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1	Learning by heart: Cardiac cycle reveals an effective time window for learning
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12 ABSTRACT

13 Cardiac cycle phase is known to modulate processing of simple sensory information. This 14 effect of the heartbeat on brain function is likely exerted via baroreceptors, the neurons 15 sensitive for changes in blood pressure. From baroreceptors the signal is conveyed all the 16 way to the forebrain and the medial prefrontal cortex. In the two experiments reported here, 17 we examined whether learning, as a more complex form of cognition, can be modulated by 18 the cardiac cycle phase. Human participants (Experiment 1) and rabbits (Experiment 2) were 19 trained in trace eyeblink conditioning, while neural activity was recorded. The conditioned 20 stimulus was presented in contingency with either the systolic or diastolic phase of the cycle. 21 The tone used as the conditioned stimulus evoked amplified responses in both humans 22 (electroencephalogram from 'vertex' (Cz)) and rabbits (hippocampal CA1 local-field potential) when its onset was timed at systole. In humans, the cardiac cycle phase did not 23 24 affect learning, but rabbits trained at diastole learned significantly better than those trained at 25 a random phase of the cardiac cycle. In summary, our results suggest that neural processing 26 of external stimuli and also learning can be affected by targeting stimuli based on the cardiac 27 cycle phase. These findings might be useful in applications aimed at maximizing or minimizing the effects of external stimulation. 28

29 NEW & NOTEWORTHY

30 It has been shown, that rapid changes in bodily states modulate neural processing of external 31 stimulus in brain. Here, we show that modulation of neural processing of external stimulus 32 and learning about it depends on the phase of the cardiac cycle. This is a novel finding that 33 can be applied to optimize associative learning.

34 Keywords: baroreceptor, classical conditioning, hippocampus, theta oscillation

35 INTRODUCTION

36 In 1884, William James reminded "brain physiologists", as he called them, that bodily states 37 affect how we experience the world. James' philosophy has the fundamental idea of the 38 consciousness being an inseparable stream of bodily and mental states. Since the 1880's science has verified in many ways that bodily states do alter the way we perceive or 39 40 experience the outer world through the inner world. Wilson (2002) suggests that episodic 41 memory consists of embodied experiences of the world. In his view, new forming episodic 42 memories are merged from contextual experiences of the environment as sensory information 43 and information of the body in different states experiencing the world and itself within it. As 44 the time passes on and these memories "become crystalized" they are not anymore modified by the bodily sensations. Therefore, if episodic memories are embodied, how is bodily 45 information merged with sensory information? 46

47 Information about the state of internal organs travels to the brain through neural and humoral pathways (Critchley and Harrison 2013). Baroreceptors, stretch and pressure -sensitive 48 49 sensory neurons found in blood vessels, activate during each heart beat as the vessel walls 50 distort. Their function is crucial in maintaining suitable blood pressure and they convey 51 information about the timing and strength of heartbeat to the nucleus of the solitary tract 52 (NTS, Critchley and Harrison 2013; Jänig 2006). The NTS is connected to the hypothalamus, 53 the parabrachial nucleus and the periaqueductal gray, which in turn are connected to forebrain 54 regions such as the amygdala, insular cortex, cingulate cortex and orbitomedial prefrontal 55 regions (Critchley and Harrison 2013). These anatomical and functional connections hint at 56 the idea that the heartbeat via baroreceptor activity could affect cognition and behavior (Lacey and Lacey 1974, 1978). 57

58 Indeed, the reported detection of a visual stimulus can be enhanced by presenting the stimuli 59 time-locked to a certain cardiac cycle phase (for T-wave, see Park et al. 2014; for P-wave, see Sandman et al. 1977). Likewise, visual evoked potentials are modulated as a function of 60 61 cardiac cycle phase (Walker and Sandman 1982). Thus, baroreceptor activity affects 62 cognition at least on the sensory level. There are indications that baroreceptor activity might 63 affect even more complex cognitive processes such as short-term memory performance 64 (Martins et al. 2014) and emotional appraisals of facial expressions (Gray et al. 2012). To our knowledge, it has not been tested whether baroreceptor activity could influence hippocampus 65 66 dependent associative learning.

67 The hippocampus is the critical hub of complex learning and episodic memory in the 68 mammalian brain (Squire 1992). Different frequencies of hippocampal electrophysiological 69 oscillatory activity reliably index different behavioral states. Theta (~4–8 Hz being peak frequency, depending on the species) is the most prominent oscillation in the hippocampus 70 71 (e.g., Buzsáki 1989). Theta is elicited by external stimuli and paced by cholinergic input from 72 the medial septum and glutamatergic input from the entorhinal cortex (Buzsáki 2002). The 73 critical role of the hippocampal theta in declarative learning is supported by a multitude of 74 experimental findings (Berry and Thompson 1978; Berry and Seager 2001; Griffin et al. 2004; Nokia and Wikgren 2010, 2014; for conflicting findings, see Múnera et al., 2001). 75 76 Overall, temporally robust hippocampal theta-band responses to the conditioned stimulus 77 predict good learning (Nokia et al. 2015).

In addition to contributing to cognitive processes via its mutual connections with the
neocortex (Buzsáki 1989), the hippocampus has connections with the hypothalamus
including supramammillary nucleus and posterior nucleus (Abrahamson and Moore 2001;
Cavdar et al. 2001; Pan and McNaughton 2004). The hypothalamus regulates the function of

the autonomic nervous system; therefore, for example, it affects blood pressure and heart beat
(Guyenet 2006), both directly and indirectly (Fanselow and Dong 2010). A few studies
actually propose that hippocampal theta oscillations are temporally aligned with the cardiac
cycle (Komisaruk 1970; Pedemonte et al. 2003) and cycles of rhythmic behavior like
mammalian sniffing (Macrides et al. 1982) and rats' whisking with their snout hairs (Grion et
al. 2016).

88 Here, to elucidate the potential connection between different phases of the cardiac cycle, 89 brain activity and associative learning, we subjected humans (Experiment 1) and rabbits 90 (Experiment 2) to trace eveblink conditioning (see Figure 1), a hippocampus-dependent task 91 considered to model declarative learning both in animals and in humans (Solomon et al. 92 1986; Holland and Bouton 1999). The onset of the conditioned stimulus (CS) was timed to 93 either the systolic or the diastolic phase of the cardiac cycle (see Figure 2). In rabbits, we also 94 included a group that was trained irrespective of cardiac cycle phase. Brain activity during 95 training was recorded from the scalp in humans (electroencephalogram, EEG) and directly 96 from the hippocampus in rabbits. In rabbits, we expected to see strong phase synchrony 97 between the ongoing hippocampal theta rhythm and the cardiac cycle. Further, we expected 98 both neural responses to the conditioning stimuli and learning at the behavioral level to be 99 different between the experimental groups in both humans and in rabbits. However, we had 100 no presumption concerning which phase, diastole or systole, would be optimal for learning.

101

102 EXPERIMENT 1: HUMAN EYEBLINK CONDITIONING AND EVENT RELATED 103 POTENTIALS

104 MATERIALS AND METHODS

105 Participants

106 Participants, recruited mainly via e-mail lists, gave an informed written consent to this study 107 and were free to discontinue participation at any point. The study was approved by the 108 University of Jyväskylä Ethical Committee. Thirty (23 female and 7 male, aged 18–32 years) 109 right-handed adults took part in the study. All participants were healthy with no history of psychiatric or neurological illnesses or medication affecting brain function. One participant 110 111 had to be excluded from all analyses because of a software malfunction. Due to technical 112 difficulties electrophysiological data from five participants were not analyzed. 113 *Experimental procedure* 114 Before the experiment, participants filled out a modified BIS/BAS personality inventory and 115 answered background questions about age, gender, height, weight, and schooling. In addition, 116 their blood pressure was measured before (and after) the experiment. 117 Participants were informed that the aim of the study was to record physiological and neural 118 responses to different types of stimuli. After recording 5 minutes of resting data, the trace 119 eyeblink conditioning procedure was started. The experiment was controlled by custom-120 written software running on an Arduino-based device that received input signal from the 121 electrocardiogram (ECG) -recording device whenever the signal exceeded the threshold set 122 roughly at 2/3 of the peak amplitude of the R-peak. The participants were randomized into 123 systole (n = 15) and diastole (n = 14) groups. In systole group, the trial onset was delayed by 124 a 100 ms from the rising slope of the R-peak, whereas in diastole group the trial onset was 125 delayed by 500 ms. The conditioned stimulus was a 200-ms, 440-Hz tone delivered via a 126 loudspeaker situated to the lower right-hand corner of the room. A 600-ms trace interval 127 separated the tone offset and the unconditioned stimulus (US) onset. The US was an airpuff

(0.4 bar source pressure, 100 ms) targeted to the corner of the left eye and it was delivered viaa plastic tube attached to modified safety goggles.

Before the actual conditioning phase, 4 air puffs alone were delivered at 5-second intervals to
accustom the participant to it. The conditioning procedure consisted of 80 trials. The first
(unpaired) and last (extinction) ten trials were CS-alone trials. The inter trial interval (ITI)
varied randomly between 9 to 19 seconds.

134 *Recordings and data analysis*

135 During the experiment, heart rate, eyeblinks and brain activity were recorded. The 136 participants were in a seated position during the experiment. Heart rate was recorded using 137 three ECG electrodes: one placed near the sternum and one over the right ribs, and the 138 grounding electrode over the left flank. Eyeblinks were recorded using two electromyography 139 (EMG) electrodes, which were placed underneath the participant's left eye. EEG data was 140 recorded using a 64-channel EEG-cap (64 BrainCap with Multitrodes, EASYCAP GmbH, 141 Woerthsee-Etterschlag, Germany). Resting state data was recorded for 5 minutes before and 142 after the actual experiment.

143 Eyeblinks

144 The EMG signal was high-pass filtered (> 60 Hz), rectified, and then low-pass filtered (< 20 Hz) off-line using Brain Vision Analyzer software. One of the authors (JW) blind to the 145 146 experimental group visually assessed all trials for conditioned responses. An eye-blink was 147 considered a CR if it occurred within a period of 500ms before the US onset. The exclusion 148 criterion was subjectively rated as excessive EMG activity during the 500ms time period 149 before the CS onset. The trials were grouped into 8 blocks of 10 for the sake of analysis and 150 the percentage of CRs per block was calculated. The learning curves of the systole and 151 diastole groups were compared using repeated measures ANOVA.

152 Brain responses

153 Valid EEG data was gathered from 24 participants (Systole: n = 12; Diastole: n = 12). BrainVision Analyzer 2.1 (Brain Products GmbH, Gilching, Germany) was used to remove 154 155 bad channels and low-pass filter (< 30 Hz) the raw data. Independent component analysis 156 (ICA) was run on the data and components related to eye blink, eye movement and heart beat artifacts were removed. The heartbeat itself is an event which induces a stereotypical activity 157 158 pattern in the electroencephalogram, called heartbeat evoked potential (HEP). It is found in 159 recordings over the somatosensory cortex (Kern et al. 2013) as well as fronto-cortical 160 (Schandry and Montoya 1996) and fronto-temporal (Montoya et al. 1993) areas. Therefore, 161 including an ECG channel in the ICA was used to remove a potential confounder from the 162 EEG data.

163 Event-related potentials (ERPs) recorded from the Cz channel were used for further analysis.

164 ERPs were calculated from a 500-ms time window at -100 to 400 ms in relation to the CS

165 onset. These epochs were first baseline-corrected by subtracting the average amplitude during

166 the 100-ms time window prior to CS onset. Then the baseline-corrected epochs were

averaged. For each participant, an average peak for N1 (minimum amplitude within 90 to 130

168 ms post-CS onset) and P2 (maximum amplitude within 150 to 230 ms post-CS onset) were

169 calculated from the paired trials (n = 60).

170 RESULTS

171 *CS evoked a larger N1 response when presented to the systole*

172 An independent samples t-test on ERPs showed that the N1 responses were larger in the

173 systole group compared to the diastole group (t (22) = 3.14, p < 0.01, Cohen's d = 1.28; see

174 Figure 3). The P2 responses were also larger in amplitude in the systole group, but the difference did not reach statistical significance (t (22) = 1.23, p = 0.23, Cohen's d = 0.52). 175 176 Due to potential confounding effect of heart-beats on ERPs, the ICA was used to remove 177 heart-related artefacts. Figure 4 depicts the effects of ICA on the EEG recorded at the Cz 178 electrode. As seen, there is a small (~1 μ v) deflection at about 350-400 ms after the R-peak in 179 the signal before the ICA. However, the corrected EEG signal is virtually flat. Thus, it can be 180 concluded that neither the heart beat related evoked potentials nor artifacts related to cardiac 181 cycle contribute to CS-evoked ERP amplitudes.

182 Cardiac cycle phase did not modulate learning in humans

183 Repeated measures ANOVA on the effects of Block (1-7, extinction block excluded) and Group (systole vs. diastole) on CR percentage revealed a significant main effect of Block (F 184 185 [6, 162] = 39.89, p < 0.001, partialEta2 = 0.57), indicating that the amount of conditioned responses increased as a function of training. Neither the main effect of group (F [1, 27] =186 0.03, p = 0.864, partialEta2 = 0.001) nor the interaction between Group and Block (F [6, 162]) 187 188 = 0.585, p = 0.742, partialEta2 = 0.021) reached significance, indicating that timing the CS 189 onset based on different phases of the cardiac cycle did not have an effect on learning trace 190 eyeblink conditioning (see Figure 5).

191 INTERIM DISCUSSION

As even relatively complex cognitive processes have been shown to be modulated by baroreceptor activity, we assumed that learning hippocampus-dependent trace eyeblink conditioning would differ between the humans trained at systole vs. diastole. Contrary to this expectation, both groups learned the task equally well. However, the systole group showed larger evoked responses (N1) to the conditioned stimulus. This suggests that cardiac cycle

- 197 phase affects sensory processing of external stimuli but these effects do not directly carry
- 198 over to learning at the behavioral level.

200 EXPERIMENT 2: RABBIT EYEBLINK CONDITIONING AND HIPPOCAMPAL 201 FIELD RESPONSES

202 MATERIALS AND METHODS

203 Subjects

204 The subjects were 25 adult female New Zealand White rabbits (Lidköpings Kaninfarm, Sweden) weighing ~2.8 kg at the time of surgery. The rabbits were housed in individual 205 206 cages in the laboratory center of the University of Jyväskylä. Food and water were freely 207 available, and room temperature and humidity were controlled. The rabbits were maintained on a 12/12-hour light/dark cycle, with lights on at 8:00 a.m. All experiments were carried out 208 209 during the light period. All experimental procedures, caretaking and handling were executed 210 in accordance with Directive 2010/63/EU of the European Parliament and the Council of 211 September 22, 2010 on the protection of animals used for scientific purposes. Animal 212 handling was done only by trained personnel and rabbits were introduced to human contact 213 and handling for a sufficient amount of time before the surgery.

214 Surgery

215 Before the surgery, rabbits were treated with subcutaneous injections (s.c.) of an anti-

216 inflammatory drug (50 mg/ml carprofen [Rimadyl vet, Pfizer Inc. Animal Health], dose: 0.1

217 ml/kg) and with 2 ml of an analgesic drug (0.3 mg/ml buprenorphine [Temgesic, Schering-

218 Plough Europe] diluted with 0.9 ml of 0.9% NaCl) to moderate acute pain after surgery. The

219 rabbits were anesthetized with an intramuscular injection (i.m.) of ketamine-xylazine cocktail

- 220 (7.8 mL of 50 mg/ml Ketaminol vet [Intervet International B.V.] mixed with 2.8 ml of 20
- 221 mg/ml Narcoxyl vet [Intervet International B.V.]). A dose of 0.8 ml/kg of the cocktail was
- 222 injected i.m. before surgery. During surgery, additional doses of either the cocktail or

ketamine alone were injected s.c. approximately every 20–30 min. or as needed. Before the
surgery, the rabbit's fur was shaved from the top of its head. Then, the rabbit was positioned
in a stereotaxic instrument (Kopf Instruments) with the bregma 1.5 mm higher than the
lambda. Eye gel was inserted into the rabbit's eyes. At this point, 2.0 ml of lidocaine (10
mg/ml Lidocain [Orion Pharma]) was injected s.c. in the area of surgery before making the
opening incision.

229 A longitudinal incision was made on the scalp and local anesthetic (2 g of lidocaine-230 hydrochloride Xylocain [AstraZeneca]) was administered to the wound. The skull was drilled 231 with holes for electrodes and four holes for the anchoring screws (5 mm anterior and 5 mm 232 lateral to the bregma; 13 mm posterior and 5 mm lateral to the bregma). Two of the screws 233 were connected together, and they were used as a reference. The other two served as the 234 ground for the electrophysiological recordings. For eight rabbits, eight monopolar recording 235 electrodes (Formvar-insulated nichrome wire; 0.05 mm bare [A-M Systems]) were 236 chronically implanted in the left dorsal hippocampus, with four electrodes aiming at the CA1 237 (4 mm posterior, 3.5–6.5 mm laterally and from the bregma; electrode tip depth from the 238 bregma 6-8 mm) and four above the hippocampal fissure (5 mm posterior, 4-7 mm laterally 239 from the bregma; electrode tip depth 6.2–8.5 mm below the bregma). For nine rabbits, eight 240 monopolar electrodes were implanted in both hippocampi (see coordinates above). For four 241 animals, a 32-channel (E32B-20-S04-L10.0-200, ATLAS Neuroengineering) adjustable four-242 shank probe was chronically implanted in the left dorsal hippocampus (5 mm posterior, 4 mm 243 laterally from the bregma) with a microdrive (nDrive xL, NeuroNexus). Wires, skull screws, 244 a preamplifier interface, one mounting screw for an air puff mount and the incision area were 245 cemented with dental acrylic. To prevent nausea after surgery, metoclopramide (0.1 ml/kg, 246 concentration 5mg/ml; Primperan [Sanofi Winthrop Industrie]) was administered s.c. and the 247 rabbit was returned to its home cage wrapped in a towel. Recovery was monitored and the

rabbits were medicated with analgesic (buprenorphine [Temgesic, Schering-Plough Europe]
diluted with 0.9 ml of 0.9% NaCl) four hours after surgery and then every eight hours for the
next 44 hours.

251 *Experimental procedure*

252 The experimental procedure is illustrated in Figure 1. After one week of recovery from 253 surgery, animals were accustomed to a Plexiglas restraining box without restraining and their 254 overall behavior was monitored. Local field potentials (LFPs) and EMG from the right eye were recorded 5 minutes prior, during and 1 minute after each session. ITI always varied 255 256 randomly between 30 and 60 s. LabVIEW (National Instruments) was used to monitor the 257 cardiac cycle and blinking on-line, to execute the experimental procedures and to present 258 stimuli. After the ITI was expired, trial presentation was always delayed for one second every 259 time the rabbit was spontaneously blinking. The percentage of learned responses performed 260 by each animal was analyzed after every session using MATLAB (The MathWorks Inc.).

During the first training session (CS-Alone), 60 tone-alone (200-ms, 5-kHz, 75-dB tone)
trials were presented regardless of cardiac cycle phase. In addition to hippocampal LFPs,
EMG from the right eye was also recorded to determine the frequency of spontaneous
eyeblinks elicited by the tone later used as a CS.

Trace eyeblink conditioning was carried out with the tone specified above as the CS and a 100-ms air puff (0.35 bar source pressure) to the right eye as an unconditioned stimulus (US). A trace period of 500 ms was used. A total of 60 training trials were presented during each session, regardless of neural state and in the absence of spontaneous blinking. The trials were timed so that the CS started either at the systolic or the diastolic phase of the cardiac cycle, or irrespective of cardiac cycle phase. A total of 14 sessions were conducted.

271 Recordings and data analysis

272 Cardiac cycle

- 273 The cardiac cycle was monitored with a pulse oximeter (Shimmer Optical Pulse Sensor,
- 274 Realtime Technologies Ltd) attached to the rabbit's shaved right earlobe.
- 275 Photoplethysmography (PPG) is a robust measure for monitoring the cardiac cycle (see
- 276 Wisely and Cook 2001). The temporal relation between ECG and PPG in rabbits was
- 277 confirmed during surgery with an anesthetized rabbit by recording the ECG with two needles
- 278 positioned on both sides of the animal, leaving the heart in between.

279 Eyeblinks

- 280 Bipolar EMG from the trained eye was recorded using stainless steel wire-hooks placed
- around the right upper and lower eyelids for the duration of the training sessions. The raw
- 282 EMG signal was conveyed to a filter-amplifier (A-M Systems Model 2100), amplified 1000x
- and band-pass filtered from 100 to 500 Hz. The EMG signal was high-pass filtered off-line
- 284 (>100 Hz) and Hilbert-transformed. An envelope curve following the peaks of the signal was
- 285 calculated. Baseline EMG activity was defined for each animal and session as the mean of the
- 286 peak EMG amplitude during a 250-ms pre-CS period (MEANpre). The mean of the standard
- 287 deviation of the EMG activity during the 500-ms pre-CS period (SDpre) was also determined.
- 288 Eyeblinks were defined as EMG activity exceeding a threshold of [MEANpre + $4 \times$ SDpre]
- for at least 10 ms. Trials with eyeblinks during the 100-ms period immediately preceding CS
- 290 onset were rejected. Eyeblinks 100ms prior to US onset were counted as conditioned

responses.

292 <u>Hippocampal local-field potentials</u>

293 For neural recordings of monopolar electrodes, a tenfold amplification was performed with a 294 preamplifier (MPA8I, MultiChannel Systems [MCS]) attached to the electrode connector in the rabbit's head. Then the signal was band-pass filtered (1-5000 Hz) with a 64-channel filter 295 296 amplifier (MCS). Lastly, the signal was further low-pass filtered (500 Hz) and digitized at a 297 rate of 2 kHz with a MCS USB-ME64 System (MC_Rack software). SPSS (IBM) and 298 MATLAB (MathWorks) were used for offline data analysis. Rabbits implanted with the 32-299 channel probes (Atlas Neuroengineering) were recorded with a wireless data acquisition 300 system (W2100-HS32-headstage, MultiChannel Systems [MCS]) with a 20 kHz sampling 301 rate.

302 To assess the temporal accuracy of the theta-band responses to the conditioning stimuli, a 303 phase-locking value (PLV) was calculated (Palva et al. 2005b). The PLV is based on 304 amplitude-normalized phase information and is thus resistant to changes or differences in 305 signal amplitude. This allows comparable measurements to be obtained from data recorded 306 over time in multiple subjects. The hippocampal LFP data were first band-pass filtered 307 between 4 and 8 Hz. Then, a Hilbert transform was run on the signal to obtain the phase 308 information, and the amplitude of the transformed signal was normalized to 1 by dividing 309 each data point by its absolute value. Finally, the PLV was obtained by averaging over 60 310 trials (one session) and taking the absolute value of the mean. The PLV varies between 0 and 1, 0 indicating no phase locking and 1 indicating perfect phase locking. For statistical 311 312 analyses, the mean of the PLV during the CS and subsequent trace-period (700 ms) was 313 derived and averaged over one session for each subject.

The phase synchrony (PS, Palva et al. 2005b) of the hippocampal theta (band-pass filtered

between 4–8 Hz) and PPG (band-pass filtered between 3–6 Hz) was analyzed next. The LFP

and PPG sweeps were selected randomly from occasions where theta ratio was high (>80%)

and the PPG signal quality was good. Both signals were Hilbert-transformed and their
amplitudes normalized to 1, as explained above. Then, the phase difference of the two signals
was calculated by multiplying the first signal with the complex conjugate of the second signal

320 (each data point of each sweep). Finally, the PS was derived by averaging the phase

321 difference matrix over sweeps, taking the absolute value.

322 Neural responses evoked by the CS in hippocampal CA1 were averaged within each session

323 per animal. Negative peak amplitudes of these event-related potentials (ERPs) were analyzed

from 25 to 60 ms after CS onset (see Figure 6B) and normalized to CS-Alone session

325 amplitudes [({session ERP amplitude – CS-Alone ERP amplitude} / CS-Alone ERP

amplitude) * 100]. The placement of the electrodes in CA1 was confirmed with histology and

in addition by inspecting sharp-wave ripples.

328 Statistical analyses

329 Repeated-measures analysis of variance (ANOVA), with training sessions (or blocks of two 330 sessions) as a within-subjects factor and group as a between-subjects factor, was used to 331 analyze changes across training and differences between experimental groups. For post hoc 332 comparisons, Bonferroni-corrected p-values are reported. One-way ANOVA or an 333 independent samples t-test was used for comparisons between groups one dependent variable 334 at a time. One-way ANOVA was used to test the difference between the groups using the 335 session from the last four training sessions for each individual animal where they achieved 336 their best performance in conditioned responses.

337 Histology

Rabbits were anesthetized with an i.m. injection of ketamine-xylazine cocktail and then
overdosed with an i.v. pentobarbital (Mebunat vet, Orion-Yhtyma Oyj) injection. Then, the
brain was perfused with physiological saline followed by 9% formalin solution through the

341 ascending aorta. The locations of the electrode tips were marked by passing a DC current

342 (200 mA, 10 s) through them. The brain was then removed and stored in formalin for several

343 days. The brain was coronally sectioned with a vibratome into 60-µm-thick slices. The slices

344 were attached to gelatinized slides, dried, and stained with Prussian blue and cresyl violet.

345 The electrode locations were determined with the help of a microscope.

346 RESULTS

347 Hippocampal theta phase was not in synchrony with cardiac cycle phase

348 Phase synchrony between hippocampal fissure LFP and the PPG signal reflecting the cardiac

349 cycle was analyzed from periods of spontaneously occurring theta oscillations (theta ratio >

350 80% during ITI). The average phase synchrony from all sessions and all rabbits was 0.17 (SD

351 = 0.07), on a scale of 0 to 1, with 1 indicating perfect phase synchrony. That is, no phase

352 synchrony between theta and the cardiac cycle was detected. The mean heart rate of the

rabbits during the sessions was ~180 beats per minute (bpm), which is within normal

variation (130 bpm to 325 bpm; see Pritchett-Corning et al. 2011).

355 *CS evoked larger hippocampal responses when presented to the systole*

356 Histological examinations confirmed that recording electrodes were in or near the

357 hippocampal CA1, as intended, in 19 animals (see Figure 6A). Note that four of the rabbits

358 had been implanted with multisite silicon probes that were adjusted constantly during the

359 experiment. Therefore they are not included in this analysis.

360 Event-related potentials to the CS recorded from the CA1 had a mean latency of 41 ms (SD =

361 4.80 ms) from CS onset. The amplitude of this response was moderated by the phase of the

362 cardiac cycle so that the amplitudes were higher in the Systole group (n = 6) compared to the

363 Diastole group (n = 7) (repeated measures ANOVA, interaction of group and session: F [12,

364	96] = 0.28, p = 0.99; main effect of session: F [6, 96] = 0.47, p = 0.83; main effect of group
365	F[2, 16] = 4.44, p < 0.05; Systole vs. Diastole, Bonferroni-corrected post-hoc comparison: p
366	= 0.027).see Figure 6C).

- 367 Phase-locked hippocampal theta-band responses to the CS were not different between the
- 368 Diastole, the Systole, and the Random group during trace conditioning (repeated measures
- ANOVA, interaction of group and session: F[12, 108] = 1.23, p = 0.27; main effect of

370 session: F[6, 108] = 1.29, p = 0.267; main effect of group F[2, 18] = 0.24, p = 0.79).

371 Rabbits trained at diastole learned better than those trained irrespective of cardiac cycle
372 phase

- 373 Twenty-one out of the 25 animals learned trace eyeblink conditioning. Learning differed
- between the Diastole (n = 10), Systole (n = 8) and the Random (n = 7) groups (repeated
- 375 measures ANOVA, interaction of group and session: F[12, 132] = 0.79, p = 0.66; main effect
- of session: F [6, 132] = 11.08, p < 0.0001; main effect of group F [2, 22] = 3.94, p < 0.05; see
- Figure 7A). Specifically, learning was better, when the CS onset was timed to the diastole
- 378 phase of the cardiac cycle compared to when it was presented in a random phase (Diastole vs.
- 379 Random, Bonferroni-corrected post-hoc comparison: p = 0.036).
- 380 There was also a significant difference between the groups in the best performance (CR %)
- they reached during the last four sessions of conditioning (one-way ANOVA: F [2, 22] =
- 4.38, p = 0.025). Post hoc comparisons indicated that the best performance in the Diastole
- group (M = 82.36, SD = 11.93) was significantly higher than that in the Random group (M =
- 52.33, SD = 32.54) (Bonferroni-corrected p = 0.022). However, the Systole group (M =

385 68.64, SD = 15.67) did not significantly differ from the other two groups (Bonferroni-

- 386 corrected post-hoc comparisons: Systole vs. Diastole: p = 0.524; Systole vs. Random: p =
- 387 0.422; see Figure 7B).

388 INTERIM DISCUSSION

389 As anticipated, the cardiac cycle phase affected both neural responses as well as behavior 390 during trace eyeblink conditioning in rabbits. Namely, hippocampal responses evoked by the 391 conditioned stimulus were larger in amplitude in the Systole group compared to the Diastole 392 group. Further, rabbits learned trace eyeblink conditioning better when the CS onset was timed to the diastole phase of the cardiac cycle. In fact, almost all (90%) of the animals in the 393 394 Diastole group reached a limit of 80% CRs per session, whereas only half in the Systole 395 group and less than third in the Random group reached this limit during the 14 sessions of 396 trace eyeblink conditioning. The animals in the Diastole group also learned exceptionally 397 well when compared to previous results from our lab using the same paradigm for trace 398 eyeblink conditioning (Nokia and Wikgren 2014; Nokia et al. 2015; Waselius et al. 2017).

400 **DISCUSSION**

401 Neural responses, as well as simple sensory phenomena have been shown to vary depending 402 on the timing of the stimuli in relation to the phase of the cardiac cycle. Here, both human 403 participants and rabbits were subjected to trace eyeblink conditioning where the onset of the 404 conditioning trial was timed either to the systolic or diastolic phase of the cardiac cycle. This 405 task revealed that neural responses (scalp EEG in humans and LFPs from hippocampal CA1 406 in rabbits) to the tone-CS differed between the Systole and Diastole groups. Namely, the 407 responses to the tone-CS were enhanced when targeted to the systolic phase. On the contrary, 408 an enhancement of the learning rate was evident in the Diastolic group in rabbits. No effect of 409 cardiac cycle phase on learning rate was found in humans.

410 Earlier studies show, that behavioral (Sandman et al. 1977; Gray et al. 2012; Martins et al. 411 2014; Park et al. 2014) and neural (Walker and Sandman 1982) responses in humans can be 412 modulated by presenting stimuli time-locked to the cardiac cycle phase. In Experiment 1 413 neural responses to the tone used as a conditioned stimulus measured with EEG differed 414 between participants trained at diastolic vs. systolic phase of the cardiac cycle. However, both 415 of the groups learned the task at the same pace. It is to be noted that single-cue trace eyeblink 416 conditioning is a relatively easy task for humans and learning occurs rapidly. Factors known 417 to affect the learning rate include, for example, awareness (Manns et al. 2000), and 418 cholinergic blockade by scopolamine (Solomon et al. 1993). While cardiac cycle might have 419 some effect on the way a stimulus is processed (as indicated by previously reported ERP and 420 sensory threshold studies), the effect might be too subtle to manifest in associative learning 421 which is a process governed by a multitude of top-down and bottom-up factors. It might be 422 that making the task a bit more demanding (e.g., increasing the trace period or lowering the 423 amplitude of the conditioned stimulus near to the detection threshold) might yield differences

424 also at a behavioral level. Running Experiment 1 again using the same parameters but with 425 elderly adults could also reveal differences in learning between groups, as it is known that 426 aging has a deteriorating effect on the ability to learn trace eyeblink conditioning (for 427 example, see Woodruff-Pak et al. 2001). In the future, it would be important to use a more 428 demanding task that should make the initial learning rate slower but eventually result in 429 progressing to a better overall performance of conditioned responses.

430 In Experiment 2, we utilized the same set-up as in Experiment 1, but conducted the study in 431 rabbits with chronically implanted recording electrodes in the hippocampus. First we tested 432 whether there is a temporal correlation between the cardiac cycle and hippocampal theta 433 oscillation (see, Komisaruk 1970; Pedemonte et al. 2003). Much to our surprise, there was no 434 phase synchrony between theta and the cardiac cycle. Next, we examined hippocampal 435 responses to the conditioned stimulus. Our previous studies indicate that hippocampal responses at the theta-band (4-8 Hz) are generally better time-locked to the CS-onset in 436 437 subjects that learn well (Nokia et al. 2015). Like in human participants in Experiment 1, 438 neural responses evoked by the tone-CS were also modulated by the cardiac phase in rabbits. 439 That is, the hippocampal CA1-evoked potentials were larger in the Systole group compared 440 to the Diastole group. However, the phase-locking of CA1 theta-band responses evoked by 441 the CS did not differ between groups. This is perhaps a consequence of the lack of synchrony 442 between theta and the cardiac cycle. Last but not least, rabbits trained at diastole learned trace 443 eyeblink conditioning better than those trained at systole.

We admit, that the timing of the US in the Experiment 2 was incoherent compared to the timing of the CS, since the heart rate varied greatly in rabbits. This could have affected learning in trace eyeblink conditioning. At the same time, we emphasize that varying the trace interval between the CS and US could have affected the learning even more and the

448 results of the experiment would have been hard to interpret. If we could have managed to 449 come around with a solution, where trace interval would have been stable and timing of CS 450 and US would have been in the same phase of the cardiac cycle, the results could have been 451 different i.e. learning rates in Systole group would have been lower. Also, we recorded neural 452 responses to CS only in the CA1 region which is in the end of the trisynaptic circuit of the 453 hippocampus. Neural responses in the CA3 and the dentate gyrus could have been modulated 454 differently i.e. responses to the CS during the diastolic phase could have been larger than 455 those elicited in the systolic phase.

456 Taken together our results suggest that the effects that the cardiac cycle phase has on neural 457 responses to a conditioned stimulus, or learning at the behavioral level, cannot be explained 458 by the connection between hippocampal theta and learning (Nokia et al. 2015; Waselius et al. 459 2017; see also Hasselmo et al. 2002). Based on our current results it would seem, that the 460 neural state affecting learning fluctuates also according to baroreceptor signaling based on the 461 pressure in arteries. This signal is conveyed to the brain via the NTS but which brain regions 462 and what mechanisms are affected by the fluctuating signal remains unclear and should be 463 studied further. It is known that input from sensory terminals arrives to the hippocampus 464 through two primary, connected pathways: the non-lemniscal (via the medial septal nucleus) and the lemniscal (through the primary auditory cortex and the entorhinal cortex) (see 465 466 Bickford et al. 2002). The function of the non-lemniscal pathway is reflected in hippocampal 467 theta activity when cholinergic input from the medial septum to the hippocampus is strong 468 (Buzsáki 2002). At the same time, the pulsatile activity of baroreceptors is constantly 469 projected to the hippocampus through the lemniscal pathway, via the neocortex. 470 Hippocampal responses to external stimuli are modulated by the functioning of these two 471 pathways and possibly by some other mechanisms as yet unknown. In the future we should 472 run a cardiac cycle phase -contingent experiment and record i.e. the activity of ventral portion

473 of the medial prefrontal cortex (vMPFC) which has inputs from baroreceptors (see Resstel et

474 al. 2004). Also, we could study peripheral sensitivity (see Edwards et al. 2009) of auditory

475 organs during different phases of the cardiac cycle.

476 CONCLUSIONS

We found that the phase of the cardiac cycle at stimulus onset affects neural responses to a behaviorally relevant external stimulus in humans and in rabbits. Furthermore, learned behavioral responding to the stimulus was modulated in rabbits. That is, very rapid changes in bodily state can affect learning. Monitoring cardiac cycle and timing of the stimulus in contingency with it might be used to optimize the effect of external stimulation and learning.

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615 FIGURE CAPTIONS

- 616 Figure 1. Trace eyeblink conditioning.
- 617 Rabbits (n = 25) and humans (n = 29) were trained in trace eyeblink conditioning with tone as
- a conditioned stimulus (CS) and an air puff aimed toward the eye as an unconditioned
- 619 stimulus (US). The trace period was 500 ms for the rabbits and 600ms for the human
- 620 participants. UR = unconditioned response, CR = conditioned response.
- 621

622 Figure 2. In both human participants (A, Experiment 1) and rabbits (B, Experiment 2) the

- 623 conditioned stimulus was timed either to the diastole or the systole.
- 624 The cardiac cycle can be divided into two phases, systole and diastole. During systole, the
- heart contracts and pumps blood to arteries. During this phase the electrocardiogram (ECG)
- shows the QRS-complex reflecting ventricular depolarization and the T-wave, reflecting
- 627 ventricular repolarization. Diastole follows systole. During diastole, the heart relaxes and fills
- 628 with blood and the P-wave is seen in ECG. It reflects atrial depolarization which occurs when
- the ventricles are almost full of blood. A) In human participants, the ECG was recorded and
- 630 used for CS timing. B) In rabbits, the oxygen saturation signal (photoplethysmogram, PPG)
- 631 was measured from the earlobe using a pulse oximeter. The PPG is at its lowest at the
- beginning of the systole phase. The highest peak in the PPG can be observed during the
- relaxation of the heart just before the atrial contraction (ECG P-wave), that is, during
- 634 diastole. The CS was timed to start either at the systole (trough, arrowhead) or the diastole
- 635 (peak, double arrowhead) or irrespective of cardiac cycle phase (Random, not shown).
- 636
- 637 Figure 3. Topographies of the event-related potentials to the tone-CS (A) and maximum
- 638 amplitudes of N1 and P2 responses measured at Cz-electrode (B) in human participants in
- 639 Experiment 1. The N1 response was significantly larger when the tone onset was contingent
- 640 with the systolic phase (n = 12) compared to diastolic phase (n = 12). The P2 response was

641 also larger in amplitude but did not reach statistical significance. In B, error bars equal 642 standard error of mean. Asterisks in B denote p-value of < 0.01. 643 644 Figure 4. (A) Effect of ICA-based artefact correction on EEG recorded at Cz electrode. The 645 EEG traces in the upper panel are grand average responses to the heart beat. The arrows mark 646 the onsets of CSs in the Systole and Diastole groups. As can be seen, there are minor 647 deflections related to cardiac cycle in the signal before ICA correction (Raw) but after that (Corrected), the signal is virtually flat. (B) ECG topography in the same time scale is plotted 648 649 in the lower panel. 650 Figure 5. Human participants in Experiment 1 learned trace eyeblink conditioning at the same 651 652 rate regardless of group (Systole vs. Diastole). Error bars equal standard error of mean. UP = unpaired, CC = conditioning, EXT = extinction. 653 654 Figure 6. In Experiment 2, relative amplitudes of event-related potentials to the conditioned 655 stimulus (CS) recorded from the rabbit hippocampal CA1 were larger in Systole group 656 657 compared to the Diastole group. A) A 60-micron cresyl violet-stained slice of a rabbit left dorsal hippocampus with the 658 locations of CA1 and pyramidal cell layer (pyr) indicated. The electrode tip location was 659 marked by passing direct current through the electrode (arrow). B) Example of a 660 representative CS-evoked ERP (average of 60 trials) in hippocampal CA1. Negative peak 661 amplitudes of ERPs were analyzed from 25 to 60 ms after CS onset (vertical gray bar). C) 662 ERPs in CA1 were averaged per session, per animal. The value from the CS-Alone session 663 was used as a baseline to calculate the relative change (%) in amplitude of the ERP during 664 subsequent conditioning sessions (see Methods). Throughout the 7 session blocks of trace 665 eyeblink conditioning the responses to CS were amplified in the Systole group and attenuated 666 667 in the Diastole group (p < 0.05). Error bars equal standard error of mean. 668 Figure 7. In Experiment 2, rabbits trained at diastole learned better than those trained 669 irrespective of cardiac cycle phase. 670 A) Learning was faster and conditioned responding remained at a higher level in the Diastole 671 group throughout trace eyeblink conditioning. Asterisk refers to repeated-measures ANOVA 672 673 indicating statistically significant difference (Bonferroni-corrected post-hoc comparisons, p < 0.05) between the Diastole group and the Random group. B) B) The best performance was 674 determined as the highest achieved performance in learned responses during one session from 675 the last four training sessions for each individual animal. There was a significant difference 676 between the Diastole and the Random groups (p < 0.05). C) The cumulative number of 677 learned responses plotted as a function of trial number in the Diastole, Systole and Random 678 679 groups indicates no initial difference in responding. Error bars equal standard error of mean.













