Biological Psychology 88 (2011) 116-123

Contents lists available at ScienceDirect



Biological Psychology

journal homepage: www.elsevier.com/locate/biopsycho

Compatibility-sequence effects in the Simon task reflect episodic retrieval but not conflict adaptation: Evidence from LRP and N2

Michiel M. Spapé^{a,b,*}, Guido P.H. Band^{a,b}, Bernhard Hommel^{a,b}

^a Leiden Institute for Brain & Cognition, Netherlands ^b Leiden University Institute of Psychology, Netherlands

ARTICLE INFO

Article history: Received 23 September 2009 Accepted 3 July 2011 Available online 19 July 2011

Keywords: Executive control Conflict adaptation Episodic retrieval Feature-integration N2 LRP Compatibility-sequence effects

1. Introduction

Psychologists and philosophers alike have long wondered how humans achieve their intended goals in the face of distractions and temptations. To study how distraction affects intentional behavior, numerous conflict-inducing tasks have been developed. Stroop (1935) found that it is difficult to ignore color words while naming incongruent colors; Simon and Rudell (1967) observed that it is difficult to ignore the location of stimuli when carrying out a spatially defined response; and Eriksen and Eriksen (1974) discovered that irrelevant flankers are difficult to ignore when responding to a central stimulus. Such effects have often been taken to demonstrate an *automatic* impact of stimuli of action control: stimuli sometimes seem to be able to evoke unwanted and interfering action tendencies against an agent's will.

At the same time, however, in none of these or other experimental tasks participants are enslaved to what stimuli tell them: even though it may take them a few more milliseconds to respond in the face of incongruent or incompatible stimuli, they are commonly able to do so. This suggests that voluntary and involuntary action tendencies compete, which has prompted researchers to conceive of action-control models that comprise of at least two processing routes—an automatic route that translates stimuli into habitually

ABSTRACT

Behavioral and psychophysiological studies on the Simon effect have demonstrated that stimuli automatically activate spatially corresponding responses, even if their location is irrelevant to the task. Interestingly, this Simon effect is attenuated after stimulus–response incompatible trials (Gratton effect or compatibility-sequence effect), a pattern that has often been attributed to online conflict adaptation, even though an account in terms of episodic binding and retrieval is just as plausible. Here we show that the compatibility-sequence effect can be eliminated and partly reversed by rotating the boxes in which stimuli are presented in between two given trials, a manipulation that is likely to affect episodic representation but not online control. Sequential modulations of electrophysiological indicators of automatic response priming were also eliminated (N2) or even reversed in sign (LRP), suggesting that these effects are due to episodic retrieval of stimulus–response bindings but not, or to only a negligible degree, to online adaptation.

© 2011 Elsevier B.V. All rights reserved.

BIOLOGICAL

acquired or otherwise associated response tendencies and a voluntary route that makes sure that the intended action comes out as planned eventually (for a review, see Hommel, 2000).

There is general agreement that so-called "automatic" routes are not entirely independent of the current intention. Seeing a color word, processing a stimulus location, and confronting a flanker stimulus does not always evoke action tendencies in people but they do so because particular experimental tasks provide a context that promotes processing of color, location, and flanker information. So in some sense, automaticity is enabled by intentions, thus creating what one may call a prepared reflex (Hommel, 2000; Woodworth, 1938).

However, recent research suggests that automaticity may be under much tighter voluntary control than suggested by this scenario. Gratton et al. (1992) found that the effect of irrelevant flankers is mediated by the congruence of flankers and targets in the previous trial: the delay of responding with incongruent as compared to congruent flankers (i.e., the flanker effect) was significantly reduced after an incongruent as compared to a congruent trial-an effect that we will refer to as the "compatibility-sequence effect" (CSE). This reduction was visible in both behavioral observations and event-related potentials (ERPs). It is well-known that responseincompatible flankers activate a lateralized readiness potential (LRP) reflecting the flanker-related response (Coles et al., 1985). This suggests that flankers indeed prime the response they are associated with. Most interestingly, however, Gratton et al. (1992) observed that these incorrect LRPs are also reduced after an incongruent trial.

^{*} Corresponding author. Present address: School of Psychology, University of Nottingham, Nottingham NG7 2RD, United Kingdom. Tel.: +44 115 82 32248.

E-mail address: michiel.spape@nottingham.ac.uk (M.M. Spapé).

^{0301-0511/\$ -} see front matter © 2011 Elsevier B.V. All rights reserved. doi:10.1016/j.biopsycho.2011.07.001

Conditions with response conflict usually also induce an increase in the N2 component of the ERP relative to conditions without conflict. The peak of this component has a fronto-central scalp distribution, presumably a source in the anterior cingulate cortex (ACC), and a latency of around 250 ms (Nieuwenhuis et al., 2003; Sasaki et al., 1993; Watanabe et al., 2002). The N2 has been claimed to reflect a general process of inhibiting erroneous responses (Kok et al., 2004) or exerting cognitive control (Folstein and Van Petten, 2008). Based on such observations, Botvinick et al. (1999) have suggested a model in which the ACC is held responsible for continuously monitoring for the occurrence of response conflict. As soon as conflict is detected, an adaptive mechanism fine-tunes control processes, thus reducing the risk of running into conflict in the future. Later modifications of this conflict-monitoring model describe conflict as an aversive stimulus (Botvinick, 2007) that triggers the avoidance of decision-making strategies that are likely to lead to its re-occurrence.

Stürmer et al. (2002) have suggested that the presence of such an error-detection/resolution mechanism may explain CSEs as observed by Gratton et al. (1992) and others. In their own study, Stürmer et al. (2002) demonstrated that the Simon effect, characterized by delayed responding to stimuli that spatially correspond to an alternative action (Simon and Rudell, 1967), is affected by manipulations of the probability of stimulus-response compatibility and by the compatibility of the previous trial. Similar to the observation of Gratton et al., Stürmer and colleagues found that, after incompatible trials, the Simon effect is smaller, absent, and sometimes (if compatible trials are more frequent) even reversed. These manipulations caused response-incompatible stimuli to induce an LRP for the incorrect response, resulting in an initial dip in the LRP before peaking in the direction of the final response. Moreover, they found that the amplitude of this dip was reduced in blocks with a high probability of incompatible responses and completely absent after incompatible trials. Stürmer et al. suggested that these observations reflect a mechanism of error detection/resolution along the lines of Botvinick et al. (1999). Experiencing response conflict leads people to suppress the automatic processing route, so that information processed by this route impacts response selection to a lesser degree or not at all. As a consequence, response selection is driven by the intentional route only and irrelevant information no longer impairs (or facilitates) performance.

It should be noted that CSEs in the Simon task (e.g. Stürmer et al., 2002; Leuthold and Schröter, 2006) provide a more process-pure indication of the possibility that response conflict is online-controlled by intentions than those based on flanker experiments do. Flanker effects are likely to have multiple causes: in part they seem to result from direct interactions between target and flanker representations (stimulus conflict) and in part from interactions between the responses that are mapped onto targets and flankers (response conflict; e.g. Fournier et al., 1997; Hommel, 1997; Rösler and Finger, 1993). In contrast, stimulus conflict can be excluded in the case of the Simon task (where there is no contradiction between any stimulus location and the relevant non-spatial stimulus feature), which leaves response conflict as the most likely culprit.

1.1. Episodic retrieval

the conditions in which stimuli and responses are repeated eliminates the adaptation-like effect. Hommel et al. (2004) have made a similar argument for the Simon effect. They point out that there is independent evidence suggesting that stimulus features and responses are spontaneously integrated and bound into episodic memory traces (Hommel, 1998, 2004). These bindings have been shown to impair performance in subsequent trials if some features and/or the response are repeated while others alternate (partial repetitions), suggesting that repeating components of a binding leads to the retrieval of the whole binding, thereby inducing code conflict if repetitions are only partial. Again, sorting trials into those following compatible versus incompatible trials leads to a confound with partial-repetition costs,¹ which may account for parts or all of what looks like adaptation effects (Hommel et al., 2004).

Two different strategies have been applied to disentangle whether CSEs in conflict tasks reflect true adaptation through cognitive control, an effect of stimulus–response binding, or a mixture of the two. Most authors advocate the use of only those conditions that are unaffected by binding processes. For instance, Akçay and Hazeltine (2007) restricted their analyses of trial-to-trial effects in the Simon task to trial transitions where not a single stimulus or response feature is repeated, and argued that this would avoid any contribution from binding-related effects.

Unfortunately, there are reasons to assume that even nonrepetitions may be affected by binding (Dutzi and Hommel, 2009). Assume, for instance, the combination of stimulus $S_{\rm b}$ and $R_{\rm v}$ is encountered right after having processed S_a and R_x , and assume that the representations of S_a and R_x were bound on this occasion. Discriminating between the two stimuli and the two possible responses requires the code of $S_{\rm b}$ to outcompete the code of $S_{\rm a}$ and the code of R_v to outcompete the code of R_x . This process can be assumed to benefit from the previously created binding between S_a and R_x : losses of S_a in its struggle against S_b would also weaken R_x in its struggle against R_v, and vice versa—a mechanism that Duncan (1996) has called integrated competition. In other words, the previous binding of stimulus and response components helps rejecting them in alternation trials. This means that binding and episodic retrieval can affect performance even if not a single stimulus or response feature is repeated, suggesting that confounds between binding effects and possible adaptation effects are impossible to avoid in principle.

Therefore, Spapé and Hommel (2011) suggested a different research strategy. Rather than trying to isolate possible contributions from binding and control, they sought for manipulations that strongly impact, and even eliminate CSEs while being unlikely to have any bearing on control. In fact, Spapé and Hommel (2011) demonstrated that a seemingly minor modification of the visual background in between two trials is sufficient to eliminate CSEs.

In the present study, we aimed to extend this observation to LRPs that produced outcomes that hitherto were taken as particularly convincing support of control accounts. In a nutshell, we compared performance in what one may consider a standard Simon task with performance in an only slightly modified version of this task where the left and right box in which the targets appeared rotated in the

Recent behavioral results have, however, shed doubt as to whether sequential conflict studies truly demonstrate a mechanism of conflict-adaptation. Mayr et al. (2003) showed that Gratton et al.'s (1992) findings can also be accounted for without referring to any higher-order mechanisms, such as conflict monitoring. Mayr et al. point to the fact that sorting trials into those following flankertarget congruence versus incongruence induces a confound with (stimulus- and/or response-) priming effects, and that excluding

¹ To see that, consider RT_{cC} , RT_{cI} , RT_{iC} , and RT_{iI} the means of reaction times in compatible trials after compatible trials, incompatible trials after compatible trials, compatible trials after incompatible trials, and incompatible trials after incompatible trials, respectively. The standard pattern that is interpreted to reflect adaptation is that the Simon effect after compatible trials ($RT_{cI} - RT_{cC}$) is larger than the Simon effect after incompatible trials ($RT_{iI} - RT_{iC}$). Now consider that partial repetitions of stimulus–response combinations (which are known to delay responding for reasons unrelated to control; Hommel, 1998) only occur in conditions where the compatibility in the present trials differs from the one in the previous trial. This would selectively increase RT_{cI} and RT_{iC} , and may thus mimic adaptation effects even in the absence of any control–related effects.

intertrial interval. In the standard condition, we expected to replicate the observations of Stürmer et al. (2002): the Simon effect should be reduced in size or even disappear after incompatible trials in both reaction times (RTs) and LRPs.

Unlike Stürmer et al. (2002), we employed a horizontal version of a sequential Simon paradigm. This is important to mention because an easily overlooked issue with studies measuring the Simon effect on the LRP concerns a particular confound in the lateralized nature of the original task (see Praamstra, 2007 for a discussion). That is, in incompatible conditions, the ERP components resulting from the visual location of the stimulus are necessarily contralateral to those resulting from the manual response. As a consequence, positive LRPs on incompatible trials may be partially due to lateralized perceptual or attentional processes, rather than to motor preparation for the incorrect response. Several studies have sought to avoid this by using vertical layouts (e.g. De Jong et al., 1994; Stürmer et al., 2002). This approach has its own difficulty, however, since it has been questioned whether the vertical Simon effect can be compared with its horizontal counterpart (Wiegand and Wascher, 2005). Fortunately, however, there is no reason to expect that these confounds limit the interpretation of the *effects* of rotation on CSEs, on which the current study focuses. That is, to the extent that attention-related lateralized effects occur and mimic a compatibility effect on the LRP, this is equal for the two conditions that we compare. The only difference between the two conditions is whether or not the placeholders for imperative stimuli are reversed by rotation in between trials. We could therefore afford to employ the more standard horizontal configuration.

Given that our minor manipulation is unlikely to affect any control process but, as we will argue, can be expected to affect binding processes, a control account is unable to explain any impact of this manipulation on CSEs. However, merely showing some effect of a binding-related manipulation is not particularly revealing. Control researchers increasingly admit that binding does play some role in attenuating conflict (e.g. Verguts and Notebaert, 2008; Notebaert and Verguts, 2008) so that modulations of CSEs may easily be attributed to the "binding portion" of these effects without invalidating the assumption of further "control portions". What would be more informative is the demonstration that CSEs can be eliminated altogether or even reversed by means of exclusively binding-related manipulations. And this is what we aimed to achieve in the present study.

1.2. Aim of present study

To test whether CSEs in the Simon task can be eliminated or even reversed in both RTs and LRPs we combined the design of Spapé and Hommel (2011), who demonstrated such an effect in RTs, with EEG recordings along the lines of Stürmer et al. (2002). CSEs were assessed by presenting pairs of trials of a rather standard Simon task. Participants responded to the shape of the visual stimuli by pressing a left versus right key, and the stimuli were presented randomly in one of the two placeholder boxes, to the left and right of the center of the screen. For the sake of clarity, we will call the first stimulus of each pair S1 and the second stimulus S2. Not only did we expect the standard Simon effect-better performance if the location of the stimulus corresponds to the location of the response (i.e. with stimulus-response compatibility) than if it does not-but a standard CSE as well: the Simon effect should be reduced, absent, or even reversed after incompatible as compared to compatible trials. In our terminology, the Simon effect for S2 should be less pronounced after an incompatible as compared to a compatible S1 trial, and this was expected for both RTs and LRPs (Stürmer et al., 2002).

This replication was expected for what we will call the *static* condition, where the two boxes in which the stimuli appeared stayed on the screen in the same positions throughout the whole experiment. However, in another condition, the *rotation* condition, the boxes were gradually rotated 180° around the screen center, as is schematically shown on the right branch of Fig. 1, which led to the reversal of the two boxes. This rotation was taking place after the response to S1 was given and before S2 was presented, so that this manipulation should not have any effect on online conflictmonitoring in the sense of Botvinick et al. (1999) or Stürmer et al. (2002). In contrast, a binding approach would lead one to expect an impact on the CSE.

Let us consider, for example, that a participant responds to circles and stars by pressing a right and a left key, respectively. In the scenario shown in Fig. 1, he or she might encounter a compatible S1 (location: right, response: right) followed by an incompatible S2 (location: right, response: left), which typically results in long RTs and many errors on S2-according to control theories the result of a lack of suppression of the automatic route (e.g. Stürmer et al., 2002). However, according to binding-theories of attention (Treisman, 1996), the rotation would cause the binding representing the right circle to be updated, so that at the time point of S2 presentation it would refer to a left circle (Kahneman et al., 1992). If CSEs reflect the benefits and costs of the repetition and alternation of feature bindings (Hommel et al., 2004), this should produce one of the two outcome patterns. On the one hand, it might be that S2 processing is affected by the updated binding only. In this case, one would expect a complete reversal of the pattern obtained in the static condition, as turning left into right, and vice versa, should render compatible transitions incompatible, and vice versa. On the other hand, it may also be that updating a binding does not overwrite the original binding altogether, so that S2 processing should reflect a mixture of compatible and incompatible transitions. As this mixture should be the same for all conditions, one would expect that CSEs are eliminated in this condition. Indeed, this is the pattern that was observed by Spapé and Hommel (2011), who therefore preferred the mixture hypothesis. They were also able to demonstrate independent contributions from original and updated bindings with rotations of 90°, which rules out the less theoretically interesting possibility that rotations simply erase existing bindings and/or the aftereffects of control processes.

Even though the elimination of the CSE by means of controlunrelated manipulations provides a strong challenge of the control account of sequential effects, converging evidence seems necessary to provide positive support for the binding account. Given that the prediction of a behavioral null effect remains somewhat unsatisfactory in principle, we sought for more insights by using EEG recordings, especially with regard to the temporal dynamics of rotation-induced effects on response tendencies as measured by LRPs. We thus computed LRPs for all compatibility sequences as a function of rotation. In line with previous findings (for an overview, see Praamstra, 2007), we predicted activation of the incorrect response as a result of incompatibility. Following Stürmer et al. (2002), this "invalid" activation was expected to be reduced after incompatible trials in the static condition. Similarly, the N2 component was predicted to be greater for incompatible following compatible trials as compared to incompatible following incompatible trials. However, both effects were expected to be reduced or even reversed in sign in the rotation condition. In order to determine whether this could be due to proactive conflict-monitoring mechanisms (cf. Botvinick, 2007), we also analyzed the S1 stimuluslocked LRPs. If the conflict-monitoring mechanism would adapt during the rotation, this was predicted to reduce amplitudes of the LRPs collected during this time-frame. On the other hand, if the LRPs of rotating trials would not differ from those of static trials up until the S2 was presented, binding-retrieval mechanisms that were prompted by the onset of S2 could be considered as the responsible mechanism.

M.M. Spapé et al. / Biological Psychology 88 (2011) 116-123



Fig. 1. Schematic depiction of the trial-sequence of two example trials. After presenting a fixation crosshair, two boxes were presented for 500 ms in the left and right of the screen, one containing the shape (S1) to which participants were required to respond. In the "static" condition (left), an inter-stimulus interval (ISI) followed in which the boxes stood still for 800 ms, whereas in the rotating condition, they rotated around the center of the screen during this ISI. In both conditions, the boxes were statically presented for another 200 ms before the second target (S2) was shown. S2 was shown for 700 ms before an inter-trial interval of 1100 ms ended the trial.

2. Methods

2.1. Participants

Sixteen students from Leiden University voluntarily participated in this experiment for a small fee or course credits. During the analysis of the LRPs over all compatible S1 conditions, four participants showed no negative LRP during responses, implying that their LRP was not diagnostic of motor preparation, so they were left out from further analysis.

2.2. Apparatus and stimuli

Stimuli were presented on a flat-screen 17" CRT monitor in 800 × 600 pixel resolution and a refresh-rate of 120 Hz. A Pentium-IV 2.60 GHz PC running E-Prime 1.2 on Windows XP SP2 controlled stimulus-presentation and recorded reactions via serial response boxes mounted on the armrests, left and right of the participant. The visual boxes that contained the targets were gray (RGB value of 128, 128, 128), black-edged squares of 60×60 pixels or an approximate visual angle of 2.4° presented against a silver (RGB value of 191, 191, 191) background. The target also measured 60×60 pixels and was either a circle or a four-pointed star. Boxes were presented 180 pixels (approximately 7.3°) left and right from the center of the screen and kept at this distance during the gradual shifts in location.

EEG was recorded at 512 Hz from seven Ag/AgCl scalp electrodes, positioned on the Fz, FCz, Cz, CPz, Pz, C3 and C4 locations, mounted in an elastic cap, using the Biosemi Active Two recording system. Common Mode Sense and Driven Right Leg electrodes (see www.biosemi.com/faq/cms&drl.htm) were used for online grounding and as initial reference, but the signal was re-referenced to the average mastoid signal off-line. Bipolar recording from approximately 1 cm above and below the left eye, and 1 cm lateral of the outer canthi of the eyes provided vertical and horizontal electrooculograms (EOGs), respectively. EEG and EOG were passed through a low-cut filter of 0.10 Hz and corrected for eye-movements using the Gratton et al. (1983) algorithm, after which they were passed through high-cut filters of 8 Hz (for LRPs) or 16 Hz (for N2). After this, an average of 8.8% (\pm 4.1%) of trials was excluded from analysis because of EEG or EOG artifacts.

2.3. Procedure

As outlined in Fig. 1, a fixation cross was presented for 500 ms, after which the two boxes were presented in the left and right of the screen, one of them containing the target shape (S1) to which participants were required to respond. After approximately 606 ms, the targets disappeared. The empty boxes stayed on the screen for another 818 ms, either in the same position (the "static" condition) or being rotated around the center of the screen at a speed of approximately 4 degrees per screen update of 44 ms (the "rotated" condition). Then the second target (S2) was presented for 700 ms before a screen with feedback informed the participant about his or her performance. This last screen disappeared after 1100 ms and then the next trial started.

Participants were instructed to ignore the location of the stimuli but react to their shapes using their index fingers. Half of the participants were to press the left response key for stars and the right key for circles, whereas the other half received the opposite stimulus–response mapping. They were required to respond as quickly and accurately as possible and during the ITI they were shown a personal score next to a high score, which they were encouraged to break. Getting points could only be done by responding both fast (1 point for each reaction below 600 ms) and accurately (1 point for each accurate reaction) and although breaking the high score was not reinforced with any kind of monetary or other incentive, most participants

did indicate being positively motivated to aim for the (fictional, computed as $3 \times$ number of trial-pairs) high score. Participants received a break after about every quarter of the trials. The experimented lasted about 100 min in total.

2.4. Design

Results were coded for S1 and S2 compatibility (vs. incompatibility) and rotation (static vs. rotated). The design was fully balanced, randomized within 18 blocks of 64 trials, which resulted from the orthogonal combination within blocks of the two S1 and S2 locations and shapes, and the two types and directions (clock- or counterclockwise) of rotation.

LRPs were calculated by averaging the differences between contralateral and ipsilateral ERP for left and right responses to S2. Magnitudes of *Gratton-dips* (after Gratton et al., 1988) – thought to reflect the stimulus-induced activation of incorrect hand responses – were measured as the maximally positive local voltage between 100 and 200 ms. To explore the effect of rotation on the LRP, and to see whether, for instance, the response activation itself might be affected by rotation, the effects of rotation on S1 response encoding were investigated by calculating LRPs for S1 and the average difference between static and rotating conditions in the entire time-window in which rotation could be present (i.e. between 604 and 1426 ms after S1 onset).

N2 amplitude was measured as the difference in Cz amplitude between S2 compatible and incompatible conditions, the onset and offset of which was measured as the negative area of the FCz, Cz and CPz electrodes between 232 and 356 ms. This window corresponded with the period of a significant (>4 SD of baseline activity) difference between the grand average ERPs of compatible and incompatible conditions.

3. Results

Responses with latencies below 50 ms and above 1000 ms were not considered, and all incorrect reactions to S1 or S2 were excluded from RT and ERP analyses. There were 6.9% errors to S1 (SD = 2.3%) and 5.5% errors to S2 following a correct response to S1 (SD = 2.7%).

3.1. Behavioral results

Table 1

Behavioral results.

11)=79.89, p < .001, $\eta^2 = .89$, and error %, F(1, 11) = 47.42, p < .001, $\eta^2 = .81$. The CSE pattern of 89 ms/8.3% that was found under static conditions (see Table 1 for calculus), broke down after rotation to values of -4 ms/-0.4%. Two ANOVAs testing the static and rotating conditions separately revealed that the S1-by-S2 compatibility interaction was only significant for RTs, F (1, 11) = 111.43, p < .001, $\eta^2 = .92$, and for error % F(1, 11) = 37.51, p < .001, $\eta^2 = .76$, of static conditions, but not for RTs, p > .4, $\eta^2 = .04$ or error %, p > .7, $\eta^2 = .02$, of rotating conditions.

3.2. S2 EEG results

In repeated measures ANOVAs on the LRP Gratton-dip magnitude with S1 compatibility, S2 compatibility and rotation as factors, neither rotation nor S1 compatibility produced a significant main effect, ps > .8. However, S2 compatibility strongly affected the Gratton-dip magnitude, F(1, 11) = 39.78, p < .001, $\eta^2 = .78$. Incompatible trials showed activations of the incorrect hand of ca. $1.4 \,\mu V$ compared to compatible trials. S1 compatibility did not interact significantly with S2 compatibility, p > .9. This appeared to be due to the interaction with the rotation factor, as evidenced by a significant three-way interaction with rotation, F(1, 11) = 13.46, p < .004, η^2 = .55. It indicated a change in sign for the static CSE (tested with a separate ANOVA as F(1, 11) = 3.90, p = .08) versus rotating CSE (separate ANOVA *F*(1, 11) = 11.40, *p* < .01). That is, in the static condition, the Gratton-dips were reduced after incompatible trials by 0.8 µV (compare the two upper panels in Fig. 2), whereas in the rotated condition, the dips increased by $0.8\,\mu\text{V}$ (compare the two lower panels in Fig. 2).

Further ANOVAs on the Cz mean amplitude of the N2 (computed as the difference between incompatible and compatible S2s) with S1 compatibility and rotation as factors showed that S1 compatibility decreased mean N2 activity, F(1, 11) = 34.48, p < .001, $\eta^2 = .76$. Rotation itself did not have a significant effect, p > .5, $\eta^2 = .03$, but did interact with S1 compatibility, F(1, 11) = 23.73, p < .001, η^2 = .68. We conducted two post-hoc tests to better understand this interaction. In post-hoc comparisons between the S1 compatible and incompatible conditions, the difference was shown to be strongly present in static conditions, t(11) = 5.61, p < .001, but not significantly in rotating conditions, t(11) = 1.10, p > .2. That is, in static trials, following incompatible S1s, the N2 difference changed sign (compatible showed larger N2 amplitudes than incompatible trials), while in rotating trials, no such change was present. Another set of post-hoc comparisons, now between the static and rotation conditions, revealed that there was no significant difference between the S1 compatible conditions, t(11) = 1.64, p > .1, but only between the S1 incompatible conditions, t(11) = 4.10, p < .005.

	S2 compatible	S2 incompatible	Simon effect	CSE
Reaction times (ms)				
Static				
S1 compatible	341 (9)	411 (11)	70	
S1 incompatible	381 (10)	362 (9)	-19	89
Rotating				
S1 compatible	356 (9)	390 (10)	34	
S1 incompatible	355 (9)	393 (10)	38	-4
Error rates (%)				
Static				
S1 compatible	9.3 (0.3)	12.3 (3.0)	4.0	
S1 incompatible	6.3 (0.9)	2.0 (1.0)	-4.3	8.3
Rotating				
S1 compatible	2.5 (0.5)	6.6 (1.7)	4.1	
S1 incompatible	2.0 (0.5)	5.7 (1.2)	3.7	-0.4

Reaction times, error rates and standard errors (in parentheses) for the second Stimulus (S2) as a function of its compatibility, preceding (S1) compatibility and rotation. Effect sizes to the right show the Simon effect and how it is affected by preceding (S1) compatibility. The CSE is measured as the degree to which the Simon effect of S2 is attenuated after incompatible S1s.

M.M. Spapé et al. / Biological Psychology 88 (2011) 116-123



Fig. 2. Effect of current (S2) compatibility on the LRP as a function of rotation and preceding (S1) compatibility. Reaction times (RT) are averaged across the two conditions in each panel. Horizontal lines show latency where initial 'incorrect' activation – or, 'Gratton dip' – of the LRP was detected. Dotted vertical lines are the RTs.

Thus, the difference between static and rotation in N2 magnitude was only found after incompatible conditions. Table 2 shows that these results in the N2 component based on the Cz measurements were generally similar in the other midline (FCz, CPz) electrodes (Figs. 3 and 4).

3.3. S1 ERP results

A repeated measures ANOVA on the mean LRP amplitude between 604 (rotation onset) and 1426 (rotation offset) with S1 compatibility and rotation as factors revealed that after incom-

Table 2

ERP results.

	Simon effect (I-C) after			CSE, conflict after C–I
	Rotation	Compatible	Incompatible	
LRP				
Gratton dip latency (ms)	Static	24	-17	41
	Rotating	64	65	-1
Gratton dip magnitude (μV)	Static	1.7	1.1	0.8
	Rotating	0.9	1.9	-0.8
May IRD latency (mc)	Static	95	-35	130
Wax LKF latency (IIIS)	Rotating	100	62	38
	N2 (I-C) component after			CSE, N2 after C–I
	Rotation	Compatible	Incompatible	
ERP				
Mean amplitude FCz (μV)	Static	-1.2	1.7	-2.9
	Rotating	-0.8	-0.7	-0.1
Mean amplitude Cz (μV)	Static	-2.6	1.4	-4.0
	Rotating	-1.0	-0.8	-0.2
Moon amplitude $P_{\mathcal{T}}(uV)$	Static	-2.8	1.1	-3.9
Mean amplitude PZ (µV)	Rotating	-1.9	-1.5	-0.4

Average effect sizes for lateralized readiness potential (LRP) and individual electrodes, stimulus-locked to the second stimulus (S2). Gratton Dip refers to the positive peak in the LRP at ca. 150 ms after stimulus, which is thought to be associated with the automatic (i.e. location-based) activation of the response. Maximum LRP latency was measured as the most negative peak, just prior to the produced response. N2 voltages were calculated as the difference between compatible and incompatible conditions in mean amplitude of the FCz, Cz and Pz electrodes between 232 and 356 ms after S2 onset. Effect of CSE was measured as the difference in compatibility as a function of preceding compatibility for both rotating and static conditions.

Author's personal copy

M.M. Spapé et al. / Biological Psychology 88 (2011) 116-123

Fig. 3. Left panel: ERPs over Cz for compatible (C) and incompatible (IC) conditions as a function of preceding compatibility (C->) and incompatibility (IC->) for both static and rotating conditions. Right panel: N2 difference waves over Cz (subtraction of compatible from incompatible conditions) as a function of preceding compatibility and rotation.

patible S1s, the incorrect response became more activated, F(1, 11)=10.52, p < .01, in a manner best described as an 'echo' of the Gratton-dip caused by S1. This is, at the present time, difficult to explain theoretically, as it could reflect, as was mentioned in the discussion on horizontal Simon effects before, both visual and/or manual deactivation after incompatibility. More importantly, rota-

Fig. 4. Effects of rotation and compatibility on the LRP of the first stimulus (S1).

tion did not affect LRP, p > .4, nor did it interact with compatibility, p > .4.

4. Discussion

The aim of this study was to test whether what looks like effects of control might be due to the impact of previously created stimulus-response bindings on present performance. We argued that if the CSEs that are taken to diagnose the impact of control could be eliminated entirely, and perhaps even reversed in sign, by means of a manipulation that is arguably unrelated to any control process; this would provide strong evidence for a binding approach to CSEs. We applied this logic to the Simon task that was previously demonstrated to produce particularly strong CSEs in both RTs and LRPs. We attempted to replicate these previous findings in our static condition and, indeed, the standard patterns were obtained: notably, the Simon effect was reduced after incompatible trials, as witnessed by reduced RT effects and less stimulus-induced activation of incorrect responses as indexed by LRPs (e.g. Stürmer et al., 2002).

Not so for rotating conditions however: Closely replicating the behavioral observations of Spapé and Hommel (2011), CSE patterns disappeared in the rotation condition. As the electrophysiological findings demonstrate, this was not due to the overwriting of control effects or the flushing of control systems (Logan and Gordon, 2001). Rather than just disappearing in the rotation condition, the 'incorrect' part of the LRP that is commonly taken to indicate the automatic, stimulus-induced activation of responses was strongly affected.

As mentioned before, some caution is required in the interpretation of the ERPs, because of the horizontal stimulus layout. The early parts of the LRP (i.e. the dip) are very likely to be contaminated by volume conducted lateralized components of the (stimulusrelated) N1 and P1. Importantly, however, the effect of rotation on the CSEs can safely be interpreted, since the contamination of the LRP by lateral stimulus presentation, if any, is identical in static and rotation conditions. Thus, any effect of rotation of the placeholders on LRP needs to be attributed to changes in the mental representation of stimuli, rather than to the physical stimuli as such.

A control account would need to explain why a task-unrelated rotation would suddenly cause non-conflicting stimuli to activate incorrect responses. As this seems rather difficult and counterintuitive, we suggest attributing the impact of trial transitions on LRPs to binding effects entirely. As pointed out by Hommel et al. (2004) and explained in Footnote 1, what looks like adaptation effects may also result from a confound of CSEs and the repetition versus alternation of stimulus and response aspects. In particular, stimulus and response features may be bound upon S1 processing and the resulting bindings be automatically retrieved upon S2 processing (Hommel, 2004). If the combination of stimulus and response features is the same in both trials, or if no feature overlaps, performance is unimpaired or even facilitated, but with partial overlap, code conflict occurs and performance suffers (Hommel, 1998).

In the rotation condition, bindings are updated to reflect the spatial change (Kahneman et al., 1992; Zacks and Swallow, 2007; Spapé and Hommel, 2010), so that "left" codes become "right" codes, and vice versa. This change can account for the reversals we have seen in the LRP data. However, the overall pattern across behavioral and psychophysiological data does not show a complete reversal, confirming the claim of Spapé and Hommel (2011) that overt performance on S2 is affected by both the original (not-updated) binding and the updated binding, resulting in a mixture of effects ranging from elimination to reversal. The existence of two bindings seems indeed mirrored in the N2 results. Whereas in static conditions, the N2 only appears after compatible trials, in rotation conditions, it appears regardless of the preceding conflict. One may thus speculate that the presence of two separate bindings results in additional, irrelevant information or response alternatives that would require a need for adaptive control.

Whether and how the two bindings interact, how strong their relative contribution to present performance is, and whether this relative strength depends on context and task requirements, remains to be determined. As long as this is so, it seems logically impossible to exclude contributions from control in principle. More detailed and more quantitative predictions from both binding and control approaches would be necessary to conclusively decide on this issue. We do emphasize, however, that none of our findings is predicted by a control approach and that our outcome does not leave much space, if any, for contributions from control processes to CSEs.

References

- Akçay, Ç., Hazeltine, E., 2007. Conflict monitoring and feature overlap: two sources of sequential modulations. Psychonomic Bulletin & Review 14, 742–748.
- Botvinick, M.M., Nystrom, L.E., Fissell, K., Carter, C.S., Cohen, J.D., 1999. Conflict monitoring versus selection-for-action in anterior cingulate cortex. Nature 402, 179–181.
- Botvinick, M.M., 2007. Conflict monitoring and decision making: reconciling two perspectives on anterior cingulate function. Cognitive, Affective, & Behavioral Neuroscience 7, 356–366.
- Coles, M.G.H., Gratton, G., Bashore, T.R., Eriksen, C.W., Donchin, E., 1985. A psychophysiological investigation of the continuous flow model of human information-processing. Journal of Experimental Psychology: Human Perception and Performance 11, 529–553.
- De Jong, R., Liang, C.-C., Lauber, E., 1994. Conditional and unconditional automaticity: a dual-process model of effects of spatial stimulus-response correspondence. Journal of Experimental Psychology: Human Perception and Performance 20, 721–750.

- Duncan, J., 1996. Cooperating brain systems in selective perception and action. In: Inui, T., McClelland, J.L. (Eds.), Attention and Performance XVI. MIT Press, Cambridge, MA, pp. 549–578.
- Dutzi, I.B., Hommel, B., 2009. The microgenesis of action-effect binding. Psychological Research 73, 425–435.
- Eriksen, B.A., Eriksen, C.W., 1974. Effects of noise letters upon the identification of a target letter in a non-search task. Perception & Psychophysics 16, 143–149. Folstein, J.R., Van Petten, C., 2008. Influence of cognitive control and mismatch on
- the N2 component of the ERP: a review. Psychophysiology 45, 152–170.
- Fournier, L.R., Scheffers, M.K., Coles, M.G.H., Adamson, A., Vila, E., 1997. The dimensionality of the Eriksen compatibility effect: a psychophysiological analysis. Psychological Research 60, 145–155.
- Gratton, G., Coles, M.G., Donchin, E., 1983. A new method for off-line removal of ocular artifact. Electroencephalography and clinical Neurophysiology 55, 468–484.
- Gratton, G., Coles, M.G.H., Donchin, E., 1992. Optimizing the use of information: strategic control of activation of responses. Journal of Experimental Psychology: General 121, 480–506.
- Gratton, G., Coles, M.G.H., Sirevaag, E.J., Eriksen, C.W., Donchin, E., 1988. Pre- and poststimulus activation of response channels: a psychophysiological analysis. Journal of Experimental Psychology: Human Perception & Performance 14, 331–344.
- Hommel, B., 1997. Interactions between stimulus-stimulus congruence and stimulus-response compatibility. Psychological Research 59, 248–260.
- Hommel, B., 1998. Event files: Evidence for automatic integration of stimulusresponse episodes. Visual Cognition 5, 183–216.
- Hommel, B., 2000. The prepared reflex: automaticity and control in stimulus-response translation. In: Monsell, S., Driver, J. (Eds.), Control of Cognitive Processes: Attention and Performance XVIII. MIT Press, Cambridge, MA, pp. 247–273.
- Hommel, B., 2004. Event files: Feature binding in and across perception and action. Trends in Cognitive Sciences 8, 494–500.
- Hommel, B., Proctor, R.W., Vu, K.-P.L., 2004. A feature-integration account of sequential effects in the Simon task. Psychological Research 68, 1–17.
- Kahneman, D., Treisman, A., Gibbs, B.J., 1992. The reviewing of object files: objectspecific integration of information. Cognitive Psychology 24, 175–219.
- Kok, A., Ramautar, J.R., de Ruiter, M.B., Band, G.P.H., Ridderinkhof, K.R., 2004. ERP components associated with successful and unsuccessful stopping in a stopsignal task. Psychophysiology 41, 9–20.
- Leuthold, H., Schröter, H., 2006. Electrophysiological evidence for response priming and conflict regulation in the auditory Simon task. Brain Research 1097, 167–180.
- Logan, G.D., Gordon, R.D., 2001. Executive control of visual attention in dual-task situations. Psychological Review 108, 393–434.
- Mayr, U., Awh, E., Laurey, P., 2003. Conflict adaptation effects in the absence of executive control. Nature Neuroscience 6, 450–452.
- Nieuwenhuis, S., Yeung, N., Van den Wildenberg, W., Ridderinkhof, K.R., 2003. Electrophysiological correlates of anterior cingulate function in a go/no-go task: effects of response conflict and trial-type frequency. Cognitive, Affective, and Behavioral Neuroscience 3, 17–26.
- Notebaert, W., Verguts, T., 2008. Cognitive control acts locally. Cognition 106, 1071–1080.
- Praamstra, P., 2007. Do's and don'ts with lateralized event-related brain potentials. Journal of Experimental Psychology: Human Perception and Performance 33, 497–502.
- Rösler, F., Finger, T., 1993. A psychophysiological analysis of outcome states and response channel activation in Eriksen's noise-compatibility task. Psychological Research 55, 20–28.
- Sasaki, K., Gemba, H., Nambu, A., Matsuzaki, R., 1993. No-go activity in the frontal association cortex of human subjects. Neuroscience Research 18, 249–252.
- Simon, J.R., Rudell, A.P., 1967. Auditory S–R compatibility: the effect of an irrelevant cue on information processing. Journal of Applied Psychology 51, 300–304.
- Spapé, M.M., Hommel, B., 2010. Actions travel with their objects: evidence for dynamic event files. Psychological Research 74, 50–58.
- Spapé, M.M., Hommel, B., 2011. Sequential modulations of the Simon effect depend on episodic retrieval. Manuscript submitted for publication.
- Stürmer, B., Leuthold, H., Schröter, H., Soetens, E., Sommer, W., 2002. Control over location-based response activation in the Simon task: behavioral and electrophysiological evidence. Journal of Experimental Psychology: Human Perception and Performance 28, 1345–1363.
- Stroop, J.R., 1935. Studies of interference in serial verbal reactions. Journal of Experimental Psychology 18, 643–662.
- Treisman, A.M., 1996. The binding problem. Current Opinion in Neurobiology 6, 171–178.
- Verguts, T., Notebaert, W., 2008. Hebbian learning of cognitive control: dealing with specific and nonspecific adaptation. Psychological Review 115 (2), 518–525.
- Watanabe, J., Sugiura, M., Sato, K., Sato, Y., Maeda, Y., Matsue, Y., Fukuda, H., Kawashima, R., 2002. The human prefrontal and parietal association cortices are involved in no-go performances: an event-related fMRI study. NeuroImage 17, 1207–1216.
- Wiegand, K., Wascher, E., 2005. Dynamic aspect of stimulus-response correspondence: evidence for two mechanisms involved in the Simon effect. Journal of Experimental Psychology: Human Perception and Performance 31, 453–464.
- Woodworth, R.S., 1938. Experimental Psychology. Holt, Rinehart and Winston, New York.
- Zacks, J.M., Swallow, K.M., 2007. Event segmentation. Current Directions in Psychological Science 16, 80–84.