# Response processing during visual search in normal aging: the need for more time to prevent cross talk between spatial attention and manual response selection

Elena Amenedo<sup>a</sup>, Laura Lorenzo-López<sup>b</sup>, Paula Pazo-Álvarez<sup>a</sup>

#### Abstract

It is still not well known whether the age-related behavioural slowing observed during visual search is due to changes in the allocation of attention, in response activation patterns, or to a combination of both. To help in clarifying it, attention-related (N2 posterior contralateral; N2pc, and N2 central contralateral; N2cc) and response-related (Motor Potential; MP, and Reafferent Potential; RAP) event-related potentials (ERPs) were obtained in healthy young and older participants executing a visual search task. Age was associated with N2pc and N2cc longer latencies, earlier MP onsets and longer MP rise times. Lower N2pc, higher MP and lower RAP amplitudes were also observed. Results suggest that older participants need more time to allocate spatial attention onto the target (N2pc) and to prevent cross talk between response selection and attention direction (N2cc), and that they are slower and need higher cortical activation when preparing and executing correctly selected responses (MP).

## Keywords

Aging; Visual search; Response-processing; Attention ERPs; Motor ERPs

<sup>&</sup>lt;sup>a</sup> Department of Clinical Psychology and Psychobiology, Faculty of Psychology, University of Santiago de Compostela, Spain

 $<sup>^</sup>b$  Gerontology Research Group, Department of Medicine, Faculty of Health Sciences, University of A Coruña, Spain

#### 1. Introduction

The visual search paradigm has been extensively used in laboratory studies to examine the basic properties of human visual selective attention (Luck and Ford, 1998; Luck and Hillyard, 1994a,b). In this paradigm, participants search for a predefined target stimulus that is randomly presented in arrays containing a variable number of bilateral distractor stimuli. In a typical single feature visual search task, like that employed in the present study, the target differs from distractors in a physical attribute (i.e. orientation change), and the participants are required to indicate whether the target is present or absent in each array of stimuli by pressing a pre-assigned response button.

Previous electrophysiological studies of visual search in young humans have revealed a posterior negative-going ERP component that appears contralateral to the visual hemifield in which the target is located 200–300 ms after the onset of a bilateral stimulus array (Eimer, 1996; Luck and Hillyard, 1994a,b; Woodman and Luck, 1999). This component was first described as the N2pc (N2-posterior-contralateral) to indicate its polarity, latency range, and scalp distribution (Luck and Hillyard, 1994a), and it has been well-validated as an electrophysiological correlate of the focusing of visuospatial attention onto a target stimulus during visual search (Eimer, 1996; Luck and Hillyard, 1994a,b; Luck and Ford, 1998; Luck et al., 1997; Woodman and Luck, 1999, 2003). The N2pc is recorded with maximum amplitude over parietal-occipital electrode sites contralateral to the attended item (Eimer, 1996; Luck and Hillyard, 1994a,b), and it is primarily generated in lateral occipital–temporal regions (Hopf et al., 2000; Lorenzo-López et al., 2011).

There is considerable evidence that older adults are behaviourally slower in tasks involving visual search (Hommel et al., 2004; Madden and Whiting, 2004; McDowd and Shaw, 2000). However, several questions remain unsolved regarding the interpretation of these findings. Mainly, it is still not well established if they represent a slowing related to deficiencies in the allocation of attention onto the target, to deficient response activation patterns, or to a combination of them. During the period between the onset of the target in the visual field and the emission of a correct response, several processes related to both attention and motor mechanisms take place. Specifically, allocation of attention onto the target and motor selection and preparation processes both contribute to the reaction time (RT) recorded during the execution of the task, and any or both of them may be affected by age.

Recent studies have helped in solving the first of the above questions by providing the first evidence that normal aging significantly affects the allocation of visuospatial attention itself during single feature visual search (N2pc component, Lorenzo-López et al., 2008a) by delaying the time needed to allocate spatial attention shifts onto the target and reducing the attention resources deployed to it. More recently, these age-related changes in N2pc parameters have been associated with a significant hypoactivation of its occipital—temporal neural sources that is more marked in the right hemisphere (Lorenzo-López et al., 2011). However, it is still not possible to conclude whether the age-related behavioural changes observed during visual search tasks are also due to changes in the processes related to the selection and preparation of motor responses to the correct detection of the target.

In recent years, an ERP component related to the prevention of cross talk between attention direction and manual response selection has been described (Oostenveld et al., 2001; Praamstra, 2006; Praamstra and Oostenveld, 2003). This component, named N2cc (N2 central-contralateral) is recorded at central electrodes contralateral to the side of presentation of the target stimuli during visuospatial attention tasks, and it presents the same polarity and latency as the above-described N2pc component. Based on results from single cell recordings on non-human primates (Crammond and Kalaska, 2000; Shen and Alexander, 1997; Wise et al., 1996, 1997), from neuroimaging data (Connolly et al., 2000; Dassonville et al., 2001) and from ERP current source analyses (Praamstra, 2006; Praamstra and Oostenveld, 2003), N2cc has been proposed to reflect activation of the dorsal premotor cortex that is invoked to prevent the selection of a manual response depending on the location of the target stimulus in the visual field, and hence to ensure that the response selection is not biased by the direction of spatial attention (Praamstra and Oostenveld, 2003). More recently, the fact that this component is reduced in amplitude when advance information of target location is provided by a cue has added support to its functional interpretation (Praamstra, 2006). In

visual search tasks, the target stimulus appears randomly interspersed with distractor stimuli in the right or left visual field in each trial. In these tasks, it is frequent to assign one hand to respond to the presence of the target and the other to its absence independently of its position in the visual field. Under such circumstances, responses executed with the hand ipsilateral to the visual hemifield where the target appears are facilitated. Thus the ability to select the correct response hand depending on the task instructions and not on the position of the target in the right or left visual field can be considered an executive function that helps inhibiting the vulnerability of response choice to be influenced by the direction of attention (Praamstra, 2006). In this regard, measuring the N2cc component makes it possible to explore the effects of aging on this executive function during visual search. Moreover, although the N2pc component has been well characterized in older samples (Lorenzo-López et al., 2008a, 2011), it was also examined in the present study in order to compare age-related effects on stimulus-related attention ERPs with those on response-related attention ERPs (Praamstra and Oostenveld, 2003).

The observation of slower RTs in older populations is not limited to visual search tasks, but is also present in almost every study on aging. In fact, the ubiquity of this age-related effect has lead to influential theories of cognitive aging and to classical debates in this research context (Salthouse, 1996; McDowd and Shaw, 2000). Some recent ERP evidence has indicated that cortical dysregulation in motor activation patterns may underlie response slowing with age in choice reaction tasks (Falkenstein et al., 2006; Yordanova et al., 2004), although to our knowledge the mechanisms underlying the behavioural slowing during visual search tasks have not been examined yet.

ERP research has shown that any task requiring an overt response to a stimulus activates electrocortical motor mechanisms immediately after or even in parallel with stimulus processing. Before a correct response is overtly executed, several ERP components associated to movement selection and preparation processes are generated in motor cortical areas. One of the most prominent ERPs associated with active limb movements is the Readiness Potential (RP; Deecke et al., 1969; Böcker et al., 1994; Shibasaki and Hallett, 2006). The RP develops through different components among which the most prominent for finger movements is a maximum negative deflection appearing just before the observable movement onset at contralateral central electrodes (Motor Potential, MP) followed by a positive deflection that is maximal around the overt movement execution (Reafferent Potential, RAP). The MP indexes motor generation of a selected response at contralateral motor cortex (Böcker et al., 1994). The functional significance of the RAP, although less well known, has been related to sensory-motor integration processes during response execution (Bötzel et al., 1997; Seiss et al., 2002; Szurhaj et al., 2006). Thus, measuring the MP and the RAP components elicited during correct responses to the target stimulus in a visual search task should allow examining motor generation and sensory-motor integration processes that take place between target onset and response execution, and their age-related changes.

Finally, in order to examine the relations between the recorded response-related electrophysiological activity and the behavioural performance during the task independently of the effects of age, partial correlations (Baron and Kenny, 1986; Perry et al., 2009; Volkow et al., 1998) were computed between the N2cc and RP (MP and RAP) parameters (latency and amplitude values) and the execution data (RTs and hit rates).

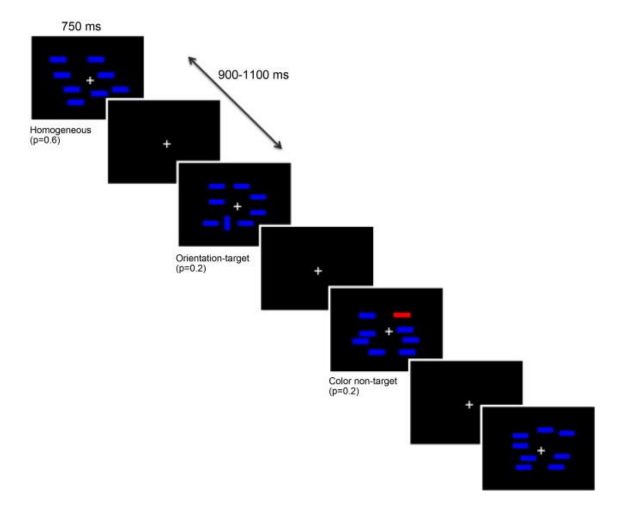
#### 2. Methods

## 2.1. Participants

The original sample consisted of 17 young (10 females,  $19.6 \pm 1.9$  years, range 18-24) and 22 older adults (11 females,  $68.5 \pm 6$  years, range 60-84). However, four young (all female) and five older (4 female) participants were discarded from the original sample because of excessively noisy EEG motor-related activity that led to low signal-to-noise ratios for ERP-averaging purposes. Finally, data from 13 young (6 females,  $20.08 \pm 2.02$  years, range 18-24), and 17 older adults (8 females,  $20.08 \pm 2.02$  years, range  $20.08 \pm 2.02$  years, range 20

## 2.2. Stimuli and experimental procedure

Recordings were made in an electrically shielded and sound attenuated room. Subjects sat in a comfortable armchair at 100 cm viewing distance from a computer screen with a black background and a continuously visible fixation white cross. They were instructed to maintain central fixation on this cross while they performed a visual search task consisting in detecting a target stimulus presented among an array of distractors that differed from them in its orientation (Fig. 1). On each trial a multi-element search array was presented composed of eight bars subtending a visual angle of 0.3° × 0.9°, which were located at random positions within an imaginary rectangle of  $9.2^{\circ} \times 6.9^{\circ}$  of visual angle around fixation cross. There were always four bars in each visual hemifield. Three types of search arrays were randomly presented: homogeneous arrays, arrays containing a target defined by a change in orientation of one of the bars, and arrays containing a non-target defined by a change in colour of one of the bars. Homogeneous arrays (p = .6) consisted of eight blue-horizontal (RGB 0, 0, 255) identical bars. Orientation target arrays (p = .2) consisted of seven blue-horizontal bars and one blue-vertical bar. Colour non-target arrays (p = .2) consisted of seven blue-horizontal bars and one red-horizontal (RGB 255, 0, 0) bar. The orientation target and the colour non-target were equally likely to appear in the right or left visual hemifield and their location was unpredictable. Each search array was presented for 750 ms, followed by a variable inter-trial interval of 900-1100 ms during which only fixation cross was present. The same feature (orientation) defined the target across all trials and the subjects were not informed about the appearance of the colour non-target. All the stimuli and search arrays were created, presented, and controlled using the Presentation software application (Neurobehavioral Systems, Inc., version 0.76). The experimental session was divided into six blocks of trials, and several training trials were run before testing to ensure a good level of performance in both age groups. Each block consisted of at least 10 arrays containing an orientation target and 10 arrays containing a colour non-target presented to each hemifield, and at least 80 homogeneous arrays, to a possible maximum of 250 arrays in total. Participants were required to indicate as rapidly and accurately as possible whether the orientation target (a vertical bar) was present or absent in each search array by pressing a button with one hand for target-present trials and another button with the other hand for target-absent trials. Thus, the colour non-target arrays required the same response hand as the homogeneous arrays (target-absent). Response buttons were counterbalanced across subjects. As a result, in 16 participants (7 young and 9 older) the right hand was assigned to respond to target-present arrays and the left hand to target-absent arrays, and in 14 participants (6 young and 8 older) the left hand was assigned to target-present arrays and the right hand to target-absent arrays.



 $\textbf{Fig. 1}. \ Graphic \ description \ of \ task \ properties.$ 

# 2.3. ERP recordings

The electroencephalogram (EEG) was recorded with a NeuroScan system using scalp electrocaps (ECI, Inc.) with 30 electrodes placed at FP1, FP2, FPz, Fz, Cz, Pz, POz, Oz, F7, F8, F3, F4, C3, C4, T3, T4, PO3, PO4, FCz, CPz, CP3, CP4, T5, T6, P3, P4, FC3, FC4, O1 and O2 (10/20 International System). All the active electrodes were referred to the nose tip and grounded with an electrode placed at nasion. Vertical and horizontal electrooculogram (EOG) activities were recorded bipolarly from above and below the left eye and from the outer canthi of both eyes. Electrode impedances were kept below  $10~\rm k\Omega$ . The EEG signals were continuously amplified (10 K) and digitized at a rate of 500 Hz/channel, and filtered on-line with a band pass of  $0.05-100~\rm Hz$ .

## 2.4. Data analysis

## 2.4.1. Behavioural data

Reaction times (RTs) were on-line recorded for all subjects to the three types of search arrays in all experimental blocks (for orientation target and colour non-target arrays, RTs to stimuli appearing in the right or left visual field were separately recorded). Only RT values associated with correct responses were considered for data analyses. Hit rates were calculated as the total percentage of correct responses with RTs no longer than 1100 ms. A common problem with using percentages as response variables is that the distribution of proportions may "pile-up" against 0 or 1, resulting in skewed data (i.e. non-normal distributions). Additionally this may lead to unequal variances. For meeting the assumptions of ANOVA, the original hit rates were firstly converted to proportions and then arcsine root transformed (Osborne, 2002). Mean correct RTs were compared across groups using a repeated measures analysis of variance (ANOVA) with age (young, older) and response hand (right, left) as the between-subjects factors, and search array (homogeneous, orientation right-left target, and colour right-left non-target) as the within-subject factor. Original and transformed hit rates were also compared across groups using ANOVAs with age (young, older) and response hand (right, left) as the between-subjects factors. An alpha level of .05 was used for all analyses.

#### 2.4.2. EEG data

All EEG data were off-line processed using Vision Analyzer software (Version 2.0). The EEG was digitally filtered with a 0.1–30 Hz band pass filter. Filtered EEG was segmented in epochs of 900 ms post-stimulus and 100 ms pre-stimulus to obtain attention-related ERPs (N2pc and N2cc, see below), and in epochs of 1000 ms post-response and 1000 ms pre-response for the motor-related RP components (MP and RAP, see below). In all segmentation files, epochs exceeding  $\pm 100~\mu V$  and those containing blinks, and horizontal or vertical eye movements were rejected and excluded from averaging, as well as epochs associated with incorrect or no responses. Because in previous research by our group no differential ERP effects related to attentional focusing were observed to colour non-target stimuli (Lorenzo-López et al., 2008a), only EEG segments associated with correct responses to orientation target stimuli were averaged for both attention ERPs (N2pc and N2cc) and motor ERPs (RP). Separate averages were obtained to orientation targets appearing in the right visual field (RVF) and to those appearing in the left visual field (LVF). This resulted in 4 attention-related and 2 response-related waveforms for each participant (see below).

2.4.2.1. Attention-related ERP data: N2pc and N2cc components. For the derivation of the N2pc component, difference waveforms for each participant were obtained by subtracting ipsilateral from contralateral ERPs relative to the target location in the right or the left visual field at posterior electrodes PO3 and PO4 respectively (see Lorenzo-López et al., 2008a,b). For the derivation of the N2cc component, difference waveforms for each participant were obtained by subtracting ipsilateral from contralateral ERPs relative to the visual field of attention following Praamstra (2006) formula that reads  $N2cc = [(C3 - C4)_{RVF \text{ attention}} + (C4 + C3)_{LVF \text{ attention}}]/2$ , where 'RVF' refers to the right visual field, and 'LVF' to the left visual field. This procedure extracts lateralized potentials collapsing the activity of both hemispheres (see Praamstra, 2006). In the resulting waveforms of both N2pc and N2cc components, mean amplitude values were obtained from 150 to 300 ms for the young adults, and from 200 to 450 ms for the older adults (these latency intervals were selected as the most representative for each group after visual inspection of the respective grand mean waveforms, see Figs. 2 and 3, and Lorenzo-López et al., 2008a,b). In order to test the effects of age on N2pc and N2cc parameters, peak amplitude, mean amplitude, and latency values of these components were entered into separate ANOVAs. For N2pc values, mixed ANOVAs with age (young, older) as the between-subjects factor and electrode (PO3, PO4) as the within-subject factor were run. For N2cc values, one-way ANOVAs with age (young, older) as the between-subjects factor were executed. An alpha level of .05 was used.

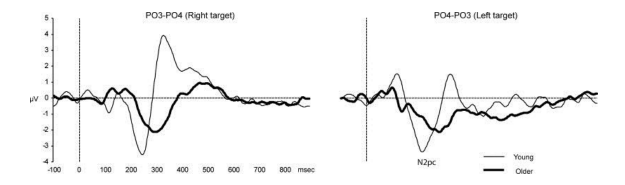


Fig. 2. N2pc component. Subtraction waveforms superimposed for young and older participants.

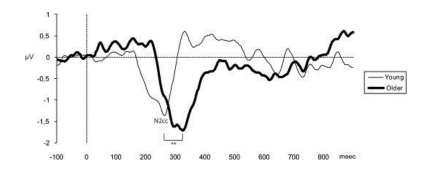


Fig. 3. N2cc component obtained by the formula 'N2cc =  $[(C3 - C4)_{RVF \ attention} + (C4 + C3)_{LVF \ attention}]/2$ '. Subtraction waveforms superimposed for young and older participants.

2.4.2.2. Response-related ERP data: MP and RAP components. RP averages to orientation targets were analysed at relevant electrodes at the contralateral motor areas (C3 for right hand responders, and C4 for left hand responders). Baseline was defined in these ERPs as a 200 ms interval between 1000 and 800 ms before the motor response (button press). Peak latency and amplitude values of the most negative displacement of the RP, the MP, were measured within the interval between stimulus presentation and response execution. Peak latency and amplitude values were also measured for the most positive displacement of the RP, the RAP, after the MP. The onset latency of the MP was calculated as the time when MP amplitude was 15% of its maximal value (cf. Mordkoff and Gianaros, 2000; Schwarzenau et al., 1998). The duration of the MP activation (MP rise time) was measured as the difference between MP peak latency and MP onset latency.

MP and RAP peak latency and amplitude values, and MP onset and rise time values were submitted to separate repeated measures ANOVAs with age (young, older) and response hand (right, left) as the between-subjects factors, and target-response compatibility (compatible target-response side, incompatible target-response side) as the within-subject factor. An alpha level of .05 was used for all statistical tests. Whenever appropriate, degrees of freedom were corrected by the conservative Greenhouse–Geisser estimate. When necessary, post hoc comparisons were performed using the Bonferroni adjustment for multiple comparisons.

2.4.2.3. ERP-behaviour analyses. To examine the relationships between behavioural execution and electrophysiological activity independently of the effects of age, partial correlation analyses controlling for age (Baron and Kenny, 1986; Perry et al., 2009; Volkow et al., 1998) were executed between both RT and hit rates, and the latency and amplitude values measured for N2cc, MP and RAP components. Because no differences between RTs to right and left targets, and no interactions between response hand, age or array type were observed (see results), the mean RT values to both target stimuli were employed to compute partial correlations between N2cc and RT. In the case of motor components (MP and RAP) the amplitude and latency values obtained to compatible target stimuli were used for the correlation analyses. Partial correlations were considered significant at p < .05 (two-tailed).

#### 3. Results

#### 3.1. Behavioural results

As can be appreciated in Table 1, the RTs were significantly slower in the older group for each array type  $(F(1,26) = 33.69, p < .0001, n_p^2 = .56)$  irrespective of the hand assigned to respond to the targets  $(F(1,26)=.38,\ p=.54,\ n_p^2=.01)$ . The effect of the array type was also significant on mean RTs  $(F(4,104) = 29.29, p < .0001, \varepsilon = .43, n_p^2 = .53)$  showing the longest response times for orientation targets, intermediate for colour non-targets, and the shortest RTs for the homogeneous arrays (see Table 1). No statistically significant interactions were observed between age and response hand (F(1,26) = 3.11, $p = .09, n_p^2 = .11$ ), between age and array type ( $F(4,104) = .58, p = .54, n_p^2 = .02$ ) or between response hand and array type (F(4,104) = .80, p = .44,  $n_p^2 = .03$ ). Moreover, target-response compatibility effects were not significant on RT values  $(F(1,28) = .61, p = .44, n_p^2 = .02)$  in either young (target-response compatible:  $500.55 \pm 51.61$  ms, incompatible:  $500.50 \pm 49.97$  ms) or older participants (target-response compatible:  $595.72 \pm 55.61$  ms, incompatible:  $604.34 \pm 42.07$  ms). Effects of age (F(1,26) = 3.77,p = .06,  $n_p^2 = .13$ ) or response hand  $(F(1,26) = .16, p = .70, n_p^2 = .006)$  were not statistically significant on hit rates (young right hand:  $97.58 \pm 2.68\%$ , left hand:  $98.55 \pm .36$ ; older right hand:  $96.51 \pm 3.23\%$ , left hand:  $94.50 \pm 5.40\%$ ). No significant interactions between age and response hand were observed  $(F(1,26) = 1.28, p = .27, n_p^2 = .05)$ . The results of the ANOVA executed on the arcsine root transformed data revealed significant effects of age that indicated worse performance levels in the older participants  $(F(1,26) = 4.73, p < .04, n_p^2 = .15)$ . No significant effects of the response hand (F(1,26) = .15, p = .70, p = .70) $n_p^2 = .006$ ) or of the interaction between age and response hand  $(F(1,26) = 1.26, p = .27, n_p^2 = .05)$  were observed on these transformed data.

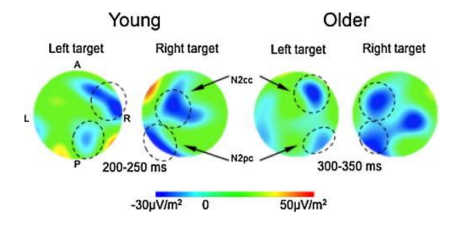
Table 1. Mean values (standard deviation) of RT (ms) across array types and assigned response hand in each age group.

	Homogeneous	Colour right	Colour left	Target right	Target left
Right hand	d				
Young	454.86	469.10	471.91	512.67	513.26
	(70.51)	(77.37)	(72.49)	(54.31)	(55.24)
Older	538.44	545.72	556.21	569.78	593.82
	(57.61)	(57.90)	(62.76)	(58.31)	(41.87)
Left hand					
Young	438.45	453.53	449.15	485.61	486.42
	(52.54)	(55.08)	(49.59)	(42.85)	(49.06)
Older	588.83	596.70	589.88	628.67	624.90
	(51.77)	(47.09)	(48.34)	(36.98)	(36.67)

## 3.2. Electrophysiological results

## 3.2.1. Attention-related ERPs: N2pc and N2cc components

Significant effects of age were observed on both N2pc (Figs. 2 and 4) and N2cc (Figs. 3 and 4) peak latencies (N2pc: F(1,28) = 37.69, p < .0001,  $n_p^2 = .69$ ); N2cc: F(1,28) = 16.56, p < .0001,  $n_p^2 = .49$ ) indicating longer values in the older group (see Table 2). N2pc peak amplitude (F(1,28) = 5.6, p < .02,  $n_p^2 = .17$ ) and mean amplitude (F(1,28) = 4.67, p < .04,  $n_p^2 = .32$ ) were significantly lower in the older participants, whereas N2cc peak amplitude (F(1,28) = .05, p = .82,  $n_p^2 = .06$ ) and mean amplitude (F(1,28) = .57, p = .46,  $n_p^2 = .01$ ) were not affected by age, although they showed a slight trend to higher values in the older group (see Table 2 and Fig. 3).



**Fig. 4.** Current source density maps (spherical spline interpolation, Perrin et al., 1989). The maps show central and posterior negative maxima corresponding to N2cc and N2pc components in each age group (highlighted with dashed black circles). At anterior; P: posterior: L: left; R: right.

Table 2. Mean values (standard deviation) of N2pc and N2cc components across age groups.

N2pc PO3–PO4 (right target)		N2pc PO4–PO3 (left target)		N2cc	
oung	Older	Young	Older	Young	Older
48.31 (16.08)	304.12 (35.11)	236.31 (26.14)	302.12 (39.42)	247.69 (25.74)	317.18 (42.27)
4.46 (3.87)	-3.10 (1.74)	-3.97 (3.27)	-2.98 (1.96)	-1.46 (1.05)	-1.72 (1.22)
1.40 (2.75)	-0.79 (1.33)	-1.93 (2.59)	-1.31 (1.36)	-0.80 (0.67)	-0.98 (0.69)
4	oung 8.31 (16.08) 4.46 (3.87)	oung Older  18.31 (16.08) 304.12 (35.11)  14.46 (3.87) -3.10 (1.74)	oung Older Young  8.31 (16.08) 304.12 (35.11) 236.31 (26.14)  8.46 (3.87) -3.10 (1.74) -3.97 (3.27)	Oung         Older         Young         Older           8.31 (16.08)         304.12 (35.11)         236.31 (26.14)         302.12 (39.42)           4.46 (3.87)         -3.10 (1.74)         -3.97 (3.27)         -2.98 (1.96)	oung         Older         Young         Older         Young           8.31 (16.08)         304.12 (35.11)         236.31 (26.14)         302.12 (39.42)         247.69 (25.74)           4.46 (3.87)         -3.10 (1.74)         -3.97 (3.27)         -2.98 (1.96)         -1.46 (1.05)

#### 3.2.2. Response-related motor ERP: MP and RAP components

The results on the motor MP component showed higher peak amplitudes, earlier onset latencies and longer durations in the older participants independently of the hand assigned to respond to the target, and to target-response compatibility effects. Specifically, the older participants showed significantly higher amplitudes of the MP component  $(F(1,26) = 4.15, p < .05, n_p^2 = .15)$  than the young participants (Table 3 and Fig. 5). MP amplitudes were significantly higher in target-response compatible trials (F(1,26) = 5.16, p < .03,  $n_p^2 = .17$ ) irrespective of the hand assigned to respond to the orientation target (F(1,26) = .97,p = .34,  $n_p^2 = .04$ ). No significant interactions were observed on this parameter ('age by response hand': F(1,26) = 1.37, p = .25,  $n_p^2 = .05$ ; 'target-response compatibility by age': F(1,26) = .81, p = .38,  $n_p^2 = .25$ .03; 'target-response compatibility by response hand': F(1,26) = .64, p = .43,  $n_p^2 = .02$ ). Similar values of MP peak latency were observed for both young and older participants (F(1,26) = .03, p = .87,  $n_p^2 = .001$ ). The spatial compatibility between target location and response hand had no significant effects on peak latency values of the MP  $(F(1,26) = .28, p = .60, n_p^2 = .01)$ . However, the hand assigned to respond influenced the results  $(F(1,26) = 4.57, p < .04, n_p^2 = .15)$  indicating that MP peaked slightly earlier in participants who responded to the target with their right hand (see Table 3, and Fig. 5). Again, no significant interactions between age and response hand  $(F(1,26) = .35, p = .87, n_p^2 = .01)$ , age and targetresponse compatibility  $(F(1,26) = .64, p = .43, n_p^2 = .02)$  or response hand and target-response compatibility  $(F(1,26) = .77, p = .39, n_p^2 = .03)$  were found. MP onset showed significant differences  $(F(1,26) = 16.84, p < .0001, n_p^2 = .39)$  between young and older participants indicating that in the older participants the MP began earlier (Table 3). Target-response compatibility effects were significant in both age groups  $(F(1,26) = 42.12, p < .0001, n_p^2 = .62)$ , and no differences as a function of the hand employed to respond to the target were observed in any age group on this parameter (F(1,26) = .50, p = .48, $n_p^2 = .02$ ). Specifically, irrespective of the hand assigned to respond, MP started earlier for targetresponse incompatible trials than for target-response compatible trials in all participants ('target-response compatibility by age' F(1,26) = .47, p = .50,  $n_p^2 = .02$ ) indicating the need for more time to prepare the correct response when target appeared in the location opposite to the response hand. No interactions were found between age and response hand  $(F(1,26) = .04, p = .84, n_p^2 = .002)$  or between response hand and target-response compatibility (F(1,26) = .79, p = .38,  $n_p^2 = .03$ ). MP rise time was also significantly affected by age  $(F(1,26) = 18.94, p < .0001, n_p^2 = .42)$  showing longer duration for the older compared to the young participants (Table 3, and Fig. 5) irrespective of the hand assigned to respond to the target  $(F(1,26) = .47, p = .50, n_p^2 = .02)$ . Moreover, MP rise times were shorter when targets appeared in the visual field compatible with the response side  $(F(1,26) = 27.73, p < .0001, n_p^2 = .52)$  in all participants ('target-response compatibility by age' F(1,27) = .06, p = .81,  $n_p^2 = .002$ ) and irrespective of the response hand ('target-response compatibility by response hand' F(1,26) = 3.10, p = .09,  $n_p^2 = .11$ ; 'age by response hand'  $F(1,26) = .71, p = .41, n_p^2 = .03$ ).

**Table 3**. Mean amplitude and latency values (standard deviation) of RP components at the contralateral electrode (C3 for right hand; C4 for left hand) across assigned response hand in each age group.

Hand	Age grou	up RP con	nponent	Compatible target	Incompatible target
Right	Young	MP	Peak amplitude (μV)	-5.21 (2.87)	-3.55 (3.09)
rugiit	roung	1111	Peak latency (ms)	-267.14 (49.43)	-287.43 (115.07)
			Onset latency (ms)	-293.71 (44.95)	-419.43 (84.41)
			Rise time (ms)	55.14 (75.35)	132.00 (57.60)
		RAP	Peak amplitude (μV)	19.18 (7.23)	14.22 (8.79)
		10.11	Peak latency (ms)	-47.14 (32.22)	-50.57 (46.40)
	Older	MP	Peak amplitude (μV)	-11.53 (6.34)	-7.91 (5.39)
	Older	1.11	Peak latency (ms)	-261.78 (66.98)	-310.89 (112.16)
			Onset latency (ms)	-389.11 (120.96)	-502.00 (76.80)
			Rise time (ms)	127.34 (87.77)	211.12 (96.58)
		RAP	Peak amplitude (μV)	6.44 (7.63)	5.82 (7.03)
		11	Peak latency (ms)	-88.23 (37.67)	-75.12 (43.95)
			•		
Left	Young	MP	Peak amplitude (μV)	-8.32 (1.91)	-7.62 (3.69)
			Peak latency (ms)	-261.34 (85.50)	-227.67 (54.62)
			Onset latency (ms)	-275.00 (93.37)	-397.34 (63.49)
			Rise time (ms)	13.67 (9.07)	166.67 (62.49)
	]	RAP	Peak amplitude (μV)	18.61 (7.41)	12.07 (15.65)
			Peak latency (ms)	-38.00 (28.40)	-27.50 (29.62)
	Older	MP	Peak amplitude (μV)	-10.32 (6.86)	-8.50 (6.05)
			Peak latency (ms)	-220.00 (128.84)	-236.50 (27.79)
			Onset latency (ms)	-337.75 (41.05)	-530.50 (104.80)
			Rise time (ms)	117.75 (120.93)	286.75 (96.27)
	]	RAP	Peak amplitude (μV)	2.92 (5.59)	3.47 (5.93)
			Peak latency (ms)	-37.75 (53.65)	-100.50 (64.48)
			* * /	,	, ,

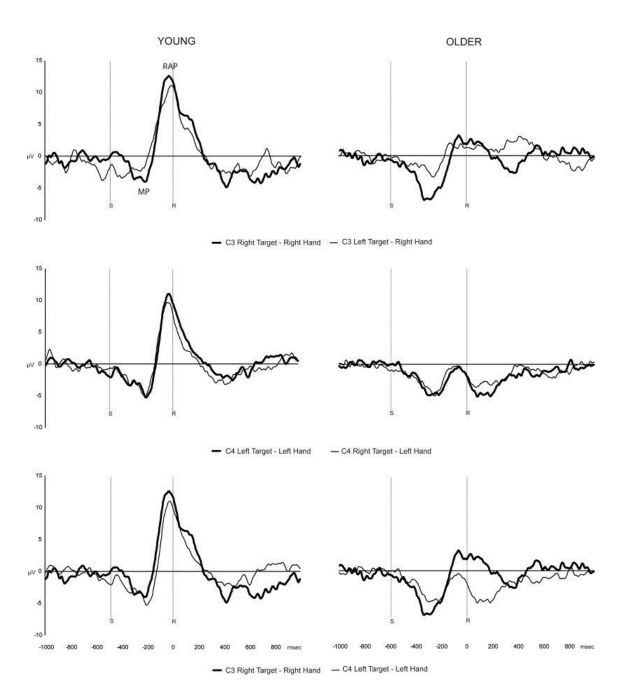


Fig. 5. Motor ERPs (MP and RAP) in young (left column) and older (right column) participants. Top and middle panels: target-response compatibility effects for right and left hand responders. Lower panel: assigned hand effects shown for compatible trials. S: target stimulus onset. R: response execution (button pressure). The waveforms were low-pass filtered at 10 Hz for visualization purposes only.

The RAP component showed effects of both age and response hand on its amplitude. Specifically, RAP amplitudes  $(F(1,26) = 32.78, p < .0001, n_p^2 = .56)$  were significantly lower in the older than in the young group (Table 3 and Fig. 5). Moreover, RAP amplitudes differed in all participants as a function of the hand used to respond to the target  $(F(1,26) = 5.56, p < .03, n_p^2 = .18)$  showing slightly larger amplitudes for participants using their right hand (Table 3). Target-response compatibility effects were not observed on the amplitude of this component  $(F(1,26) = .03, p = .86, n_p^2 = .001)$ , and no interactions were observed ('age by response hand' F(1,26) = .18, p = .68,  $n_p^2 = .007$ ; 'age by target-response compatibility' F(1,26) = .29, p = .59,  $n_p^2 = .01$ ; 'response hand by target-response compatibility' F(1,26) = 2.67, p = .11,  $n_p^2 = .09$ ). Finally, no significant effects of age  $(F(1,26) = 3.41, p = .08, n_p^2 = .09)$ .12), response hand  $(F(1,26) = .94, p = .34, n_p^2 = .03)$  or target-response compatibility  $(F(1,26) = .49, n_p^2 = .03)$ p = .49,  $n_p^2 = .02$ ) were observed on the latency of the RAP component, and no interactions between any of the factors were found ('age by response hand' F(1,26) = .002, p = .97,  $n_p^2 = .00$ ; 'age by targetresponse compatibility' F(1,26) = .79, p = .38,  $n_p^2 = .03$ ; 'response hand by target-response compatibility'  $F(1,26) = .92, p = .34, n_p^2 = .03$ ).

## 3.3. ERP-behaviour relationships

Partial correlation analyses showed that, once the effects of age were taken into account significant relationships emerged between behavioural (RT and hit rates) and response-related electrophysiological (N2cc, MP and RAP) measures (Table 4). Specifically, RT was positively correlated to N2cc peak latency, N2cc peak amplitude and MP peak amplitude, and negatively correlated to N2cc mean amplitude. On the other hand, hit rate was negatively correlated to MP and RAP peak latencies, and MP rise time (Table 4). Regarding RT-N2cc correlations, these results indicated that longer RT values were associated with longer N2cc latencies, higher N2cc peak amplitudes, and lower N2cc mean amplitudes. The correlation between RT and MP peak amplitude showed that slower responses to targets were associated with higher amplitudes of the MP. Finally, partial correlations indicated that higher hit rates were related to earlier MP latencies and shorter MP rise times, and to earlier RAP peak latencies (Table 4).

Table 4. Partial correlation values between behavioural and electrophysiological measures with age as control variable.

	N2cc latency (ms)	N2cc peak amplitude (μV)	N2cc mean amplitude (μV)	$\begin{array}{c} MP \ peak \\ amplitude \ (\mu V) \end{array}$	MP peak latency (ms)	MP rise time (ms)	RAP peak latency (ms)
RT (ms) Hit rate	0.36*	0.49*	-0.55**	0.37*			
(%)					-0.57**	-0.44*	-0.78***

p < .01 (27 two-tailed degrees of freedom).

<sup>\*\*</sup> p < .001 (27 two-tailed degrees of freedom).

p < .0001 (27 two-tailed degrees of freedom).

#### 4. Discussion

In the present study, stimulus and response-related ERPs were recorded while healthy young and older adults searched for a singleton feature target defined by an orientation change relative to bilateral surrounding distractor stimuli. The main objective was to complement previous research on age-related changes in visual search (Lorenzo-López et al., 2008a, 2011) by examining the possibility that electrocortical activity related to motor and sensory-motor processing is contributing to the slowing of behavioural response observed in visual search tasks. Specifically, to address age-related changes in attention-related ERPs associated with the prevention of manual responses determined by the direction of attention the N2cc component was examined (see Praamstra, 2006). To address movement preparation and execution-related ERP activity, the MP and RAP components of the RP were analysed (Shibasaki and Hallett, 2006). Moreover, the possible relationships between the above referred electrocortical activity and behavioural performance (RT and hit rate) were also examined independently of the effects of age.

#### 4.1. Behavioural results

Behavioural results revealed an age-related delay in RTs, which is in line with those of our previous (Lorenzo-López et al., 2008a, 2011) and other behavioural studies showing that older participants are slower than young participants in tasks involving visual search (Hommel et al., 2004; Madden and Whiting, 2004; McDowd and Shaw, 2000). There was an effect of age on hit rates once the original data were arcsine root transformed. This result, according to previous studies (Hommel et al., 2004; Lorenzo-López et al., 2008a,b, 2011; Madden and Whiting, 2004; McDowd and Shaw, 2000), indicates that older adults executed the search task at lower accuracy levels than young participants.

## 4.2. Electrophysiological results

The electrophysiological results on the attention-related components showed that age was significantly associated with longer latencies of both N2pc and N2cc. Moreover, in agreement with previous results (Lorenzo-López et al., 2008a) lower peak amplitudes and mean amplitudes of N2pc were also observed in the elderly group of the present study. N2cc amplitude values, although showing a slight trend to be higher in the older participants, were not significantly affected by age. Regarding the results on the motor ERPs, the older participants showed higher amplitudes, earlier onsets and longer rise times of MP, along with lower amplitudes of RAP. Moreover, these effects of age on MP and RAP components were independent of the hand assigned to respond to the target, and of target-response compatibility effects.

## 4.2.1. Attention-related ERPs: N2pc and N2cc components

In visual spatial attention tasks, correctly detected targets appearing at lateral locations elicit two contralateral negative deflections at the same latencies, one that is recorded over parietal-occipital electrodes (the N2pc, Luck and Ford, 1998; Luck and Hillyard, 1994a,b) and another that is recorded over central electrodes (the N2cc, Oostenveld et al., 2001; Praamstra and Plat, 2001). The N2pc is a well-established electrophysiological index of the deployment of visuospatial shifts of attention onto the location of a target stimulus that is generated in extrastriate occipital–temporal cortical areas (Hopf et al., 2004; Lorenzo-López et al., 2011). The present results on N2pc indicate, in line with our previous findings (Lorenzo-López et al., 2008a,b, 2011), that the older participants shift their attentional focus slower, and deploy less attentional resources onto the location of the target stimulus during visual search.

The N2cc deflection has been described more recently (Oostenveld et al., 2001; Praamstra and Plat, 2001) and it has been proposed as an attention-related motor component that reflects the activation of dorsal premotor cortex to suppress or prevent the emission of incorrect responses promoted by targets appearing at locations ipsilateral to the responding hand (Praamstra and Oostenveld, 2003; Praamstra,

2006). The inhibition of preeminent responses determined by spatial stimulus-response compatibility effects is considered an important part of the executive control system that promotes correct and flexible responding (Miller and Cohen, 2001; Ridderinkhof et al., 2004). The longer N2cc peak latencies observed in the older participants of the present study, considering the functional significance proposed for this component, indicated a slower attention-related activation of dorsal premotor areas in order to prevent the emission of incorrect responses promoted by the spatial location of the target stimuli in the visual field (i.e. right responses to right targets in right-hand responders, and left responses to left targets in left-hand responders). In this regard, this finding would suggest an age-related need of additional time to inhibit automatic location-based response activation due to poorer visuospatial motor processing. Moreover, although it did not reach statistical significance, the N2cc tended to have higher amplitudes in the older participants, suggesting a trend to a higher premotor activation to help in preventing such incorrect responses. Van der Lube and Verleger (2002) compared young and older adults in a visual Simon task and found a trend to lower N2pc-like and higher N2cc-like amplitudes (in their study Posterior Contralateral Negativity, PCN, and pre-Lateralized Readiness Potential, pre-LRP, respectively) in the older group. Based on previous results on Parkinsońs Disease (see Praamstra and Plat, 2001) they suggested that this age-related trend to lower PCN and larger pre-LRP might be due to an increased direct visuomotor transmission from posterior to anterior brain areas that would decrease inhibitory control of incorrect responses along a direct visuomotor pathway. In the present study, lower amplitudes of N2pc along with a trend to higher amplitudes of N2cc have been observed in the older group. These results complement those by Van der Lube and Verleger (2002) and therefore suggest a poorer attentional sensory-motor mapping during the visual search that could promote the selection of responses ipsilateral to the location of the target independently of being the stimulus associated whit the correct response. This trend would hinder the selection of correct responses, manifest in the present study by a slowness of the N2cc component that may be underlying the longer RTs observed in the aged participants.

It could be argued that the task employed in the present study hindered obtaining the N2cc, because this component has been observed to be attenuated when the task does not involve competing response alternatives to the target stimulus defined at opposite hands (Praamstra and Oostenveld, 2003) as in the present task, where only one hand was assigned to respond to the target. However, an N2cc component with similar amplitudes to that obtained in response-choice tasks (Praamstra, 2006; Praamstra and Oostenveld, 2003) was observed in the present study in all subjects. The reason may be that the design of the present task implied compatible and incompatible target-response mappings during the visual search process (i.e. in each trial the target could appear equiprobably in the right or left visual field, and depending on the hand assigned to respond to the target, it might appear at locations spatially compatible or incompatible to the response side). Moreover, participants had to select and prepare a correct response to a target surrounded by several distractor stimuli appearing randomly in different locations of the visual field. In such situation, it is essential to be able to effectively combine different sources of information for accurate and appropriate responses. One of those sources of information comes from the place where the target appears in each trial in the visual field, which promotes the shift of attention to its location. Another source is related to the motor significance of the target, that is, the target instructs a response that must be executed with the hand assigned to it independently of its location in the visual field. Therefore, under such circumstances, a proper inhibition of incorrect responses based on the location of the target is needed, which would promote dorsal premotor cortex activation and N2cc generation (see Praamstra, 2006).

## 4.2.2. Response-related motor ERPs: MP and RAP components

The results on the motor MP component showed higher peak amplitudes, earlier onset latencies and longer durations in the older participants independently of the hand assigned to respond to the target, and to target-response compatibility effects. The MP is generated in the contralateral primary motor cortex, and it is related to the preparation and initiation of the motor command resulting in an overt response (Böcker et al., 1994; Deecke et al., 1969; Shibasaki and Hallett, 2006). Previous reports on motor ERPs have showed higher age-related amplitudes of this component in choice-reaction tasks (Falkenstein et al., 2006; Yordanova et al., 2004), and studies employing fMRI have found additional age-related activation

in the contralateral motor cortical areas during the performance of simple movements (Mattay et al., 2002; Ward and Frackowiak, 2003) and in areas associated to suppression of preeminent response tendencies and inhibitory cognitive control both during simple and complex movements (Heuninckx et al., 2005). According to the above reports, the age-related changes observed in the amplitude, onset latency and rise time of the MP component in this study indicate that older adults need higher, earlier and longer activation of the contralateral motor cortex to prepare and execute correctly selected responses to the targets during the visual search task, thus complementing the scarce existing data on response-related brain activity on aging.

A new finding in the present study was that age also affected the amplitude of the RAP component that was markedly reduced in the older participants independently of the hand employed to respond and to the target-response compatibility effects. To our knowledge, this is the first study having tested agerelated changes in this motor component. The origins and functional significance of this component are still not well known although recent reports have shown evidence of a relationship between RAP and sensory-motor integration during movement execution. Specifically, more classical studies had suggested that the RAP was related to post-movement reafferent sensory-motor processes generated in somatosensory cortical areas (Bötzel et al., 1997; Shibasaki et al., 1980). This interpretation, however, received little support (for a review see Shibasaki and Hallett, 2006) until recent years when studies employing intracortical recordings in humans found clear relationships between RAP amplitude and gamma event-related synchronization (ERS) in the 40-60 Hz band at contralateral sensorimotor cortices (Szurhaj et al., 2005, 2006). Initial descriptions of gamma band oscillations related them to multiple sensory and cognitive processes, but recent studies associate 40-60 Hz rhythms in sensorimotor cortical regions with the somesthetic reafferentation from the muscles and joints necessary for controlling the ongoing movement (see Szurhaj and Derambure, 2006). Szurhaj et al. (2005) interpreted the relationship between the amplitude of RAP component and gamma ERS as indicating that this component is functionally related to the facilitation of afferences from muscles and joints involved in the movement to the motor cortico-spinal cells, necessary for controlling the ongoing movement (Szurhai et al., 2005, 2006). The lower RAP amplitudes observed in the older group of the present study would therefore suggest a deficient sensory-motor integration and hence a poorer control of movement during the execution of the correct responses in the visual search task.

## 4.2.3. Target-response compatibility effects

Stimulus-response compatibility has been extensively studied. In general, it consists in a shortening of RTs when target stimulus and response are spatially compatible (Hommel, 1993). Even when the spatial location of the target stimulus is irrelevant to the task this effect is clearly seen. Simon and Rudell (1967) described this special compatibility effect with auditory stimulation, and since then it has been called Simon effect. In the present study the relevant feature to the task was target orientation, therefore the compatibility effects here can be attributed to a Simon-like effect. We observed similar RTs in all participants for both target-response compatible and incompatible trials. Moreover, no interactions with age or response hand were observed. The absence of Simon-like effects on RTs in the present study may probably be related to the fact that only one hand was assigned to respond to both right and left targets, reducing the choice component that matches the spatial dimensions of stimulus and response and resulting in similar RTs for both compatible and incompatible trials. In this sense, the Simon effect is normally reduced when the task demands blur the match between the spatial dimensions of stimulus and response (Hommel, 1993; Juncos-Rabadán et al., 2008). Moreover, the lack of age-related differences in compatibility effects may be associated with the stimulus configuration in the present task. In this sense, it has been found that when tasks include irrelevant, spatially varying secondary stimuli accompanying the relevant stimulus (as in the visual search task used in the present study) there are no effects of age on stimulus-response compatibility effects on RTs (Proctor et al., 2005a,b).

Although no effects of target-response compatibility were observed on the RT values, the MP component showed significant modulations as a function of spatial target-response compatibility independently of age and response hand. When target and response were spatially incompatible (left target-right response and vice versa) MP showed lower amplitude, it began earlier and consequently its duration was longer indicating lower and slower electrocortical activation in contralateral motor areas to prepare and execute the correct response in this trials. This result partially agrees with previous reports on stimulus-response compatibility effects observed on lateralized motor ERPs obtained by averaging the contralateral minus the ipsilateral electrocortical activity for the responses executed with both hands (the Lateralized Readiness Potential, LRP, see Coles et al., 1988). In general, in stimulus-response incompatible trials lower amplitudes of the LRP are observed that have been traditionally explained as the result of the automatic initiation of electrocortical activity at the ipsilateral but incorrect hemisphere (see Gratton et al., 1988). The compatibility effects observed on the MP motor component were independent of the age of participants and of the hand assigned to respond to the targets. To our knowledge, only a previous study has explored age-related changes in the Simon effect on motor ERPs. In this study, Van der Lube and Verleger (2002) obtained the LRP during the execution of a visual Simon Task in young and older adults and they did not find effects of age on this ERP component. Although in the present study it was not possible to obtain the LRP as only one hand was assigned to the target presence in the visual field, our results are in line with the absence of age-related differences in the compatibility effects on the LRP observed previously by these authors (Van der Lube and Verleger, 2002).

## 4.2.4. Hand assignment effects

The hand assigned to respond to the target showed significant effects only on MP peak latency and on RAP amplitude, which peaked earlier and presented higher amplitude respectively when participants were assigned the right hand to detect the presence of the target in the visual field. This result, probably related to the handedness of the participants (all right-handed; see Knösche et al., 1996; Kutas and Donchin, 1974) may indicate facilitated motor preparation and sensory-motor integration during response execution when movements were performed with the dominant hand (Knösche et al., 1996). However, there were no significant interactions between response hand and age or target-response compatibility, therefore indicating that earlier MP latencies and larger RAP amplitudes were present in both young and older participants responding with their right hand to all targets (independently of they appearing in the left or in the right visual field).

# 4.3. Response ERPs-behaviour relationships

The results of partial correlation analyses indicated that, independently of age, longer RT values were associated with longer N2cc latencies, higher N2cc peak amplitudes, and lower N2cc mean amplitudes. Moreover, the correlations between RT and MP peak amplitudes showed that slower responses to targets were associated with higher amplitudes of the MP. Finally, partial correlations indicated that higher hit rates were related to earlier MP latencies and shorter MP rise times, and to earlier RAP peak latencies. In general, the significant correlations observed between RTs and response-related ERPs help in explaining the significant age-related effects observed on the response-related components analysed in the present study. Firstly, the association 'slower N2cc - slower RT' makes the explanation given for the age-related longer RTs and later N2cc latencies observed in the present study more feasible since it indicates that slower behavioural responses are related to the need for more time to prevent the selection of responses determined by the location of the target stimulus in the visual field during a single feature visual search task. Secondly, the correlations between RT and N2cc amplitudes indicate that in general those participants responding with slower RTs presented higher and more phasic activation in dorsal premotor areas to successfully inhibit incorrect responses when targets appeared in the same side than the response hand, which would add partial support to the slight trend to higher N2cc amplitudes observed in the older group. Thirdly, the association 'slower RTs - higher MP amplitudes' gives support to the explanation of age-related behavioural slowing as partly due to higher motor cortical activation during the preparation of the selected correct responses (Falkenstein et al., 2006; Yordanova et al., 2004).

#### 5. Conclusions

The results of the present study complement our previous findings on the electrophysiological patterns that underlie the age-related behavioural slowing observed in visual search tasks by showing that slower responses to correctly detected targets in these tasks are also due to (a) slower activation in dorsolateral premotor cortex to inhibit incorrect responses promoted by the target location when selecting the correct ones determined by task instructions, (b) higher and longer activation patterns in motor areas when preparing the correctly selected responses, and (c) poorer sensory-motor mapping during the execution of the correct responses during the task. Moreover, the conjoint examination of attention-related ERP components associated to stimulus (N2pc) and response (N2cc) during the visual search task gives support to a possible age-related increased direct visuomotor transmission from posterior to anterior brain areas that would decrease inhibitory control of incorrect responses.

#### Disclosure statement

There are no actual or potential conflicts of interest that could inappropriately influence this work. Informed consent was obtained from all participants in the experiment and their rights were protected.

#### Acknowledgements

This study was supported by grants from the Spanish MICINN (PSI2010-21427) and Xunta de Galicia (10PXIB211220PR).

#### References

- Baron, R.M., Kenny, D.A., 1986. The moderator-mediator variable distinction in social psychological research: conceptual, strategic, and statistical considerations. Journal of Personality and Social Psychology 51, 1173–1182.
- Böcker, K.B.E., Brunia, C.H.M., Cluitman, P.J.M., 1994. A spatio-temporal dipole model of the readiness potential in humans. I. Finger movement. Electroencephalography and Clinical Neurophysiology 91, 275–285.
- Bötzel, K., Ecker, C., Schulze, S., 1997. Topography and dipole analysis of reafferent electrical brain activity following the Bereitschaftspotential. Experimental Brain Research 114, 352–361.
- Coles, M.G.H., Gratton, G., Donchin, E., 1988. Detecting early communication: using measures of movement-related potentials to illuminate human information processing. Biological Psychology 26, 69–89.
- Connolly, J.D., Goodale, M.A., Desouza, J.F.X., Menon, R.S., Vilis, T., 2000. A comparison of frontoparietal fMRI activation during anti-saccades and anti-pointing. Journal of Neurophysiology 84, 1645–1655.
- Crammond, D.J., Kalaska, J.F., 2000. Prior information in motor and premotor cortex: activity during the delay period and effect on pre-movement activity. Journal of Neurophysiology 84, 986–1005.
- Dassonville, P., Lewis, S.M., Zhu, X.H., Ugurbil, K., Kim, S.G., Ashe, J., 2001. The effect of stimulus–response compatibility on cortical motor activation. NeuroImage 13, 1–14.
- Deecke, L., Scheid, P., Kornhuber, H.H., 1969. Distribution of readiness potential, premotion positivity, and motor potential of the human cerebral cortex preceding voluntary finger movements. Experimental Brain Research 7, 158–168.
- Eimer, M., 1996. The N2pc component as an indicator of attentional selectivity. Electroencephalography and Clinical Neurophysiology 99, 225–234.
- Falkenstein, M., Yordanova, J., Kolev, V., 2006. Effects of aging on slowing of motor-response generation. International Journal of Psychophysiology 59, 22–29.
- Folstein, M.F., Folstein, S.E., McHugh, P.R., 1975. Mini-mental state. A practical method for grading the cognitive state of patients for the clinician. Journal of Psychiatric Research 12 (3), 189–198.
- 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 and Performance 14, 331–344.

- Heuninckx, S., Wenderoth, N., Debaere, F., Peeters, R., Swinnen, S.P., 2005. Neural basis of aging: the penetration of cognition into action control. The Journal of Neuroscience 25, 6787–6796.
- Hommel, B., 1993. The relationship between stimulus processing and response selection in the Simon task: evidence for a temporal overlap. Psychological Research 55, 280–290.
- Hommel, B., Li, K.Z., Li, S.C., 2004. Visual search across the life span. Developmental Psychology 40 (4), 458–545.
- Hopf, J.M., Luck, S.J., Girelli, M., Hagner, T., Mangun, G.R., Scheich, H., Heinze, H.J., 2000. Neural sources of focused attention in visual search. Cerebral Cortex 10, 1233–1241.
- Hopf, J.M., Boelmans, K., Schoenfeld, A.M., Luck, S.J., Heinze, H.J., 2004. Attention to features precedes attention to locations in visual search: evidence from electromagnetic brain responses in humans. Journal of Neuroscience 24 (8), 1822–1832.
- Juncos-Rabadán, O., Pereiro, A.X., Facal, D., 2008. Cognitive interference and aging: insights from a spatial stimulus-response consistency task. Acta Psychologica 127, 237–246.
- Knösche, T., Praamstra, P., Stegeman, D., Peters, M., 1996. Linear estimation discriminates midline sources and a motor cortex contribution to the readiness potential. Electroencephalography and clinical Neurophysiology 99, 183–190.
- Kutas, M., Donchin, E., 1974. Studies of squeezing: handedness, responding hand, response force, and asymmetry of readiness potential. Science 186, 545–548.
- Lorenzo-López, L., Amenedo, E., Cadaveira, F., 2008a. Feature processing during visual search in normal aging: electrophysiological evidence. Neurobiology of Aging 29, 1101–1110.
- Lorenzo-López, L., Amenedo, E., Pascual-Marqui, R.D., Cadaveira, F., 2008b. Neural correlates of age-related visual search decline: a combined ERP and sLORETA study. NeuroImage 41, 511– 524
- Lorenzo-López, L., Gutiérrez, R., Moratti, S., Maestú, F., Cadaveira, F., Amenedo, E., 2011. Agerelated occipito-temporal hypoactivation during visual search: relationships between mN2pc sources and performance. Neuropsychologia 49, 858–865.
- Luck, S.J., Ford, M.A., 1998. On the role of selective attention in visual perception. Proceedings of the National Academy of Sciences of United States of America 95 (3), 825–830.
- Luck, S.J., Hillyard, S.A., 1994a. Spatial filtering during visual search: evidence from human electrophysiology. Journal of Experimental Psychology: Human Perception and Performance 20, 1000–1014.
- Luck, S.J., Hillyard, S.A., 1994b. Electrophysiological correlates of feature analysis during visual search. Psychophysiology 31 (3), 291–308 Luck, S.J.,
- Girelli, M., McDermott, M.T., Ford, M.A., 1997. Bridging the gap between monkey neurophysiology and human perception: an ambiguity resolution theory of visual selective attention. Cognitive Psychology 33, 64–87.
- Madden, D.J., Whiting, W.L., 2004. Age-related changes in visual attention. In: Costa, P.T., Siegler, I.C. (Eds.), Recent Advances in Psychology and Aging. Elsevier, Amsterdam, pp. 41– 88.
- Mattay, V.S., Fera, F., Tessitore, A., Hariri, A.R., Das, S., Callicott, J.H., Weinberger, D.R., 2002. Neurophysiological correlates of age-related changes in human motor function. Neurology 58, 630–635.
- McDowd, J.M., Shaw, R.J., 2000. Attention and aging: a functional perspective. In: Craik, F.I.M., Salthouse, T.A. (Eds.), The Handbook of Aging and Cognition., 2nd ed. Lawrence Erlbaum Associates Publishers, Mahwah, NJ, US, pp. 221–292. Miller, E.K., Cohen, J.D., 2001. An integrative theory of prefrontal cortex function. Annual Review of Neuroscience 24, 167–202.
- Mordkoff, J.T., Gianaros, P.J., 2000. Detecting the onset of the lateralized readiness potential: a comparison of available methods and procedures. Psychophysiology 37, 347–360.
- Oldfield, R.C., 1971. The assessment and analysis of handedness: the Edinburgh inventory. Neuropsychologia 9, 97–113.
- Oostenveld, R., Praamstra, P., Stegeman, D.F., van Oosterom, A., 2001. Overlap of attention and movement-related activity in lateralized event-related brain potentials. Clinical Neurophysiology 112, 477–484.
- Osborne, J., 2002. Notes on the use of data transformations. Practical Assessment. Research & Evaluation 8. 6. 1–9.
- Perrin, F., Pernier, J., Bertrand, O., Echallier, J.F., 1989. Spherical splines for scalp potential and current density mapping. Electroencephalography and Clinical Neurophysiology 72, 174–187 (posterior correction in 1990 in Electroen- cephalography and Clinical Neurophysiology 76, 565)
- Perry, M.E., McDonald, C.R., Hagler Jr., D.J., Gharapetian, L., Kuperman, J.M., Koyama, A.K., Dale, A.M., McEvoy, L.K., 2009. White matter tracts associated with set-shifting in healthy aging. Neuropsychologia 47, 2835–2842.

- Praamstra, P., 2006. Prior information of stimulus location: effects on ERP measures of visual selection and response selection. Brain Research 1072, 153–160.
- Praamstra, P., Oostenveld, R., 2003. Attention and movement-related motor cortex activation: a high-density EEG study of spatial stimulus–response compatibility. Cognitive Brain Research 16, 309–322.
- Praamstra, P., Plat, F.M., 2001. Failed suppression of direct visuomotor activation in Parkinson's disease. Journal of Cognitive Neuroscience 13, 31–43.
- Proctor, R.W., Pick, D.F., Vu, K.L., Anderson, R.E., 2005a. The enhanced Simon effect for older adults is reduced when the irrelevant location information is conveyed by an accessory stimulus. Acta Psychologica 119, 21–40.
- Proctor, R.W., Vu, K.L., Pick, D.F., 2005b. Aging and response selection in spatial choice tasks. Human Factors. The Journal of the Human Factors and Ergonomics Society 47, 250–270.
- Ridderinkhof, K.R., van den Wildenberg, W.P.M., Segalowitz, S.J., Carter, C.S., 2004. Neurocognitive mechanisms of cognitive control: the role of prefrontal cortex in action selection, response inhibition, performance monitoring, and reward- based learning. Brain and Cognition 56, 129–140.
- Salthouse, T.A., 1996. The processing-speed theory of adult age differences in cognition. Psychological Review 103 (3), 403–428.
- Schwarzenau, P., Falkenstein, M., Hoorman, J., Hohnsbein, J., 1998. A new method for the estimation of the onset of the lateralized readiness potential (LRP). Behavioral Research Methods, Instrumentation and Computation 30, 110–117.
- Seiss, E., Hess, C.W., Drane, S., Oostenveld, R., Wing, A.M., Praamstra, P., 2002. Propioception-related evoked potentials: origin and sensitivity to movement parameters. NeuroImage 17, 461–468.
- Shen, L., Alexander, G.E., 1997. Preferential representation of instructed target location versus limb trajectory in dorsal premotor area. Journal of Neurophysiology 77, 1195–1212.
- Shibasaki, H., Hallett, M., 2006. What is Bereitschaftspotential? Clinical Neurophys- iology 117, 2341–2356.
- Shibasaki, H., Barrett, G., Halliday, E., Halliday, A.M., 1980. Cortical potentials following voluntary and passive finger movements. Electroencephalography and Clinical Neurophysiology 50, 201–213.
- 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.
- Szurhaj, W., Derambure, P., 2006. Intracerebral study of gamma oscillations in the human sensorimotor cortex. In: Neuper, C., Klimesch, W. (Eds.), Event-Related Dynamics of Brain Oscillations. Progress in Brain Research, vol. 159. Elsevier, Amsterdam, pp. 297–310.
- Szurhaj, W., Bourriez, J.L., Kahane, P., Chauvel, P., Mauguière, F., Derambure, P., 2005. Intracerebral study of gamma rhythm reactivity in the sensorimotor cortex. European Journal of Neuroscience 21, 1223–1235.
- Szurhaj, W., Labyt, E., Bourriez, J.L., Kahane, P., Chauvel, P., Mauguière, F., Derambure, P., 2006. Relationship between intracerebral gamma oscillations and slow potentials in the human sensorimotor cortex. European Journal of Neuroscience 24, 947–954.
- Van der Lube, R.H.J., Verleger, R., 2002. Aging and the Simon task. Psychophysiology 39, 100-
- Volkow, N.D., Gur, R.C., Wang, G.J., Fowler, J.S., Moberg, P.J., Ding, Y.S., Hitzemann, R., Smith, G., Logan, J., 1998. Association between decline in brain dopamine activity with age and cognitive and motor impairment in healthy individuals. American Journal of Psychiatry 155, 344–349.
- Ward, N.S., Frackowiak, R.S.J., 2003. Age-related changes in the neural correlates of motor performance. Brain 126, 873–888.
- Wise, S.P., Di Pellegrino, G., Boussaoud, D., 1996. The premotor cortex and nonstandard sensorimotor mapping. Canadian Journal of Physiology and Pharmacology 74, 469–482.
- Wise, S.P., Boussaoud, D., Johnson, P.B., Caminity, R., 1997. Premotor and parietal cortex: corticocortical connectivity and combinatorial computations. Annual Review of Neuroscience 20, 25–42.
- Woodman, G.F., Luck, S.J., 1999. Electrophysiological measurement of rapid shifts of attention during visual search. Nature 400, 867–869.
- Woodman, G.F., Luck, S.J., 2003. Serial deployment of attention during visual search. Journal of Experimental Psych: Human Perception and Performance 29, 121–138.
- Yordanova, J., Kolev, V., Hohnsbein, J., Falkenstein, M., 2004. Sensorimotor slowing with ageing is mediated by a functional dysregulation of motor-generation processes: evidence form high-resolution event-related potentials. Brain 127, 351–362