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Pattern analyses reveal separate experience-based fear memories in the human right amygdala

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Abstract

30 Learning fear via the experience of contingencies between a conditioned stimulus (CS) and an
31 aversive unconditioned stimulus (US) is often assumed to be fundamentally different from
32 learning fear via instructions. An open question is whether fear-related brain areas respond
33 differently to experienced CS-US contingencies than to merely instructed CS-US contingencies.
34 Here, we contrasted two experimental conditions where subjects were instructed to expect the
35 same CS-US contingencies while only one condition was characterized by prior experience with
36 the CS-US contingency. Using multi-voxel pattern analysis of fMRI data, we found CS-related
37 neural activation patterns in the right amygdala (but not in other fear-related regions) that
38 dissociated between whether a CS-US contingency had been instructed and experienced versus
39 merely instructed. A second experiment further corroborated this finding by showing a
40 category-independent neural response to instructed and experienced, but not merely instructed,
41 CS presentations in the human right amygdala. Together, these findings are in line with
42 previous studies showing that verbal fear instructions have a strong impact on both brain and
43 behaviour. However, even in the face of fear instructions, the human right amygdala still shows
44 a separable neural pattern response to experience-based fear contingencies.

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Significance statement

48 In our study we addressed a fundamental problem of the science of human fear learning
49 and memory, namely whether fear learning via experience in humans relies on a neural pathway
50 that can be separated from fear learning via verbal information. Using two new procedures and
51 recent advances in the analysis of brain imaging data, we localize purely experience-based fear
52 processing and memory in the right amygdala, thereby making a direct link between human and
53 animal research.

54

55

Introduction

56 As a product of evolution, animals are equipped with the ability to learn relations that
57 impact their survival. For example, by recognizing contingencies between certain stimuli in the
58 environment (CSs) and harmful events (USs), animals can learn to anticipate these events in the
59 future, a process which is thought to underlie Pavlovian fear conditioning (Maren, 2001;
60 Pavlov, 1927). Pavlovian learning is often distinguished from other forms of fear acquisition
61 (e.g., via instructions or observation; Olsson & Phelps, 2007) as it necessitates first-hand
62 experiences of paired events rather than information transfer from an instructor or model. The
63 experience-based nature and strong evolutionary conservation of Pavlovian conditioning led
64 many theorists to think that conditioning in humans happens relatively automatically and
65 independently from verbal processing or even awareness (Dolan & Vuilleumier, 2003; Grillon,
66 2009; LeDoux, 2014; Mineka & Öhman, 2002; Olsson & Phelps, 2004; Schultz & Helmstetter,
67 2010). Accordingly, it has been proposed that there is an evolutionary old fear module in the
68 human brain, centered around the amygdala, that contributes to the acquisition and expression
69 of Pavlovian fear by specifically mediating its putative non-verbal, experience-based element
70 (Öhman & Mineka, 2001). In its strongest form, this theory postulates the amygdala-centered
71 fear module to be “encapsulated”, i.e., impenetrable to conscious cognitive control (Öhman &
72 Mineka, 2001).

73 If there is a brain module responsible for purely experience-based Pavlovian fear learning,
74 it must operate relatively independently from verbally-mediated or instructed fear learning.
75 Therefore, to test whether Pavlovian fear learning can operate independently from verbally-
76 mediated fear learning, previous studies tried to isolate neural correlates of Pavlovian learning
77 by evidencing conditioning in the absence of CS awareness, that is, with backward-masked or
78 subliminally presented CSs. These studies often pointed towards the amygdala (Critchley et al.,
79 2002; Knight et al., 2009; Morris et al., 1998; Tabbert et al., 2011) as the neural substrate of
80 Pavlovian fear learning, but also received substantial criticism based on methodological
81 (potential residual CS awareness, Mitchell et al., 2009) and statistical grounds (Vadillo et al.,

82 2016). Even evidence for fear conditioning in non-verbal human children (Watson & Rayner,
83 1920) still leaves open the main question that motivated our research: Once a human becomes
84 verbal, can these verbal processes override learning pathways via experience, or do we keep
85 separable pathways for experience-based fear conditioning instead? In other words, it remains
86 unclear whether the human brain reserves space for the unique impact of actually experiencing
87 CS-US pairings, in the face of explicit fear instructions.

88 If we want to demonstrate a truly independent, separable neural response to experience-
89 based Pavlovian conditioning, we must contrast it to verbally-mediated instruction-based
90 learning. Therefore, rather than trying to exclude conscious or language-based processing, as in
91 previous studies, we here developed an experiment where conditions were optimal for verbally-
92 mediated language-based processing to override the hypothesized separate experience-based
93 component to fear learning. To this end, we compared neural pattern responses to two CSs that
94 were both part of explicitly instructed contingencies but of which one was (CS⁺P, CS⁺ Paired)
95 and the other was not (CS⁺U, CS⁺ Unpaired) previously paired with the US (Mertens et al.,
96 2016; Raes et al., 2014). Hence, whereas both stimuli were expected to activate the same
97 instruction-based fear memory during the memory testing phase, only the previously paired CS
98 (CS⁺P) should additionally activate experience-based memory elements, which we here call the
99 Pavlovian trace. This way, we studied a unique, experience-based component of Pavlovian fear
100 conditioning in humans. In a second experiment, we aimed to replicate and further extend this
101 finding by testing whether a similar neural signature could be observed when comparing
102 entirely novel (i.e., merely instructed) to old (i.e., instructed and experienced) CS presentations.

103

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Material and Methods

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General Method

106

In our first and main experiment, participants were first instructed that two visual stimuli
107 (CS⁺P, CS⁺U) could be followed by a painful electrocutaneous stimulus (US). A third stimulus (CS⁻
108) was introduced as a control stimulus that would never be paired with the US. Participants were

109 then told that, in a preparatory training phase, only one of the two CS⁺s (CS^{+P}) would be
110 occasionally paired with an electrical stimulus, whereas the other CS⁺ (CS^{+U}) would be
111 occasionally followed by a placeholder (drawing of a lightning bolt) – under the false pretense
112 that this limitation of the absolute number of electrical stimulation given would allow the
113 subjects to gradually adjust to the aversive task conditions. During this training phase, we
114 randomly presented each CS nine times; three out of nine CS^{+P} presentations were followed by
115 a US, making subjects experience the CS^{+P}-US contingency, while three out of nine CS^{+U}
116 presentations were followed by the placeholder (Figure 1). Before the subsequent test phase,
117 participants were re-instructed that, from then on, both CS⁺s could be followed by a US. In fact,
118 no more USs were delivered, keeping this critical phase of the experiment free of any Pavlovian
119 fear learning. Because instructions in the test phase were identical for both CS⁺s, neural activity
120 patterns associated exclusively with the CS^{+P} during the test phase would reflect a Pavlovian
121 trace of actual CS-US pairings. This procedure has been tested before by Raes and colleagues
122 (2016) and Mertens and colleagues (2016) but without simultaneous MRI recordings. Therefore,
123 the behavioral data will be discussed in close comparison to those studies.

124 A second experiment was performed to examine whether our main finding could also be
125 observed in a different, but conceptually similar procedure. As an additional motivation, this
126 second experiment also allowed us to examine whether a higher visual category-independent
127 neural response to instructed and experienced CS presentations could be observed, as compared
128 to merely instructed CS presentations. This experiment was different in design from Experiment
129 1, but nonetheless allowed for a similar comparison between instructed and experienced versus
130 merely instructed fear. Specifically, we used a large range of different CSs (house or face
131 pictures), and participants were instructed on the relevant CS-US contingencies before each CS
132 presentation (Figure 8a). That is, each trial of this experiment started with the presentation of an
133 instruction screen that defined the CS⁺ and the CS⁻, followed by the presentation of the CS⁺ or
134 CS⁻. Each CS⁺ was in its turn followed by a US presentation. This way, participants saw many
135 different CS instructions. Importantly, however, whereas some contingency instructions (and

136 the experience thereof) recurred multiple times throughout the experiment (i.e., old CS stimuli),
137 others were always new (i.e., new CS stimuli). Hence, some contingencies had been "instructed
138 and experienced" whereas, for new CSs, this was not the case. The new CSs could therefore be
139 considered "merely instructed".

140 In contrast to Experiment 1, Experiment 2 also allowed for a comparison within CS
141 conditions but across visual stimuli or visual categories. Specifically, the different CS
142 instructions could be further subdivided into those that used pictures of faces as CSs, and those
143 that used pictures of houses as CSs. This way, we could compute the similarity in neural
144 activation patterns at the time of CS presentation between faces versus houses separately (see
145 Figure 8b) for each of the four different CS presentation conditions: CS⁺old, CS⁻old, CS⁺new,
146 and CS⁻new. By computing similarities between responses to faces and houses, this study
147 allowed us to investigate to which extent fear-related regions showed a neural response that was
148 independent from CS object category, and thus specific to whether a CS carried a representation
149 of threat (CS⁺ vs. CS⁻) or whether it was old or novel (CSold vs. CSnew). Following up on the
150 findings of the first experiment, we zoomed in on the pattern similarities in the left and right
151 amygdala. In particular, we were interested in whether the right amygdala would show a higher
152 similarity between face and house evoked patterns when a CS⁺ was old, which would indicate a
153 representation of threat sensitive to whether a contingency had been experienced before.

154 **Participants**

155 Twenty participants took part in Experiment 1 (twelve women and eight men, mean age =
156 24, SD = 2.5, range = 19 - 28), and another twenty in Experiment 2. One participant from
157 Experiment 2 was excluded from analyses due to self-reported nausea and inattention to the
158 task, so the final sample of Experiment 2 contained 19 participants (ten women and nine men,
159 mean age = 24, SD = 3.7, range = 18 - 34). All participants had normal or corrected to normal
160 vision, and were right-handed as assessed by the Edinburgh Handedness Inventory. They gave
161 their informed written consent and reported no current or history of neurological, psychiatric or

162 major medical disorder. Every participant was paid 35€ for participating. The work has been
163 completed with the approval of the Ghent University Hospital Ethical Committee.

164 **Stimuli and Procedure: Experiment 1**

165 *Stimuli.* The conditioned stimuli consisted of three dissociable blue fractal figures
166 (snowflakes) that were presented on a white background. Counterbalanced across participants,
167 one of the fractals served as a conditioned stimulus (CS) that would never be followed by an
168 electro tactile pain stimulus (CS⁻), another served as a CS that could occasionally be followed by
169 a pain stimulus (CS^{+P}), and a third as a CS that subjects were told could be followed by a pain
170 stimulus, but was actually only occasionally followed by a placeholder (CS^{+U}). The
171 electro tactile pain stimulus that served as the aversive stimulus (unconditioned stimulus, US)
172 consisted of a train of twelve square-wave pulses of 2 ms duration each (interval 18 ms). The
173 US was delivered through a surface electrode with platinum pin (Specialty Developments,
174 Bexley, UK) onto the right leg over the retromalleolar course of the sural nerve using a DS5
175 electrical stimulator (Digitimer, Welwyn Garden City, UK). The intensity was determined
176 through a standard work-up procedure prior to the experiment (more info below). The
177 placeholder consisted of a centrally presented yellow drawing of a lightning bolt.

178 *Ratings.* Self-reported CS fear and US expectancy were assessed for all CSs in separate
179 rating blocks interspersed between conditioning trials. These ratings were performed on screen.
180 On a typical rating trial, the CS was presented centrally, while the question on fear or US
181 expectancy was situated on top of the CS and a rating scale was presented below. Before each
182 rating phase, participants were instructed to respond to the questions that would appear at the
183 top of the screen through selecting the response possibility that felt most appropriate to them.
184 Furthermore, it was stressed that these questions pertained to their most recent encounter with
185 the CSs during the foregoing (conditioning) phase. In addition, participants were instructed that
186 “Whenever you are asked about your expectancy of an electrical stimulus, we refer to the actual
187 stimulation, not to the picture of the lightning bolt”. The questions that appeared were “How
188 much fear did you experience when looking at this figure?” (self-reported CS fear) and “To

189 what extent did you expect an electrical stimulation while looking at this figure?" (US
190 expectancy). Participants responded verbally on a 9-point Likert scale. Numbers 1, 3, 5, 7, and 9
191 of this scale carried a response label that was presented right above the number (with 1 = none
192 at all/certainly not; 3 = very little/rather not; 5 = uncertain; 7 = quite some/to some extent; 9 =
193 very much/most certainly).

194 *Procedure.* The experimenter attached the electrodes to the participant, who was positioned
195 on the scanner table, right before the table was inserted into the scanner. Next, the tolerance
196 level of the pain stimulus was determined for each participant individually, by means of an
197 adapted interleaved staircase procedure. This procedure consisted of 20 trials where USs were
198 presented and participants rated the subjectively experienced pain intensity on a scale from zero
199 (not painful at all) to ten (extremely painful). In order to increase reliability of the threshold
200 procedure, the 20 trials were divided into two separate sequences of ten trials differing in
201 current amplitude on their first trial, which was randomly drawn from either .5 to 1.0 mA, or 1.0
202 to 1.5 mA, respectively. After the first trial of each sequence, the current amplitude of the
203 following trial depended on the participant's rating in the previous trial of the respective
204 sequence. If the rating was below five, current amplitude would increase by 0.1 mA in the
205 following trial of that sequence, whereas it would decrease by 0.1 mA in case of a rating above
206 five, or stay the same if the rating equaled five. Thus, ratings from both sequences would
207 approach five on the rating scale for each participant. The two sequences were presented
208 intermixed and the final electrical current amplitude was then calculated as the mean of the final
209 values from both sequences. Participants were instructed to rate an intensity five when it was
210 experienced as unpleasant, but not intolerable, and informed that the pain stimuli used
211 throughout the experiment would not surpass this value.

212 After this preparation phase, the participants were instructed to wait for five minutes,
213 during which an anatomical scan would be administered. Upon completion of the anatomical
214 scan, the participants were presented with the instructions. Participants were informed that three
215 fractal figures (see Figure 1a) would appear successively for 8 seconds each. Following the

216 presentation of all three fractals on a white background, the participants were instructed that two
217 of the fractals would sometimes be followed by an electrical stimulation, whereas the third
218 fractal would never (in capital letters) be followed by an electrical stimulation. Subsequently,
219 participants were informed that these fractal-pain contingencies would be clearly displayed and
220 were encouraged to closely attend to these contingencies. After this instruction, a slide
221 containing both CS⁺ fractals and the text “+ electrical stimulation!” was presented during 8
222 seconds. This was followed by another 8 second presentation of a slide containing the CS⁻
223 fractal and the text “This figure will never be followed by a stimulation”.

224 Following these general instructions, the participants were informed that they would first
225 be allowed to familiarize themselves with the stimuli and the procedure in an initial training
226 phase. The training phase was said to be very similar to the test phase that would follow, except
227 that some of the electrical stimulations would be replaced by a picture of a lightning bolt (i.e.,
228 the placeholder stimulus, see Figure 1a). Participants were told that this was to prevent them
229 from getting too many pain stimuli before the real experiment actually started and asked to keep
230 in mind that whenever a lightning bolt would be presented, this meant that in the actual test
231 phase, a real electrical stimulation would occur. This was followed by a slide presenting the
232 CS⁺P (with electrical stimulation) and CS⁺U (with lightning bolt) contingencies during 8
233 seconds. The final page of instructions informed participants that they would be asked to
234 perform fear and US expectancy ratings at regular intervals during the upcoming phase. They
235 were told that no stimuli would be administered during the ratings and asked to remember the
236 most recent encounter with the fractals while answering the questions.

237 The actual training phase consisted of 27 conditioning trials (9 for each CS) interspersed
238 with blocked ratings. Each conditioning trial started with a 4 second presentation of a fixation
239 cross followed by a CS presentation for 8 seconds, with an inter-trial interval of 13, 15, or 17
240 seconds (see Figure 1a). On reinforced trials, the US or placeholder was presented at CS⁺ offset.
241 The US was presented for 300 ms. The placeholder remained on screen for a duration of 500
242 ms. The CSs were presented in “triplets” of three CS presentations so that each CS-type had

243 been presented once before the next triplet started. Trial order was randomized within triplets.
244 Blocked ratings of fear and US expectancy were presented after 9, 18 and 27 conditioning trials
245 (3, 6 and 9 triplets) respectively. As such, three mini-blocks were created within the training
246 phase, each containing 3 trials of CS⁺P, CS⁺U and CS⁻ presentation. Three out of nine CS⁺P and
247 CS⁺U presentations were reinforced during the training phase. For half of the participants, the
248 first, third, and penultimate presentation of the CS⁺P was followed by the US and the first,
249 second, and last presentation of the CS⁺U was followed by the placeholder. The other half of the
250 participants had counterbalanced reinforcement schedules (e.g., the first, second and last
251 presentation of the CS⁺P would be followed by the US).

252 Each block of ratings contained six ratings (two for each CS). The order of rating trials
253 within each rating block was fully randomized. However, due to technical difficulties, the CS
254 presentation and question type were independently randomized within each rating block for the
255 first five participants, resulting in repeated measurements or empty cells for some of the
256 questions. For this reason, the rating analyses are reported for the last 15 subjects only. Brain-
257 behavior correlation analyses, however, which focus on averaged CS ratings for an entire phase
258 (across the three rating blocks), were possible to perform on the entire set of participants. Before
259 the start of each rating block, it was stressed that by electrical stimulation (expectancy), we
260 referred solely to real electrical stimulations (i.e., not placeholders).

261 After the training phase, participants again received on-screen instructions. They were
262 informed that now the test phase would start, meaning that all electrical stimulations would be
263 presented for real and no placeholders would be used. Participants were instructed that the test
264 phase would evolve similarly to the training phase in all other respects. The course of the test
265 phase was very similar to that of the test phase, with 27 trials and 3 rating blocks in between.
266 However, no USs or placeholders were presented during this phase.

267 **Stimuli and Procedure: Experiment 2**

268 *Stimuli.* The stimuli for Experiment 2 were gray-scaled images selected from a database of
269 252 images of faces and 252 images of houses as previously used by Muhle-Karbe and

270 colleagues (2016). For each subject separately, 128 pictures (64 from each category) were
271 randomly selected and assigned to sixty-four pairs of pictures with the only restriction that both
272 pictures should be from the same category. When the category was faces, we further assured
273 that both pictures were from the same gender (and an equal amount of pairs were formed per
274 gender), to avoid that participants could dissociate the pictures based on gender. Per pair, one
275 picture served as a conditioned stimulus (CS) that would never be followed by a electrical
276 stimulation (CS⁻), while another served as a CS that would always be followed by a electrical
277 stimulation (CS⁺). While eight pairs of CS⁻ and CS⁺-pictures would re-occur throughout the
278 experiment (four pairs of houses, two pairs of male faces, and two pairs of female faces), the 56
279 remaining pairs only appeared once, hereafter referred to as "old" and "new" stimuli,
280 respectively. The electrotactile stimulus (US) consisted of the same sequence of pulses, and was
281 applied to the same location, as in Experiment 1. Different from Experiment 1, however, the
282 intensity could either be low or high (see below). The instructions would indicate the intensity
283 of the potentially following electrotactile stimulus by showing a grey intensity meter which
284 either pointed to the left or the right indicating a low or high intensity, respectively (see Figure
285 8a).

286 *Procedure.* Before entering the scanner, the participants were shown an example trial of the
287 experiment (without electrotactile stimulation) and were instructed about the general procedure
288 of the experiment. Namely, participants were informed that they would encounter several
289 instruction screens where two pictures were presented above one another on the left side of the
290 screen and an intensity meter on the right side of the screen, next to one of the two pictures (see
291 Figure 8a). It was further explained to the participants that after these instruction screens, one of
292 the two pictures would be presented (i.e., CS presentation) in the center of the screen. If this
293 picture was presented next to the intensity meter during the instructions, they would receive an
294 electrotactile stimulation shortly afterwards. The intensity of this stimulation was dependent on
295 which direction the intensity meter pointed to. To ensure that participants paid attention to the
296 task, one out of eight CS presentations (or one out of four in the practice block) were replaced

297 by a catch question where participants were shown either one of the two CSs, or a third picture
298 that they had never seen before (of the same category). On these trials, their task was to indicate
299 whether this picture was instructed to be followed by a stimulation, not followed by a
300 stimulation, or never presented before (all participants performed above chance level on these
301 catch questions; mean = 87.1%, SD = 10.6 %, minimum = 67%, maximum = 100%). Last, on
302 some trials participants would also see a centrally presented intensity meter, which would
303 indicate that an electroactile stimulus could follow with a 50% probability. These trials are
304 hereafter referred to as control trials. Briefly, this last condition was designed to control for
305 general US expectancy effects during instruction presentation, but is not important for the
306 current focus of analysis.

307 Next, participants were placed on the scanner table and after attaching the electrode, the
308 tolerance level was determined for each subject separately. The two intensities were determined
309 through a work-up procedure where gradually increasing current amplitudes were presented to
310 the participant. Participants were asked to determine which amplitude was the first noticeable,
311 and when we should stop increasing the amplitude, upon which the work-up procedure
312 automatically ended. The first noticeable current level was used as the low intensity, the last
313 tolerable as the high intensity. We assured subjects that only those two intensities could be used
314 in the remainder of the experiment.

315 After the anatomical scan, we presented the instructions to the participant once more. They
316 were further informed that they would receive a practice block half the length of the following
317 three experimental blocks. This practice block was to ensure that participants were familiarized
318 with all the "old" pairs, and was not included in the analyses.

319 During each block, the participant was presented with an equal amount of all three possible
320 trial types: old instructed trials, new instructed trials, and control trials. The general structure of
321 each of those trial types was that they started with 2500 ms instruction presentation, followed by
322 a 2000 to 4200 ms instruction-CS interval, a 1000 ms CS presentation, 2000 to 4200 ms CS-US
323 interval, a 300 ms US presentation (if the CS was a CS⁺), and, finally, a 2500 to 4700 ms inter-

324 trial interval. For instructed trials, instruction presentation consisted of a presentation of the CS⁻
325 and CS⁺ picture above one another on the left side of the screen (location of CS-type was
326 randomly determined per trial), and an intensity meter next to the CS⁺ picture. On control trials,
327 instruction presentation consisted of a centrally presented intensity meter. CS presentation on
328 instructed trials consisted of one of the two CSs (CS⁻ or CS⁺) centrally presented on screen.
329 During control trials, this was replaced by the presentation of a central fixation cross.

330 Each test block consisted of 16 new instructed trials, 16 old instructed trials (each specific
331 pair was presented twice per block), and 16 control trials. All trial types were further subdivided
332 in eight high intensity and eight low intensity trials. The instructed trial types also showed an
333 equal amount of faces and houses (and an equal amount of male and female pictures among the
334 faces). The different trials were presented in a random order.

335 **fMRI data acquisition**

336 In both experiments, participants were positioned headfirst and supine and instructed not
337 to move their heads to avoid motion artefacts. Images were collected using a 3T Magnetom Trio
338 MRI scanner system (Siemens Medical Systems, Erlangen, Germany) with a standard thirty-
339 two-channel radio-frequency head coil. First, a 3D high-resolution anatomical image of the
340 whole brain was acquired for co-registration with the functional images using a T1-weighted 3D
341 MPRAGE sequence (TR = 2530 ms, TE = 2.58 ms, TI = 1100 ms, acquisition matrix = 256 ×
342 256 × 176, sagittal FOV = 220 mm, flip angle = 7°, voxel size = 0.9 × 0.86 × 0.86 mm). Next,
343 whole brain functional images were collected using a T2*-weighted EPI sequence, sensitive to
344 BOLD contrast (TR = 2000 ms, TE = 28 ms, image matrix = 64 × 64, FOV = 224 mm, flip
345 angle = 80°, slice thickness = 3 mm, distance factor = 17%, voxels resized to 3.0 × 3.0 × 3.0
346 mm, 34 axial slices). The number of images per run varied depending on response speed during
347 the rating blocks (Experiment 1) or catch questions (Experiment 2).

348 **Experimental Design and Statistical Analysis**

349 *Behavioral data analyses for Experiment 1.* We carried out two ANOVAs with the three
350 within-subjects factors phase (training versus test), CS type (CS⁺P, CS⁺U and CS⁻), and block

351 (first, second, or third rating block) for each rating scale separately. Further ANOVAs focused
352 on specific CS contrasts (e.g., CS⁺P vs. CS⁺U) to allow for a more detailed picture of the
353 significant interactions observed in the main ANOVA. Due to incomplete data collection for the
354 first five subjects, we performed all behavioral ANOVAs on fifteen subjects. Analyses
355 excluding the factor block allowed us to study nineteen subjects and all twenty subjects for the
356 fear and US expectancy data (one subject did not have any fear ratings for the CS-),
357 respectively. Including these subjects did not change the significance of our findings, but,
358 naturally, are limited to analyses excluding the factor block. Therefore, we will only report the
359 overall ANOVA. We did, however, use these general ratings (across blocks) to investigate
360 brain-behavior correlations.

361 *Introduction to Representational Similarity Analyses.* A traditional mass univariate voxel-
362 wise comparison of CS-related activations in canonical fMRI analyses can inform about gross
363 differences in the recruitment of brain regions at the macroscopic scale but is blind to
364 differences in the recruitment of separable neural ensembles *within* a brain region. More
365 recently, multivariate multi-voxel pattern analysis (MVPA) techniques have permitted to
366 investigate intra-regional spatial patterns of neuronal activation (Haxby et al., 2001;
367 Kriegeskorte et al., 2008; Norman et al., 2006), including in fear conditioning (Bach et al.,
368 2011; Dunsmoor et al., 2013; Hauner et al., 2013; Li et al., 2008; Visser et al., 2011; 2013), with
369 high sensitivity, often going beyond conclusions that can be derived from univariate analyses
370 (notably, the latter did not show any significant differences in CS⁺P versus CS⁺U activity during
371 the test phase, neither in a whole-brain corrected contrast, nor when restricting the analysis to
372 any of the below-defined regions). We therefore investigated (dis)similarities in neural
373 processing of different CSs or of the same CS at different time points of the experiment by
374 extracting and comparing multi-voxel activation pattern data from previously selected regions
375 of interest (ROIs) event-locked to the presentation of the different CSs (Kriegeskorte et al.,
376 2008).

377 *fMRI data analyses.* Data processing and analyses were performed using the SPM8 Matlab-
378 package software (<http://www.fil.ion.ucl.ac.uk/spm/software/spm8/>). The first four volumes of
379 each run in which no stimulation occurred were discarded before estimating statistical models.
380 Anatomical images were spatially normalized to the SPM T1-template image and resliced to a
381 voxel size of 1 x 1 x 1 mm. Functional data were slice-time corrected and spatially realigned to
382 the first volume of the task. Next, EPIs were spatially normalized based on the T1-derived
383 normalization parameters and a temporal high-pass filter of 128 s was applied to remove low-
384 frequency drifts. No spatial smoothing was used.

385 In Experiment 1, BOLD responses were modelled with boxcar functions at CS onset till CS
386 offset (eight seconds) or with delta functions for all other regressors, which were then
387 convolved with a standard hemodynamic response function (HRF). Event-related regressors
388 were created corresponding to the onset of the CS and defined by CS-type (CS⁻, CS^{+P}, or
389 CS^{+U}), to the onset of US and defined by US-type (electrotactile stimulation or placeholder), or
390 to the onset of a ratings question. Specifically, the model included three (or more, see below)
391 regressors denoting the CS-type, one for the ratings, two for the US-type (in the training phase
392 only), and six movement parameters derived from the realignment procedure, for the two runs
393 separately (which corresponded with the training and test phase, respectively). The statistical
394 parameter estimates were computed separately for each voxel for all columns in the design
395 matrix. Three different first-level models were fitted. The CS-type regressors per phase were
396 either further split up in a single regressor for each trial separately (i.e., the trial-based model), a
397 regressor for each mini-block separately (i.e., the mini-block model), or a regressor per phase
398 (i.e., the phase model). We analyzed the resulting data by computing pair-wise Pearson
399 correlations between all CS event-related spatial patterns of activation. For a visual depiction of
400 the resulting similarity matrices from these three types of models in the anterior cingulate cortex
401 (ACC), see Figure 2. The strength of these correlations was used as a metric of similarity. The
402 correlations were Fisher-transformed for statistical analyses. Different types of correlations
403 were selected for the analyses of interest and analyzed in ANOVAs using Statistical Package for

404 the Social Sciences (version 22, SPSS). Each analysis started with an overall ANOVA including
405 the factor region, whose six levels corresponded to the six regions of interest (ROIs) identified
406 below. Only when effects interacted with the factor region, the effects were further studied for
407 each region separately. All reported p -values are two-sided and a Greenhouse-Geisser correction
408 was applied whenever the assumption of sphericity was violated, but uncorrected degrees of
409 freedom are reported for ease of reading.

410 Task events in Experiment 2 were similarly modelled as in Experiment 1, only now we also
411 added separate regressors for instruction presentations (also modelled with boxcar functions
412 from instruction onset till instruction offset, 2500 milliseconds). In total, 31 event-related
413 regressors were created. Eight regressors corresponded to the onset of instruction presentation
414 and were defined by instruction-type (coding for either old versus new instructions, house
415 versus face stimuli, and low versus high US intensity), sixteen to the onset of CS presentation
416 defined by CS-type (similarly determined by the same conditions as instruction-type and further
417 coding for being a CS⁻ or CS⁺ presentation), four to the onset of control instructions and the
418 subsequent fixation cross presentation (further determined by low or high intensity), two to the
419 onset of US presentations and defined by being either of low or high intensity, and, finally, one
420 to the onset of catch questions. Although instruction presentations, US presentations, and catch
421 trials were modelled separately, they were not included in our contrasts and considered
422 regressors of non-interest. For the analysis of this dataset, we only focused on high intensity
423 trials, where the CS⁺ was instructed and expected to be followed by a high intensity electrical
424 stimulation. Specifically, we focused on voxel patterns evoked by CS presentation in any of the
425 six below-defined ROIs. We again analyzed the resulting data by computing Fisher-transformed
426 pair-wise Pearson correlations. In particular, we were interested in the correlations between
427 house and face presentations of the same trial type (see Figure 8b). This way, we created four
428 possible correlations of interest: the correlations between multi-voxel patterns of activity to face
429 presentation versus house presentation, for CS⁻old trials, CS⁺old trials, CS⁻new trials, and
430 CS⁺new trials, separately. These values were interpreted as the degree to which a certain region

431 showed a response independent of visual category. Next, these values were analyzed using two
432 by two ANCOVAs with the factors fear relevance (CS⁺ versus CS⁻) and novelty (old versus
433 new), and, to control for differences in preferred US intensity, the standardized covariate US
434 intensity (baseline-corrected by dividing the high intensity value by the low intensity value), for
435 each ROI separately.

436 *ROI analyses.* We extracted β estimates from the separate voxels of anatomically defined
437 ROIs, known to be relevant in fear conditioning (the same fear related regions as were used in
438 Visser et al., 2013). Specifically, in addition to the amygdala, we focused on the anterior
439 cingulate cortex (ACC), superior frontal gyrus (SFG), insula, and ventromedial prefrontal cortex
440 (vmPFC), areas previously implicated in fear conditioning (Visser et al., 2013; Fullana et al.,
441 2015; Mechias et al., 2010). We extracted the mean voxel activation from those regions, as
442 obtained from the Harvard-Oxford cortical and subcortical structural atlases (Harvard Center for
443 Morphometric Analysis), thresholded at 25%, and included a factor "region" in each of our
444 analyses to identify potential between-region differences.

445

446 Results

447 Experiment 1: Rating data

448 In the rating data, we tested whether we could replicate the behavioral results by Raes and
449 colleagues (2014), and Mertens and colleagues (2016), by analyzing the mean fear and US
450 expectancy ratings per phase and block. There was a clear main effect of CS type for both
451 ratings, both $ps < .001$, which interacted with phase, both $ps < .001$ (see Figure 1b and 1c).
452 Marginal significant three-way interactions between phase, CS type, and block for both the fear,
453 $F(4, 11) = 3.4, p = .050$, and US expectancy ratings, $F(4, 11) = 2.7, p = .088$, hinted at a
454 differential evolution of CS ratings over time depending on the phase.

455 Next, we investigated these interactions for each CS type comparison and phase separately,
456 by running separate block \times CS type ANOVAs. In the training phase, both the CS⁺P and CS⁺U
457 elicited a higher US expectancy and fear rating, relative to the CS⁻; all $ps < .001$. Moreover, the

458 CS⁺P elicited higher ratings on both scales relative to the CS⁺U, both $ps < .005$. In the testing
459 phase, again, both the CS⁺P and CS⁺U elicited higher ratings relative to the CS⁻, all $ps < .001$.
460 The CS⁺P, however, was no longer significantly different from the CS⁺U, on both the fear, $F(1,$
461 $14) = 1.3, p = .272$, as well as the US expectancy scale, $F(1, 14) = 1.2, p = .277$, suggesting
462 similar fear responses.

463 For all CS⁺ to CS⁻ comparisons, on both phases, there was a significant interaction between
464 CS type and block, all $ps < .001$, suggesting that CS⁺ ratings did, while CS⁻ ratings did not,
465 decay over time, irrespective of the phase (see Figures 1b and 1c). Interestingly, however,
466 similar interactions between CS type and block between CS⁺P and CS⁺U were absent in the
467 training phase, $F_s < 1.7, ps > .21$, but present in the testing phase, with a significant interaction
468 for the fear ratings, $F(2, 13) = 12.0, p = .001$, and a marginally significant interaction in the US
469 expectancy data, $F(2, 13) = 1.2, p = .052$. These interactions demonstrated that while there was
470 no overall difference between both CS⁺ ratings in the test phase, there were some initial
471 differences in the first mini-blocks of the test phase (see Figures 1b and 1c). Specifically, the
472 fear rating for CS⁺P relative to CS⁺U was significantly higher in the first block, $t(14) = 2.22, p =$
473 $.044$, marginally significant in the second block, $t(14) = 1.87, p = .082$, and absent in the third
474 block, $t(14) = .76, p = .458$. The US expectancy rating for CS⁺P relative to CS⁺U was marginal
475 significantly higher in the first block, $t(14) = 1.79, p = .095$, but not significantly different in the
476 second or third block, $t(14) = 1.08, p = .301, t(14) = 0.52, p = .610$, respectively.

477 In sum, these results clearly replicate the results by Raes and colleagues (2014) and
478 Mertens and colleagues (2016). Most importantly, the CS⁺P elicited slightly but reliably higher
479 fear ratings than the CS⁺U, especially at the beginning of the test phase (Figure 1; for similar
480 findings in fear potentiated startle responses, but not skin conductance responses, see Mertens et
481 al., 2016; Raes et al., 2014). This indicates a dissociable contribution of prior actual CS-US
482 pairings to the fear reaction to the CS⁺P in the testing phase.

483 **Similarity analyses: Experiment 1**

484 Different pattern similarity analyses are reported to allow for a comprehensive picture of
485 the data. Importantly, each of those analyses were motivated by specific hypotheses, which are
486 detailed below when discussing each analysis separately. We will first discuss an analysis that
487 tested whether the different CSs were responded to similarly within the training and test phases.
488 That is, did the ROIs respond more similarly to the CS⁺P and the CS⁺U, than, for example, the
489 CS⁺P and the CS⁻? Thereafter, we report an analysis that looked at the internal consistency of the
490 separate patterns to each CS within a phase. Namely, did certain regions respond in a more
491 consistent manner to one CS as opposed to another CS? Third, and most importantly, we tested
492 whether the pattern response to CS⁺P during the training phase was a better predictor of itself
493 during the test phase, than it was to CS⁺U or CS⁻. As a post-hoc analysis following up on this
494 hypothesized result, we also tested whether regions that show this differential processing of the
495 CS⁺P also show a relation with the difference in fear ratings. That is, we wanted to explore
496 whether the difference in subjective fear experience as observed in the present study (replicating
497 previous findings by Mertens et al., 2016; Raes et al., 2014) can be linked back to a neural trace
498 of experience-based fear. Last, we will report a targeted pattern-informed connectivity analyses
499 by investigating parallels in trial-to-trial pattern similarities between the ACC, and left and right
500 amygdala, depending on CS type, as well as a broader connectivity analysis involving all six
501 regions. Note that every analysis started with an omnibus ANOVA that included the factor
502 region, to detect between-region differences.

503 *Inter-CS similarities per phase.* We first confirmed that the selected ROIs process learned
504 stimulus qualities (rather than merely processing the perceptual properties of the CSs).
505 Specifically, we observed that the trial-averaged multi-voxel activation patterns evoked by the
506 CS⁺P and the CS⁺U were more similar to each other (CS⁺P-CS⁺U inter-CS similarity, green bars
507 in Figure 3) than each of them was to the trial-averaged patterns evoked by the CS⁻ (comparison
508 with CS⁻-CS⁺P inter-CS similarity: $F(1,19) = 11.08, p = .004$, blue bars in Figure 3; with CS⁻-
509 CS⁺U inter-CS similarity: $F(1,19) = 18.59, p < .001$, red bars; for statistical procedures, see
510 Method). This was the case during both training and test (effects of phase: both $F_s < 1$), but,

511 intriguingly, the effect differed significantly between regions (both $ps < .004$). Specifically in
512 the left and right amygdalae, the CS⁺P, which is the only CS whose acquired qualities result (in
513 part) from experience, evoked activation patterns that were not more similar to CS⁺U patterns
514 than to CS⁻ patterns (both $F_s < 1$). This analysis suggests that while participants were instructed
515 to treat the CS⁺P and CS⁺U similarly (and most fear-related regions also seem to reflect this),
516 the amygdala could be sensitive to whether the fear value results from experience, and therefore
517 did respond to them differently. This difference in responding was even comparable to the
518 difference in responding to CS⁺P versus CS⁻. Whereas we cannot prove that this effect is not a
519 floor effect, we would like to note that the other analyses below do show reliable differences in
520 the amygdala (sometimes even exclusively in the amygdala), speaking against the idea that
521 activity in the amygdala regions was simply too variable or too noisy to detect reliable
522 differences.

523 *Intra-CS similarities per phase.* Areas responsible for processing acquired as opposed to
524 mere perceptual stimulus qualities are thought to exhibit more consistent activation patterns to
525 each of the CS⁺s than to the CS⁻ from one trial to the next of the experiment (Visser et al., 2011;
526 2013). However, note that one could also expect the opposite. Namely, regions responsive to
527 dynamic trial-to-trial changes in a certain condition might show a smaller internal consistency.
528 Therefore, these analyses only allow us to conclude that regions which show differences in
529 these internal consistency measures between conditions must respond to these two conditions
530 differently. Such a result would be another confirmation that such regions encode acquired
531 stimulus qualities. For simplicity and robustness, we grouped trials into three mini-blocks per
532 phase and computed intra-CS similarities from one mini-block to the next. As expected, intra-
533 CS⁺P similarity was significantly higher than intra-CS⁻ similarity (main effect of CS type:
534 $F(1,19) = 18.11, p < .001$; Figure 4), and this difference decreased in the test phase relative to
535 the training phase, consistent with the observed extinction in fear ratings in this phase ($F(1,19)$
536 $= 4.49, p = .047$; no interactions with region, $F_s(5,95) < 1.95, ps > .135$). In stark contrast, the
537 intra-CS⁺U versus intra-CS⁻ similarity analysis (main effect of CS type: $F(1,19) = 7.31, p =$

538 .014) exhibited an interaction effect with region ($F(5,95) = 2.82, p = .020$; but no interactions
539 with phase, $F_s < 1$). The amygdalae were the only ROIs not showing significantly higher intra-
540 CS⁺U than intra-CS⁻ similarities (both $F_s(1,19) < 1.25, p_s > .278$). That is, an analysis based on
541 the temporal consistence of neural activation patterns (Visser et al., 2011; 2013) found no
542 evidence that the amygdala processes the learned qualities of a merely instructed stimulus, the
543 CS⁺U.

544 Although all our analyses reflect between-region differences in different forms of fear
545 processing between six identified fear-related ROIs, one could also investigate whether
546 completely unrelated ROIs show a fear response. Although it is hard to select completely
547 unrelated regions, we investigated whether the superior temporal gyrus (i.e., auditory cortex)
548 shows a similar main effect of fear conditioning (as suggested by an independent reader).
549 Specifically, we zoomed in on the most reliable effect across all cortical fear-related regions: the
550 enhanced correlation between patterns of responses to the instructed and experienced stimulus
551 (CS⁺P) versus the neutral stimulus (CS⁻). While this effect did reach significance in each of
552 those regions, even after correcting for multiple comparisons, it did not reach significance in the
553 superior temporal gyrus, $F(1,19) = 2.52, p = .129$.

554 *Inter-CS similarities across phase.* Both the CS⁺P and the CS⁺U, but not the CS⁻, carry a
555 representation of threat. Fear-related regions should therefore show CS⁺ evoked activation
556 patterns that are more similar to each other than to CS⁻ evoked activation patterns. Intriguingly,
557 the above set of similarity analyses indicated that the amygdala's neural activation pattern to the
558 CS⁺P was not more similar to the CS⁺U than it was to CS⁻, whereas all other fear-related regions
559 did show a higher similarity between their responses to CS⁺P and CS⁺U than between any of the
560 two CS⁺s and the CS⁻ (Figure 3). Moreover, while all fear-related regions, including the
561 amygdala, showed a higher consistency (within phase, but across mini-blocks) in their neural
562 pattern response to the CS⁺P than the CS⁻, only the non-amygdala fear-related regions also
563 showed a higher consistency in their response to the CS⁺U than the CS⁻ (Figure 4). Together,

564 these results are already suggestive of the idea that the amygdala is involved differently in the
565 processing of an instructed and experienced versus a merely instructed fear contingency.

566 However, the central question in our analyses was whether brain regions involved in fear
567 learning would respond differently to both CS⁺s in the test phase, despite the fact that the same
568 verbal information was given about both CS⁺s. A neural trace of experience-based Pavlovian
569 fear learning should be apparent from similarities between CS evoked activation patterns during
570 test and the activation pattern evoked by the CS⁺P during training (CS⁺Ptr), i.e., where the
571 contingency was experienced and the Pavlovian fear memory was formed. Specifically, one
572 should expect higher similarities between CS⁺Ptr patterns and the patterns evoked by the same
573 CS during testing (i.e., CS⁺Pte). If those similarities were larger than between CS⁺Ptr and
574 CSU⁺te (as well as between CS⁺Ptr and CS⁺te), they would indicate a Pavlovian memory trace.
575 By contrast, if CS⁺PtrCS⁺Pte similarities were no more pronounced than CS⁺PtrCS⁺Ute
576 similarities, this would indicate a more generalized representation of threat during testing that
577 does not retain a specific experience-based memory element. In this critical analysis, we
578 observed a significantly higher similarity between CS⁺Ptr and CS⁺Pte (CS⁺PtrCS⁺Pte) as well as
579 CS⁺Ptr and CS⁺Ute (CS⁺PtrCS⁺Ute), than between CS⁺Ptr and CS⁺te (CS⁺PtrCS⁺Pte vs.
580 CS⁺PtrCS⁺te similarity: $F(1,19) = 19.36, p < .001$; CS⁺PtrCS⁺Ute vs. CS⁺PtrCS⁺te similarity:
581 $F(1,19) = 14.27, p = .001$; Figure 5). Importantly, CS⁺PtrCS⁺Pte did not differ from
582 CS⁺PtrCS⁺Ute pattern similarity, $F(1,19) = .397, p = .536$. These results suggest both CS⁺s
583 evoked similar threat-related processing during test. However, there was an interaction with
584 region ($F(10,190) = 3.32, p = .007$; also when comparing the left and right amygdala only:
585 $F(2,18) = 3.31, p = .047$). Namely, the right amygdala exhibited CS⁺Pte patterns that were
586 significantly more similar to CS⁺Ptr patterns than were both CS⁺Ute and CS⁺te patterns
587 (CS⁺PtrCS⁺Pte vs. CS⁺PtrCS⁺Ute similarity: $t(19) = 2.204, p = .040$; CS⁺PtrCS⁺Pte vs.
588 CS⁺PtrCS⁺te similarity: $t(19) = 3.990, p = .001$; Figure 5f). The CS⁺Ute and CS⁺te patterns did
589 not differ in their similarity to the CS⁺Ptr pattern, $t(19) = 1.013, p = .324$. Importantly, in all
590 other regions, the patterns of CS⁺Pte and CS⁺Ute were not significantly different in their

591 similarity to the pattern of CS⁺Ptr, all $r_s(19) < 1$. Hence, this analysis isolated a threat-related
592 neural response during testing in the right amygdala that was exclusively evoked by the CS⁺P,
593 as opposed to the CS⁺U, meeting our criterion for a Pavlovian trace of actually experienced CS-
594 US pairings. Other regions appeared to register a merely instructed threat (CS⁺U) in the same
595 way as a threat that is not only instructed but also previously experienced (CS⁺P).

596 The observation that activation in the right amygdala is more similar for CS⁺Ptr and CS⁺Pte
597 than for CS⁺Ptr and CS⁺Ute might also reflect the fact that CS⁺Ptr and CS⁺Pte are visually more
598 similar than CS⁺Ptr and CS⁺Ute. However, if this alternative explanation is correct, then
599 activation for CS⁺Utr and CS⁺Ute should also be more similar than activation for CS⁺Utr and
600 CS⁺Pte. No such difference was observed, $t(16) = .727, p = .476$. In fact, the similarity between
601 CS⁺Ptr and CS⁺Pte was higher than that between the CS⁺Utr and CS⁺Ute, $t(19) = 3.502, p =$
602 $.002$.

603 Further supporting the encoding of this Pavlovian fear trace in the right amygdala, we
604 observed a relation between CS⁺P specific right amygdala neural responses and the observed
605 differential (CS⁺P > CS⁺U) fear response, in that the difference between CS⁺PtrCS⁺Pte and
606 CS⁺PtrCS⁺Ute pattern similarities as depicted in Figure 5f predicted the difference in CS⁺Pte
607 and CS⁺Ute fear ratings during test across participants (*Spearman's* $\rho = .465, p = .039$; Figure
608 5g). This post-hoc analysis should of course be treated with caution because our study (and its
609 sample size) was not in first instance set up to study inter-subject correlations.

610 *Inter-region similarities in Intra-CS similarities.* If the right amygdala processes
611 experience-based threat in a way that can be dissociated from its processing of instruction-based
612 threat, it might also preferentially exchange that information with other threat areas.
613 Specifically, we wondered whether CS⁺P related functional connectivity of the right amygdala
614 with the ACC would differ from that of the left amygdala, for comparison. The ACC is the
615 region that is most prominently and consistently activated during fear conditioning studies
616 (Fullana et al., 2015; Mechias et al., 2010) and also exhibits strong structural and functional
617 connectivity with the amygdalae (e.g., Bissière et al., 2008; Carlson et al., 2013; Etkin et al.,

618 2006; Van Marle et al., 2010; Williams et al., 2006; for a review, see Kim et al., 2011). To this
619 end, we opted to carry out voxel-pattern-informed connectivity analyses, which have recently
620 been demonstrated to be more sensitive and reliable than standard connectivity analysis
621 (Geerligs & Henson, 2016). To perform a similarity-based functional connectivity analysis, we
622 used the trial-based model (Figure 2a) and Spearman correlated the trial-by-trial intra-CS
623 similarities per CS and phase (Figure 6a) between regions for each subject separately, thereby
624 indexing the similarities between these three regions in how a CS pattern relates to itself across
625 time (Figure 6b-c, Kriegeskorte et al., 2008). We found that intra-CS⁺P relative to intra-CS⁺U
626 similarity time courses from the right, but not the left amygdala, showed a higher correlation
627 with corresponding ACC time courses (interaction between CS type and amygdala side: $F(1,19)$
628 $= 5.68, p = .028$; follow-up comparison of correlations between intra-CS⁺P and intra-CS⁺U
629 similarity time courses from right amygdala and ACC: $t(19) = 2.698, p = .014$; from left
630 amygdala and ACC: $t(19) = .767, p = .453$; Figure 6b-c). These results further corroborate the
631 conclusion that the right amygdala reacts differently to experienced and instructed fear, but also
632 point towards an extended role of a larger network (Okon-Singer et al., 2015; Pessoa &
633 Adolphs, 2010), by showing an increased functional connectivity with the ACC for
634 communicating this experience-based component of fear learning.

635 To further illustrate this idea, and allow for a more comprehensive picture of the data, we
636 also computed inter-region similarity matrices depicting each possible region-to-region
637 connectivity, for each CS-type separately (see Figure 7a). As a point of reference, we further
638 included two occipital regions, namely the lateral occipital cortex (LO) and occipital pole (Occ).
639 Multi-dimensional scaling analyses on Figure 7c depict the relations between the six fear-related
640 regions for both CS⁺s in the test phase. We used two-dimensional solutions (using PROXSCAL,
641 SPSS), as these offered the most optimal stress levels relative to the number of dimensions (i.e.,
642 the elbow in the scree plot). Most importantly, these analyses visualize how the connectivity
643 patterns change depending on the CS type processing. For example, Figure 7c suggests a more
644 integrated role for the right amygdala when it comes to processing the CS⁺P. In fact, when

645 comparing the overall connectivity between all six fear related regions per CS (averaging all 15
646 possible connections; Figure 7b), it appeared to be enhanced for CS⁺Pte relative to CS⁺Ute,
647 $t(19) = 2.633, p = .016$. When testing this for each region separately (its average connectivity
648 with all other five regions for CS⁺Pte relative to CS⁺Ute), the only two regions showing
649 significantly stronger connectivity during CS⁺Pte processing were the right amygdala and ACC,
650 $t(19) = 2.780, p = .012, t(19) = 2.541, p = .020$, respectively (all other regions, $t(19) < 1.91, p >$
651 $.071$), again suggesting that these two regions and their interaction play an important role in the
652 learning or expressing of experience-based fear.

653 **Similarity analyses: Experiment 2**

654 In the second experiment, we used a different, but conceptually similar procedure to
655 investigate whether we could replicate the observation that the right amygdala (in comparison to
656 the left amygdala) dissociated between instructed and experienced (i.e., old) versus merely
657 instructed (i.e., new) fear contingencies. Moreover, this experiment employed stimuli belonging
658 to different visual categories (houses versus faces) across different trials of the same condition,
659 allowing us to study the similarity between pattern responses to the presentation of a house
660 versus a face as a CS. The results, shown in Figure 8, hinted at a main effect of fear relevance,
661 CS⁺ versus CS⁻, $F(1,17) = 4.00, p = .062$, suggesting that object category independence (i.e.,
662 house-face pattern similarity) indexed activation of a threat representation. More importantly,
663 there was a three-way interaction between CS type, CS novelty, and amygdala side, $F(1,17) =$
664 $5.38, p = .033$, that was qualified by a two-way interaction between CS type and novelty in the
665 right amygdala, $F(1,17) = 5.36, p = .033$, but not the left amygdala, $F(1,17) = .133, p = .720$.
666 More specifically, the right amygdala again differentiated between the processing of novel (i.e.,
667 merely instructed) and old (i.e., previously instructed and experienced) fear contingencies
668 (Figure 8c-h). Namely, the similarity between faces and houses was higher for CS⁺old than CS⁻
669 old, $t(18) = 2.653, p = .016$, but not for CS⁺new than CS⁻new, $t(18) = -.217, p = .831$. This result
670 further supports our conclusion that the right amygdala carries a trace of the CS-US contingency
671 experience made during Pavlovian fear conditioning. No other fear-related regions (displayed in

672 Figure 8 for illustrative purposes only) showed a similar CS type by novelty interaction (all $F_s <$
673 $1.782, p_s > .200$), again suggesting that the right amygdala was most sensitive in encoding a
674 Pavlovian trace.

675

676

Discussion

677 Demonstrating a purely experience-based element in human Pavlovian fear conditioning
678 and identifying its neural correlates has been a major goal of learning research over the past
679 decades. While the strong phenomenological and functional homologies between human and
680 non-human fear conditioning have always suggested language-independent processing in
681 humans, too, previous efforts have not yielded conclusive evidence. Our new approach is not
682 dependent on controversial methods to exclude verbal or conscious processing (Mitchell et al.,
683 2009) and exploits recent advances in the multivariate analysis of neural signatures of fear
684 learning and memory (Bach et al., 2011; Dunsmoor et al., 2013; Hauner et al., 2013; Li et al.,
685 2008; Visser et al., 2011; 2013). Because of these unique features, we are able to provide much
686 sought evidence for experience-based threat processing. Specifically, using pattern similarity
687 analyses, our two experiments demonstrated that the right amygdala was the only fear-related
688 region whose neural activation pattern to experienced CSs was different from its neural pattern
689 to merely instructed CSs. Of course, our data do not imply that the right amygdala does not
690 process instructed threat information. Rather, it appears to respond in a unique way, perhaps in
691 the form of specialized neural ensembles, to experience-based threat information. More
692 generally, the amygdala has also been implicated in other processes besides fear processing
693 (Okon-Singer et al., 2015; Pessoa & Adolphs, 2010). Therefore, future studies should determine
694 whether other separable experience-dependent neural traces in the right amygdala can be
695 identified for other forms of learning as well.

696 The present findings seem to converge on those of previous fear conditioning studies that
697 were set up to single out unconscious fear conditioning. However, as noted above, the present
698 study was not designed to study unconscious or implicit fear conditioning. Instead, the

699 manipulations in the present study were made very explicit: participants were very much made
700 aware of the instructions and the actual CS-US pairings. This way, our study tried to create
701 conditions for instructed fear learning to override a hypothesized experience-based component
702 to fear learning. Our results show that these instructions were successful in evoking a similar
703 neural response to the merely instructed as compared to the instructed and experienced stimulus
704 in most fear related regions, except in the right amygdala. It is possible that the present
705 experience-based trace in the right amygdala is related to the one identified in previous
706 unconscious fear conditioning studies. However, the present study cannot (and did not aim to)
707 prove the hypothesized unconscious nature of this experience-based trace (Öhman & Mineka,
708 2001). Therefore, it does not distinguish between an experience-based memory trace that is
709 generated fully automatically and unconsciously versus one that relies on conscious contingency
710 knowledge.

711 The present observation of apparent hemispheric differences in threat processing in the
712 amygdala also adds to another important question in fear research, more specifically, concerning
713 the presence of amygdala lateralization (Baker & Kim, 2004; Sergerie et al., 2008). Our results
714 clearly suggest that the right amygdala shows a separable neural response to actual CS-US
715 pairings, whereas the left amygdala appears to be more susceptible to fear instructions. These
716 findings are concordant with previous (lesion) studies suggesting that the right amygdala is
717 associated with a fear response to experiencing negative events, while the left amygdala is more
718 responsive to the verbally-mediated cognitive representation of fear (Funayama et al., 2001;
719 Phelps et al., 2001).

720 Last, our findings also fit well with those of another recent study by Atlas and colleagues
721 (2016). In this study, Atlas and colleagues used two CSs which were both predictive of the US,
722 but in different phases of the experiment. Crucially, they contrasted a condition in which this
723 reversal in contingencies was always instructed to a condition where it was not, and observed
724 that the amygdala was more responsive to the actual (changes in) contingencies rather than the
725 instructions that preceded those. Interestingly, their analyses did not show a hemispheric

726 difference, in contrast to earlier findings suggesting that the left amygdala can be responsive to
727 instructions (present results; Funayama et al., 2001; Phelps et al., 2001). However, as also
728 argued in their discussion, Atlas and colleagues (2016) focused on the effects of instruction in
729 changing environments and examined dynamic learning-related responses, whereas our study
730 employed a manipulation where the contingencies were not changing, and instructions were
731 given full opportunity to override the hypothesized Pavlovian trace.

732 In sum, across two experiments, we investigated neural pattern responses in fear related
733 regions to experience-based fear processing in the face of verbal fear instructions. Our results
734 show that verbal instructions were successful in evoking a similar neural response in fear-
735 related regions to merely instructed versus instructed and experienced fear stimuli, except for
736 the right amygdala. Instead, the human right amygdala showed a Pavlovian trace, suggesting it
737 to be more sensitive to the actual experience of CS-US contingencies.

738

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Author Contributions

740 S.B., J.D.H., R.K., & M.B. designed the experiment. S.B. & J.D. programmed and
741 conducted the experiments. S.B., J.D., K.S.L.Y., & R.K. analyzed the data. S.B., J.D.H., J.D.,
742 R.K., & M.B. wrote the paper.

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856 **Figure 1.** Procedure Experiment 1 and behavioral results. A. Experiment trials and rating screens. Each
857 trial started with the presentation of a fixation cross, followed by the presentation of a CS. In the training
858 phase, the CS⁺P was occasionally followed by a US (electrical stimulation), the CS⁺U by a US
859 placeholder (picture of a lightning bolt), and the CS⁻ was never followed by either a US or placeholder. In
860 the test phase, none of the CSs was followed by a US or placeholder. Rating screens assessed participant's
861 fear experience and US expectancy associated with each of the CSs. B. Experimental procedure and
862 instructions. Before training, subjects were instructed that only the CS⁺P could be followed by the US,
863 while the CS⁺U could only be followed by the placeholder. Before testing, subjects were told to expect
864 USs after both CS⁺s. Both phases consisted of three mini-blocks (where every CS was randomly
865 presented three times), each followed by a rating block. Three out of nine CS⁺ presentations were
866 followed by either the US or the US placeholder. C. Mean fear ratings. D. Mean US expectancy ratings.
867 The error bars are ± 1 standard error of the mean (SEM). tr=training; te=test.

868
869 **Figure 2.** An example of the different types of similarity matrices in the anterior cingulate cortex
870 (Experiment 1). The similarity matrices represent color-coded average Pearson correlation coefficients
871 across subjects, for every possible correlation between all different CS-presentation regressors in the trial-
872 based model (A), mini-block model (B), or the phase model (C). The vertical bars adjacent to each matrix
873 indicates its color coding depending on the correlation coefficient.

874
875 **Figure 3.** Inter-CS similarities per region and experimental phase (Experiment 1), based on the mini-
876 block model (see Figure 2). More similar multi-voxel activation patterns between the two CS⁺s than
877 between a CS⁺ and the CS⁻ indicate processing of learned stimulus value in ACC, SFG, Insula, and
878 vmPFC. The amygdala does not exhibit higher similarity between the instructed and experienced CS⁺
879 (CS⁺P) and the merely instructed CS⁺ (CS⁺U) relative to the similarity between these CS⁺s similarities
880 and the CS⁻. The error bars are ± 1 SEM.

881
882 **Figure 4.** Intra-CS similarities from mini-block to mini-block per CS type, region and experimental phase
883 (Experiment 1), based on the mini-block model (see Figure 2). Higher temporal consistency in intra-CS⁺

884 than intra-CS similarities indicates processing of learned stimulus qualities in a given ROI. This is not
885 observed for the merely instructed CS⁺ (CS⁺U) in the amygdalae. The error bars are ± 1 SEM.

886

887 **Figure 5.** Comparison of inter-CS similarities between each of the three different CS types from the test
888 phase (CS⁺te, CS⁺Pte, CS⁺Ute) and the CS⁺P pattern from the training phase (CS⁺Ptr), based on the phase
889 model (see Figure 2), reveals CS⁺P specific processing of threat-related information in the right amygdala
890 (Experiment 1): CS⁺P associated multi-voxel activation patterns during test (CS⁺Pte) are more similar to
891 CS⁺P associated patterns during training (CS⁺Ptr) than CS⁺U associated patterns during test (CS⁺Ute) (F).
892 Other regions do not show such differentiation (A-E). The error bars are ± 1 SEM. G. Individual
893 differences in the difference between CS⁺PtrCS⁺Pte and CS⁺PtrCS⁺Ute inter-CS similarities in the right
894 amygdala were correlated with differences in fear ratings between CS⁺P and CS⁺U in the test phase.

895

896 **Figure 6.** Inter-region similarity analyses between the ACC and left and right amygdala (Experiment 1).
897 A shows trial-by-trial intra-CS similarity matrices from the trial-based models for each region and CS
898 type. On this basis, Spearman correlation coefficients were calculated between each combination of the
899 resulting trial-by-trial ACC and left and right amygdala intra-CS similarity time courses, separately for
900 each CS. The right amygdala showed a higher inter-region similarity with the ACC for the CS⁺P
901 specifically (B), relative to the left amygdala where no such effect was observed (C). The error bars are
902 ± 1 SEM.

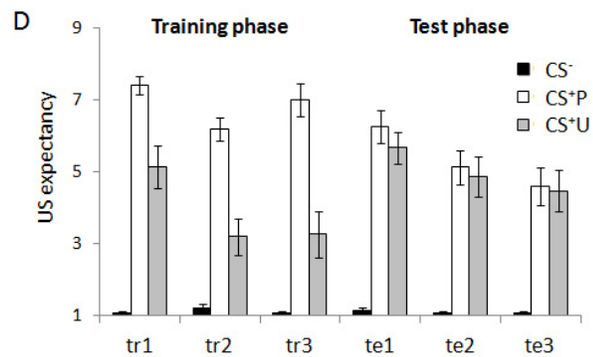
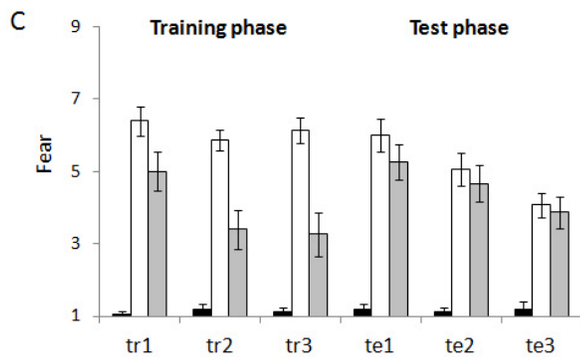
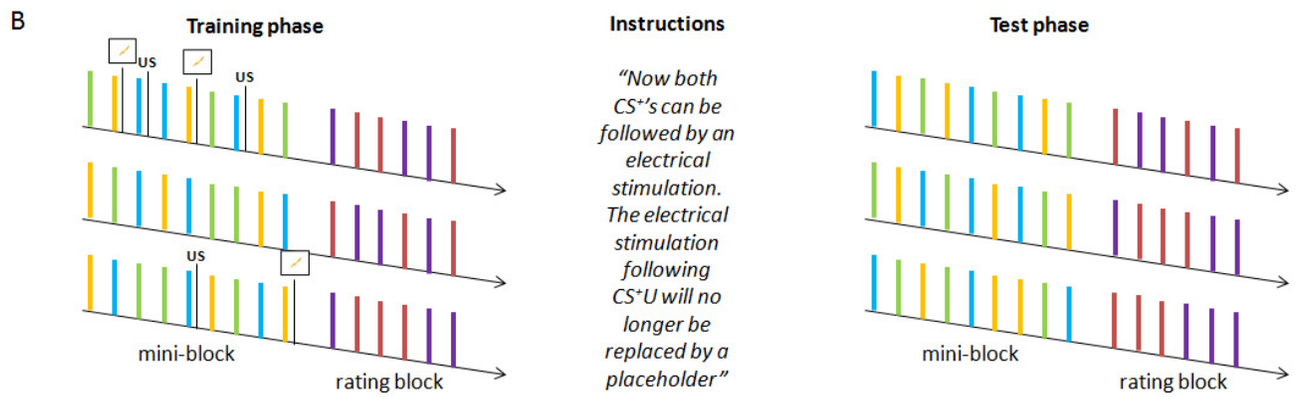
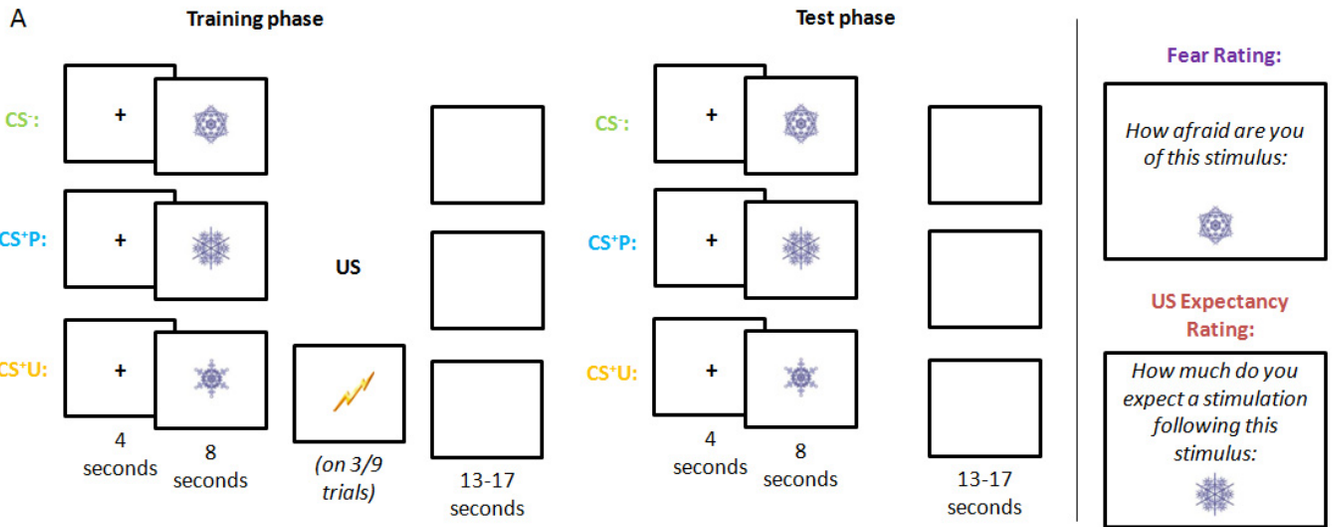
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904 **Figure 7.** Inter-region similarity analyses per different CS-type and experimental phase. A. Correlations
905 were calculated between each combination of two regions' intra-CS similarity matrices from the trial-
906 based models (see Figure 2), per CS and phase separately, as explained in Figure 6. B. The correlations
907 across all fear-related regions were averaged and are presented per CS type and phase separately. C.
908 Visual two dimensional scaling depiction of the similarities between different regions for each CS⁺ in the
909 test phase separately.

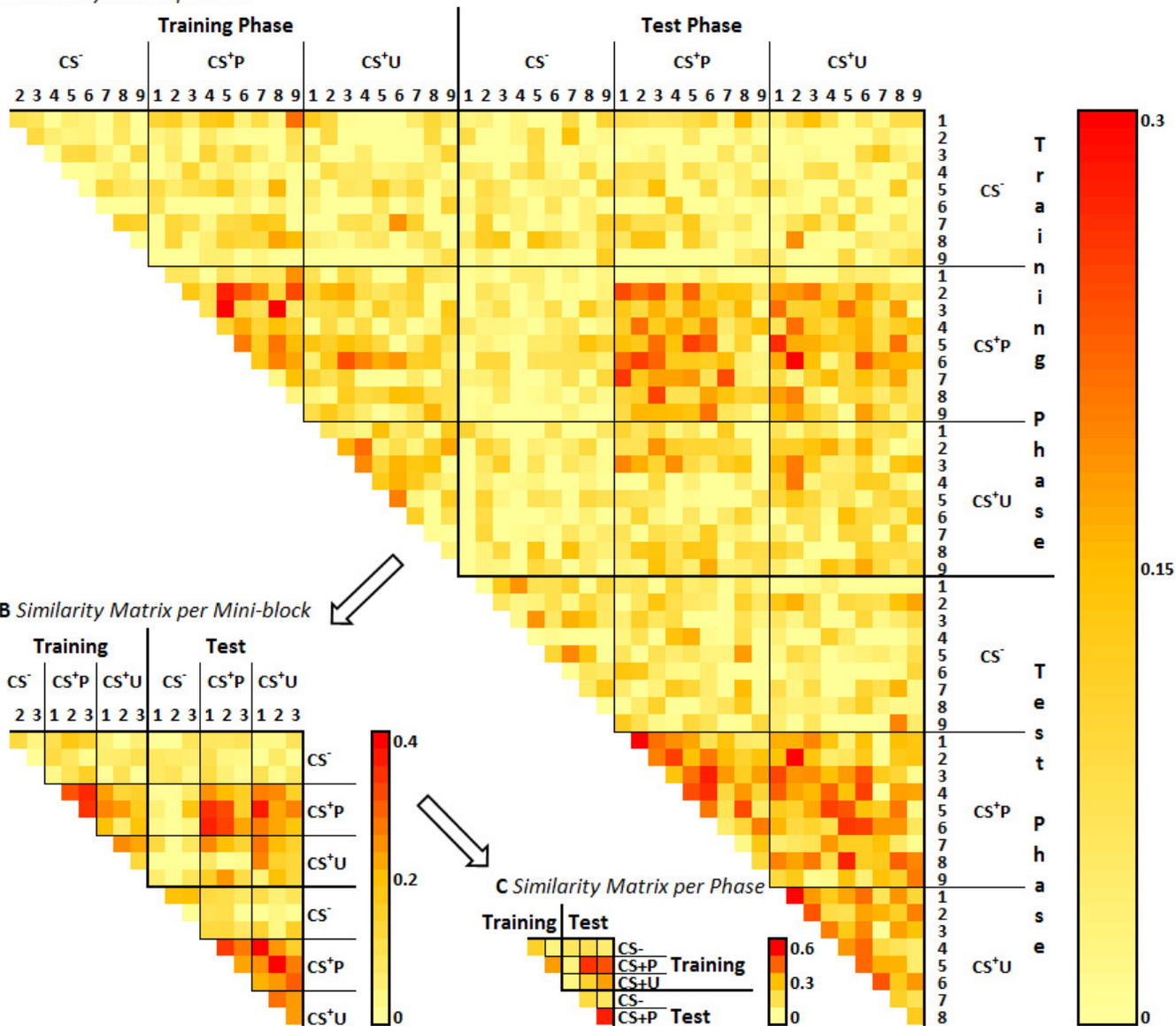
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911 **Figure 8.** Procedure Experiment 2 and results. A. Each trial consisted of a fear contingency instruction
912 and a CS presentation. The instruction indicated which of two pictures (CS⁺) would be followed by an

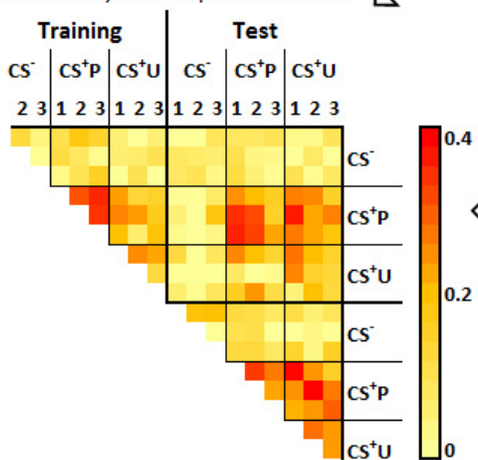
913 electrical US by presenting an intensity meter next to that picture. As illustrated on the lower left part of
914 A, some CSs and instructions were recurring (OLD), others were always novel (NEW). Orthogonal to
915 this, some instructions and subsequent CS presentation employed pictures of houses, others of faces. CS⁺
916 presentation was always followed by a US presentation. On a small subset of trials, CS presentation was
917 replaced by a catch question, to assure participants paid attention to the experiment. B. The similarity
918 analyses focused exclusively on pattern similarities between face and house pictures during CS
919 presentations, for each CS type separately (CS⁺old, CS⁻old, CS⁺new, and CS⁻new). C-H. These analyses
920 revealed that the right, but not the left, amygdala showed a differential response to fear relevance as a
921 function of novelty. Namely, the pattern response for houses and faces were more similar when these
922 denoted a CS⁺ than when they indicated a CS⁻. However, this difference was only present when the CSs
923 had been instructed and experienced before (old CSs), but disappeared when they were novel. The error
924 bars are ± 1 SEM.
925
926



A Similarity Matrix per Trial



B Similarity Matrix per Mini-block



C Similarity Matrix per Phase

