

Visual Probe Tasks (VPTs) have been extensively used to measure spatial attentional biases, but as usually analysed, VPTs do not consider trial-to-trial carryover effects of probe location: Does responding to a probe on, e.g., the location of a threat cue affect the bias on the subsequent trial? The aim of the current study was to confirm whether this kind of carryover exists, using a novel task version, the diagonalized VPT, designed to focus on such trial-to-trial interactions. Two versions of the task were performed by a sample of college students. In one version cues were coloured squares; in the other, cues were threat-related and neutral images. Both versions included partially random positive or negative response feedback and varying Cue-Probe Intervals (200 or 600 ms). Carryover effects were found in both versions. Responding to a probe at the location of a cue of a given colour induced an attentional bias on the subsequent trial in the direction of that colour. Responding to a threat-related cue induced an attentional bias towards threat on the subsequent trial. The results provide evidence that trial-to-trial carryover effects on spatial attentional bias indeed exist. A methodological implication is that previous probe location could be considered in analyses or re-analyses of spatial visual attention tasks.

1 Trial-to-trial Carryover Effects on Spatial Attentional Bias

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18

19 **Abstract**

20 Visual Probe Tasks (VPTs) have been extensively used to measure spatial attentional biases,  
21 but as usually analysed, VPTs do not consider trial-to-trial carryover effects of probe  
22 location: Does responding to a probe on, e.g., the location of a threat cue affect the bias on  
23 the subsequent trial? The aim of the current study was to confirm whether this kind of  
24 carryover exists, using a novel task version, the diagonalized VPT, designed to focus on such  
25 trial-to-trial interactions. Two versions of the task were performed by a sample of college  
26 students. In one version cues were coloured squares; in the other, cues were threat-related and  
27 neutral images. Both versions included partially random positive or negative response  
28 feedback and varying Cue-Probe Intervals (200 or 600 ms). Carryover effects were found in  
29 both versions. Responding to a probe at the location of a cue of a given colour induced an  
30 attentional bias on the subsequent trial in the direction of that colour. Responding to a threat-  
31 related cue induced an attentional bias towards threat on the subsequent trial. The results  
32 provide evidence that trial-to-trial carryover effects on spatial attentional bias indeed exist. A  
33 methodological implication is that previous probe location could be considered in analyses or  
34 re-analyses of spatial visual attention tasks.

35 **Keywords:** Carryover; Visual Probe; Colour; Threat; Attentional Bias; Spatial Attention

36

## 37 1. Introduction

38 The ability to select relevant information for further processing and response selection is  
39 essential for efficient, adaptive behaviour. Visual spatial attention is an important form of this  
40 ability, in which information is selected from regions of the visual field. This process  
41 involves bottom-up or intrinsic visual features versus top-down or task-dependent signals,  
42 together creating a spatial map of saliency (Soltani & Koch, 2010). Saliency maps are also  
43 affected by attentional biases involving emotional or motivational stimuli (Mogg & Bradley,  
44 2016). Such biases involve effects on selection or inhibition that are not due to intrinsic visual  
45 features, but that are nevertheless automatic rather than controlled and in that sense bottom-  
46 up. Attentional biases are commonly studied using dot-probe or visual probe tasks (VPTs)  
47 (MacLeod, Mathews, & Tata, 1986). In these tasks, emotional cue stimuli are presented on  
48 screen, and their appearance affects the saliency map as measured by responses to probe  
49 stimuli appearing at their location versus away from their location (Cisler & Koster, 2010;  
50 Mogg & Bradley, 2016; Notebaert, Crombez, Van Damme, De Houwer, & Theeuwes, 2011)  
51 or predicted location (Gladwin, Möbius, Mcloughlin, & Tyndall, 2019). Attentional approach  
52 versus avoidance of emotional cues is inferred from faster versus slower responses to probes  
53 at their location, relative to responses to probes at the location of non-emotional cues.  
54 Attentional biases, in terms of both attentional approach and avoidance, have been connected  
55 to a wide range of clinical disorders, including anxiety (for review, see Mogg & Bradley,  
56 2016), aggression (e.g., Kimonis, Frick, Fazekas, & Loney, 2006) and post-traumatic stress  
57 disorder (for review, see Aupperle, Melrose, Stein, & Paulus, 2012).

58 VPTs may however also contain information in the trial-to-trial variability that would long  
59 have been considered noise. That is: the bias towards or away from a certain stimulus  
60 category could change from one trial to the next, or over relatively brief periods of time  
61 within a task session. This variability of the attentional bias to and from salient stimuli over

62 trials has received recent research interest, although questions have been raised about the  
63 interpretation of most measures of attentional bias variability (Kruijt et al., 2016).  
64 Nevertheless, attentional bias variability has been related to, e.g., trauma (Iacoviello et al.,  
65 2014), anxiety (Zvielli, Bernstein, & Koster, 2014), and conflicting positive and negative  
66 alcohol-related associations (Gladwin & Vink, 2018).

67 One as yet rarely explored source of attentional bias variability could be trial-to-trial  
68 *carryover effects* (Gladwin, 2017; Hill & Duval, 2016). This refers to effects caused by the  
69 probe appearing on the emotional versus non-emotional location that are observed on the  
70 subsequent trial. Say, for instance, that on trial N the probe appears at the location of the  
71 emotional cue. The question is whether the attentional bias on trial N + 1 is different from if  
72 the probe had appeared at the location of the non-emotional cue on trial N. Analogous effects  
73 have been found to affect non-spatial attentional biases in the emotional Stroop task (Cane,  
74 Sharma, & Albery, 2009; Clarke, Sharma, & Salter, 2014; Waters, Sayette, Franken, &  
75 Schwartz, 2005; Wilson, Sayette, Fiez, & Brough, 2007) and spatial attentional biases  
76 carrying over between different tasks (Thompson & Crundall, 2011). The rationale for  
77 translating the carryover concept to trial-to-trial effects in spatial visual probe tasks is that  
78 responding to probes at the location of the emotional versus non-emotional cue could cause a  
79 state that affects attentional bias on the subsequent trial. Such a state could be described using  
80 a generalized concept of binding (Roelfsema, Engel, König, & Singer, 1997; Treisman &  
81 Gelade, 1980) in which the stimulus feature “threat” is bound to an attentional function. If  
82 this binding remains active on the subsequent trial, it would cause an attentional bias towards  
83 the location of the cue corresponding to the previous probe’s location. Some evidence for  
84 carryover effects has been found for threat VPTs (Gladwin, 2017): Responding to probes at  
85 the location of threat cues caused lower overall accuracy on the subsequent trial (but no  
86 change in bias towards or away from threat), and subclinical post-traumatic stress disorder

87 symptoms were associated with this effect. Further, symptoms appeared to be associated with  
88 a time-dependent carryover effect on bias, in which responding to threat on a trial induced a  
89 bias towards threat on the next trial, expressed by increased errors when the probe appeared  
90 on the neutral cue location. Such effects would be missed without considering previous trial  
91 cue location as a factor in analyses. However, it remains to be firmly established that trial-to-  
92 trial carryover exists as a phenomenon in spatial attentional bias tasks.

93 The aim of the current study was therefore to confirm the hypothesis that trial-to-trial  
94 carryover effects exist in visual probe tasks. We used a variant of the VPT, the diagonalized  
95 VPT (dVPT), optimized to study such effects. This task is designed in such a way as to  
96 reduce trial-to-trial interference other than the type of carryover effect of interest. Essentially,  
97 neither response keys nor stimulus locations were ever repeated. In task version 1 (the Colour  
98 task), the cues concern a basic visual feature (the colour of cues), while in task version 2 (the  
99 Threat task), the cues concern an emotional-motivational feature (threatening versus non-  
100 threatening scenes). An additional, more exploratory question involved the use of random  
101 feedback on responses. This was based on the theoretical perspective that the adaptive  
102 activation of cognitive responses to stimuli must depend on prior reinforcement processes (de  
103 Wit & Dickinson, 2009; Gladwin & Figner, 2014; Hazy, Frank, & O'Reilly, 2007). Just as  
104 how motor responses are learned and subsequently selected, likely involving dopaminergic  
105 signals in the basal ganglia, cognitive responses and even executive functions are determined  
106 by whether they were previously reinforced (Bunge, 2004; Lanciego, Luquin, & Obeso,  
107 2012). We therefore hypothesized that trial-to-trial carryover would depend on whether  
108 positive or negative feedback occurred on the previous trial, even if this feedback was task-  
109 independent. If positive versus negative feedback occurred, carryover was expected to be  
110 stronger, as positive feedback would reinforce the most recently performed cognitive action  
111 (i.e., attending to a location associate with a given cue category). Finally, the Cue-Probe

112 Interval (CPI), the duration of the interval between the cues (the stimuli expected to induce an  
113 attentional shift) and the probe (the stimulus requiring a response), was manipulated, as  
114 temporal dynamics are known to play an important role in attentional biases (Mogg, Bradley,  
115 Miles, & Dixon, 2004). There was no specific a priori hypothesis concerning CPI and  
116 variability, but using multiple CPIs allows potential time-dependent effects to be detected.

## 117 2. Methods

### 118 2.1. Participants

119 Participants were students who enrolled for participation credits ( $N = 163$ , analytical sample  
120 of 144 after removing subjects who showed low overall accuracy (below .8) or incomplete  
121 data; 119 female and 25 male, mean age 20,  $SD = 4$ ).

### 122 2.2. Diagonalized Visual Probe Task

123 Two versions of the dVPT were used (Figure 1). In both versions, trials started with the  
124 presentation of two cue stimuli. In the Colour version of the task, the cues were a yellow and  
125 a blue square. In the Threat version of the task, the cues were neutral and threatening pictures  
126 drawn from a subset of 14 images from the International Affective Pictures Set (Lang,  
127 Bradley, & Cuthbert, 2008). Threatening pictures included attacking animals and scenes with  
128 physical violence such as a pointed gun. Neutral pictures included non-threatening animals  
129 and sports scenes. Pictures never repeated from one trial to the next. The positioning of the  
130 two cue stimuli changed per trial, alternating between the diagonals of locations on a two by  
131 two grid. That is, they either appeared at the top-left and bottom-right locations, or at the  
132 bottom-left and top-right locations. The cues remained on-screen for a CPI of either 200 or  
133 600 ms, with equal probability. During cue presentation and throughout the trial, the current  
134 score was shown in white (if the score was non-negative) or red (if the score was negative)  
135 digits at the top of the screen. Following the CPI, the probe stimulus appeared. The probe  
136 consisted of two symbols: The target symbol >><< which replaced one of the two cues, and a

137 non-target symbol  $\vee$  or  $\wedge$  on the other location. The task was to press the button  
138 corresponding to the location of the target. The keyboard response buttons were R, F, J, and I;  
139 note that these had a strong stimulus-response compatibility in terms of spatial locations (e.g.,  
140 “upper-left”, “lower-left”, etc). The task continued after a response was given. Following an  
141 incorrect response, a red “-1” was presented as negative feedback, and the score was  
142 decreased. Following a correct response, a red “-1” or a green “+1” could appear, with equal  
143 probability, while the score was in the range -2 to +2. Outside this range, there was a  
144 tendency for the score to be pushed back towards zero. If the score was lower than -2 and the  
145 initial random feedback was negative, there was a .4 chance for the random feedback to  
146 become positive. If the score was higher than +2 and the initial random feedback was  
147 positive, there was a .4 chance for the random feedback to become negative. The score was  
148 updated according to the feedback. Participants were instructed that the feedback was  
149 random, but that incorrect responses were always followed by negative feedback. It was  
150 therefore still optimal to provide correct responses. The intertrial interval was 250 ms.

151 The Colour dVPT consisted of 9 blocks of 35 trials per block. The Threat dVPT consisted of  
152 16 blocks of 35 trials per block. The difference in block numbers was due to the expectation  
153 that fewer trials would be needed to detect effects involving the simple Colour cues due to the  
154 simpler categories and the lack of variation of cues per category.

155 Importantly for the current study, by using the diagonalized locations and these response  
156 keys, neither stimulus locations nor response keys were repeated from one trial to the next.  
157 This removed these sources of trial-to-trial influence.

### 158 2.3. Procedure

159 The study was performed online. Participants received information via a webpage, clicked on  
160 a clearly marked button to indicate informed consent, and then received an invitation by



161 email with a link to participate. Participants performed the Colour and Threat versions of the  
162 dVPT, always starting with the Colour version. Participants also filled in questionnaires and  
163 performed other tasks and subsequent sessions unrelated to the current study.

#### 164 2.4. Data pre-processing and statistical analyses

165 Analyses were performed in Matlab (The Mathworks, 2015). The first four trials of the task,  
166 the first trial of each block, trials with incorrect responses and trials following incorrect  
167 responses and trials with RTs above 3000 ms were removed as being likely noisy. Further  
168 pre-processing concerned the removal of trial data that was logged more than once (due to a  
169 feature of the software that re-logged data when the connection was slow, to avoid data loss)  
170 and the removal of data of task performance that was repeated or restarted. Repeated  
171 measures ANOVA was used to test effects of the within-subject factors of Current Probe  
172 Location (Blue or Yellow for the Colour version; Threat or Neutral for the Threat version),  
173 Previous Probe Location (Probe Location on the previous trial), CPI (200 versus 600 ms), and  
174 Previous Feedback (Negative or Positive). Higher-order interactions were explored using  
175 post-hoc tests which performed lower-order interactions per level of one of the variables of  
176 the higher-order interaction. The dependent variable was median RT, as this removes effects  
177 of outliers and the need to set arbitrary RT criteria for defining outliers.

178 Data and scripts are available on request.

#### 179 3. Results

180 Descriptive statistics are provided in Table 1. Overall accuracy was good, .96 for the Colour  
181 task and .96 for the Threat task. Figure 2 illustrates the main findings. For the Colour version,  
182 the primary test—the interaction between Current Probe Location and Previous Probe  
183 Location—was significant,  $F(1, 143) = 91, p < .0001, \eta_p^2 = 0.39$ : On the trials following a  
184 response to a probe at the location of a blue cue, responses were faster for probes on blue

185 than on yellow cue locations,  $t(143) = -3.44, p = .00076, d = 0.29$ . On the trials following a  
186 response to a probe at the location of a yellow cue, responses were slower for probes on blue  
187 than on yellow cue locations,  $t(143) = 7.66, p < .0001, d = -0.64$ . This interaction was not  
188 further moderated by CPI or Previous Feedback. There were also effects of Previous Probe  
189 Location (responses were faster following responses to yellow than to blue locations:  $t(143) =$   
190  $-2.22, p = .028, d = -0.18$ ), and of CPI (responses were faster following the longer (600 ms)  
191 than the shorter (200 ms) CPI:  $t(143) = -5.54, p < .0001, d = -0.46$ ).

192 For the Threat version, the interaction between Current Probe Location and Previous Probe  
193 Location was also significant,  $F(1, 143) = 8.5, p = .0042, \eta_p^2 = 0.056$ . On trials following  
194 respond-to-threat trials, responses to the threat location were faster than responses to the non-  
195 threat location,  $t(143) = -2.92, p = .0041, d = -0.24$ . On trials following respond-to-non-threat  
196 trials, there was no significant difference between probes at the threat versus non-threat  
197 location,  $t(143) = -0.63, p = .53, d = 0.0027$ . There was no further moderation of the  
198 interaction. There was a main effect of CPI, with faster responses following the longer than  
199 the shorter CPI,  $t(143) = -27.89, p < .0001, d = -2.32$ ; and an effect of Previous Probe  
200 Location, with slower responses following probes at the threat versus non-threat location,  
201  $t(143) = 2.00, p = .048, d = .17$ .

#### 202 4. Discussion

203 The results confirmed the primary hypothesis: Carryover effects were found in both task  
204 variants. In the Colour task, responses were faster on probes appearing at the location of the  
205 same Colour-cue as where the previous trial's probe had appeared, versus on probes  
206 appearing at the location of the other cue. In the Threat task, an attentional bias to threat was  
207 only found following a trial with a response to a probe on the threat location. This was  
208 previously interpreted in terms of a kind of binding (Roelfsema et al., 1997; Singer et al.,  
209 1996) between the function of attentional selection and the stimulus category associated with

210 the position to which attention is shifted (Gladwin, 2017). Questions clearly remain on the  
211 precise processes underlying carryover effects. Whether effects occur at the level of the  
212 saliency map or involve later processing involving response selection cannot yet be  
213 determined. However, the current carryover effects fit the binding interpretation, or stated  
214 somewhat differently the model of a task set (Monsell, 2003) of stimulus – response  
215 mappings, with cue categories as imperative stimuli and attentional shifting as the responses  
216 to which the stimuli are mapped. That is, it appears that by responding to a probe at the  
217 location of a given cue, a mapping is established between that cue category and the covert  
218 cognitive response of shifting attention to that cue’s location (or potentially, away from the  
219 non-attended location’s cue).

220 We note that while the carryover effect was found in both tasks, it was stronger in the Colour  
221 than in the Threat task. The effect size of the interaction was greater in the Colour task, and  
222 the effect in Threat task was limited to trials following a probe-on-threat trial. There are a  
223 number of reasons that could have played a role in this. First, the colour cues were highly  
224 visually salient and there was no variation between cues. In contrast, threat versus non-threat  
225 stimuli were complex and varied, requiring more visual processing to determine the  
226 categories and presumably also varying in how threatening different exemplars were. This  
227 would be expected to lead to more noise in the Threat task. Further, the limitation of the  
228 effect to post-threat trials may be a true effect: perhaps responding to neutral trials does not  
229 induce a bias in the way that attending to threat trials does. Speculatively, this would make  
230 evolutionary sense, in that becoming attuned to threat and downregulating unthreatening  
231 information could aid survival.

232 A limitation of the current study is that the results concern a novel task variant, specifically  
233 designed to answer the theoretical question of whether carryover effects exist in spatial  
234 attentional bias. While it appears difficult to explain these effects in a different way than an

235 attentional bias, whether similar effects can be found in classical dot-probe tasks remains to  
236 be determined by future research. Some current task-variations involving feedback, such as  
237 the changing colour of the score, may be unnecessary or suboptimal for future work. Less  
238 abstract positive and negative feedback could yet prove to influence carryover, for instance  
239 angry faces or electric shock. A second limitation is the use of true randomization per trial  
240 rather than precisely counterbalanced trials. However, analyses of trial numbers showed the  
241 expected averaging to very similar numbers for comparable conditions; there did not seem to  
242 be any possible way random variations in trial numbers could result in systematic RT  
243 differences. Nevertheless, future work could consider controlling the trial numbers per  
244 condition, per participant. Third, the possibility was raised during review of a different kind  
245 of carryover, namely of CPI – could effects involve differences involving the same versus  
246 different CPI being used on consecutive trials? We note that there was no systematic  
247 relationship between CPI-carryover and the type of carryover, Category-carryover, that was  
248 the focus of the current study. However, future work could restrict the design to a single CPI  
249 to remove any effect of this type of carryover. Fourth, the stimulus categories of threat versus  
250 non-threat could be further decomposed, in particular in terms of being negative and  
251 arousing. In the current study, threat stimuli would be both more negative and more arousing  
252 than the control stimuli. Future work could determine whether carryover effects are also  
253 found while controlling for either dimension. Fifth, the order of the Colour and the Threat  
254 tasks was not counterbalanced, so that comparisons between the tasks are confounded by  
255 order and time on task.

256 In conclusion, trial-to-trial carryover effects were found in spatial attentional bias tasks  
257 involving colour and threat cues. Including previous probe location as a factor in future  
258 analyses may contribute to the understanding of trial-to-trial variability and reveal previously  
259 undetected effects and relationships.

## 260 Compliance with Ethical Standards

261 The authors declare no conflict of interest. The study was approved by the institutional  
262 ethical review board (Ethics Committee of the Radboud University Nijmegen, application  
263 ECSW2016-1710-422). Participants provided informed consent before performing the  
264 experiment.

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366 **Table 1. RTs per condition**

367 1A. Colour variant

		Blue				Yellow			
		Neg		Pos		Neg		Pos	
		200	600	200	600	200	600	200	600
Blue		563	540	562	553	567	561	582	563
		(121)	(124)	(126)	(142)	(114)	(118)	(118)	(124)
Yellow		576	563	572	563	550	535	552	544
		(122)	(130)	(114)	(128)	(115)	(127)	(110)	(142)

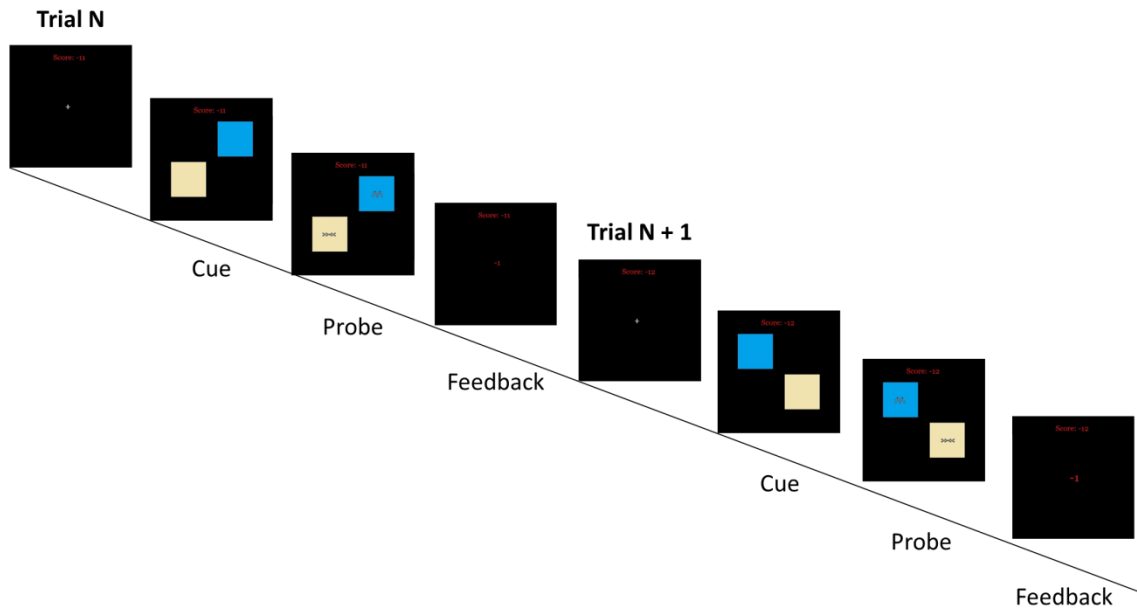
368

369 1B. Threat variant

		Neutral				Threat			
		Neg		Pos		Neg		Pos	
		200	600	200	600	200	600	200	600
Neutral		582	529	584	526	592	531	593	533
		(95)	(99)	(93)	(87)	(102)	(89)	(94)	(102)
Threat		588	528	588	524	588	525	583	532
		(93)	(88)	(85)	(83)	(101)	(88)	(79.7)	(109)

370 *Note.* The Table shows the mean RT per condition, with standard deviations in brackets, of  
 371 the Colour and Threat variants of the dVPT. Standard deviations are given for the between-  
 372 subject data, i.e., without removal of the subject means. Rows show the probe locations on  
 373 the current trial. Columns show the probe location on the previous trial, feedback on the  
 374 previous trial (Negative or Positive), and Cue-Probe interval (200 or 600 ms). The overall  
 375 accuracy was .96 in the Colour task and .96 in the Threat task.

376 Figure 1. Illustration of the diagonalized Visual Probe  
 377 Task

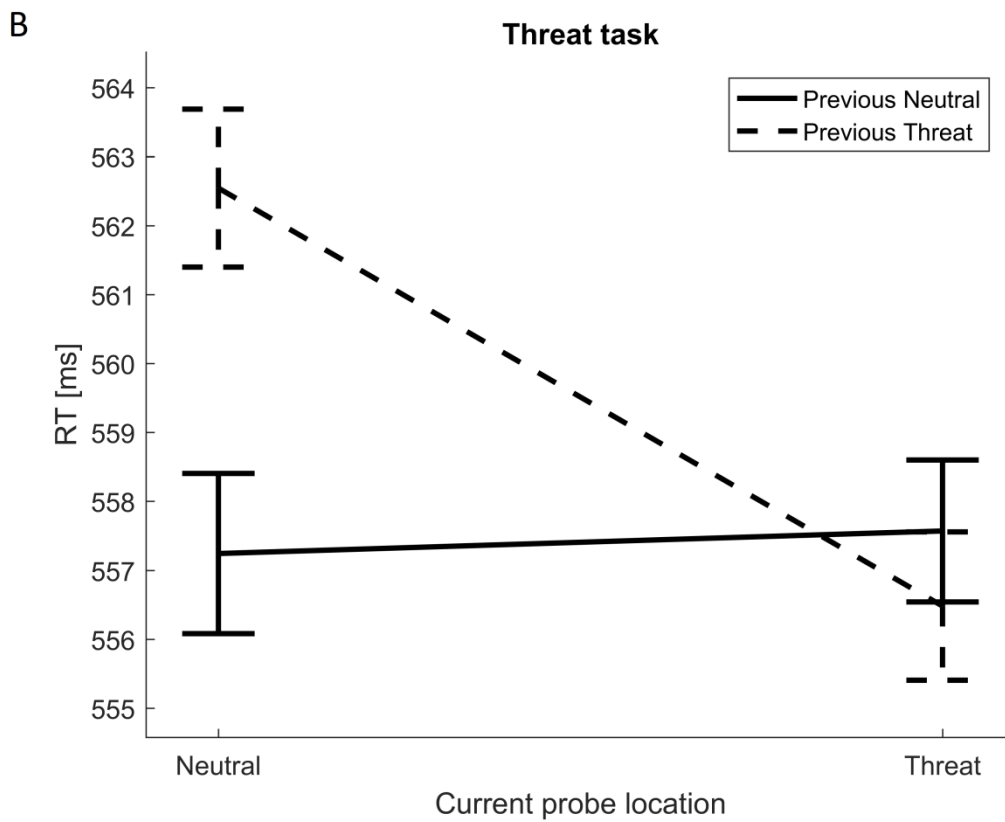
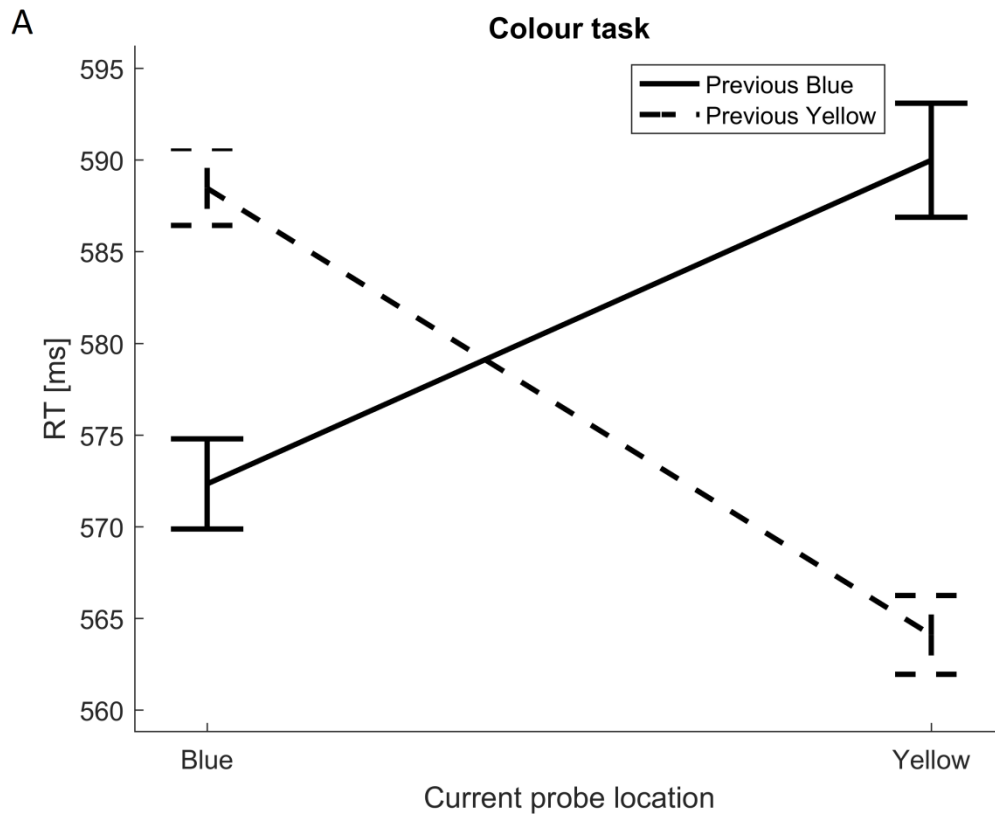


378

379 *Caption.* Trials consisted of a cue, which remained on screen for 200 or 600 ms. In the  
 380 Colour version of the task, cue stimuli were a yellow and a blue box. In the Threat version of  
 381 the task, a neutral and a threatening picture were used. A probe stimulus then appeared  
 382 requiring a button press indicating the location of a target stimulus. Correct responses were  
 383 followed by random positive or negative feedback. Incorrect responses were always followed  
 384 by negative feedback only. The diagonal on which the two elements of the cue appeared  
 385 alternated over trials so that spatial location and response button were never repeated.

386

387 **Figure 2. Carryover effects**



388

389 *Caption.* The figures illustrate the main findings involving carryover. The x-axis represents  
390 the location of the probe on the current trial. The lines are separated based on the location of  
391 the probe on the previous trial. The error bars are  $+1/-1$  standard errors based on the data after  
392 removal of the subject means, as effects concerned within-subject factors (Cousineau, 2005;  
393 O'Brien & Cousineau, 2016). In both task versions, attentional bias was affected by the probe  
394 location on the previous trial. In the Colour task (A), an attentional bias was induced in the  
395 direction of the cue associated with probe location on the previous trial. In the Threat task  
396 (B), an attentional bias to threat, expressed as slower responses when the probe appeared  
397 away from the threat cue, was found only following trials when the probe appeared at the  
398 location of the threat cue.