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1	The evolutionary ecology of circadian rhythms in infection
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3	Mary L Westwood <sup>1*</sup> , Aidan J O'Donnell <sup>1</sup> , Charissa de Bekker <sup>2</sup> , Curtis M Lively <sup>3</sup> , Marlene Zuk <sup>4</sup> ,
4	Sarah E Reece <sup>1</sup>
5	
6	<sup>1</sup> Institute of Evolutionary Biology and Institute of Immunology and Infection Research, School of
7	Biological Sciences, University of Edinburgh, Charlotte Auerbach Road, Edinburgh EH9 3FL,
8	United Kingdom.
9	<sup>2</sup> Department of Biology, University of Central Florida, 4111 Libra Drive, Orlando, Florida 32816,
10	United States of America.
11	<sup>3</sup> Department of Biology, Indiana University, Bloomington, 1001 East Third Street, Indiana 47405,
12	United States of America.
13	<sup>4</sup> Department of Ecology, Evolution and Behavior, University of Minnesota, 1479 Gortner Ave, St.
14	Paul, Minnesota 55108, United States of America.
15	
16	* Corresponding author: Mary Westwood (mary.westwood@ed.ac.uk)
17	
18	Biological rhythms coordinate organisms' activities with daily rhythms in the environment. For
19	parasites, this includes rhythms in both the external abiotic environment and the within-host biotic
20	environment. Hosts exhibit rhythms in behaviours and physiologies, including immune responses,
21	and parasites exhibit rhythms in traits underpinning virulence and transmission. Yet, the
22	evolutionary and ecological drivers of rhythms in traits underpinning host defence and parasite
23	offence are largely unknown. Here, we explore how hosts use rhythms to defend against infection,
24	why parasites have rhythms, and whether parasites can manipulate host clocks to their own ends.
25	Harnessing host rhythms or disrupting parasite rhythms could be exploited for clinical benefit; we
26	propose an interdisciplinary effort to drive this emerging field forward.
27	
28	Circadian rhythms have long been taken for granted by science. Indeed, the first observation of a
29	clock-controlled behaviour (leaf opening and closing in Mimosa pudica) was not recorded until the
30	18th century <sup>1</sup> . Following the fundamental observation that organisms can adaptively anticipate daily
31	rhythms in their environment, the field of "chronobiology" took off in the mid-20th century with a
32	focus on evolutionary and ecological questions. However, the advent of genetic tools a few decades
33	later shifted the remit to determining the molecular and genetic workings of circadian clocks. Yet,

## 34 despite their assumed major impact on fitness, circadian rhythms remain overlooked in evolutionary

35 ecology<sup>2–4</sup>. Here, we propose that the integration of chronobiology and evolutionary ecology return

to its roots to tackle a topic of growing and applied interest; the role of rhythms in host-parasite
interactions. Note that we use the term "parasite" to collectively refer to all agents of infection (e.g.
single-celled and multicellular eukaryotes, bacteria, viruses).

39

One of the most fundamental ecological interactions is that between hosts and parasites. Research from diverse taxa (plants, mammals, and insects) reveals that host clocks drive daily rhythms in immune defences, disease severity and spread<sup>5,6</sup>. Parasites display daily rhythms in traits underpinning within-host survival and between-host transmission<sup>7,8</sup>. Rhythms in parasite activities and in host responses to infection could provide an advantage to parasites, hosts, both, or neither. To what extent parasites and hosts are in control of their own and/or each other's rhythms is also poorly understood.

47

48 Understanding the evolution (and possibly, coevolution) of rhythms may enable vaccines and drugs 49 to take advantage of rhythmic vulnerabilities in parasites or harness host rhythms to improve efficacy and reduce drug toxicity. For such interventions to be robust to parasite evolution, 50 51 understanding how host-parasite interactions shape rhythms in hosts and parasites is necessary<sup>7</sup>. 52 Key questions include how rhythms in diverse host traits contribute to defence, how parasites cope 53 with exposure to their host's rhythms, and whether hosts and parasites can manipulate each other's 54 rhythms for their own benefit. We discuss these three scenarios, identify systems to explore them, 55 and offer ways in which this knowledge can be exploited to improve health. An evolutionary 56 ecologist's introduction to chronobiology is provided in Boxes 1 and 2.

57

#### 58 **Rhythms in host defence**

59 The most patent defence against infection is the immune response, and a wealth of evidence reveals that circadian clocks play a role in orchestrating immune defences<sup>5</sup>. Circadian clock genes are 60 61 expressed in many types of immune cell, and the immune and circadian systems are connected in multiple ways<sup>9,10</sup>. For instance, the clock gene *Bmal1* mediates the balance between pro- and anti-62 63 inflammatory responses<sup>11</sup>. Rhythmic production of the pro-inflammatory cytokines TNF- $\alpha$  and IL-6 by macrophages is clock controlled<sup>12</sup>, and mobilization of inflammatory monocytes is also 64 65 regulated by the clock<sup>10</sup>. This phenomenon, termed "anticipatory inflammation", appears uncoupled to metabolic rhythms and may defend against incoming parasites<sup>13</sup>. Similarly, in humans, 66 67 proinflammatory cytokines peak in circulation during the day (active phase)<sup>14</sup>, whereas hematopoietic stem and progenitor cells, and most mature leukocytes, peak at night<sup>14,15</sup>. In 68 69 nocturnal mammals, an inverse rhythm is often observed, with innate defences peaking at night 70 (active phase) and repair mechanisms peaking during the day (resting phase)<sup>9</sup>.

71

72 Observations of immune rhythms have given rise to the notion that organisms invest in defence 73 during the active phase when parasite encounter is assumed most likely, and repair during the 74 resting phase<sup>16</sup>. Temporal segregation of immune responses may thus solve problems caused by 75 having immune defences continually tuned to maximal (e.g. collateral damage via 76 immunopathology<sup>17</sup>). Also, energetic demands imposed by activity and metabolism may trade-off 77 against immune defence<sup>18</sup>. Intuitively, "defence only during the active phase" suggests the host is 78 achieving the most "bang for the buck" by ensuring activities that are energetically costly, or likely 79 to cause collateral damage, are only performed when most useful. However, this intuition may be 80 naïve. First, it ignores the potential for constraints imposed by the need to temporally couple (or de-81 couple) certain immune rhythms with other internal rhythms<sup>7</sup>. This includes separating the timing of 82 metabolism from defensive actions within immune cells themselves<sup>5,16</sup>. Second, it assumes that a 83 parasite encounter is rhythmic and predictably occurs in the active phase. This is clearly the case for 84 food-borne parasites, but ingestion is not the only route into a host. Rather, the immune system 85 functions within a broad set of energetic demands in which parasite defence is just one of many 86 requirements. For example, rhythmic stomatal opening for gas exchange during the day is a wellused route into plants by bacterial pathogens<sup>19</sup>. Consequently, *Arabidopsis* is better able to detect 87 and defend against parasites in the morning than evening<sup>20,21</sup>. Given the wealth and diversity of data 88 89 (illustrated in Table 1), meta-analyses are needed to test whether the timing (phase) of rhythms in 90 immune effectors relates to nocturnal vs diurnal lifestyles and whether they function in front-line or 91 secondary defences, or healing.

92 Infection in the active vs resting phase for diverse hosts (flies, plants, mammals) dramatically 93 affects disease severity and mortality rates (Table 1), suggesting that the phase of immune rhythms 94 upon infection matters. Most studies performed in plants (Table 1) point towards infection during 95 the active phase resulting in greater resistance to infection and less damage to the plant. But the 96 degree to which immune rhythms result in time-of-day differences in parasite control can be 97 counter-intuitive. For example, mice mount higher clock-controlled proinflammatory responses 98 against Salmonella enterica Typhimurium when challenged in their rest phase, but bacterial load is also higher and hosts have worse symptoms<sup>22</sup>. Furthermore, Leishmania parasites infect host 99 100 neutrophils and macrophages, and the clock-controlled secretion of chemoattractants by these 101 immune cells facilitates their infection, making parasite invasion more successful at night when immune activity is highest<sup>23</sup>. Thus, whether immune rhythms are sufficient to entirely explain 102 103 divergent outcomes of time-of-day of infection is unclear (Table 1). Studies that separate the effects 104 of immune rhythms on preventing infection from their role in dealing with ongoing infection will

reveal the extent to which immune rhythms are beneficial and when they should be overruled to deal with a major threat. Additionally, most time-of-day immune challenges have used either bacteria or chemicals, raising the question of whether a more diverse array of challenges are needed to establish general patterns.

109

110 That host circadian clocks impact on infection via traits other than immune responses has been largely overlooked. Rhythmicity in host activity may determine when hosts provide the best 111 resources to their parasites and offer the most opportunities for onwards transmission<sup>24–26</sup>. For 112 113 example, a recent study of the intestinal helminth Trichuris muris demonstrates the role of host 114 rhythms in foraging. Mice infected in the morning (resting phase) expel worms sooner and have a 115 stronger T-helper 2 response than dusk-infected (active phase) mice, and this effect is reversed when mice are fed only in the day, in an immune-independent manner<sup>27</sup>. Host feeding rhythms are 116 117 relevant to gut microbiota, and a two-way feedback between host and microbe rhythms has been proposed<sup>28</sup>. Daily rhythms in host reproductive behaviours may make hosts vulnerable to infection. 118 119 For example, the crepuscular and nocturnal singing activity of the cricket *Teleogryllus oceanicus* allows the acoustically-orienting parasitoid fly Ormia ochracea to locate hosts, but the flies are best 120 able to hunt when darkness is incomplete<sup>29</sup>. A rhythmically expressed reproductive behaviour 121 (singing) got the host into this mess, and it appears that natural selection has found two solutions 122 123 (see Box 3).

124

In addition to immune responses, infected hosts often exhibit adaptive sickness behaviours 125 consisting of endocrine, autonomic, and behavioural changes that perturb circadian rhythms<sup>30,31</sup>. For 126 example, wild red colobus monkeys (Procolobus rufomitratus tephrosceles) decrease energetically 127 costly activities, and rest frequently, while shedding whipworm eggs<sup>32</sup>. Fever, another common 128 sickness behaviour, is sufficiently advantageous to offset the 10-12.5% increase in metabolic rate 129 required for each 1°C increase in temperature<sup>33</sup> and has been conserved throughout more than 600 130 million vears of vertebrate evolution<sup>34</sup>. Fever enhances an organisms chance of survival by creating 131 a hostile environment for parasites and a more active immune response<sup>34–37</sup>. Under normal 132 133 circumstances, the so-called central (SCN) clock controls body temperature rhythms, but how the 134 SCN and inflammation interact to control temperature is unknown. Though many behaviours 135 altered during infection are clock-controlled during health, the extent to which organisms become 136 too sick to maintain normal behaviour or adaptively disrupt their rhythms is unclear. Additionally, 137 clock-control could facilitate recovery of rhythms during the return to health.

- 139 Viewing the host as a collection of traits connected by the circadian system has the potential to
- 140 uncover novel strategies to resist infection and reveal the circumstance in which immune rhythms
- 141 reflect constraints or adaptations. Indeed, rhythmic metabolism of xenobiotic substances (e.g. drugs
- 142 and vaccines) influences efficacy and toxicity in a time-of-day dependent manner<sup>38</sup>. For example,
- 143 halothane (a commonly used anaesthetic) administered to mice in the daytime results in low
- 144 mortality (5%), but mortality increases (76%) if administered at night<sup>39</sup> and half of the best-selling
- 145 drugs in the USA for humans target the products of genes that are rhythmically expressed (in
- <sup>146</sup> mice)<sup>40</sup>. A better understanding of host rhythms could be harnessed to make drugs and vaccines
- 147 more effective, as well as mitigating the negative effects of modern lifestyles that involve shift work
- 148 and jet lag. However, for such interventions to be sustainable in the face of parasite evolution,
- 149 understanding the ecology of rhythms from the perspective of parasites is also required.
- 150

#### 151 **Rhythms in parasite offence**

152 Scheduling activities to take advantage of daily rhythms in transmission opportunities could be a 153 general explanation for rhythms in parasites. The most well-known example concerns the transmission forms (microfilariae) of different species of filarial worms. They move from the host's 154 155 organs to the capillaries during the day or night, depending on whether they are transmitted by dayor night-biting insect vectors<sup>41</sup>. In addition to the activity patterns of vectors, rhythmic interactions 156 with hosts also matter. For example, the larval stage of the blood fluke Schistosoma japonicum 157 158 emerge from their invertebrate host to seek a mammalian host at different times of day. Flukes 159 emerge in the afternoon when the preferred host is nocturnal or in the morning if seeking a diurnal host<sup>42</sup>. Parasites that have free-living stages are also subject to rhythms in the abiotic environments. 160 The coccidian parasite *Isospora* sheds from its host in the late afternoon to minimise UV exposure 161 162 and desiccation risk whilst undergoing a developmental transition necessary to infect new hosts<sup>43</sup>. 163 However, key questions remain about the adaptive nature of these rhythms. For example, why 164 aren't microfilariae located in the peripheral capillaries all day long? Is a cost associated with this 165 location, which is only worth paying at times of day when vectors are active?

166

In contrast to the role of parasite rhythms in transmission, their role in within-host survival has received less attention. Many host rhythms (in addition to immune rhythms) present opportunities and constraints for parasites. *Trypanosoma brucei* (which cause sleeping sickness) display circadian clock-driven rhythms in the expression of metabolic genes<sup>8</sup>. These rhythms correlate with time-ofday sensitivity to oxidative damage, thereby suggesting the need to cope with redox challenges caused by rhythmic digestion of food by hosts. In contrast, rhythms in the development of asexually replicating malaria parasites capitalise on daily variation in the nutritional content of blood caused

- by host immune responses and feeding patterns<sup>44,45</sup>. Whether malaria parasites cannot complete
- their developmental cycle until the host makes nutrients available, and/or use nutrients rhythms as a
- 176 time-of-day cue to set the pace of their development, is unknown<sup>46</sup> (see Box 3).
- 177

178 Clocks in parasites or hosts could have fitness consequences for one or both parties, or neither. 179 Fitness consequences for both hosts and parasites suggests that clocks could coevolve. Clock coevolution is suspected for the plant-pollinator system Petunia axillaris and Manduca sexta<sup>47</sup>, in 180 181 which nocturnal scent emission by *P. axillaris* coincides with foraging activity in the hawkmoth *M.* 182 sexta. Both traits are clock-controlled, and appear so well synchronized that, even in the absence of floral scent emission, *M. sexta* exhibits a burst in foraging activity at the same time that floral scent 183 184 emission is expected to be greatest. However, foraging behaviour also remains sensitive to the 185 environment, as evidenced by absence of activity when the moth is subjected to light at night. If rhythms in different organisms do coevolve, then they should use the same Zeitgeber, but how 186 187 robust should their timing systems be to fluctuations in the environment? If the rhythm of one party is more readily disrupted (masked) by environmental change, or faster at tracking seasonal changes 188 189 in photoperiod, then the relationship may be disrupted to the gain of hosts or parasites. Exploring 190 the degree and consequences of plasticity in rhythms is pertinent because climate change is interfering with the ability of interacting species to synchronise<sup>48</sup>. 191

192

193 The situation is further complicated when interactions between both host and parasite clocks shape 194 disease trajectories. For example, in a plant-fungus system (Arabidopsis thaliana and Botrytis 195 *cinerea*, respectively), when both parties are in the same photoperiod schedule, primary plant 196 defences peak in the morning, and the fungus produces the biggest lesions when inoculated at 197 dusk<sup>49</sup>. The authors were able to separate the contributions to pathogenicity by host and parasite 198 clocks using reverse lighting schedules for fungus and plants: fungus at dusk produced more severe 199 infections than fungus at dawn, regardless of time-of-day for recipient plants<sup>49</sup>. Furthermore, this 200 suggests B. cinerea anticipates and exploits weaknesses in plant defence at dusk rather than 201 attempting to overwhelm dawn defences (see section "Rhythms in host defence"). Separately 202 assigning the contributions of rhythms in hosts/vectors and parasites to virulence and transmission 203 is necessary to understand whose genes control which rhythms, and hence how they can be shaped 204 by selection.

205

If parasite rhythms are adaptive, then disrupting them could reduce disease severity as well as transmission. However, understanding the timing mechanisms of parasite rhythms is necessary to disrupt them<sup>7</sup>. Unravelling how parasite rhythms are controlled is a considerable challenge. 209 Parasites might allow the host to inadvertently schedule their activities for them, in which case the 210 genes encoding parasite timing mechanisms belong to hosts. Alternatively, parasites might keep 211 time using a circadian clock (with the properties described in Box 1), as demonstrated for *T. brucei* 212 and *B. cinerea*. Given the diversity in clock genes across taxa, searching genomes for known clock 213 genes often yields "absence of evidence" not "evidence of absence." Instead, round-the-clock 214 transcriptomics or proteomics, paired with bioinformatics approaches to mine for known core 215 clock-related functional domains and sequence patterns may find candidates. However, simpler time-keeping strategies exist, though they do not necessarily have the advantages of temperature 216 217 compensation or anticipation. For example, cell division cycles are often controlled by hourglass mechanisms that rely upon threshold concentrations of substances, independently of periodic 218 219 phenomena<sup>50</sup>. Alternatively, organisms can react directly (via "tracking") to temporal changes in the environment. Note, this differs from masking, a chronobiological phenomenon in which the 220 221 expression of a clock-controlled rhythm is suppressed by a change in the environment without 222 having a direct effect on the period or phase of the underlying rhythm<sup>51</sup>. A response that directly tracks time-of-day cues may suit parasites with multi-host lifecycles if each host type provides a 223 224 different time-cue.

225

226 Given that rhythms in *T. brucei* metabolism and plasticity in development during the asexual cycle 227 of *Plasmodium spp.* enables these parasites to tolerate drugs, there is an urgent need for proximate 228 and ultimate explanations of their rhythms. The T. brucei clock is entrained by temperature cycles, 229 but if other parasites use Zeitgebers to set their clocks, or respond directly to time-of-day cues, that 230 are readily perturbed, it should be possible to reduce parasite fitness by interfering with their rhythms. Further, reports of changes to the biting time of mosquito populations that transmit 231 232 malaria suggests that insecticide-treated bed nets are imposing selection on vector rhythms<sup>8,52,53</sup>. Given that rhythms of parasites and mosquitoes each affect malaria transmission in lab 233 234 experiments<sup>54,55</sup>, what are the likely epidemiological consequences? Recent work suggests that mosquitoes are more susceptible to infection when they feed in the daytime and parasites are more 235 236 infectious at night<sup>54</sup>. Thus, day-biting could increase the prevalence, but not burden, of malaria in 237 mosquitoes. However, in the longer term, if parasites evolve to invert their rhythm but mosquitoes 238 do not, both prevalence and burden may increase.

239

#### 240 Parasite manipulation of host rhythms

Rhythms in host processes offer opportunities that parasites could exploit. Could parasite fitness beincreased by coercing hosts into altering their rhythms? Although many striking examples of

243 parasite manipulation of host phenotypes (i.e. changes to host traits that benefit parasites) are known<sup>56</sup>, the notion of "parasite manipulation of host clocks" is largely unexplored<sup>57</sup>. A pre-244 245 requisite for parasite manipulation is that a phenotypically plastic host trait is targeted; and 246 circadian clocks are flexible. Because clocks control much of the host's behaviour and physiology<sup>58</sup> 247 and clocks throughout a given host involve the same players in the canonical clock (the TTFL), 248 manipulation of the host's time-keeping may be an efficient way to simultaneously alter many aspects of the within-host environment. Alternatively, parasites interests may be served by 249 250 bolstering circadian rhythms of their hosts during sickness to ensure they forage and interact with 251 conspecifics, as usual.

252

253 As outlined in the section "Rhythms in host defence," separating the effects of being sick per se 254 from host defence and parasite manipulation is challenging. Recently, a combination of culture and 255 comparison of infection models has revealed that T. brucei alters expression rhythms of clock genes 256 in host mice<sup>59</sup>. Specifically, infected hosts are more active in the resting phase (phase-advanced) because the clock runs faster (shorter period). Effects at organismal, cellular, and molecular levels 257 suggests the behaviour is not just a result of sickness<sup>59</sup>. However, it is not clear how *T. brucei* 258 259 achieves this, and whether the parasite benefits from altering host rhythms. One target of circadian 260 disruption by viral parasites is the gene *Bmal1*, a core clock gene. Herpes and influenza A virus replication and dissemination within the host is enhanced in infections where *Bmal1* is knocked 261 out<sup>60</sup>. However, it remains unclear if virus replication is maximised by simply disturbing 262 rhythmicity in host cell cycles or if this is a case of immune manipulation since *Bmal1* appears 263 involved in innate host defence<sup>60</sup>. Having observed changes to host clocks, the proceeding step is to 264 decipher the ecological context behind these effects. 265

266

267 The above examples lend proof-of-principle to the idea that parasites can manipulate host clocks 268 and could be a general explanation for examples of host manipulation. Hairworms (Nematomorpha) 269 are a well-known case of temporally linked behavioural manipulation. They infect various 270 arthropods, notably crickets, and cause the host to wander in an erratic manner until a body of water 271 is encountered. The host commits suicide by jumping in water, and the adult hairworm emerges. 272 Infected hosts are found wandering only in the early part of the night<sup>61</sup>, and uninfected hosts are rarely motivated to jump into water. Infected crickets differentially express an array of proteins, 273 274 some of which are linked to visual processes and circadian clocks<sup>62</sup>. Culturing isolated host cells 275 with parasite products and quantifying the expression of clock genes (following Rijo-Ferreira 2018) 276 could illuminate this case of parasite manipulation. For systems without relevant insect cells lines, 277 or cases where manipulation is likely to be tissue/cell type specific, a transcriptomics approach may

be useful<sup>63</sup>. Round the clock expression data can be mined for putative core clock genes and their
phase, amplitude and period assessed in control and manipulated hosts. This however, is likely to be
extremely challenging for host species whose timekeeping does not rely on a canonical circadian
clock.

282

283 Another putative case for clock manipulation concerns the New Zealand freshwater snail (Potamopyrgus antipodarum) infected with Microphallus trematodes<sup>64</sup> (Trematoda: 284 Microphallidae). Uninfected adult snails forage primarily at night on the upper surfaces of rocks in 285 286 the shallow-water margins of lakes. These snails retreat to under rocks at sunrise, which likely 287 reduces their risk of predation by waterfowl, which are the definitive host for Microphallus. 288 Infected snails, however, show delayed retreating, potentially making them more likely to be 289 consumed<sup>25</sup>. Crucially, the apparent manipulation only occurs when the parasite is mature. Snails 290 infected with immature (non-transmissible) stages exhibit the same risk-averse retreating behaviour 291 as uninfected snails<sup>25</sup>. In addition, snails infected with other species of sterilizing trematodes, which 292 are not trophically transmitted, do not exhibit the same risky behaviour as those infected with *Microphallus*<sup>65</sup>, thereby eliminating the possibility that the *Microphallus*-induced behavioural 293 294 change is a simple artefact of parasitic castration. Finally, Microphallus-infected snails spend more 295 time foraging on the top of rocks, even when food was removed whereas uninfected snails retreated to shelter<sup>65</sup>. Taken together, the data suggest that *Microphallus* induce a change in snail behaviour 296 297 that increases trophic transmission, potentially via manipulation of clock-controlled activity 298 rhythms.

299

300 There are many ways that parasites could interfere with clock-controlled host behaviours. A blunt 301 instrument would be to alter perception/detection of the Zeitgeber that sets the time of the host's 302 clock, which is usually light. For example, Microphallus could interfere with photoreception to 303 reduce the sensitivity of snails to dawn, causing their clocks to phase delay and forage at higher 304 light intensities than un-manipulated snails. Alternatively, parasites could induce the host to ignore 305 its clock (mask) or alter clock regulation of hormones that relay time-of-day information around the host. For example, baculoviruses appear to perturb the circadian rhythms of their caterpillar hosts 306 307 by disrupting hormones that control climbing behaviour. In the baculovirus (Lymantria *dispar* nucleopolyhedrovirus), a single gene inactivates 20-hydroxyecdysone<sup>66</sup> (a host hormone 308 309 regulated by a circadian oscillator), motivating the caterpillar to climb high atop their host plants. 310 Here, they liquefy and disseminate the virus to caterpillars below, as well as infecting birds who consume the corpses<sup>67</sup>. Similar to the manipulation of caterpillar hosts, many species of parasitic 311

fungi (*Ophiocordyceps spp.* and *Pandora spp.*) alter the daily behavioural rhythm of a variety of ant
species<sup>68,69</sup> (See Box 3).

314

Parsing out whether temporal disruption is a host response or clock manipulation is nearly, if not entirely, impossible without uncovering the mechanism of manipulation. The lack of insight into the mechanisms parasites use to interfere with their hosts has stalled progress in the field of "host manipulation by parasites"<sup>70</sup>. This gap could be filled by harnessing the tools and conceptual framework developed in chronobiology. Many of the examples above have employed an ecological approach, yet a chronobiological approach can help elucidate both proximate and ultimate explanations.

322

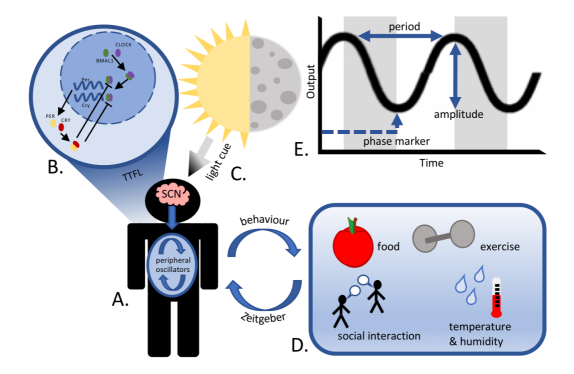
### 323 Conclusion

324 Over the past few decades, the focus of chronobiology has been to elucidate the mechanistic 325 underpinnings of biological rhythms. We propose that now is the time to integrate this knowledge into parasitology, evolutionary ecology, and immunology (see Box 2). Indeed, the role of biological 326 rhythms in infectious disease is a growing topic that holds promise for improving human and 327 328 animal health. History clearly illustrates that attempts to control parasites are usually met with counter-evolution (in the form of drug resistance, vaccine escape, and host shifts). A comprehensive 329 understanding of how rhythms affect parasite invasion and exploitation of a host (or vector) offers 330 331 novel ways to disrupt the chain of transmission and treat disease. Further, clock coevolution may 332 occur in host-parasite-vector interactions, resulting in complex arms races best understood through 333 the lens of chronobiology coupled with evolutionary ecology. Chronobiology supplies a myriad of tools to help elucidate rhythmic phenotypes and reveal to what extent host and parasite genes are 334 335 responsible for rhythms in disease phenotypes. Adding an evolutionary ecology framework will 336 ensure this information is generalisable and used to make interventions as evolution-proof as 337 possible.

- 338 Box 1. What are circadian rhythms? Biological rhythms are deemed to be controlled by circadian 339 clocks if they meet several criteria<sup>71</sup>. First, their duration (period) must be approximately 24 hours. Second, they must persist (free-run) in conditions without time-of-day cues, which is usually 340 341 assessed by observation in constant light or dark. Third, the phase of the oscillator or outputs are set 342 (entrained) by a time-of-day cue (Zeitgeber) which is usually light. Fourth, unlike the rate of many 343 chemical reactions, the speed of a circadian clock varies little over a biologically realistic range of 344 environmental temperatures (temperature compensation). Together, these criteria allow organisms 345 to fulfil a key feature of circadian rhythms: anticipatory, rather than reactionary, behaviour. For instance, plants ready photosynthetic machinery in anticipation of sunlight<sup>72,73</sup> and animals exhibit 346 food-anticipatory activity (e.g. increases in core temperature, activity, serum corticosterone, and 347 duodenal disaccharides) prior to foraging<sup>74</sup>. The workings of circadian clocks are sufficiently 348 349 flexible to allow organisms to cope with gradual changes in photoperiod across seasons, but not 350 flexible enough to instantly cope with changes in time zones (which is why travellers experience jet
- 351

lag).

- 352
- 353 The mammalian circadian system is composed of the "central" clock in the brain (suprachiasmatic
- nucleus; SCN) and "peripheral clocks" in other organs and tissues (A). Clocks in nucleated cells are
- run by transcription-translation feedback loops (TTFL). For example, in animals the proteins
- 356 CLOCK and BMAL1 act as activators and members of the PER and CRY families are repressors<sup>75</sup>
- 357 (B). Retinal photoreceptors receive light cues which are carried through the hypothalamic optic tract
- and transmitted to the SCN, resulting in its synchronization/entrainment (C). Clocks in organs and
- 359 tissues (peripheral clocks) can be entrained by feeding rhythms, and in
- taxa other than mammals, exercise, social cues, and abiotic rhythms in temperature and humidity
  may entrain clocks (D). Rhythms are often characterised by their period, amplitude, and markers for
  phase (E; grey bars illustrate night time for a rhythmic trait measured over 48 hours). They are
  described in relation to the time since the Zeitgeber (ZT) occurred (e.g. ZT6 refers to 6 hours after
  dawn) which usually differs from the actual time-of-day (Circadian Time; CT).
- 366 \*we suggest that the image [Box\_1] be placed here.
- 367
- 368
- 369
- 370
- 371
- 372





#### Box 1 image.

Box 2. Why have circadian rhythms evolved? Circadian clocks appear so advantageous that nearly all eukaryotes have a circadian system in most cells<sup>76</sup>. Circadian clocks may confer two kinds of fitness benefit: coordinating behaviours with rhythms in the external environment (extrinsic adaptive value), and temporally compartmentalising incompatible processes (intrinsic adaptive value)<sup>2</sup>. For instance, intrinsic benefits are conferred when cell division in yeast is temporally constrained to the reductive phase of metabolism, minimising rates of genetic mutation<sup>77</sup>. However, most studies of the fitness consequences of circadian rhythms have focussed on the benefits of synchronizing activities with rhythms in the abiotic environment: matching the period of day-night rhythms enables cyanobacteria to outcompete strains whose clocks run faster or slower<sup>78</sup> and enhances the survival of *Arabidopsis*<sup>73</sup>. Rhythms in the biotic environment<sup>2</sup> matter too. For example, the sea urchin Centrostephanus coronatus avoids predatory sheephead wrasse (Pimelometopon pulchrum) by foraging at night and retreating to shelter prior to the onset of wrasse activity<sup>79</sup>. 

Despite the diversity of extrinsic rhythms that could select for the scheduling of diverse processes, there are surprisingly few demonstrations that circadian clocks actually affect fitness. For example, fitness is greater in wild-type mice than mutant mice with shortened periods<sup>80</sup>, flies with clock mutations die more rapidly than wild types after infection with bacteria<sup>81,82</sup>, and circadian knockout plants flower later and are less viable than wild-type plants<sup>3</sup>. However, depending on ecological context, rigidly scheduling activities according to day and night is not always the best strategy. For example, nocturnal mice boost energy efficiency by switching to diurnality when challenged with cold and hunger<sup>83</sup>. Nursing honeybees, that remain in the hive are arrhythmic, because round-the-clock care is necessary for larvae; and, if needed, diurnal foraging bees can revert to arrhythmic nursing behaviour<sup>84</sup>. Shorebirds also display considerable plasticity in activity rhythms during breeding, likely explained by predator avoidance strategies<sup>85</sup>. 

The above examples illustrate the gains to be made from integrating chronobiology with evolutionary ecology in general<sup>4</sup>. We propose that such an approach offers a novel advance to the study of host-parasite interactions and coevolution. Coupling the well-developed conceptual frameworks for unravelling how circadian oscillators operate, and probing the costs and benefits of phenotypically plastic traits that are relevant to infection, will explain why rhythms in immune defences and parasite traits occur.

455 Box 3. Case studies illustrating the role of circadian rhythms in parasite offence, host defence,

## 456 and host manipulation

- Host-parasite system: *Teleogryllus oceanicus* (Pacific field cricket) & *Ormia ochracea* (parasitoid
   fly)
- 459 What we know: O. ochracea deposit larvae which burrow into the host and emerge 7-10 days later,
- 460 resulting in host death. A flatwing morph that is physically incapable of calling has evolved to 461 evade the risk of parasitism by acting as a silent, satellite male<sup>24</sup>.
- 462 A more nuanced form of parasite evasion? In addition to the flatwing morph, natural selection
- 463 may have found another solution. Some males condense singing activity to the darkest part of the
- 464 night<sup>29</sup> which may hamper the fly's ability to use visual cues to home in on hosts. Parasite evasion
- 465 (via a flatwing phenotype or phase-shifted calling) trades off against attracting females, potentially 466 constraining selection on these strategies. Moreover, multiple activities need to be coordinated for
- 466 constraining selection on these strategies. Moreover, multiple activities need to be coordinated for 467 successful reproduction (e.g. locomotion, foraging, spermatophore production). Given that many of
- 468 these traits are clock-controlled, could altering the timing outputs of the clock be a streamlined way
- 469 of phase-shifting all related activities and minimizing the costs of parasite evasion? [associated
- 470 image = cricket\_fly.png] Photo credit: Norman Lee
- 471



472 473

474 Host-parasite system: Carpenter ants & *Ophiocordyceps* spp. and *Pandora* spp. (fungi)

475 What we know: O. unilateralis s.l. induces workers of its carpenter ant host, ordinarily active

476 during the night-time, to wander out of the ant nest during the day-time. Hosts then summit

477 vegetation and adopt a mandibular death-grip in elevated positions. This manipulated behaviour is

478 highly time-of-day and species-specific and occurs within a 3-hour window at dawn or in the mid-

late morning, depending on the species<sup>68,86</sup>. Clinging to vegetation, the ant dies whilst the fungus
 completes its life cycle by growing a spore-producing stalk out of the dorsal region of the ant's

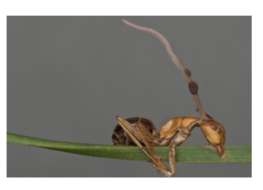
481 thorax<sup>86</sup>.

482 A case for coevolution and ecosystem specificity? The jigsaw puzzle of how the fungus controls
483 the ant is still being pieced together. Clocks may play a central role because infection alters the
484 expression of host clock homologues *period* and *cycle*<sup>68</sup>. Host manipulation also appears to involve

485 altering host chemosensory abilities, potentially via rhythmic secretion of enterotoxins<sup>87</sup>, all

486 achieved from the fungus's primary location in muscle tissues<sup>88</sup>. [associated image = ant\_fungi.png]

- 487 Photo credit: Miles Zhang
- 488



- 491 Host-parasite system: Mammals & Plasmodium spp. (malaria parasites)
- 492 What we know: Malaria parasites synchronously burst from the host's blood cells every 24, 48, or
- 493 72 hours depending on the parasite species<sup>89</sup>. When out of synch with the host's circadian rhythms,
- 494 parasites incur an approximately 50 percent reduction in the densities of both asexual stages
- 495 (necessary for in-host survival), and sexual stages (responsible for transmission)<sup>90</sup> before they 406 has many head to be in some head to be stage that the stage of the
- 496 become rescheduled to be in synch with host feeding rhythms<sup>44,45</sup>.
- **Three worlds collide: a complex system of interactions?** Why aligning the phase of parasite
- 498 rhythms with the host's rhythms is important remains mysterious, but recent work suggests that 499 parasites are also selected to coordinate with the time-of-day their mosquito vectors are active<sup>54,55</sup>
- 500 (see Rund et al. 2011 for information on *Anopheles* circadian rhythms). If differently phased
- 501 rhythms for asexual replication are required to provide the best matches to host and vector rhythms,
- 502 parasites face a trade-off between maximizing in-host survival and between-host transmission. Such

- a tension could be exploited by novel drug treatments to coerce parasites into a loss of fitness.
- 504 Further, mosquito nets have induced a shift in *Anopheles gambiae* biting activity, ultimately
- resulting in a change in host-parasite timing<sup>8,52,53</sup>. The epidemiological consequences of this are unknown. [associated image = mosquito malaria.png] Photo credit: Sinclair Stammers



535 Table 1. Impact of immune challenge during the rest and active phases of hosts. A selection of studies identified as time-of-day immune challenges from PubMed searches for ""time of day" plus "immune and 536 537 infection" and ""circadian rhythm" plus "immune and infection". Articles were included if the study involved a time-of-day immune challenge; those without a time-of-day immune challenge were not included in the table. 538 539 Time-of-day (ToD) is given as hours since lights on (ZT) for organisms in entrainment conditions, and as 540 subjective day/night for those in constant light or dark conditions (i.e. corresponding to the light or dark portion 541 of the cycle before experiencing constant conditions). Unless otherwise stated, entrainment conditions are 12 542 hour light:dark. Outcomes of challenge in the rest phase (daytime for nocturnal organisms, nighttime for 543 diurnal organisms) are compared to challenge in the active phase in terms of virulence metrics and immune 544 effectors measured.

Host spp.	Challenge	ToD	Outcome in rest versus active phase	Ref
	Salmonella typhmurium	ZT4/16	Greater inflammation and bacterial load when infected in the rest phase	22
	Leishmania major	Subjective day/night	Lower parasite burden and lower severity when infected in the rest phase	23
	Lipopolysaccharide (LPS) endotoxin	Subjective day/night	Lower concentrations of cytokines when infected in the rest phase	91
		ZT11/19	Higher mortality when challenged in the rest phase	92
Mus musculus – house mouse		Subjective day/night	Greater inflammatory responses and lower bacterial burden when challenged/infected in the rest phase	93
(nocturnal)	Streptococcus pneumoniae	ZT0/12		
	Murid Herpesvirus 4	ZT0/10	Greater viral replication when infected in the rest phase	60
	Helicobacter pylori	ZT1/7/13	Lower lymphocyte numbers when infected in the rest phase	94
	Vesicular stomatitis virus	ZT0/12	Higher mortality when infected in the rest phase	95
	Pseudomonas aeruginosa	ZT1/5/9/13 /17/21/1	Lowest mortality when infected in the rest phase (especially ZT21)	82
Drosophila		Subjective day/night	Lowest bacterial burden when infected in the rest phase	
<i>melanogaster</i> – fruit fly	Streptococcus pneumoniae	ZT7/19	Slowest rate of mortality when infected in the rest phase	81
(diurnal)	Escherichia coli	ZT0/6/12 /18	Infection at all ZT induces sleep the morning after infection and sleep was more prolonged after infection in the rest phase	96
Anopheles stephensi - Asian malaria mosquito (nocturnal)	Escherichia coli	Morning/ evening	Lower bacterial growth and lower mortality when infected in the rest phase	97
Arabidopsis	Pseudomonas syringae	ZT0/4/10 /16	Immune defences are highest when inoculation occurs early in the active phase Note photoperiod is 9 hours light:15 hours dark	98
thaliana –	Botrytis cinerea	Dawn/dusk	Larger lesions when inoculated in the rest phase	49
thale cress (diurnal)		ZT0/3/6/9/ 12/15/18 /21/24	Greater susceptibility when inoculated in the rest phase	21
	Pseudomonas syringae	Subjective day/night	Lower infiltration of bacteria when infected in the rest phase	99

			Subjective morning /evening	Greater suppression of bacterial growth at the start of the rest phase when spray-inoculated, and greater suppression of bacterial growth at the start of the active phase when syringe-infiltrated	20
		Hyaloperonospora arabidopsidis	Dawn/dusk	Highest percentage of leaves with sporangiophores when infected in the start of the rest phase	100
	Danio rerio zebrafish (diurnal)	Salmonella typhimurium	ZT4/16	Lower survival when infected in the rest phase	101
	<i>Oreochromis</i> <i>niloticus</i> – Nile tilapia (mostly diurnal)	LPS	ZT3/15	Greater humoral immune response when infected in the rest phase	102
	Phodopus sungorus - Siberian hamster (nocturnal)	LPS	ZT1/16	Shorter febrile response and more persistent locomotor activity when infected in the rest phase. Note, photoperiod is 16 hours light:8 hours dark	103
54 548 548 548 559 550 552 552 552 552 552 552 552	3         3         4         5         6         7         3         4         5         6         7         3         4         5         6         7         3         4         5         6         7         3         4         5         6         7         3         4         5         6         7         3         4         5         6         7         3         4         5         6         7         3         9         9         9          9          9          9          9          9          9          9          9          9 <td< td=""><td></td><td></td><td></td><td></td></td<>				

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- 846

### 847 Author information

### 848 Affiliations

- 849 Institute of Evolutionary Biology and Institute of Immunology and Infection Research, School of
- 850 Biological Sciences, University of Edinburgh, Edinburgh, United Kingdom
- 851 Mary L Westwood, Aidan J O'Donnell, & Sarah E Reece
- 852
- 853 Department of Biology, University of Central Florida, Orlando, FL, USA
- 854 Charissa de Bekker
- 855
- 856 Department of Biology, Indiana University, Bloomington, Indiana, USA
- 857 Curtis M Lively
- 858
- 859 Department of Ecology, Evolution and Behavior, University of Minnesota, St. Paul, MN, USA
- 860 Marlene Zuk
- 861

## 862 **Contributions**

- 863 SER conceived the study, MLW and SER drafted the manuscript, and all authors provided 864 substantial input into ideas and the writing of subsequent drafts.
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## 866 **Competing interests**

- 867 The authors declare no competing interests.
- 868
- 869 **Corresponding author**
- 870 Correspondence to Mary L Westwood (mary.westwood@ed.ac.uk)