Deep sequencing analysis of the circadian transcriptome of the jewel wasp Nasonia vitripennis

Nathaniel J. Davies and Eran Tauber[†]

Dept.	of Genetics	s, Universit	y of Leicester	, University	y Road.	Leicester LE	1 7RH.	, UK

Running title: Sequencing the Nasonia circadian transcriptome

Keywords: Circadian clock; transcriptomics; Nasonia; RNAseq

[†] For correspondence. Dept. Genetics, University of Leicester, University Road, Leicester LE1 7RH, United Kingdom. Email: et22@le.ac.uk

2 **Abstract** 3 The study of the circadian clock has benefited greatly from using *Drosophila* as a 4 model system. Yet, accumulating evidence suggests that the fly might not be the 5 canonical insect model. Here, we have analysed the circadian transcriptome of the Iewl wasp *Nasonia vitripennis* by using RNA-seq in both constant darkness 6 7 and constant light (in contrast to flies, the wasps are rhythmic under continuous 8 with). We identify approximately 6% of the transcriptome as cycling under 9 constant conditions, revealing a bimodal distribution of phases and low cycling 10 amplitude. We examine the functions under circadian control in *Nasonia*, 11 identifying clock control of functions such as metabolism, light response, and a 12 variety of neural processes, drawing comparisons between Nasonia and 13 *Drosophila.* We characterise the transcriptional differences underlying 14 phenotypic differences in free-running circadian behaviour in constant darkness 15 and constant light, revealing significant down-regulation of catabolic processes 16 in constant light. We also profile levels of opsins transcription, gaining insight 17 into how Nasonia responds to light, which is a key question in circadian research 18 in the Hymenoptera. Although there was little similarity between cycling genes 19 in Drosophila and Nasonia, the functions fulfilled by cycling transcripts were 20 similar in both species. Of the known *Drosophila* core clock genes, only *pdp1e*, 21 shaggy and Clock showed a significant cycling in Nasonia, underscoring the 22 importance of studying the clock in non-model organisms.

23

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

Introduction The circadian clock regulates fundamental biological processes such as sleep (Huang, et al. 2011), metabolism (Huang, et al. 2011), and the immune system (Scheiermann, et al. 2013), and has implications for a wide range of human diseases. Notable examples of diseases linked to the circadian clock include cancer (Kelleher, et al. 2014), Alzheimer's disease (Musiek, et al. 2015), cardiovascular disease (Takeda and Maemura, 2011), obesity (Maury, et al. 2010), diabetes (Maury, et al. 2010), and depression (Quera Salva, et al. 2011). A primary output of the clock is circadian regulation of transcription, a trait which has been demonstrated in mammals (Hughes, et al. 2009), insects (McDonald and Rosbash, 2001a), plants (Schaffer, et al. 2001), and even bacteria (Woelfle and Johnson, 2006). Therefore, analysing transcriptional oscillations in clockcontrolled genes (CCGs) is a key step in understanding how the daily rhythms produced by the clock are ultimately linked to behavioural phenotypes. The genetic mechanisms underlying the animal circadian clock were first elucidated through studies of model animals; primarily the fruit fly *Drosophila*. The first clock gene to be identified, *period (per)*, was discovered through mapping the genetic basis of *Drosophila* mutants with aberrant locomotor and eclosion rhythms (Konopka and Benzer, 1971). The discovery of period was followed by the discovery of its heterodimeric partner timeless (tim) (Sehgal, et al. 1994). These two genes are joined by a roster of other genes working together to produce robust internal rhythms. The discoveries made in *Drosophila* have been instrumental for understanding the mechanisms of the circadian clock in mammals (Yu and

Hardin, 2006). As the principal insect model, *Drosophila* has been used to great

50

51

52

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

effect to model circadian phenomena in humans (Rosato, et al. 2006). However, as circadian research into non-drosophilid insects has advanced, several alternative clock models have been proposed (Yuan, et al. 2007), some of which may better model aspects of the mammalian clock than *Drosophila*. For example, a major difference between the various clock models in insects concerns the light input pathway. The main light input to the clock in *Drosophila* is mediated through *cryptochrome* (*cry1*) which is activated in response to light (Ceriani, et al. 1999), binds to and promotes the degradation of tim (Busza, et al. 2004), ultimately resulting in the degradation of per (Ko, et al. 2002, Grima, et al. 2002). In contrast, mammalian-like cryptochrome (cry2) is not light-sensitive (Yuan, et al. 2007), but is a part of the core transcriptional feedback loop suppressing its own transcription (and that of per) by interfering with the actions of the CLK-BMAL1 heterodimer (Kume, et al. 1999, Jin, et al. 1999). Mammals also lack a homolog for timeless, possessing only a homolog of the Drosophila gene timeout (Benna, et al. 2000), a gene whose potential role in the clock is less clear and less crucial than that of timeless (Gustafson and Partch, 2015, Benna, et al. 2010). The Lepidoptera harbour both types of *cryptochrome* (*Drosophila*-like cry1 and mammal-like cry2) (Tomioka and Matsumoto, 2010), as well as homologs of timeless and timeout (Tomioka and Matsumoto, 2015). The two cryptochromes have been shown to act in a similar way to their *Drosophila* and mammal counterparts; cry1 functions as a light receptor and cry2 serves as a transcriptional repressor (Zhu, et al. 2008). Of the major insect orders, the Hymenoptera arguably possess the most mammalian-like core clock architecture, possessing cry2 and timeout but neither

mammalian-like core clock architecture, possessing *cry2* and *timeout* but neither

75

76

77

78

79

80

81

82

83

84

85

86

87

88

89

90

91

92

93

94

95

96

97

cry1 nor timeless (Tomioka and Matsumoto, 2015, Yuan, et al. 2007). In addition to these molecular similarities, there is evidence that the transcriptional profiles of these genes match more closely the mammalian model than the *Drosophila* model (Rubin, et al. 2006). Light-entrained circadian rhythms have been demonstrated in the Hymenoptera, but the question of light detection in the Hymenopteran clock remains an open one. Nasonia vitripennis is a parasitoid wasp, which as a research model offers advantages over other hymenopterans, including a fully sequenced genome (Werren, et al. 2010), systemic RNAi (Lynch and Desplan, 2006), a robust and well-characterised circadian response (Bertossa, et al. 2013), a fully functional DNA methylation kit (Park, et al. 2011), and a history as a model for photoperiodism (Saunders, 1969). In this study, we advance *Nasonia* as an alternative circadian model by using RNA-seq to profile whole-transcriptome gene expression in the *Nasonia* head. As the Nasonia clock free-runs in both constant darkness and constant light (Figure 1), we profiled both of these conditions to examine how the two circadian transcriptomes differ. To our knowledge, this is the first circadian RNA-seq study performed in an insect other than *Drosophila*, and the first study to profile the circadian transcriptome oscillating under constant light. Results Identifying rhythmic transcription We first performed an unbiased clustering analysis to ascertain the kinds of expression patterns present in the data. To this end, Mfuzz (Kumar and E Futschik, 2007) was used to carry soft c-means clustering, a method which is less

rutschik, 2007) was used to carry soft c-means clustering, a method which is less

99

100

101

102

103

104

105

106

107

108

109

110

111

112

113

114

115

116

117

118

119

120

121

122

sensitive to biological noise than traditional clustering (Futschik and Carlisle, 2005). After filtering (see Methods), thirty clusters were generated for each condition (Supplementary figures S1 and S2), revealing a variety of potentially rhythmic and non-rhythmic expression trends. Potential asymmetric wave forms were detected in LL (e.g. Supplementary figure S2, clusters 22 and 26). To identify rhythmic transcripts, we used the RAIN algorithm (Thaben and Westermark, 2014). At false discovery rate (FDR) threshold of 0.1 we identified 1,057 rhythmic transcripts in DD and 929 in LL (Table S1, S2). Rhythmic transcripts (q < 0.1) were sorted by phase, peak shape, and significance, and plotted (Figure 2A). Examining the phase distribution (Figure 2B), it is apparent that the majority of transcripts show peak expression early in the subjective morning/afternoon or in the subjective night, with fewer transcripts peaking at intermediate times. This disparity in phase is greater in the transcripts which show rhythmic expression in both DD and LL; less than 12% of transcripts in DD and less than 5% in LL show peak expression at intermediate times (Figure 2B). The majority of these transcripts (~87%) exhibit a similar (+-4 hrs) phase in LL to their phase in DD. Similarly to Drosophila (Hughes, et al. 2012) and mammals (Hughes, et al. 2009), the majority of transcripts show only small cyclic changes in expression amplitude over the day; over 80% of reliably quantified (see Methods) transcripts in both conditions have amplitudes (peak expression divided by trough expression) of 2 or less. In both DD and LL, transcripts with exceptionally high amplitudes (> 4) are transcripts with unusually low or high measurements at isolated time-points with no obvious specific shared function. This is in contrast with results in *Drosophila* and mammals, where some core clock genes

124

125

126

127

128

129

130

131

132

133

134

135

136

137

138

139

140

141

142

143

144

145

146

147

exhibit very high amplitude oscillations (Hughes, et al. 2009, Hughes, et al. 2012, Li, et al. 2015). Canonical clock genes and comparison with Drosophila The canonical clock genes were examined for rhythmicity both at the transcript level and via an additional RAIN analysis at the gene level. The q-values (FDR adjusted p-values) for the canonical clock genes are shown in supplementary table S3. We found a rather limited evidence for rhythmicity in these genes which included pdp1e ($q \sim 0.1$, LL and DD), shaqqv (q < 0.1, DD), and Clok ($q \sim$ 0.1, LL). At a less stringent FDR (q< 0.2), per, cyc, Dbt and cwo were rhythmic in DD, while cry and cyc, were oscillating in LL. For comparison between splice variants and conditions, median expression levels of the canonical clock genes and their transcripts for both DD and LL are shown in supplementary table S4. We compared the transcripts identified as cycling in Nasonia heads with the transcripts identified as cycling in *Drosophila* heads. For these purposes, we used a list of genes identified in a meta-analysis study of *Drosophila* circadian microarray data as being rhythmically expressed in either LD or DD (Keegan, et al. 2007). Of 173 genes identified as rhythmic in *Drosophila*, 33 genes (Supplementary table S5) were found to also be rhythmic in *Nasonia* (either in LL or DD, q < 0.1), no more than would be expected by chance (p = 0.11, hypergeometric test). **Functions of rhythmic genes** To capture the general functions that rhythmic genes may fulfil in *Nasonia*, we tested a broader set of rhythmic genes (FDR < 0.2 in RAIN) for GO term

149

150

151

152

153

154

155

156

157

158

159

160

161

162

163

164

165

166

167

168

169

170

171

172

overrepresentation (Davies and Tauber, 2015a), revealing 94 GO terms overrepresented for genes rhythmic in DD (including 'response to light stimulus', 'proteasome complex', and 'generation of neurons', Supplementary table S6) and 123 terms for genes rhythmic in LL (including 'locomotion', 'proteasome complex', and 'response to external stimulus', Supplementary table S7), 25 of which were shared between both conditions (Figure 3). Shared terms include terms related to neurons, signal transmission, and responses to stimuli. Notably, all four Nasonia opsins were found to exhibit similar transcriptional profiles in LL and DD, with low expression in the morning and high expression in the evening. It has previously been demonstrated that the timing of different (or indeed opposing) biological processes can be controlled through the circadian regulation of groups of genes (Sancar, et al. 2015, Zhang, et al. 2014). Unsupervised clustering methods have previously been established as a useful method for functional characterisation of circadian genes (Nguyen, et al. 2014). To establish whether temporal separation of functions occurs in Nasonia, we therefore returned to the expression clustering analysis. Firstly, we employed hypergeometric tests to identify clusters with an overrepresentation of rhythmic genes (Figure 4, Supplementary table S8 and S9). Clusters which were found to have a significant rhythmic component (q < 0.05, supplementary tables S8 and S9) were analysed for overrepresented G0 terms. Examples of clusters with enriched functions include clusters DD7 and LL20 which are significantly enriched for catalytic activity GO terms, especially genes involved in the proteasome, and clusters DD24 and LL6 which are both involved in circadian and neural processes. Other clusters (DD1 and DD2) did not turn up any

and heurar processes. Other clusters (DDT and DD2) the not turn up any

174

175

176

177

178

179

180

181

182

183

184

185

186

187

188

189

190

191

192

193

194

195

overrepresented GO terms and are thus likely comprised of genes with a wide range of functions. Transcriptional differences between constant darkness and constant light To examine whether differences in circadian period seen in locomotor activity between DD and LL could also be detected in transcriptional rhythms, we fitted parametric models with a range of periods to transcripts rhythmic in both conditions (q < 0.1). For those transcripts with statistically significant fits to the model in both conditions (q < 0.1, see Methods), we took the period with the best fit and compared these periods between conditions. Overall, transcripts in LL showed a significantly (p < 3.9e-09, Wilcoxon rank sum test) shorter (median 24) period than those in DD (median 25.4), mirroring the behavioural differences in period. We have also tested for differential expression between DD and LL. In the absence of biological replicates, we analysed differential expression using a foldchange approach. We used 1.5 fold change as a cut-off for differential expression (Dalman, et al. 2012), yielding 1,488 genes expressed higher in DD than LL and 971 genes expressed higher in LL than DD (Figure 5). Genes more highly expressed in DD were significantly enriched (q < 0.01) for genes involved in various forms of catalytic activity (Supplementary table S10), including the vast majority of proteasome genes (>75%). Genes more highly expressed in LL were enriched for a small number of terms including 'plasmalemma' and 'sequence-

specific DNA binding' (Supplementary table S11).

197

198

199

200

201

202

203

204

205

206

207

208

209

210

211

212

213

214

215

216

217

218

219

220

Discussion This study provides the first insights into global transcriptional oscillation in *Nasonia*. With RNA-seq, we profiled the circadian transcription of >26,000 transcripts in Nasonia in either DD or LL. At a relatively stringent FDR (q < 0.1), we identified 1,057 cycling transcripts in DD and 929 cycling transcripts in LL. These transcripts correspond to a cycling fraction of 6.7% and 5.9% of all transcripts analysed in DD and LL respectively. These figures are comparable to cycling fractions reported in various organisms and tissues, generally between 2% and 10% of the transcriptome (Michael and McClung, 2003). In both conditions, cycling transcripts were found to cycle at low amplitudes (mostly < 2 fold) and with a limited, bimodal, range of phases. This is in contrast to microarray/RNA-seq studies in *Drosophila*, where transcripts were found to cycle with a broader range of phases (Rodriguez, et al. 2013) and studies in both mammals and Drosophila, which have identified a group of highamplitude (> 4-fold) cycling genes among the low-amplitude majority (Akhtar, et al. 2002). High amplitude cyclers typically include clock genes (Akhtar, et al. 2002, Hughes, et al. 2012). The low oscillations of the Nasonia head transcriptome render the expression profiles of the canonical clock genes difficult to resolve (Covington, et al. 2008). This issue may also contribute to the discordance between the various circadian microarray studies in *Drosophila* (Keegan, et al. 2007). An emerging property of the circadian transcriptome in *Nasonia* is the temporal separation of function by phase (Fig 2). Notably, genes involved in catalytic activity were strongly overrepresented in morning-peaking transcripts.

This is in line with other studies which show catalytic activity confined to the

222

223

224

225

226

227

228

229

230

231

232

233

234

235

236

237

238

239

240

241

242

243

244

morning in fungi (Sancar, et al. 2015), in agreement with a general observation that an important (or even primary) function of circadian clocks (Hurley, et al. 2015) is to temporally separate catabolism and anabolism. Although we did not detect an overrepresentation of anabolic genes within the cyclic transcripts, expression clusters DD10 and LL24 (Supplementary figures S1 and S2) did show strong overrepresentation (Supplementary tables S12 and S13) for genes involved in cytosolic ribosomal genes (q < 3.e-56) and cellular anabolism (q < 2e-6) 06). These clusters exhibit an antagonistic expression pattern to the expression clusters containing the catabolic genes, suggesting that catabolism and anabolism are indeed separated by the circadian clock in Nasonia. The comparison of expression between LL and DD reveals that a majority of genes involved in the proteasome and a broader set of genes involved in catabolism, are more highly expressed in DD than LL. As turnover rates of clock proteins have shown to be coupled with changes in the circadian period (Syed, et al. 2011, He and Liu, 2005), up-regulation of the proteasome may provide an explanation for differences in period observed between DD and LL. Although the genes which cycle in *Drosophila* largely differ from those cycling in *Nasonia*, the functions fulfilled by CCGs in *Nasonia* are similar to the functions filled by CCGs in *Drosophila*. Examples of functions shared by CCGs in the Drosophila and Nasonia heads are: various aspects of metabolism (Rodriguez, et al. 2013, Ueda, et al. 2002, Ceriani, et al. 2002, Claridge-Chang, et al. 2001), phototransduction (Ueda, et al. 2002, Rodriguez, et al. 2013), synaptic/nervous functions (McDonald and Rosbash, 2001b, Ceriani, et al. 2002, Claridge-Chang, et al. 2001), oxidoreductase activity (Claridge-Chang, et al. 2001), mating behaviour

246

247

248

249

250

251

252

253

254

255

256

257

258

259

260

261

262

263

264

265

266

267

268

(Rodriguez, et al. 2013), and immunity (McDonald and Rosbash, 2001b, Ceriani, et al. 2002). We identified cycling of genes involved in response to light, particularly all four Nasonia opsins. These opsins, along with associated gPCRs, cycle with a similar phase and are all more highly expressed in LL than in DD (Supplementary figure S6). Daily and circadian changes in opsin expression have been demonstrated in other organisms (e.g. mice (Bowes, et al. 1988), zebrafish (Li, et al. 2005), honeybee (Sasagawa, et al. 2003)), and opsin expression is generally found to be up-regulated in response to light (Yan, et al. 2014). Characterising the opsins in *Nasonia* is likely to provide insights into the light input pathway into the clock, particularly as *Nasonia* does not possess other obvious light input candidate genes such as Drosophila-like CRY1 (Bertossa, et al. 2014) or Pteropsin (Velarde, et al. 2005) (Supplementary figure S6). Data availability We have made the expression profile for each transcript in both conditions available on WaspAtlas (Davies and Tauber, 2015b). Data have been archived in the NCBI short read archive (SRA), with accession number PRJNA318159. Methods Maintenance and sample collection Stocks of *Nasonia vitripennis* (strain AsymCX) were maintained at 25°C on blowfly pupal hosts in 12:12 light:dark cycles. To obtain male wasps for experiments, groups of eight females were isolated at the yellow pupal stage and transferred onto fresh hosts upon eclosion. The resulting male progeny were

collected upon eclosion and moved onto vials with a 30% sucrose agar medium,

270

271

272

273

274

275

276

277

278

279

280

281

282

283

284

285

286

287

288

289

290

291

292

293

in groups of 20. During entrainment (four full days in an LD 12:12 cycle) and collection, wasps were kept in four light boxes in the same incubator at 19°C. Starting at CT1, wasps were collected every four hours and snap-frozen in liquid nitrogen and immediately transferred to -80°C. Wasps were collected sequentially from light box to light box every four hours to minimise disturbance of wasps, and so that wasps were collected from each light box once every 16 hours, thereby minimising the effect of variations within light boxes. Temperature and light recordings were taken during the experiment, and can be viewed in Supplementary file S2. To verify that wasps entrained correctly to the experimental conditions and that free-running behaviour was as expected, individual male wasps were isolated and locomotor activity was monitored. Behavioural recordings of individual male wasps in experimental conditions can be seen in Supplementary figure S7, ruling out behavioural differences caused by inter light box variations in light intensity in LL, though not transcriptional differences. RNA extraction, sequencing, and read mapping RNA was extracted from pooled groups of 50 heads for each sample, using Trizol RNA extraction protocol, and followed by clean-up using the RNAeasy spin column kit (Qiagen). Samples were poly A selected and sequenced at Glasgow Polyomics (University of Glasgow, United Kingdom) on the Illumina NextSeq500 platform, resulting in approximately 20 million 75bp paired-end reads per sample. Read mapping was achieved with Tophat2 (v2.1.0) (Trapnell, et al. 2012) against the Nasonia Nvit_2.1 NCBI annotation. As the purpose of this study was

295

296

297

298

299

300

301

302

303

304

305

306

307

308

309

310

311

312

313

314

315

316

317

not to identify novel splice variants or improve on existing annotation, novel junction detection was disabled for accurate quantification of known transcripts. Mean mapping efficiency was above 90% for both conditions (Supplementary table S14). Read quantification was performing using the DEseq normalisation method (Anders and Huber, 2010). All 24 samples from both conditions were grouped together to allow comparison between as well as within conditions. **Expression profile clustering** Isoform expression profiles were first filtered to include only those isoforms with no missing values at any time-point in either condition. Expression values were standardised using the 'Standardise' function in Mfuzz (Kumar and E Futschik, 2007). The 'cselection' function in Mfuzz was used to select an appropriate c-value for the c-means clustering (default parameters; m=1.25). Based on this analysis, thirty fuzzy clusters were generated for each condition using the fuzzification parameter m=1.25. Rhythmic expression analysis RAIN (Thaben and Westermark, 2014) was used on all filtered isoforms (i.e. those with no missing values at any timepoint) in either condition to detect rhythmic isoforms at a period of 24 hours. As a non-parametric method, RAIN only facilitates detection of rhythmic isoforms with periods which are a multiple of the sample resolution (in this case 4 hr). The p-values produced by RAIN were corrected to q-values using the Benjamini-Hochberg method (Benjamini and Hochberg, 1995). This method was repeated using expression values for genes

319

320

321

322

323

324

325

326

327

328

329

330

331

332

333

334

335

336

337

338

339

340

341

rather than transcripts for the clock gene analysis (i.e. the summed expression values for all known transcripts of a particular gene). Maximum fold changes in expression were calculated by normalising percondition expression values by the median value and calculating the ratio from the lowest expression over 48 hours to the highest. Reliably quantified transcripts are defined as those those transcripts where the absolute FPKM value is 5 or above at all timepoints, the threshold for this set at a similar level to other analyses (Hughes, et al. 2012). To analyse the period of rhythmic transcripts, we fitted parametric waveforms with a variety of periods (20 to 28 hrs in steps of 0.2 hrs) to all transcripts identified as rhythmic (q < 0.1) in both conditions. This FDR threshold is in line with, or more strict, than thresholds chosen in other similar studies (Hughes, et al. 2012, Huang, et al. 2013, Keegan, et al. 2007). Those transcripts (85 in total) which showed a significant (q < 0.1) fit to the model in both conditions were analysed in terms of their best fitting period. GO term overrepresentation was performed in WaspAtlas (Davies and Tauber, 2015b) using the Nvit_2.1 NCBI annotation dataset. All hypergeometric tests were performed within R using the 'phyper' function. Clusters with rhythmic components were identified by collapsing the fuzzy clusters into hard clusters using the 'cluster' property of the Mfuzz object, performing hypergeometric tests to identify clusters with enrichment for rhythmic transcripts. Thirty tests were performed for each condition (i.e. for all clusters), and were corrected per-condition using the Benjamini-Hochberg method in R (R

Development Core Team, 2008).

343

344

345

346

347

348

349

350

351

352

353

354

355

356

357

358

359

360

361

362

363

For comparison to microarray studies, orthologs for *Drosophila* melanogaster were obtained from a meta-study of circadian microarray data (Keegan, et al. 2007). The 214 obtained FlyBase identifiers were converted to the latest identifiers using the validation tool, resulting in 218 unique identifiers (the increase in identifiers can be attributed to previous identifiers referring to multiple genes in the current annotation). Orthologs for these *Drosophila* genes were obtained through WaspAtlas, retrieving orthologs for 135 genes which mapped to 173 unique Nasonia genes due to gene duplications, etc. This set of 173 genes was compared with the number of genes with rhythmic transcripts that would be expected by chance using a hypergeometric test. Phylogenetic analysis of opsin genes Opsin genes were searched for using NCBI BLASTP using six species; Apis mellifera, Bombyx mori, Drosophila melanogaster, Mus musculus, Nasonia vitripennis, and Homo sapiens, using the Nasonia Lop1 protein sequence as a query. BLAST results were inspected and 7e-19 was chosen as an appropriate cut-off to include all opsin sequences. Sequences were aligned by ClustalW in MEGA (Tamura, et al. 2007) and a maximum likelihood tree generated using default parameters. Duplicated sequences were manually removed, and sequences renamed for display on the tree. Full protein name to shortened display name translations can be found in supplementary table S15.

364 References 365 Akhtar RA, Reddy AB, Maywood ES, Clayton JD, King VM, Smith AG, Gant TW, 366 Hastings MH, Kyriacou CP. 2002. Circadian cycling of the mouse liver 367 transcriptome, as revealed by cDNA microarray, is driven by the 368 Suprachiasmatic Nucleus. Current Biology, 12: 540-550. 369 Anders S and Huber W. 2010. Differential expression analysis for sequence count 370 data. Genome Biol. 11: R106-2010-11-10-r106. Epub 2010 Oct 27. 371 Benjamini Y and Hochberg Y. 1995. Controlling the False Discovery Rate: A 372 Practical and Powerful Approach to Multiple Testing. Journal of the Royal 373 Statistical Society. Series B (Methodological) 57: 289-300. 374 Benna C, Bonaccorsi S, Wulbeck C, Helfrich-Forster C, Gatti M, Kyriacou CP, Costa 375 R, Sandrelli F. 2010. Drosophila timeless 2 is required for chromosome stability 376 and circadian photoreception. Curr. Biol. 20: 346-352. 377 Benna C, Scannapieco P, Piccin A, Sandrelli F, Zordan M, Rosato E, Kyriacou CP, 378 Valle G, Costa R. 2000. A second timeless gene in Drosophila shares greater 379 sequence similarity with mammalian tim. *Curr. Biol.* **10**: R512-3. 380 Bertossa RC, van de Zande L, Beukeboom LW, Beersma DG. 2014. Phylogeny and 381 oscillating expression of period and cryptochrome in short and long 382 photoperiods suggest a conserved function in Nasonia vitripennis. Chronobiol. 383 Int. 31: 749-760.

384 Bertossa RC, van Dijk J, Diao W, Saunders D, Beukeboom LW, Beersma DG. 2013. 385 Circadian rhythms differ between sexes and closely related species of Nasonia 386 wasps. *PLoS One* **8**: e60167. 387 Bowes C, van Veen T, Farber DB. 1988. Opsin, G-protein and 48-kDa protein in 388 normal and rd mouse retinas: developmental expression of mRNAs and proteins 389 and light/dark cycling of mRNAs. Exp. Eye Res. 47: 369-390. 390 Busza A, Emery-Le M, Rosbash M, Emery P. 2004. Roles of the two Drosophila 391 CRYPTO CHROME structural domains in circadian photoreception. *Science* **304**: 392 1503-6. 393 Ceriani MF, Darlington TK, Staknis D, Mas P, Petti AA, Weitz CJ, Kay SA. 1999. 394 Light-dependent sequestration of TIMELESS by CRYPTOCHROME. Science 285: 395 553-6. 396 Ceriani MF, Hogenesch JB, Yanovsky M, Panda S, Straume M, Kay SA. 2002. 397 Genome-Wide Expression Analysis in Drosophila Reveals Genes Controlling 398 Circadian Behavior. J. Neurosci. 22: 9305-9319. 399 Claridge-Chang A, Wijnen H, Naef F, Boothroyd C, Rajewsky N, Young MW. 2001. 400 Circadian regulation of gene expression systems in the Drosophila head. Neuron 401 **32**: 657-671 402 Covington MF, Maloof JN, Straume M, Kay SA, Harmer SL. 2008. Global 403 transcriptome analysis reveals circadian regulation of key pathways in plant 404 growth and development. Genome Biol. 9: R130.

grower and development, denome Biol. 7. K130.

405 Dalman MR, Deeter A, Nimishakavi G, Duan ZH. 2012. Fold change and p-value 406 cutoffs significantly alter microarray interpretations. BMC Bioinformatics 13 407 **Suppl 2**: S11-2105-13-S2-S11. 408 Davies NJ and Tauber E. 2015a. WaspAtlas: a Nasonia vitripennis gene database 409 and analysis platform. *Database (Oxford)* **2015**: bav103. Print 2015. 410 Davies NJ and Tauber E. 2015b. WaspAtlas: a Nasonia vitripennis gene database 411 and analysis platform. Database (Oxford) 2015: 10.1093/database/bav103. Print 412 2015. 413 Futschik ME and Carlisle B. 2005. Noise-robust soft clustering of gene expression 414 time-course data. J. Bioinform Comput. Biol. 3: 965-988. 415 Grima B, Lamouroux A, Chelot E, Papin C, Limbourg-Bouchon B, Rouyer F. 2002. 416 The F-box protein slimb controls the levels of clock proteins period and timeless. 417 Nature **420**: 178-82 418 Gustafson CL and Partch CL. 2015. Emerging models for the molecular basis of 419 mammalian circadian timing. *Biochemistry* **54**: 134-149. 420 He O and Liu Y. 2005. Degradation of the Neurospora circadian clock protein 421 FREQUENCY through the ubiquitin-proteasome pathway. *Biochem. Soc. Trans.* 422 **33**: 953-956 423 Huang W, Ramsey KM, Marcheva B, Bass J. 2011. Circadian rhythms, sleep, and 424 metabolism. J. Clin. Invest. 121: 2133-2141.

metabolism. J. Chn. Mvest. 121. 2133-2141.

425 Huang Y, Ainsley JA, Reijmers LG, Jackson FR. 2013. Translational profiling of 426 clock cells reveals circadianly synchronized protein synthesis. *PLoS Biol.* **11**: 427 e1001703. 428 Hughes ME, DiTacchio L, Hayes KR, Vollmers C, Pulivarthy S, Baggs JE, Panda S, 429 Hogenesch JB. 2009. Harmonics of circadian gene transcription in mammals. 430 *PLoS Genet.* **5**: e1000442. 431 Hughes ME, Grant GR, Paquin C, Qian J, Nitabach MN. 2012. Deep sequencing the 432 circadian and diurnal transcriptome of Drosophila brain. Genome Res. 22: 1266-433 1281. 434 Hurley JM, Loros JJ, Dunlap JC. 2015. The circadian system as an organizer of 435 metabolism. Fungal Genet. Biol. 436 Jin X, Shearman LP, Weaver DR, Zylka MJ, de Vries GJ, Reppert SM, 1999. A 437 molecular mechanism regulating rhythmic output from the suprachiasmatic 438 circadian clock. Cell 96: 57-68. 439 Keegan KP, Pradhan S, Wang JP, Allada R. 2007. Meta-analysis of Drosophila 440 circadian microarray studies identifies a novel set of rhythmically expressed 441 genes. PLoS Comput. Biol. 3: e208. 442 Kelleher FC, Rao A, Maguire A. 2014. Circadian molecular clocks and cancer. 443 Cancer Lett. **342**: 9-18 444 Ko HW, Jiang J, Edery I. 2002. Role for Slimb in the degradation of Drosophila 445 Period protein phosphorylated by Doubletime. *Nature* **420**: 673-678.

Konopka R and Benzer S. 1971. Clock mutants of Drosophila melanogaster. Proc. 446 447 Natl. Acad. Sci. USA 68: 2112-2116. 448 Kumar L and E Futschik M. 2007. Mfuzz: a software package for soft clustering of 449 microarray data. Bioinformation 2: 5-7. 450 Kume K, Zylka MJ, Sriram S, Shearman LP, Weaver DR, Jin X, Maywood ES, 451 Hastings MH, Reppert SM. 1999. mCRY1 and mCRY2 are essential components of 452 the negative limb of the circadian clock feedback loop. *Cell* **98**: 193-205. 453 Li, J., Grant, G. R., Hogenesch, J. B., and Hughes, M. E. 2015. Chapter Sixteen -454 Considerations for RNA-seq Analysis of Circadian Rhythms. In *Methods in* 455 Enzymology (ed. Amita Sehgal), pp. 349-367. Academic Press, . 456 Li P, Temple S, Gao Y, Haimberger TJ, Hawryshyn CW, Li L. 2005. Circadian 457 rhythms of behavioral cone sensitivity and long wavelength opsin mRNA 458 expression: a correlation study in zebrafish. *J. Exp. Biol.* **208**: 497-504. 459 Lynch JA and Desplan C. 2006. A method for parental RNA interference in the 460 wasp Nasonia vitripennis. Nat. Protoc. 1: 486-494. 461 Maury E. Ramsey KM, Bass J. 2010. Circadian Rhythms and Metabolic Syndrome: 462 From Experimental Genetics to Human Disease. Circulation Research 106: 447-463 462. 464 McDonald MJ and Rosbash M. 2001a. Microarray analysis and organization of 465 circadian gene expression in *Drosophila Cell* **107**: 567-578.

466 McDonald MJ and Rosbash M. 2001b. Microarray analysis and organization of 467 circadian gene expression in Drosophila. *Cell* **107**: 567-578. 468 Michael TP and McClung CR. 2003. Enhancer trapping reveals widespread 469 circadian clock transcriptional control in Arabidopsis. Plant Physiol. 132: 629-470 639. 471 Musiek ES, Xiong DD, Holtzman DM. 2015. Sleep, circadian rhythms, and the 472 pathogenesis of Alzheimer disease. Exp. Mol. Med. 47: e148. 473 Nguyen TT, Mattick JS, Yang Q, Orman MA, Jerapetritou MG, Berthiaume F, 474 Androulakis IP. 2014. Bioinformatics analysis of transcriptional regulation of 475 circadian genes in rat liver. *BMC Bioinformatics* **15**: 83-2105-15-83. 476 Park J, Peng Z, Zeng J, Elango N, Park T, Wheeler D, Werren JH, Yi SV. 2011. 477 Comparative analyses of DNA methylation and sequence evolution using 478 Nasonia genomes. Mol. Biol. Evol. 28: 3345-3354. 479 Ouera Salva MA, Hartley S, Barbot F, Alvarez JC, Lofaso F, Guilleminault C. 2011. 480 Circadian rhythms, melatonin and depression. Curr. Pharm. Des. 17: 1459-1470. 481 R Development Core Team. 2008. R: A Language and Environment for Statistical 482 Computing. R Foundation for Statistical Computing, Vienna, Austria. 483 Rodriguez J. Tang CH, Khodor YL, Vodala S, Menet JS, Rosbash M. 2013. Nascent-484 Seq analysis of Drosophila cycling gene expression. Proc. Natl. Acad. Sci. U. S. A. 485 **110**: E275-84

55 **110.** E275-04.

486 Rosato E, Tauber E, Kyriacou CP. 2006. Molecular genetics of the fruit-fly 487 circadian clock. Eur. J. Hum. Genet. 14: 729-38. 488 Rubin EB, Shemesh Y, Cohen M, Elgavish S, Robertson HM, Bloch G. 2006. 489 Molecular and phylogenetic analyses reveal mammalian-like clockwork in the 490 honey bee (Apis mellifera) and shed new light on the molecular evolution of the 491 circadian clock. Genome Res. 16: 1352-1365. 492 Sancar C, Sancar G, Ha N, Cesbron F, Brunner M. 2015. Dawn- and dusk-phased 493 circadian transcription rhythms coordinate anabolic and catabolic functions in 494 Neurospora. *BMC Biol.* **13**: 17-015-0126-4. 495 Sasagawa H, Narita R, Kitagawa Y, Kadowaki T. 2003. The expression of genes 496 encoding visual components is regulated by a circadian clock, light environment 497 and age in the honeybee (Apis mellifera). Eur. J. Neurosci. 17: 963-970. 498 Saunders DS. 1969. Diapause and photoperiodism in the parasitic wasp Nasonia 499 vitripennis, with special reference to the nature of the photoperiodic clock. *Symp.* 500 Soc. Exp. Biol. 23: 301-29 501 Schaffer R, Landgraf J, Accerbi M, Simon V, Larson M, Wisman E. 2001. 502 Microarray analysis of diurnal and circadian-regulated genes in Arabidopsis. 503 Plant Cell 13: 113-123. 504 Scheiermann C, Kunisaki Y, Frenette PS. 2013. Circadian control of the immune 505 system. Nat. Rev. Immunol. 13: 190-198.

506 Sehgal A, Price J, Man B, Young M. 1994. Loss of circadian behavioral rhythms 507 and per RNA oscillations in the Drosophila mutant timeless. Science 263: 1603-508 1606. 509 Syed S, Saez L, Young MW. 2011. Kinetics of doubletime kinase-dependent 510 degradation of the Drosophila period protein. J. Biol. Chem. 286: 27654-27662. 511 Takeda N and Maemura K. 2011. Circadian clock and cardiovascular disease. J. 512 Cardiol. 57: 249-256. 513 Tamura K, Dudley J, Nei M, Kumar S. 2007. MEGA4: Molecular Evolutionary 514 Genetics Analysis (MEGA) software version 4.0. Mol. Biol. Evol. 24: 1596-1599. 515 Thaben PF and Westermark PO. 2014. Detecting rhythms in time series with 516 RAIN. J. Biol. Rhythms 29: 391-400. 517 Tomioka K and Matsumoto A. 2010. A comparative view of insect circadian clock 518 systems. Cell Mol. Life Sci. 67: 1397-1406. 519 Tomioka K and Matsumoto A. 2015. Circadian molecular clockworks in non-520 model insects. Current Opinion in Insect Science; Insect genomics * Development 521 and regulation 7: 58-64. 522 Trapnell C, Roberts A, Goff L, Pertea G, Kim D, Kelley DR, Pimentel H, Salzberg SL, 523 Rinn JL, Pachter L. 2012. Differential gene and transcript expression analysis of 524 RNA-seq experiments with TopHat and Cufflinks. *Nat. Protoc.* **7**: 562-578.

525 Ueda HR, Matsumoto A, Kawamura M, Iino M, Tanimura T, Hashimoto S. 2002. 526 Genome-wide Transcriptional Orchestration of Circadian Rhythms in Drosophila. 527 J. Biol. Chem. 277: 14048-14052. 528 Velarde RA, Sauer CD, O. Walden KK, Fahrbach SE, Robertson HM. 2005. 529 Pteropsin: A vertebrate-like non-visual opsin expressed in the honey bee brain. 530 Insect Biochem. Mol. Biol. 35: 1367-1377. 531 Werren JH, Richards S, Desjardins CA, Niehuis O, Gadau J, Colbourne JK, Nasonia 532 Genome Working Group, Werren JH, Richards S, Desjardins CA, et al. 2010. 533 Functional and evolutionary insights from the genomes of three parasitoid 534 Nasonia species. Science **327**: 343-348. 535 Woelfle MA and Johnson CH. 2006. No promoter left behind: global circadian 536 gene expression in cyanobacteria. *J. Biol. Rhythms* **21**: 419-431. 537 Yan S, Zhu J, Zhu W, Zhang X, Li Z, Liu X, Zhang Q. 2014. The expression of three 538 opsin genes from the compound eye of Helicoverpa armigera (Lepidoptera: 539 Noctuidae) is regulated by a circadian clock, light conditions and nutritional 540 status. *PLoS One* **9**: e111683. 541 Yu W and Hardin PE. 2006. Circadian oscillators of Drosophila and mammals. J. 542 Cell. Sci. 119: 4793-4795. 543 Yuan Q, Metterville D, Briscoe AD, Reppert SM. 2007. Insect Cryptochromes: 544 Gene Duplication and Loss Define Diverse Ways to Construct Insect Circadian 545 Clocks, Mol. Biol. Evol. msm011.

CIOCKS. Mol. Blol. Evol. IIISIII011.

Zhang R, Lahens NF, Ballance HI, Hughes ME, Hogenesch JB. 2014. A circadian gene expression atlas in mammals: implications for biology and medicine. *Proc. Natl. Acad. Sci. U. S. A.* 111: 16219-16224.
Zhu H, Sauman I, Yuan Q, Casselman A, Emery-Le M, Emery P, Reppert SM. 2008.
Cryptochromes define a novel circadian clock mechanism in monarch butterflies that may underlie sun compass navigation. *PLoS Biol.* 6: e4.

Figure legend

Figure 1. Free-run behavioural rhythm in *Nasonia*. Representative actograms of individual *Nasonia* males in DD (left) and LL (right). Activity counts were sorted into 30 minute bins and plotted in blue. Yellow and grey backgrounds indicate lights on and lights off respectively. Gray and black bars below the actogram indicate the 12 hr subjective day and night.

Figure 2. Circadian transcriptional rhythms. (A) Heatmap of median-normalised expression of rhythmic (q < 0.1) transcripts in both constant darkness and constant light. (B) Histograms and heatmap of phases of rhythmic transcripts (q < 0.1 in both conditions), showing bimodal phase distribution and overlap between the two conditions.

Figure 3. Enrichment of GO terms among cycling transcripts. (**A**) Bar plot of 10 top overrepresented GO terms (by gene proportion) for both DD and LL rhythmic genes. (**B**) Euler diagram showing the overlap of overrepresented terms in DD (blue) and LL (red).

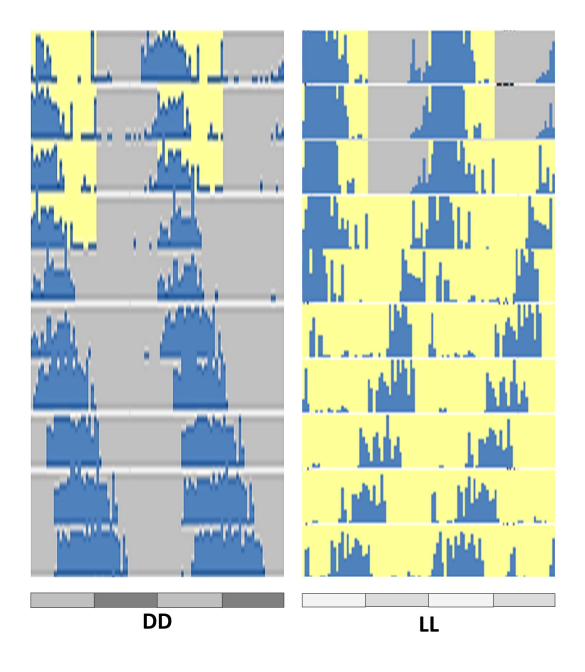
Figure 4. Normalised expression of clusters with significant (q < 0.01) overrepresentations of rhythmic genes. Each transcript profile in each cluster is coloured by that gene's membership of the cluster.

Figure 5. Comparison of the DD and LL transcriptomes. (A) FPKM (log2) expression of transcripts in DD (x axis) and LL (y axis), showing genes classified

(> 1.5 median fold change) as differentially expressed up in DD (blue) and up in LL (red). (B) Selected overrepresented (q < 0.01) GO terms for genes more highly expressed in DD. (C) Heatmap showing median-normalised expression for differentially expressed transcripts, in DD (left) and in LL (right), sorted by fold change.

cnange.

Figure 1



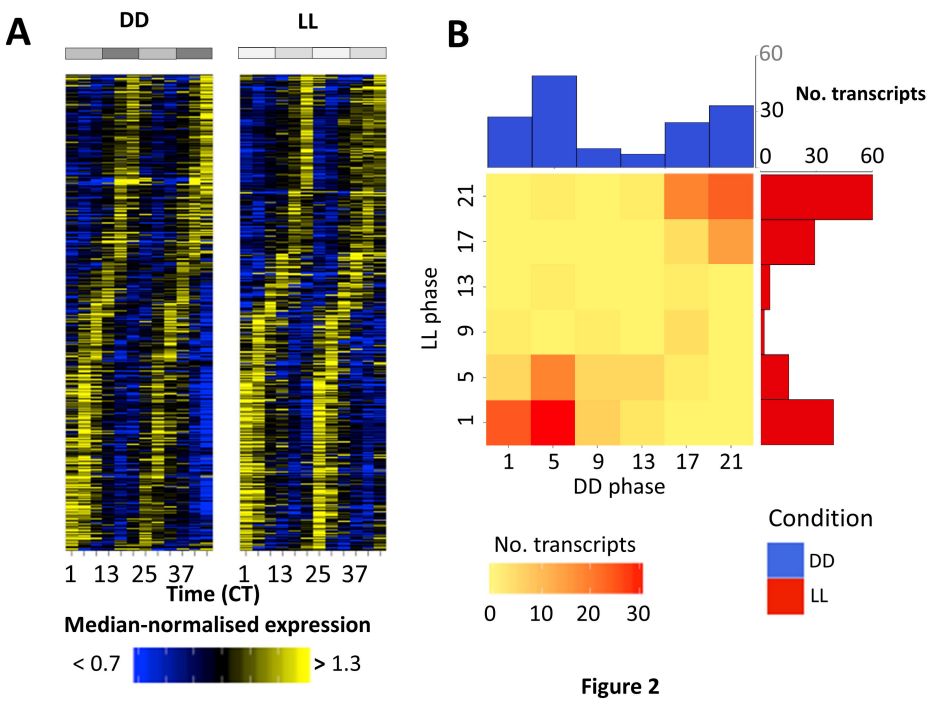


Figure 3

