a mortar and pestle. Total
789 RNA was extracted with TRIzol (Invitrogen # 15596026) and the Direct-zol RNA MicroPrep kit
790 (Zymo Research # R2060) according to the manufacturer's instructions and including the on-
791 column DNase I treatment step (Roche # 04 716 728 001, 10 U/μl stock, 30 U used per
792 sample). Total RNA samples were prepared for Northern Blotting, 3' End Sequencing, or stored
793 at -80°C.

794 Northern blotting was performed as previously described (Kelliher et al., 2020a) with
795 slight modifications. Equal amounts of total RNA (7 μg) were loaded per lane of a 0.8% w/v
796 agarose gel (Supplementary Figure 5). For blot visualization, anti-Digoxigenin-AP Fab
797 fragments was purchased from Sigma (Roche # 11 093 274 910) and used at 1:10,000 (75
798 mU/ml).

799 Total RNA was submitted to the Dartmouth Genomics Shared Resource (GSR) for 3'
800 end library preparation and sequencing. 75 bp single-end (SE) strand-specific libraries were
801 prepared using the Lexogen QuantSeq 3' REV kit, multiplexed, and sequenced on an Illumina
802 Mini-Seq. 6.92 ± 0.30 million reads were obtained for each sample, and read quality was
803 confirmed using FastQC. Raw FASTQ files were aligned to the *Neurospora crassa* OR74A
804 NC12 genome (FungiDB version 45 accessed October 25, 2019) using STAR (Dobin et al.,
805 2013). 91.5 – 93.5% of the reads mapped uniquely to the NC12 genome. Because 3' end
806 libraries generate only 1 sequencing read at the extreme 3' end of a given mRNA transcript
807 (directly before its poly(A) tail), gene expression was quantified by counting reads assigned to
808 each genetic locus using HTSeq-count (Anders et al., 2015). Gene count normalization by
809 library size between samples was performed using a custom R script. 3' End Sequencing data
810 have been submitted to the NCBI Gene Expression Omnibus (GEO;
811 <https://www.ncbi.nlm.nih.gov/geo/>) under accession number GSE201901.

812 RNA-Sequencing datasets from 4 other studies were mined in the analyses presented.
813 To examine wild-type *Neurospora* gene expression under carbon starvation (Supplementary
814 Figure 4, Supplementary Table 2), RNA-seq data were taken from a study where liquid cultures
815 (25°C, LL) were grown for 16 hours in 1X Vogel's 2% sucrose minimal medium and shifted to
816 either 0% or 2% glucose 1X Vogel's medium for 60 minutes (Wang et al., 2017) (GSE78952).
817 To examine gene expression in the Δ set-1 mutant background (Figure 5B), RNA-seq data were
818 taken from a study where liquid cultures (25°C, DD24) were grown in 2% glucose Liquid Culture
819 Medium for 48 hours of total growth (Zhu et al., 2019) (GSE121356). To examine gene
820 expression in the Δ set-2 mutant (Figure 5C), RNA-seq data were taken from a study where

821 liquid cultures (32°C) were grown in 1X Vogel's 1.5% sucrose medium (Bicocca et al., 2018)
822 (GSE82222 and GSE118495). These three RNA-Seq datasets were re-processed exactly as
823 previously described (Kelliher et al., 2020a), and FPKM gene expression values were used in
824 the analyses presented. To examine and compare wild-type poly(A) tail locations
825 (Supplementary Table 3), 2P-Seq data were taken from a previous study where nuclear
826 fractions were isolated from 1X Vogel's 2% glucose liquid medium (Zhou et al., 2018) (SRA
827 PRJNA419320). Raw 2P-Seq data were filtered according to custom a Perl script from the
828 original study ("Step 1": <https://github.com/elifeosciences-publications/poly-A-seq>). After read
829 filtering, duplicate 2P-Seq FASTQ files were processed in exactly the same manner as the new
830 3' End Sequencing dataset generated in this study.

831 NC12 mapped reads from 3' End Sequencing (this study) and 2P-Seq (Zhou et al.,
832 2018) data were sorted and indexed using Samtools (BAM file outputs) and then visualized
833 using IGV. Read pileups denoted the location of poly(A) tails in both datasets. To map locations
834 of poly(A) tails genome wide, the ChIP-Seq peak calling algorithm MACS2 was re-purposed
835 (Zhang et al., 2008). The relevant MACS2 parameters used to identify poly(A) peaks were:
836 effective genome size (-g 4.014e7), retention of duplicate reads in pileups (--keep-dup all),
837 summit and subpeak identification (--call-summits), fragment size estimation and shifting turned
838 off (--nomodel --extsize 75), and a false discovery rate cutoff for significant peaks (-q 0.01).
839 MACS2 peaks were assigned to the corresponding gene 3' UTR region using a custom R script.
840 The *Neurospora crassa* NC12 transcriptome annotation remains partially incomplete with only
841 7,793 out of 10,591 unique NCU IDs having 3' UTRs annotated. As a result, ~14-18% of all
842 MACS2 peaks were pruned from consideration due to missing annotations. Importantly, there
843 are also examples of under-annotated 3' UTR regions, where the poly(A) read pileup signal is
844 clearly located outside of the 3' UTR annotation. One such critical example occurs at the
845 frequency locus (positive / Watson strand gene), where the predominant poly(A) peak is
846 centered at LG VII coordinate 3,136,633, and its longest 3' UTR annotation ends at coordinate
847 3,136,464. The *Neurospora* NC12 transcriptome annotation was last updated in March 2015
848 before migration from the Broad Institute to the FungiDB database (Basenko et al., 2018). The
849 3' End Sequencing analyses presented here can be updated upon release of an improved
850 transcriptome annotation. Furthermore, there are 621 instances of overlapping coordinates
851 within the 3' UTRs of tail-to-tail oriented genes, and any poly(A) peaks falling in these gene
852 assignment ambiguous regions were also removed from consideration (~9% of MACS2 peaks).
853 The remaining MACS2 peaks (~6,600 unique poly(A) peaks per sample) were assigned to the
854 corresponding gene 3' UTR region and analyzed using custom R scripts. Alternative

855 Polyadenylation (APA) events were defined as instances of more than one distinct MACS2 peak
856 assigned to a single 3' UTR region. poly(A) tail read pileups from genes of interest were
857 extracted using the igvtools count function, and genome coordinate plots were generated using
858 the R Bioconductor package Gviz (Figure 4C). 3' UTR heatmaps were generated using a
859 custom R script (Figure 4B). All custom R scripts for gene expression analyses and alternative
860 polyadenylation analyses are available at: <https://github.com/cmk35>.

861

862 **Mammalian cell culture, synchronization, and siRNA knockdown reagents.**

863 U2OS cells were stably transfected under puromycin selection using a construct
864 containing the mouse *BMAL1* promoter (Sato et al., 2006; Gamsby et al., 2009) driving
865 destabilized luciferase (Ueda et al., 2002). U2OS-m*BMAL1-dLuc*-Puro (referred to as: *Bmal1-*
866 *dLuc*) cells were maintained at 37°C and 6% CO₂ in 25 mM (high) glucose DMEM (Thermo
867 Fisher # 11995-065 with 1 mM sodium pyruvate; or Thermo Fisher # 11965-092 without
868 pyruvate) supplemented with 10% v/v FBS (Thermo Fisher # 10437-036, LOT # 2199672RP)
869 and 1.5 µg/ml of puromycin (Sigma # P9620, 10 mg/ml stock).

870 For control Nutritional Compensation assays (Supplementary Figure 6), *Bmal1-dLuc*
871 cells were subcultured from the same 100-mm dish and grown to 95-100% confluence in 35-mm
872 dishes (Corning # 430165) containing 2 ml of DMEM 25 mM glucose, 10% FBS, and puromycin.
873 Confluent cells were washed once in warm 1X PBS pH 7.4 (Corning # 21-040-CV). The medium
874 was changed to either 2 ml of DMEM 25 mM high glucose (Thermo Fisher # 11995-065 with 1
875 mM sodium pyruvate) or 2 ml of DMEM 5.56 mM low glucose (Thermo Fisher # 11885-084 with
876 1 mM sodium pyruvate). Both synchronization-release medium formulations were pre-warmed
877 and each contained: 10% v/v FBS, 1.5 µg/ml puromycin, 0.1 mM luciferin (GoldBio, 0.1 M
878 stock), and 0.1 µM dexamethasone (Sigma # D2915, 1 mM stock). Dexamethasone is used to
879 reset cells to the same circadian phase and initiate the circadian free run for recording.

880 For siRNA knockdown assays (Figure 6), *Bmal1-dLuc* cells were subcultured from the
881 same 100-mm dish and grown to 60-80% confluence in 35-mm dishes containing DMEM 25 mM
882 glucose, 10% FBS, and puromycin. Cells were washed once in 1X PBS, and the medium was
883 changed to 2 ml of Opti-MEM (Thermo Fisher # 31985-070) with 5% v/v FBS. Cells were
884 transfected with the indicated siRNAs (15 pmol of total siRNA per 35-mm dish) (Baggs et al.,
885 2009) using the Lipofectamine 3000 transfection reagent (Thermo Fisher # L3000) and
886 according to the manufacturer's instructions for 6-well plates. Although the Opti-MEM medium
887 formulation is not publicly available, one study reported the Opti-MEM glucose concentration as
888 2.5 g/L or 13.88 mM (Young et al., 2004). siRNAs were obtained from Qiagen: AllStars Negative

889 Control siRNA (Qiagen # 1027280); human *SETD2* siRNA (Qiagen # 1027416, FlexTube
890 GeneSolution GS29072, 4x siRNAs used at 3.75 pmol each); human *CPSF6* siRNA (Qiagen #
891 1027416, FlexTube GeneSolution GS11052, 4x siRNAs used at 3.75 pmol each); human *CRY2*
892 siRNA (Qiagen # 1027416, FlexTube GeneSolution GS1408, 4x siRNAs used at 3.75 pmol
893 each) (Lee et al., 2019). Cells were incubated for 2 days before removing the siRNA
894 transfection medium and proceeding with RNA extraction or with circadian recordings.

895 RT-qPCR was used to validate siRNA knockdown efficiencies. 2-day transfected cells
896 were washed once in 1 ml of ice-cold 1X PBS. Cells were harvested by scraping in 1 ml of
897 TRIzol (Invitrogen), and total RNA extraction was performed according to the manufacturer's
898 instructions. 500 ng of mRNA was converted into cDNA using the oligo(dT) method from the
899 SuperScript IV First-Strand synthesis kit (Invitrogen # 18091–050). RT-qPCR was performed
900 using SYBR green master mix (Qiagen # 204054) and a StepOne Plus Real-Time PCR System
901 (Applied Biosystems). C_t values were determined using StepOne software (Life Technologies)
902 and normalized to the *GAPDH* gene (ΔC_t). The $\Delta\Delta C_t$ method was used to determine mRNA
903 levels relative to non-transfected negative control samples. Relevant RT-qPCR primer
904 sequences are: h*GAPDH*: 5' TGCACCACCAACTGCTTAGC 3' and 5'
905 ACAGTCTTCTGGGTGGCAGTG 3'. h*CPSF6*: 5' GATGTGGTAAAGGAGCAG 3' and 5'
906 CTTCATCTGTTGTCCACCA 3'. h*SETD2*: 5' CTTTCTGTCCCACCCCTGTC 3' and 5'
907 CCTTGCACCTCTGATGGCTT 3'.

908 2-day transfected cells were washed in warm 1X PBS and prepared for circadian
909 synchronization. Synchronization-release medium was pre-warmed and contained 1% v/v FBS,
910 1.5 μ g/ml puromycin, 0.1 mM luciferin, and 0.1 μ M dexamethasone. siRNA assays in DMEM
911 were conducted using 2 ml of DMEM (Thermo Fisher # 11966-025) supplemented with 10 mM
912 (low glucose) or 30 mM (high glucose) from a D-glucose stock solution (Sigma # G8644, 100
913 g/L stock). DMEM base medium contains more total nutrients than MEM—approximately 2-fold
914 higher levels of the 13 essential amino acids, about 4-fold higher levels of the 8 vitamins, and
915 includes the non-essential amino acids (Gly and Ser) in its formulation. siRNA assays in MEM
916 were conducted using 2 ml of MEM (Thermo Fisher # 11095-080) containing 5.56 mM (low
917 glucose) or supplemented up to 25 mM (high glucose) from a D-glucose stock solution (Sigma).
918 Unlike *Neurospora*, complete glucose starvation medium did not support cell viability in
919 preliminary experiments using DMEM medium containing 10% v/v FBS but zero additional
920 glucose. "Low" 5 – 10 mM glucose was defined by manufacturer formulations as well as
921 physiological levels of fasting serum glucose in humans.

922

923 **Mammalian luciferase reporter detection and period length calculations.**

924 Immediately prior to bioluminescent recording, *Bmal1-dLuc* cells in 35-mm dishes were
925 covered with 40-mm circular microscope cover glass (Fisher Scientific # 22038999 40CIR-1)
926 and sealed using high-vacuum silicone grease (Dow Corning # Z273554). Luciferase data were
927 collected in a LumiCycle 32 (ActiMetrics) luminometer every 10 minutes for 5 – 6 days. Raw
928 luciferase traces in bioluminescence counts / second units were exported using LumiCycle
929 analysis software (ActiMetrics, version 2.56). Data from individual plates were manually
930 combined and converted to hours post synchronization using Microsoft Excel. Period lengths for
931 each luciferase trace were calculated using 3 different methods and averaging the period results
932 with equal weights. For the first method, signal peaks and troughs were extracted from days 1 –
933 3 of raw data, and period was estimated by subtracting consecutive peaks or troughs as
934 described (Chen et al., 2020). Second, the WaveClock algorithm was implemented in R (Price
935 et al., 2008). Finally, the suite of *Neurospora* period length tools was used as previously
936 described (Kelliher et al., 2020a). For *Bmal1-dLuc* luciferase trace data visualization purposes
937 (Figure 6, Supplementary Figure 6), raw counts per second values sampled within the same
938 hour were averaged together (i.e. data were down-sampled from 10-minute measurement
939 intervals to 1-hour measurement intervals).

940

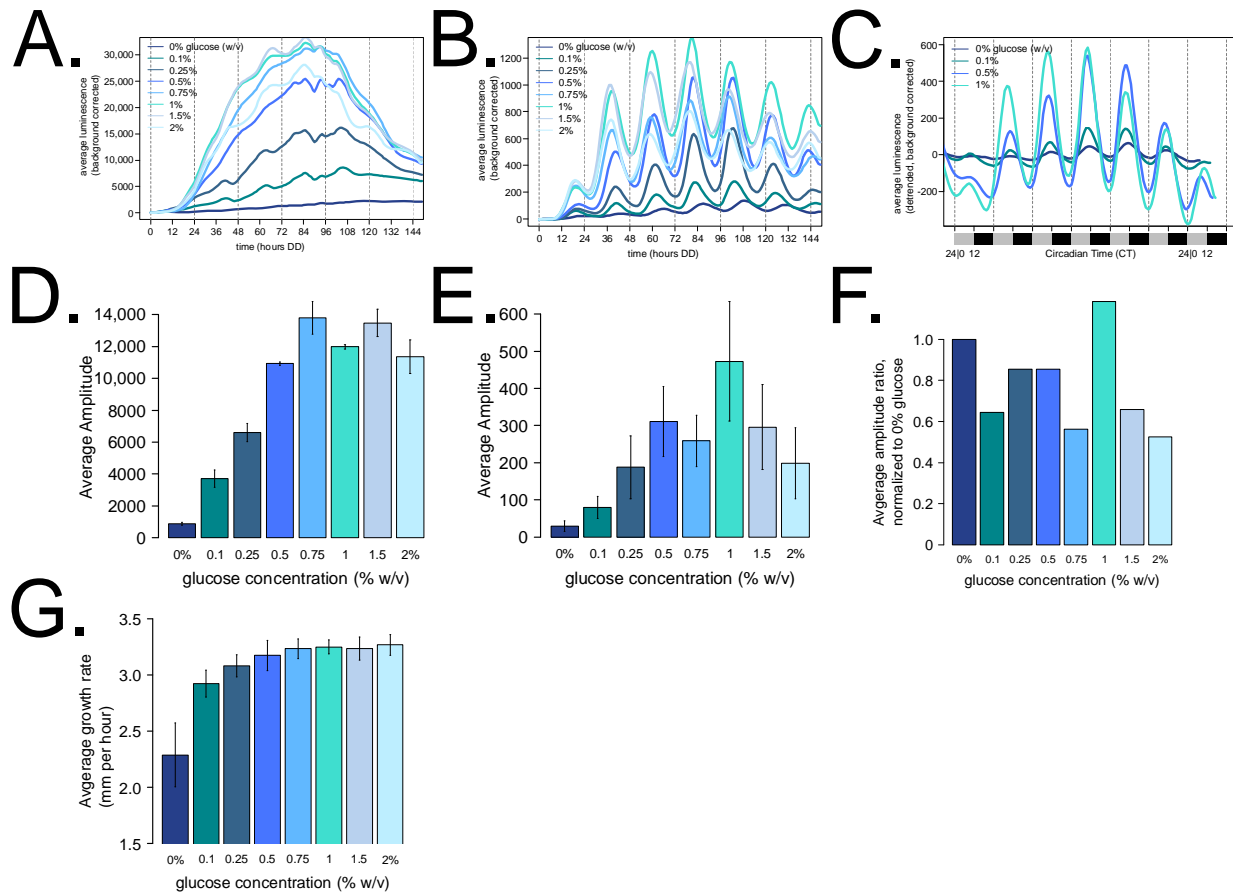
941 **Data visualization.**

942 All figures were plotted in R, output as scalable vector graphics, formatted using
943 Inkscape, and archived in R markdown format. Data represent the mean of at least three
944 biological replicates with standard deviation error bars, unless otherwise indicated. All statistical
945 tests were performed in R.

946

947 **Supplementary Figures, Tables, and Movies**

948



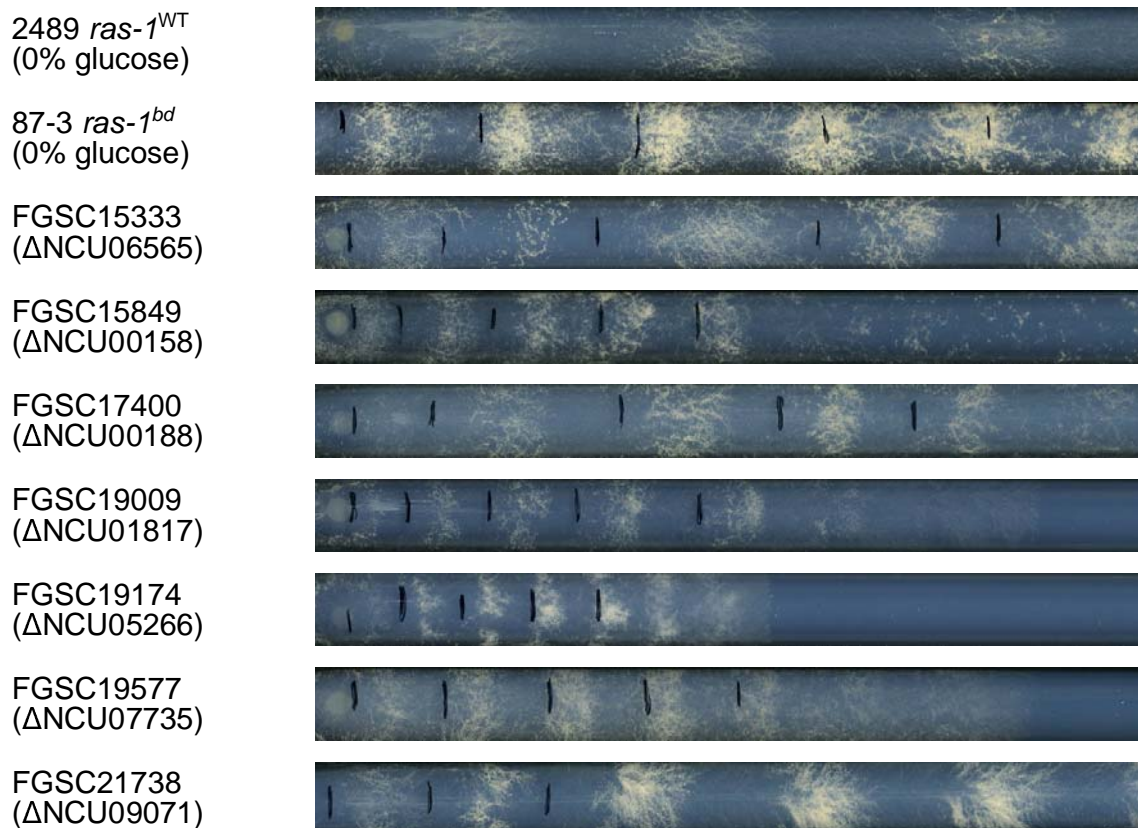
949

950 **Supplementary Figure 1. Additional properties of Nutritional Compensation in**

951 ***Neurospora crassa*.** A fungal biomass control was implemented to ask whether the amplitude
 952 of the core clock transcriptional reporter changes with glucose levels (e.g. Figure 1B). The *gpd*
 953 promoter from *Cochliobolus heterostrophus* driving constitutive *luciferase* was used as a
 954 reporter for fungal biomass. Not surprisingly, biomass increases as a function of glucose (N = 3
 955 race tubes per glucose concentration). Many traces showed a decrease in bioluminescence at
 956 ~100 hours, and this correlates with the fungal growth front reaching and surpassing the end of
 957 the device's recording area (A) (see: Supplementary Movie 1B). Averaged replicates are shown
 958 for the *frq* clock box transcriptional reporter across all glucose levels (N = 6; expanded Figure
 959 1B) (B). Detrended clock reporter traces were plotted on a circadian time (CT) scale to
 960 normalize for the slight period differences (Figure 1A). Circadian phase is consistent across
 961 glucose levels (C). Amplitude was computed for each individual biomass reporter trace using
 962 data from hours 25 – 108 (amplitude calculation: [maximum value – minimum value] / 2) (D).
 963 Average amplitude was computed for each individual core clock reporter trace using data from
 964 days 2 – 5 (hours 25 – 112) to extract 4 peak and 4 trough values. The first day (hours 0 – 24)
 965 was omitted due to low fungal biomass and consequently low luciferase signal during the first

966 recording day (**E**). Core clock amplitudes were normalized to biomass by computing the
967 amplitude ratio at the respective glucose concentrations. There is no clear increasing or
968 decreasing trend of normalized core clock amplitude as a function of glucose, and therefore the
969 higher magnitude oscillations observed at high glucose concentrations are most likely a function
970 of increased biomass only (**F**). Growth rates were computed from biomass reporter experiments
971 by estimating the linear growth rate at 5 consecutive 12-hour intervals from 36 – 84 hours in
972 constant darkness (**G**). No arginine was added to the race tube medium for any experiment
973 shown.

974



975

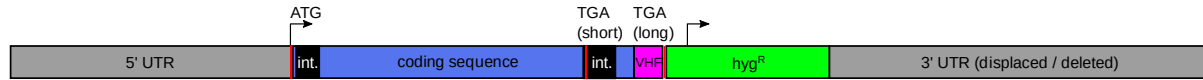
976 **Supplementary Figure 2. A subset of FGSC knockout strains were identified with strong**
977 **conidial banding phenotypes on glucose starvation medium.** A representative race tube
978 from the primary genetic screen is shown. 6 out of 7 knockout strains with strong banding
979 phenotypes have a wild-type circadian period length and normal compensation, except for
980 FGSC15333 (Figure 3). Like the *ras-1*^{bd} (NCU08823) point mutant, all knockout strains have a
981 reduced linear growth rate compared to the wild-type control FGSC2489.

982

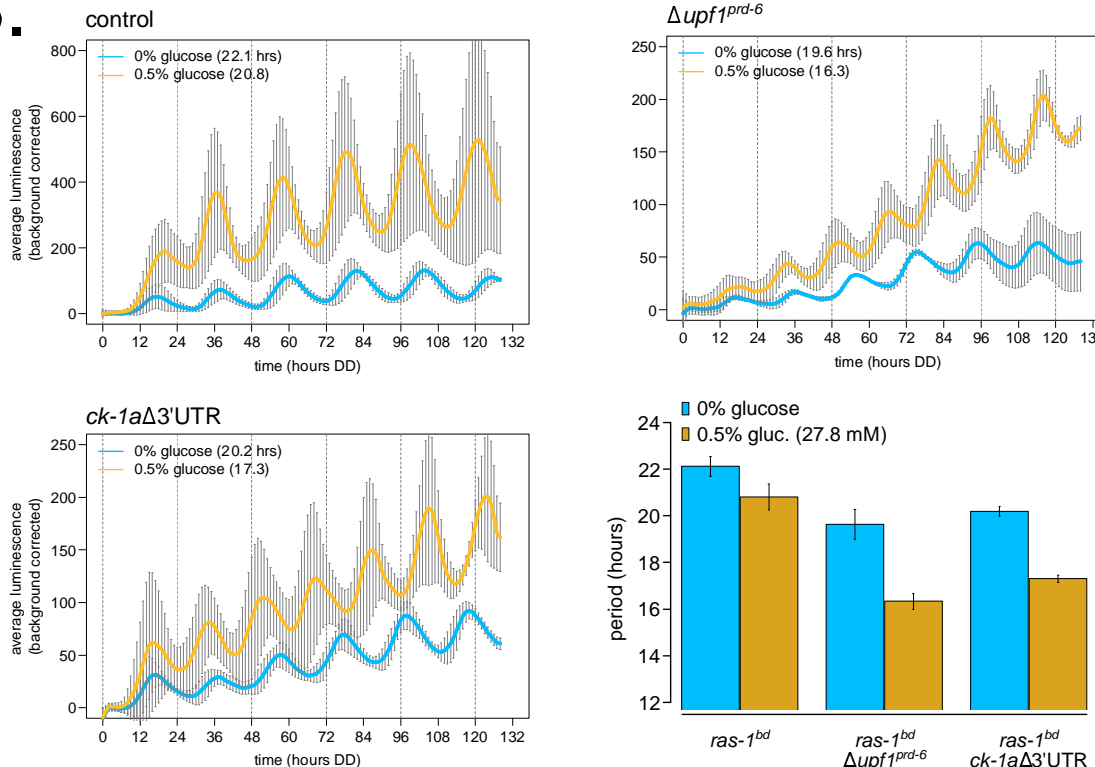
A. *ck-1a* (NCU00685) locus:



***ck-1a*Δ3'UTR mutant:**



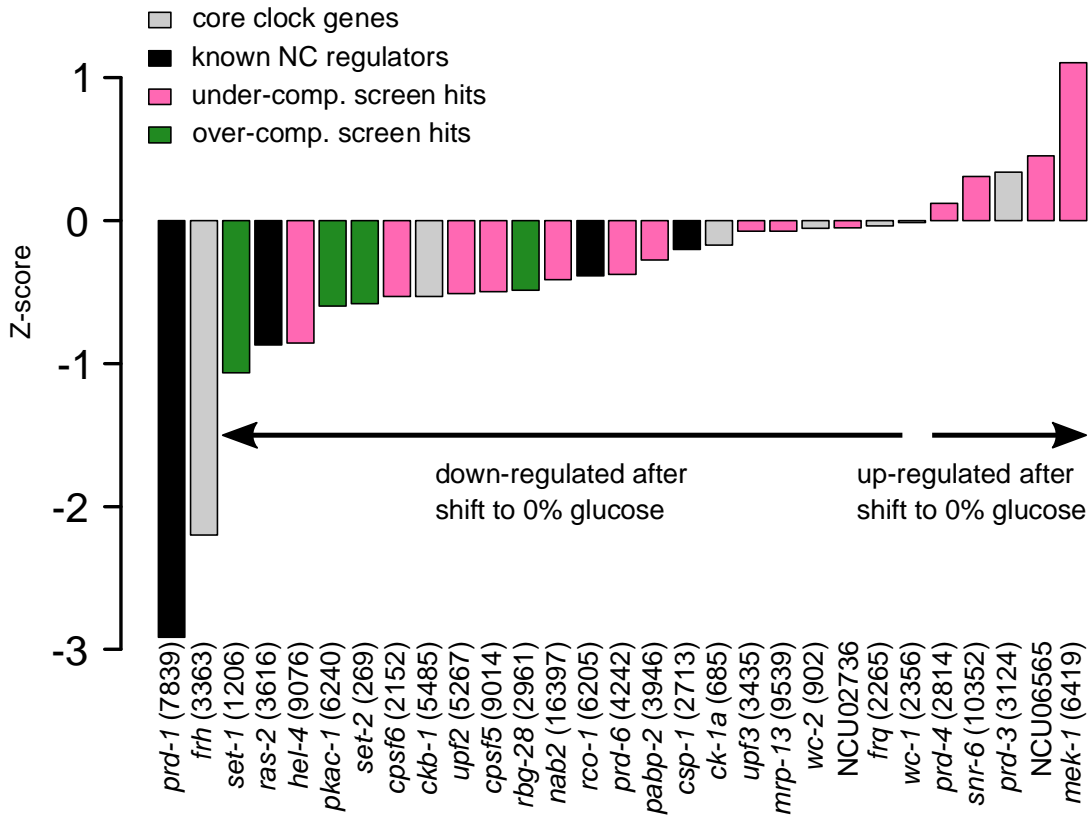
B.



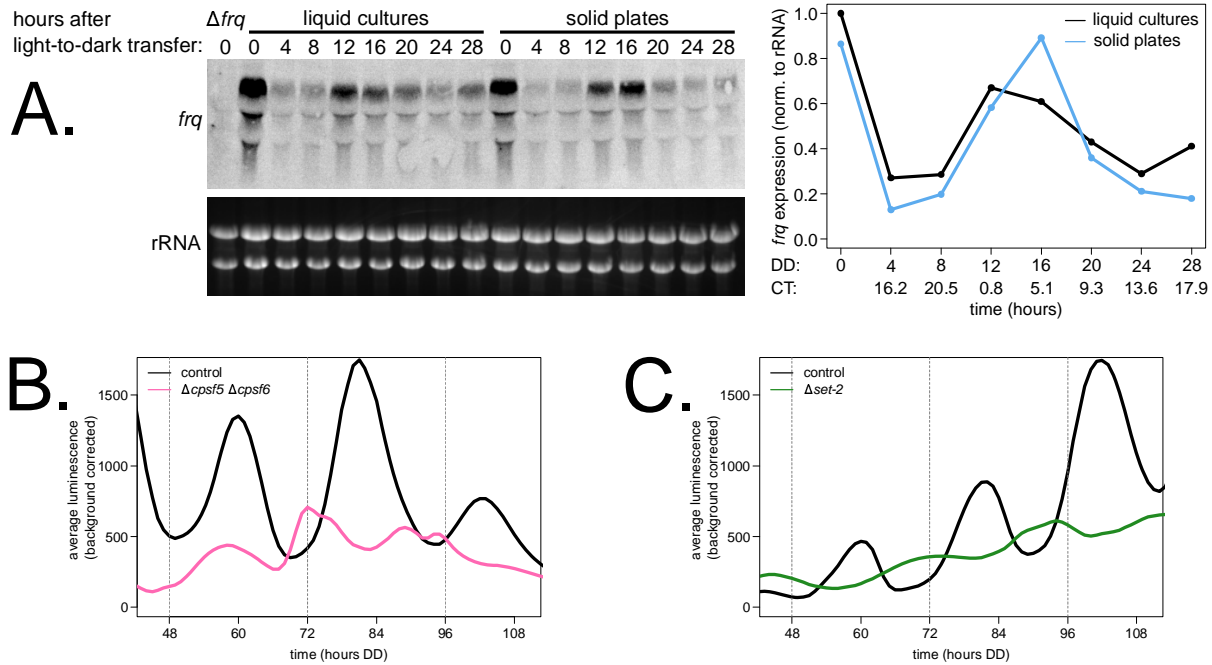
983

984 **Supplementary Figure 3. Deletion of the casein kinase 1 3' UTR phenocopies the short**
 985 **period length of nonsense-mediated decay (NMD) mutants, and partially explains the**
 986 **nutritional under-compensation phenotype of NMD mutants.** The CK1 3' UTR mutant was
 987 constructed by C-terminal epitope-tagging (V5-6xHis-3xFLAG) the LONG isoform of NCU00685
 988 at the endogenous locus and displacing 1,543 bps of the annotated 3' UTR region. Mutant strain
 989 construction is shown with a cartoon diagram to scale (A). Circadian bioluminescence was
 990 recorded from race tube cultures of the indicated genotypes. High nutrient medium (yellow lines)
 991 contained 0.5% w/v glucose 0.17% w/v arginine, and zero nutrient medium (blue lines)
 992 contained 0% glucose 0% arginine. Period lengths were computed (N = 3 - 4 biological
 993 replicates per nutrient concentration) and summarized in a bar graph (B).

994



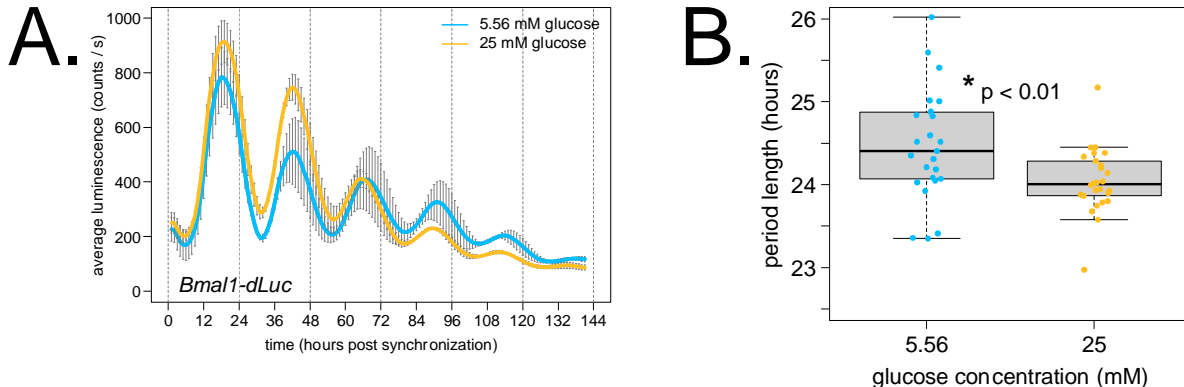
995
996 **Supplementary Figure 4. Core clock and compensation gene expression upon glucose**
997 **starvation.** RNA-Sequencing data were mined from a previous study (Wang et al., 2017)
998 (Materials and Methods), where liquid cultures were either maintained in 2% glucose or shifted
999 to glucose starvation for 60 minutes. Duplicate transcriptomes from the 2% glucose condition
1000 were compared to 0% glucose starvation replicates, Z-scores were computed for the 8,796
1001 expressed genes in the dataset, and *prd-1* (NCU07839) and *frh* (NCU03363) were found among
1002 the top 220 genes (top 2.5%) in the entire transcriptome down-regulated after glucose
1003 starvation.
1004



1005

1006 **Supplementary Figure 5. Solid medium cellophane plate cultures maintain circadian**
1007 **function and recapitulate Nutritional Compensation phenotypes of interest.** Liquid cultures
1008 containing wild-type fungal plugs (1.8% glucose) and cellophane plate cultures inoculated with
1009 wild-type conidia (0.5% w/v glucose, 0.17% w/v arginine) were set up concurrently. Liquid and
1010 solid cultures were entrained in constant light at 25°C for at least 16 hours, and serial light-to-
1011 dark transfers were performed to sample 1.5 cycles of circadian time points from DD4 to DD28
1012 (4-hour sampling density, 44 – 48 hour total culture ages). Total RNA was isolated from each
1013 time course sample, and *frq* mRNA rhythms were examined by Northern blot (N = 1 time course
1014 replicate). RNA levels were quantified using ImageJ densitometry, normalized, and plotted as
1015 line graphs. The circadian clock is clearly functional in both growth regimes (A). Circadian
1016 bioluminescence was recorded from cellophane plate cultures of the indicated genotypes grown
1017 on high nutrient medium (0.25% w/v glucose, 0.17% w/v arginine). One representative
1018 luciferase trace is shown from N = 2 biological replicates per strain. Period lengths were
1019 calculated, and results agree with Nutritional Compensation phenotypes derived from the race
1020 tube screen (Supplementary Table 1): control: 20.6 ± 0.3 hrs; $\Delta cpsf5 \Delta cpsf6$ double mutant:
1021 17.5 ± 0.4 hrs (B). Circadian bioluminescence was recorded from cellophane plate cultures of
1022 the indicated genotypes grown on zero nutrient medium (0% glucose, 0% arginine). One
1023 representative luciferase trace is shown from N = 2 biological replicates per strain. Period
1024 lengths were calculated, and results agree with Nutritional Compensation phenotypes derived

1025 from the race tube screen (Supplementary Table 1): control: 22.0 ± 0.6 hrs; Δ set-2: 20.5 ± 0.1
1026 hrs (C).
1027



1028
1029 **Supplementary Figure 6. *Bmal1-dLuc* cells are slightly under-compensated from 5.56 mM**
1030 **low glucose to 25 mM high glucose.** U2OS cells were synchronized using dexamethasone
1031 and released into DMEM low or high glucose medium. Representative luciferase traces are
1032 shown from one biological replicate with $N = 4$ technical replicates per glucose concentration
1033 (standard deviation error bars) (A). Period lengths were calculated for 25 – 26 total replicates
1034 per glucose concentration and plotted as a boxplot. As observed in *Neurospora*, the human
1035 circadian period length is under-compensated and shortens slightly with increasing glucose ($p <$
1036 0.01, student's t-test). Average period lengths were: 24.5 ± 0.7 hrs (low glucose) and 24.0 ± 0.4
1037 hrs (high glucose) (B).

1038
1039 **Supplementary Table 1. Genetic screen for Nutritional Compensation defects.** Results are
1040 presented from the 3-phase genetic screen for Nutritional Compensation defects among
1041 *Neurospora* knockout strains. Period lengths are shown for each knockout strain. Knockout
1042 strains were retained through each phase of the screen if circadian period changes were
1043 observed relative to the wild-type control. Nutritional under-compensation mutants were defined
1044 by ratios of high-to-zero glucose period lengths ≤ 0.90 (pink font). Nutritional over-compensation
1045 mutants had period length ratios ≥ 1.02 (green font).

1046
1047 **Supplementary Table 2. Informatic description of *Neurospora* genes screened for**
1048 **Nutritional Compensation defects.** Circadian rhythmicity at the gene and protein level among
1049 knockouts screened was determined from previous studies (Hurley et al., 2014, 2018). Promoter
1050 binding by the WCC positive arm transcription factors was determined from previous studies

1051 (Smith et al., 2010; Hurley et al., 2014; Sancar et al., 2015). Light-regulated gene activation or
1052 repression was determined from previous studies (Chen et al., 2009; Wu et al., 2014; Sancar et
1053 al., 2015). Gene expression Z-scores from carbon starvation conditions were also reported
1054 (Wang et al., 2017) (Supplementary Figure 4).

1055

1056 **Supplementary Table 3. Consensus list of *Neurospora* genes with Alternative**
1057 **Polyadenylation (APA) in 3' UTRs.** 843 genes contain multiple poly(A) sites within 3' UTR
1058 regions from the intersection of this study and previous work (Zhou et al., 2018). 9 / 843 genes
1059 with 3' UTR APA are highlighted as core clock genes or compensation screen hits. Annotated
1060 MACS2 poly(A) peak results are shown for the 843 genes with Alternative Polyadenylation from
1061 each dataset as individual tabs.

1062

1063 **Supplementary Table 4. Alternative Polyadenylation (APA) events altered in the Δ CFIm**
1064 **knockout mutant compared to controls.** 1,447 total genes contain multiple poly(A) sites
1065 within 3' UTR regions in wild-type control and/or Δ CFIm mutant data from this study. 940 / 1,447
1066 genes display APA in all 4 datasets. 123 / 1,447 instances are recorded where a single poly(A)
1067 peak in control expands to multiple APA events in mutant (dark green highlight). 193 / 1,447
1068 instances are reported of a single poly(A) peak in mutant expanding to multiple APA events in
1069 wild-type controls (light green highlight). 155 / 1,447 APA events occur where the location of the
1070 predominant poly(A) peak was significantly changed in the mutant background (orange
1071 highlight). Annotated MACS2 poly(A) peak results are shown for the APA genes from each
1072 dataset as individual tabs.

1073

1074 **Supplementary Movie 1. Nutritional Compensation properties of wild-type *Neurospora***
1075 ***crassa*.** Circadian bioluminescence of exemplar wild-type race tubes (Figure 1) is shown for
1076 zero nutrient medium (blue line; 0% glucose 0% arginine) (A) compared to high nutrient medium
1077 (yellow line; 0.5% w/v glucose 0% arginine) (B). Average period lengths are 21.9 ± 0.5 hours for
1078 0% glucose and 20.8 ± 0.2 hours for 0.5% w/v glucose (average \pm SD).

1079

1080 **Supplementary Movie 2. Nutritional Compensation defect in the Δ cpsf6 mutant.** Circadian
1081 bioluminescence of one exemplar Δ cpsf6 race tube (Figure 4A) is shown for whole-tube
1082 quantification of high amino acid medium growth (yellow line; 0% glucose 0.17% w/v arginine;
1083 average period = 19.4 ± 0.3 hrs) (A) compared to old tissue quantification (blue line; 0% glucose

1084 0.17% w/v arginine; period = 17.3 ± 0.3 hrs) (B). This ~2-hour period difference indicates that
1085 Δ cpsf mutants must undergo a transition from high-to-low amino acid levels to reveal the
1086 Nutritional Compensation defect.

1087

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1106

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