The neural correlates of conceptual and perceptual false recognition

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False recognition, broadly defined as a claim to remember something that was not encountered previously, can arise for multiple reasons. For instance, a distinction can be made between conceptual false recognition (i.e., false alarms resulting from semantic or associative similarities between studied and tested items) and perceptual false recognition (i.e., false alarms resulting from physical similarities between studied and tested items). Although false recognition has been associated with frontal cortex activity, it is unclear whether this frontal activity can be modulated by the precise relationship between studied and falsely remembered items. We used event-related fMRI to examine the neural basis of conceptual compared with perceptual false recognition. Results revealed preferential activity in multiple frontal cortex regions during conceptual false recognition, which likely reflected increased semantic processing during conceptual (but not perceptual) memory errors. These results extend recent reports that different types of false recognition can rely on dissociable neural substrates, and they indicate that the frontal activity that is often observed during false compared with true recognition can be modulated by the relationship between studied and tested items.
underlie the differential effect of age on conceptual versus perceptual false recognition (for evidence that conceptual recognition can be disproportionately impaired compared with perceptual recognition, see also Schacter et al. 1997b, Budson et al. 2003). Moreover, it makes good sense to postulate that conceptual false recognition would be more reliant on frontal processes than perceptual false recognition; frontal activity can reflect semantic processing (e.g., semantic retrieval, semantic elaboration, semantic relational memory) (Wagner et al. 2001; Kirchhoff et al. 2005; Prince et al. 2005), which would presumably accompany conceptual false recognition more often than perceptual false recognition.

Although no prior study has compared the neural processes associated with conceptual and perceptual false recognition, recent work has shown that different types of false recognition can rely on distinct neural substrates. For example, the neural correlates of false recognition can vary depending on the context in which one believes to have studied an item previously (Kahn et al. 2004). Moreover, related false recognition (i.e., "old" responses to related but new items) and unrelated false recognition (i.e., "old" responses to unrelated, new items) have been shown to rely on distinct patterns of neural activity (Garoff-Eaton et al. 2006), which indicates that the relationship between a falsely remembered item and previously studied items must be considered when exploring the neural basis of memory errors. Therefore, it is plausible that the precise nature of the relationship (i.e., conceptual vs. perceptual) between falsely remembered and studied items could also affect the neural correlates of memory errors.

The primary goal of the present study was to examine whether frontal activity would be equally strong during conceptual and perceptual false recognition or whether the frontal cortex would be differentially engaged during conceptual false recognition. To distinguish between these alternatives, we conducted an event-related fMRI study that compared the neural activity associated with conceptual memory errors with the activity associated with perceptual memory errors. During study, participants viewed a series of word triplets. The words in each triplet were related to each other either conceptually (i.e., based on their meaning) or perceptually (i.e., based on their spelling and sound). (It should be noted that we use the term “perceptual” to refer to both visual and phonological properties of the stimuli.) At test, participants viewed a series of single words and decided whether or not they had studied those words previously. These tested words included: (1) old, studied words from conceptual or conceptual triplets; (2) lure words that were related to studied conceptual or perceptual triplets; and (3) new, unrelated words (Fig. 1). Related false recognition occurred when participants endorsed lures as old, and the classification of these errors as either conceptual or perceptual was determined by the relationship of the lure to the related, studied triplet.

**Results**

**Behavioral results** Participants were able to successfully discriminate between studied and nonstudied words on the recognition test (Table 1). Participants were significantly more likely to give either a “remember” or a “know” response to an old conceptual word (0.88) or an old perceptual word (0.74) than they were to give either of those responses to an unrelated new word (0.37; both P < 0.0001). Furthermore, participants were also significantly more likely to give either a “remember” or a “know” response to a conceptual lure (0.62) or a perceptual lure (0.49) than they were to give those responses to an unrelated new word (both P < 0.0001), indicating that the relatedness of the lures did affect participants’ responses in both the conceptual and the perceptual conditions. It is important to note that the old conceptual and conceptual lure conditions elicited significantly more “remember” responses than the old perceptual and perceptual lure conditions, respectively (i.e., “remember”/old conceptual > “remember”/old perceptual, “remember”/conceptual lure > “remember”/perceptual lure; both P < 0.0005). These differences remained even when conditioned scores were compared (i.e., proportion of “remember” responses, given that an item was not called “new” [“remember”/ (“remember” + “know”)]; both P < 0.0005). These results replicate earlier findings that “remember” responses tend to accompany conceptual false recognition more often than perceptual false recognition (Schacter et al. 1997b; Watson et al. 2003). Thus, given that remembering and knowing can rely on distinct patterns of neural activity (Wheeler and Buckner 2004), we chose to limit our fMRI analysis to “remember” responses so that our neuroimaging comparisons of conceptual versus perceptual memory processing would not be contaminated by differences in remembering versus knowing. (Note that because conceptual true and false recognition were accompanied by “remember” responses more often than perceptual true and false recognition, the perceptual conditions are composed of fewer trials than the conceptual conditions.) “Remember” responses also are relatively straightforward to interpret (e.g., a “remember” response to a lure item is a clear indication of a false memory; the person is claiming that they remember the instance in which they were presented with the word), whereas “know” responses may be more likely to include guesses or responses based upon familiarity with related studied items. Importantly, “remember” responses were given to conceptual and perceptual old words and lures more often than they were given to unrelated new words (all P < 0.001). Because most participants gave an insufficient number of “remember” responses to unrelated new words (i.e., unrelated false recognition), we did not include that condition in any fMRI analyses. Furthermore, the set of 14 participants with a

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**Figure 1.** Schematic of the experimental procedure. During the study task, participants viewed word triplets that were either conceptually or perceptually related. Participants were instructed to decide how similar the word in each triplet were to each other while also remembering the words for the subsequent recognition test. During the recognition test, participants viewed five types of words: old conceptual words, old perceptual words, conceptual lures, perceptual lures, and unrelated new words. They gave a “remember,” “know,” or “new” response to each tested word. The condition in which words were tested was counterbalanced across participants by changing the word triplets shown during the study task (e.g., yell could be tested as a perceptual lure [shown here]; as a conceptual lure after studying shout, cry, squeal; as an old perceptual word after studying dell, bell, yell; as an old conceptual word after studying shout, yell, squeal; or as an unrelated new word after studying unrelated triplets).
Table 1. At the top, proportion (mean [SE]) of items given a "remember," "know," or "new" response as a function of item type; at the bottom, response latencies (mean [SE]) associated with each response type as a function of item type.

| Response type       | Old word (conceptual) | Old word (perceptual) | Lure (conceptual) | Lure (perceptual) | New word (unrelated) |
|---------------------|-----------------------|-----------------------|-------------------|-------------------|---------------------|
| **Accuracy**        |                       |                       |                   |                   |                     |
| "Remember"         | 0.71 (0.05)           | 0.37 (0.04)           | 0.35 (0.03)       | 0.17 (0.02)       | 0.09 (0.02)         |
| "Know"             | 0.17 (0.03)           | 0.36 (0.04)           | 0.27 (0.03)       | 0.33 (0.02)       | 0.28 (0.02)         |
| "New"              | 0.10 (0.03)           | 0.26 (0.02)           | 0.37 (0.03)       | 0.49 (0.02)       | 0.62 (0.03)         |
| **Response latencies (msec)** |                   |                       |                   |                   |                     |
| Remember            | 1267 (75)             | 1243 (62)             | 1434 (58)         | 1495 (94)         | 1735 (98)           |
| Know                | 1798 (133)            | 1766 (91)             | 1740 (105)        | 1694 (105)        | 1666 (109)          |
| New                 | 1633 (115)            | 1732 (107)            | 1676 (87)         | 1572 (88)         | 1526 (76)           |

sufficient number of "remember" responses to studied words and lures was not the same set of participants with a sufficient number of "know" responses to studied words and lures; therefore, we will not report any fMRI comparisons between "remember" and "know" responses here.

Focusing on true and false recognition (i.e., "remember" responses to old words and lures), participants were faster to respond to conceptual or perceptual old words than they were to respond to conceptual or perceptual lures, respectively (i.e., conceptual true recognition < conceptual false recognition, perceptual true recognition < perceptual false recognition, based on average median response latencies; both P < 0.005). Critically, there was no difference in participants’ median response latencies to conceptual versus perceptual true recognition (old conceptual vs. old perceptual) or false recognition (conceptual lure vs. perceptual lure; both P > 0.2) (see Table 1).

fMRI results

A frontoparietal network of brain regions was active during both conceptual and perceptual true recognition ("remember"/old conceptual and "remember"/old perceptual, respectively) compared with correct rejections ("new"/unrelated new), which is consistent with prior reports of neural activity patterns accompanying remembering (e.g., Buckner et al. 1998; Buckner and Wheeler 2001). A similar network of frontal and parietal regions was also active during both conceptual and perceptual false recognition ("remember"/conceptual lure and "remember"/perceptual lure, respectively) compared with correct rejections. These results are consistent with past reports of overlapping activity associated with true and false recognition in frontal and parietal regions (e.g., Cabeza et al. 2001; Slotnick and Schacter 2004; Garoff-Eaton et al. 2006), and they indicate that similarities do exist between conceptual and perceptual memory processing. Moreover, consistent with previous reports of frontal cortex activity observed during false recognition (e.g., Schacter et al. 1996b, 1997a; Cabeza et al. 2001; Goldmann et al. 2003; Slotnick and Schacter 2004), false compared with true recognition, regardless of information type ("remember"/conceptual lure and perceptual lure > "remember"/old conceptual and old perceptual), was associated with activity increases in multiple frontal cortex regions, including the middle, medial, and inferior frontal gyri (Table 2). By contrast, we did not observe any activity increases during all true recognition compared with all false recognition ("remember"/old conceptual and old perceptual > "remember"/conceptual lure and perceptual lure).

Although these analyses suggest that at least some memory processes are unaffected by the type of information being retrieved, we were particularly interested in differential activity associated with conceptual versus perceptual memory processing. Collapsing across true and false recognition, conceptual memory compared with perceptual memory ("remember"/old conceptual and conceptual lure > "remember"/old perceptual and perceptual lure) was accompanied by activity increases in multiple regions, including the frontal (e.g., superior, middle, and medial frontal gyrus) and parietal cortex (e.g., precuneus and superior parietal lobule) (see Table 2). By contrast, the opposite analysis ("remember"/old perceptual and perceptual lure > "remember"/old conceptual and conceptual lure) revealed activity increases during perceptual memory (compared with conceptual memory) in only two regions, the left inferior frontal gyrus (BA 44/9) and the left inferior parietal lobule (BA 40) (see Table 2).

Turning our focus to false recognition, we used a masking function (see Materials and Methods) to identify neural activity increases associated with false (compared with true) recognition that were unique to either conceptual or perceptual information. First, we identified those regions that were more active during false than true conceptual recognition but were not more active during false than true perceptual recognition (i.e., those regions

Table 2. Neural activity observed during false vs. true recognition (regardless of information type) and conceptual vs. perceptual recognition (regardless of accuracy)

| Region               | BA   | x    | y    | z    |
|----------------------|------|------|------|------|
| All false > all true |      |      |      |      |
| Left middle frontal gyrus | 9    | −46  | 29   | 30   |
| Left middle frontal gyrus | 8/9  | −44  | 9    | 33   |
| Left inferior frontal gyrus | 43/46 | −44 | 28   | 17   |
| Left inferior frontal gyrus | 44/9  | −46 | 5    | 24   |
| Left inferior frontal gyrus | 47   | −32  | 21   | −11  |
| Right medial frontal gyrus | 8/6  | 16   | 31   | 39   |
| Left medial frontal gyrus | 9/8/6 | −4  | 31   | 33   |
| Right anterior cingulate | 32/24 | 8   | 30   | 19   |
| Right insula | 13   | 32   | 20   | 6    |
| All conceptual > all perceptual |      |      |      |      |
| Left superior frontal gyrus | 8    | −40  | 18   | 49   |
| Left middle frontal gyrus | 6    | −40  | 14   | 49   |
| Left medial frontal gyrus | 11/10 | −10 | 50   | −11  |
| Left supramarginal gyrus | 40/39 | −50 | −59  | 31   |
| Left cingulate gyrus | 31   | −10  | −55  | 29   |
| Left posterior cingulate | 23/30 | −4  | −53  | 19   |
| Left superior parietal lobule | 7    | −40  | −70  | 46   |
| Left precuneus | 7    | −6   | −68  | 33   |
| Left/right precuneus | 23/31 | −2  | −61  | 18   |
| Left precuneus | 31/7  | −2   | −51  | 32   |
| Right precuneus | 31   | 2    | −65  | 29   |
| Left cuneus | 30/23  | −60  | −60  | 9    |
| Right cuneus | 18   | 4    | −71  | 16   |
| Left cuneus | 18/19 | −2   | −78  | 28   |
| Left lingual gyrus | 18   | 12   | −68  | 3    |
| Right middle occipital gyrus | 18  | 10   | −91  | 12   |
| Right pons | −14   | −40  | −28  |      |
| Left pons | −10   | −35  | −30  |      |
| Right cerebellum | 8    | −42  | −18  |      |
| Left cerebellum | −10  | −45  | −16  |      |
| All perceptual > all conceptual |      |      |      |      |
| Left inferior frontal gyrus | 44/9  | −51 | 9    | 24   |
| Left inferior parietal lobule | 40   | −50  | −35  | 46   |

BA refers to Brodmann area, and coordinated (x, y, z) are reported in Talairach space.
that were active in the “remember”/conceptual lure > “remember”/old conceptual contrast but were not active in the “remember”/perceptual lure > “remember”/old perceptual contrast. This analysis revealed widespread activity throughout the frontal cortex (BA s 6, 8, 9, 44, 45, 46, 47), as well as activity in the precentral gyrus and precuneus (Table 3; Fig. 2). By contrast, the reverse analysis (i.e., identifying those regions that were active in the “remember”/perceptual lure > “remember”/old perceptual contrast but were not active in the “remember”/conceptual lure > “remember”/old conceptual contrast) revealed only one cluster of activity in the caudate (see Table 3). Complementary analyses were performed to identify activity increases during true (compared with false) recognition that were unique to either conceptual or perceptual information. Only one cluster of activity spanning the left insula and claustrum (x = -36, y = -18, z = -1; BA 13) was more active during true than false conceptual (but not perceptual) recognition (“remember”/conceptual > “remember”/perceptual > “remember”/old perceptual), and no regions were more active during true than false perceptual (but not conceptual) recognition (“remember”/perceptual > “remember”/old perceptual > “remember”/old conceptual) and vice versa. It is unlikely that activity observed in these analyses was due to differences in effort, given that the same reaction time difference existed in both the included and the excluded contrast (i.e., activity associated with lengthier response latencies during false compared with true recognition should be present in both the contrast of interest and in the contrast used to create the mask, thus excluding activity associated only with longer reaction times from the final analysis).

To further investigate the differential activity observed in the frontal cortex during false compared with true recognition of conceptual (but not perceptual) information, we extracted event-related timecourses from each cluster of frontal activity defined by the “remember”/conceptual lure > “remember”/old conceptual > “remember”/perceptual lure > “remember”/old perceptual > “remember”/old conceptual > “remember”/perceptual > “remember”/old conceptual lure > “remember”/old perceptual lure > “remember”/old perceptual > “remember”/old conceptual lure > “remember”/old conceptual lure > “remember”/old perceptual > “remember”/old conceptual > “remember”/conceptual lure > “remember”/old conceptual > “remember”/perceptual lure > “remember”/old perceptual > “remember”/old conceptual). This analysis resulted in nine clusters of activity in the frontal cortex. More than nine frontal regions are listed in Table 3 because several clusters spanned more than one region or Brodmann area. Event-related timecourses were extracted from each cluster of activity rather than each region so that all clusters were spatially independent, a requirement for computing the joint probability reported here (see Materials and Methods). Although the random effects analysis requires a larger difference in activity between false and true conceptual recognition than between false and true perceptual recognition, it does not require a specific relationship between conceptual false recognition and perceptual false recognition (i.e., event-related timecourses are necessary to determine whether activity observed during conceptual false recognition is greater than, less than, or equal to activity observed during perceptual false recognition in the frontal cortex regions defined by the random effects analysis). Critically, the event-related timecourses revealed that, in all nine clusters of frontal cortex activity, greater activity (as assessed at 4–8 sec after stimulus onset) was associated with conceptual false recognition than perceptual false recognition (see Fig. 2). This difference reached statistical significance in two regions of left middle frontal gyrus (both P < 0.05). Importantly, the joint probability of observing this same pattern of differential activity across all nine of these clusters was significant (joint P-value <0.005). The same pattern of results was observed when event-related activity assessed at 6 sec after stimulus onset was analyzed (joint P-value <0.03).

Additional analyses that did not rely on masking procedures were also performed in order to further explore neural activity that was modulated by both accuracy and information type. Specifically, an interaction analysis revealed that the left inferior frontal gyrus (BA 45/47) showed a greater difference in activity between conceptual false and true recognition than between perceptual false and true recognition (“remember”/conceptual lure > “remember”/old conceptual > “remember”/perceptual lure > “remember”/old perceptual) (see Table 3; Fig. 3). Event-related timecourses extracted from this active cluster revealed significantly greater activity associated with conceptual false recognition than perceptual false recognition (P < 0.04) (see Fig. 3). By contrast, activity in the right inferior parietal lobule extending into the post-central gyrus (x = 55, y = -30, z = 20; BA 40), and in a region bordering the right inferior/middle frontal gyr (x = 22, y = 35, z = -5; BA 47/11). By contrast, increased activity in only one cluster of activity bordering the left cingulate gyrus (x = -20, y = -43, z = 35; BA 31) was observed during perceptual false recognition (compared with conceptual false recognition).

Table 3. Neural activity observed during false recognition compared with true recognition for either conceptual (but not perceptual) or perceptual (but not conceptual) information

| Region                              | BA  | x    | y    | z    |
|-------------------------------------|-----|------|------|------|
| Conceptual false > conceptual true  |     |      |      |      |
| (not perceptual false > perceptual true) |     |      |      |      |
| Left superior frontal gyrus         | 8   | -6   | 14   | 47   |
| Left superior frontal gyrus         | 6   | -2   | 18   | 53   |
| Left middle frontal gyrus           | 9   | -44  | 9    | 33   |
| Left middle frontal gyrus           | 6   | -26  | 12   | 45   |
| Left middle frontal gyrus           | 6   | -38  | -2   | 44   |
| Left middle frontal gyrus           | 46  | -51  | 26   | 24   |
| Right inferior frontal gyrus        | 44/45| 55   | 11   | 22   |
| Right inferior frontal gyrus        | 45  | -48  | 35   | 6    |
| Left inferior frontal gyrus         | 47  | -48  | 36   | -9   |
| Left inferior frontal gyrus         | 47  | -32  | 23   | -10  |
| Right medial frontal gyrus          | 9   | 9    | 43   | 21   |
| Right medial frontal gyrus          | 9   | 4    | 44   | 33   |
| Left medial frontal gyrus           | 9   | -2   | 40   | 31   |
| Left medial frontal gyrus           | 8   | -4   | 27   | 41   |
| Left medial frontal gyrus           | 6   | -4   | 31   | 37   |
| Right precentral gyrus              | 6   | -28  | -19  | 54   |
| Left precuneus                      | 7   | -6   | -62  | 51   |
| Left/right midbrain                 | -2  | -16  | -4   |      |
| Perceptual false > perceptual true  |     |      |      |      |
| (not conceptual false > conceptual true) |     |      |      |      |
| Right caudate                       | 12  | 12   | 23   |      |
| (Conceptual false > conceptual true) |     |      |      |      |
| (not perceptual false > perceptual true) |     |      |      |      |
| Left inferior frontal gyrus         | 45/47| -46  | 37   | 2    |
| Right middle temporal gyrus         | 21  | 55   | -48  | -3   |

BA refers to Brodmann area, and coordinates (x, y, z) are reported in Talairach space.

Discussion

The results from the current study indicate that although similarities between conceptual and perceptual memory processes ex-
The neural correlates of false recognition

The frontal cortex is preferentially active during conceptual false recognition. Specifically, similar standard patterns of neural activity (including activity in frontal and parietal regions) (Buckner et al. 1998; Buckner and Wheeler 2001) were observed during true and false recognition for both conceptual and perceptual information compared with correct rejections. Furthermore, consistent with past comparisons of true and false recognition (Schacter et al. 1996b, 1997a; Cabeza et al. 2001; Goldmann et al. 2003; Slotnick and Schacter 2004), increased frontal cortex activity accompanied false (compared with true) recognition regardless of information type. Importantly, subsequent analyses qualified that finding, indicating that the increased activity observed in certain frontal cortex regions during false recognition may be most pronounced when there is a conceptual (rather than a perceptual) relationship between the studied and the falsely recognized items. For instance, when the neural correlates of false and true recognition were compared separately for each information type, false recognition for conceptual (but not perceptual) information was supported by increased activity in multiple frontal cortex regions. In fact, event-related timecourses illustrated that in all of these frontal regions, greater activity accompanied conceptual than perceptual false recognition. By contrast, there was no indication of frontal cortex activity that supported perceptual rather than conceptual false recognition. Thus, our results suggest that conceptual false recognition is particularly reliant upon frontal lobe activity, and that the neural underpinnings of conceptual and perceptual false recognition are at least partially distinct.

These results complement previous behavioral research. Specifically, our results suggest that frontal activity is preferentially involved in conceptual false recognition, which could explain the different effects of childhood development (Holliday and Weeke 2006) and aging (Koutstaal et al. 2003) on conceptual versus perceptual false recognition, given the pronounced frontal lobe changes that occur during these developmental periods (Anderson and Craik 2000; Raz 2000; Diamond 2002). In fact, using the same behavioral paradigm as was used in this fMRI study, conceptual false recognition was especially reliant upon frontal lobe activity, which could imply that the neural underpinnings of false recognition are at least partially distinct.

Figure 2. Neural activity associated with conceptual false compared with conceptual true recognition but not with perceptual false compared with perceptual true recognition (“remember/conceptual lure > "remember/old conceptual" excluding activity in "remember/perceptual lure > "remember/old perceptual"). In the center, activity identified by this analysis is projected onto the left lateral surface of a three-dimensional template brain. On each side, activity is projected onto coronal slices of the group mean anatomic image, and each of the nine clusters of frontal activity identified by this analysis is demarcated by a circle. In the bottom panel, event-related activity (4–8 sec after stimulus onset) extracted from these active clusters is shown.

(a, inferior frontal gyrus; b, medial/superior frontal gyri; c, medial frontal gyrus; d, inferior frontal gyrus; e, middle frontal gyrus; f, middle frontal gyrus; g, middle frontal gyrus; h, inferior frontal gyrus; i, middle frontal gyrus)

Figure 3. Interaction analysis identifying regions that show a greater difference in activity between conceptual false and true recognition than between perceptual false and true recognition ("remember/conceptual lure > "remember/old conceptual" > "remember/perceptual lure > "remember/old perceptual"). On the right, activity is projected onto a coronal slice of the mean anatomic image, and on the left, the event-related activity (4–8 sec after stimulus onset) extracted from the active cluster within the left inferior frontal cortex (BA 45/47) is shown.

* p < 0.04
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Older adults compared with young adults show increased rates of conceptual false recognition, but equivalent rates of perceptual false recognition, perceptual true recognition, and conceptual true recognition (R. J. Garoff-Eaton, E. A. Kensinger, and D. L. Schacter, unpubl.). According to the results from the current study, it could be hypothesized that this differential effect of aging on conceptual and perceptual memory errors is due to altered frontal cortex activity in older adults, where conceptual false recognition is more reliant on frontal cortex activity and thus is more affected by the frontal lobe changes that accompany aging; further research would be necessary to test that hypothesis.

Our results indicate that the frontal cortex is preferentially engaged during conceptual compared with perceptual false recognition, but what cognitive processes does this frontal lobe activity reflect? Past interpretations of frontal activity during false recognition have often focused on post-retrieval monitoring processes (e.g., Schacter et al. 1996b; Cabeza et al. 2001; Goldmann et al. 2003; for review, see Schacter and Slotnick 2004). Under this view, the monitoring or verification processes that take place after retrieval (e.g., that conceptual false recognition involves the frontal cortex) are more extensive and/or effortful when the retrieved information was never studied (i.e., during false recognition) than when it was studied previously (i.e., during true recognition). This interpretation is consistent with previous reports of increased prefrontal cortex activity during memory tasks that have heightened evaluation or monitoring demands (e.g., tasks requiring source or associative memory in addition to item memory) (Rugg et al. 1996; Henson et al. 1999; Rugg et al. 2003; Achim and Lepage 2005). When considering the role of post-retrieval monitoring in false recognition, one might assume that monitoring should occur regardless of the precise relationship between the studied and the falsely recognized items. For instance, it seems likely that monitoring would be required during both conceptual and perceptual false recognition to evaluate or verify the accuracy of the familiar information being retrieved. Indeed, some of the overlapping frontal cortex activity we observed during both conceptual and perceptual false recognition may reflect post-retrieval monitoring; however, it is important to note that this overlapping activity was primarily left lateralized, and some have argued that it is the right (not left) prefrontal cortex that is preferentially involved in post-retrieval monitoring of familiarity (e.g., Dobbins et al. 2004) or evaluation/verification monitoring processes (e.g., Cabeza et al. 2003).

Furthermore, it seems unlikely that the primarily left lateralized regions found to be more active during conceptual than perceptual false recognition reflect post-retrieval monitoring alone. Instead, this frontal activity may reflect additional cognitive processes that are likely to be preferentially involved in conceptual memory. For instance, regions of the frontal cortex (especially the left inferior frontal cortex) are particularly active during successful semantic processing (Demb et al. 1995), word generation (Petersen et al. 1988), recovery of meaning/semantic retrieval (Wagner et al. 2001), verbal recall (Cabeza et al. 2003), semantic retrieval (associated with perceptual) relational memory (Prince et al. 2005), and semantic elaboration (Kirschhofer et al. 2005). In the current study, false recognition of a conceptual lure is likely to occur if a participant elaborates on the meaning of the lure and retrieves other related words (e.g., those that were actually seen at study) or additional semantic information about the lure (e.g., a broader category to which it belongs). Although one might assume that conceptual false recognition must rely on this extensive semantic elaboration process, conceptual true recognition may result from other cognitive processes (e.g., recognition of item-specific details). Furthermore, whereas elaborative processes may often accompany conceptual false recognition, perceptual false recognition should not be as dependent on this type of semantic processing; indeed, in the case of perceptual lures, there are no conceptually related words to retrieve from the study lists. Thus, at least some of the frontal activity observed during false compared with true recognition for conceptual (but not perceptual) information likely reflects some combination of the semantic processes previously reported to engage the left frontal cortex (e.g., semantic elaboration, semantic retrieval, etc.). However, further work will be necessary to determine the precise role(s) of the frontal cortex in conceptual false recognition.

Whereas conceptual false recognition may result from more extensive semantic elaboration than conceptual true recognition (as reflected by frontal lobe increases during conceptual false compared with true recognition), perceptual false recognition should occur when the visual and/or phonological features that overlap between the lure and previously studied words are recognized, which may be quite similar to the process of remembering a previously studied word in the case of perceptual true recognition. Indeed, very few regions showed a greater increase in activity during false compared with true recognition for perceptual (but not conceptual) information, which suggests that the neural correlates of false and true perceptual recognition may be more similar than those of conceptual false and true recognition. Moreover, very little activity (i.e., only one cluster bordering the cingulate gyrus) was revealed when perceptual false recognition was directly compared to conceptual false recognition. However, it is important to note that the direct comparison of conceptual and perceptual false recognition was also insensitive to the frontal cortex activity increases during conceptual (but not perceptual) false recognition identified via comparisons involving both false and true recognition. Thus, it is possible that subthreshold activity increases associated with perceptual (but not conceptual) false recognition exist (perhaps within visual or phonological processing regions), but that our current paradigm lacked the power to reveal that differential activity. Future research is necessary to determine whether there are patterns of neural activity that are unique to perceptual false recognition in the same way that certain patterns of frontal cortex activity seem to be unique to conceptual false recognition.

Previous work has suggested that activity in the anterior left inferior frontal cortex (BA 45/47) consistently accompanies conceptual/semantic processing, whereas phonological processing is more often associated with posterior left inferior frontal cortex (BA 44) activity (Poldrack et al. 1999; Poldrack and Wagner 2004). In the current study, activity increases in anterior left inferior frontal cortex (BA 45/47) occurred during conceptual (but not perceptual) false compared with true recognition; however, we did not observe posterior inferior frontal cortex activity during perceptual false (compared with true) recognition. As discussed above, one interpretation of these results is that conceptual false recognition may rely on significantly more semantic processing (e.g., elaboration, retrieval of related words) than conceptual true recognition, whereas similar levels of perceptual processing may be necessary for both false and true perceptual recognition. Although our comparisons did not reveal any frontal cortex activity during perceptual false recognition, we did observe posterior left inferior frontal cortex (BA 44/9) activity when we compared all perceptual memory to all conceptual memory, collapsing across accuracy (i.e., true and false perceptual recognition > true and false conceptual recognition). Thus, our results suggest that although perceptual memory in general does benefit more than conceptual memory from the phonological processes supported by the left posterior inferior frontal cortex, perceptual false recognition may not recruit these processes any more than perceptual true recognition does. By contrast, conceptual false recognition does seem to engage the semantic processes sup-
ported by the left anterior inferior frontal cortex to a greater degree than conceptual true recognition does. Previous work exploring priming and accurate memory has also identified differential neural activity accompanying the retrieval of conceptual versus perceptual information. In a study of repetition priming, transcranial magnetic stimulation (TMS) was used to disrupt activity in the left frontal cortex during the initial viewings of visual objects (Wig et al. 2005). Subsequent priming-related activity reductions for those objects were disrupted in left inferior frontal cortex (as were priming-related decreases in response latencies on the behavioral semantic classification task), while neural priming was spared in sensory-processing regions like the middle and inferior occipital cortex. In a study of accurate recollection, multiple regions within the left frontal cortex were preferentially active during the retrieval of conceptual details about previously studied objects, whereas the retrieval of perceptual details of those objects engaged right frontal and bilateral occipito-temporal regions (Dobbins and Wagner 2005). Our results complement these demonstrations that conceptual versus perceptual priming and accurate memory retrieval can rely on distinct neural substrates (specifically that conceptual memory tends to rely on left frontal cortex activity), and we extend these findings to the domain of memory distortion by illustrating that conceptual false recognition is associated with increased left frontal cortex activity. As discussed above, there are several possible explanations for why we did not observe unique patterns of neural activity during perceptual false recognition in the current study. However, given that sensory processing regions in the occipital cortex seem to be preferentially involved in both perceptual priming (Wig et al. 2005) and accurate perceptual retrieval (Dobbins and Wagner 2005), exploring activity in these sensory regions during perceptual false recognition may be an interesting avenue for further investigation.

In summary, although there was substantial overlapping neural activity associated with conceptual and perceptual false recognition compared with correct rejections, conceptual (but not perceptual) false compared with true recognition relied on increased frontal cortex activity (e.g., left inferior and middle frontal cortex), which likely reflected semantic processing. Our results complement previous behavioral work illustrating differential effects of aging and development on conceptual versus perceptual false recognition (Koutstaal et al. 2003; Holliday and Weekes 2006), as well as neuroimaging work reporting preferential engagement of the left frontal cortex in conceptual compared with perceptual priming and accurate retrieval (Dobbins and Wagner 2005; Wig et al. 2005). Moreover, previous research has shown that the neural basis of false recognition can depend on the relatedness of a falsely remembered item to previously studied items (Garoff-Eaton et al. 2006), as well as the context in which one believes to have seen a falsely remembered item (Kahn et al. 2004). Our findings extend this work by indicating that the precise relationship between falsely remembered and previously encountered items can also influence the neural activity that accompanies memory errors. Specifically, this study indicates that the frontal activity that is often observed during false compared with true recognition can be modulated by the relationship between studied and tested items, and it suggests that at least some of this frontal activity may reflect semantic processing during conceptual false recognition, which is distinct from previous arguments that frontal activity during false recognition reflects monitoring processes (e.g., Schacter et al. 1996b; Cabeza et al. 2001).

Materials and Methods
Participants
Twenty healthy, right-handed native English speakers with normal or corrected-to-normal vision and no history of serious neu-
"new" decision, differences between conceptual and perceptual false memories may have arisen due to differences in the qualitative nature of the retrieved memories. By asking participants to make "remember" versus "know" responses, we were able to compare conceptual to perceptual false recognition for items that were retrieved with the same qualitative nature (i.e., for items associated with a "remember" response).

Each target word was shown for 3 sec followed by a variable duration of fixation ranging from 1–17 sec. Each recognition test lasted 5 min 56 sec. Thus, one complete study-test cycle was completed every study task. After the study task, filler tasks were used in general, participants completed five full study-test cycles. Data from a single study phase for two participants and a full study-test cycle for one additional participant were excluded because of equipment malfunctions or spiking in the fMRI images. One participant completed only three full study-test cycles due to time constraints.

A particularly important aspect of this paradigm is that a single target can be tested in all five conditions (old conceptual, old perceptual, conceptual lure, perceptual lure, and unrelated new) between participants. All participants were tested on the same set of target words, while the condition in which those words were tested was counterbalanced across participants by manipulating the word triplets shown at study. For each studied word triplet, participants were tested either on an old word from that triplet or on a lure related to that triplet; thus, participants were never tested on multiple words related to the same triplet. In the new conceptual and old conceptual conditions, targets were studied as part of the appropriate (conceptual or perceptual) study triplet, and they were placed in each list position (i.e., top, middle, or bottom) an approximately equal number of times. In the conceptual lure and perceptual lure conditions, targets were not studied, but their appropriate (conceptual or perceptual) study triplets were. And finally, in the unrelated new condition, neither of the study triplets associated with the target was studied. For example, the target word yell could be tested in any of the five conditions by using the following word triplets at study: old conceptual—shout, yell, squeal; old perceptual—fell, bell, yell; conceptual lure—shout, cry, squeal; perceptual lure—fell, bell, yell; unrelated new—no studied triplet.

Imaging methods

Images were acquired on a three Tesla head-only Siemens Allegra MR imaging system at the Research Imaging Center at the University of Texas Southwestern Medical Center. The images were acquired using a T2-weighted echo planar imaging (EPI) sequence (TR = 30 msec, TE = 3.3 msec, 128 slices, 1 × 1 × 3.3 mm voxels). Functional images were acquired using a T2*-weighted echo planar imaging (EPI) sequence (TR = 2000 msec, TE = 30 msec, 34 slices, 4 × 4 × 4 mm voxels). Although images were acquired during both the study tasks and the recognition tests, here we report analyses conducted on images acquired during the recognition tests only.

Unless otherwise noted, SPM99 (Wellcome Department of Cognitive Neurology) was used for all data preprocessing and analysis. Standard preprocessing was performed on the functional data. Images were first slice-time corrected and motion corrected. Then, a temporal high-pass filter was applied to each run to remove linear, quadratic, cubic, and quartic components (using custom software written in MATLAB). Next, all study task runs were concatenated, and all recognition test runs were concatenated. Finally, images were normalized to the SPM99 EPI template (resampling at 2 × 2 × 2 mm resolution) and spatially smoothed (using a 4-mm full-width half-maximum [FWHM] Gaussian kernel).

A general linear model approach was used for the event-related analysis. For each participant, activity associated with each event type was modeled using a canonical hemodynamic response function. For the recognition tests, there were 16 possible event types; event types were defined by crossing the three possible response types ("remember," "know," and "new") with the five possible conditions (old conceptual, old perceptual, conceptual lure, perceptual lure, and unrelated new), resulting in 15 event types, with an additional event type defined as any item that received no response. All participants had a minimum of five events in each event type of interest. For each voxel, a β-weight (i.e., model amplitude) associated with each event type was computed by entering each event type's hemodynamic response model into a general linear model. The difference in magnitude between β-weights associated with two event types of interest were computed and then compared across participants using one-sample t-tests (i.e., t-test and recognition test cycle analysis). Thus, these analyses identified regions that were more active during one event type than another. It is important to note that although it is generally assumed that regions that are more active during a certain task are also more involved in that task, the opposite can also be true (e.g., certain processes, like priming, can be associated with activity decreases) (Schacter and Buckner 1998; Schacter et al. 2007).

Monte Carlo simulations (http://www2.bc.edu/~slotnics/scripts.htm) were used to define an individual voxel and cluster extent threshold that would yield results corrected for multiple comparisons (e.g., Forman et al. 1995; Slotnick et al. 2003; Slotnick and Schacter 2004, 2006; Garoff-Eaton et al. 2006). In this study, an individual voxel threshold of P < 0.01 was used in combination with a cluster extent threshold of 40 resampled voxels, which yielded results corrected for multiple comparisons (P < 0.05). For a detailed description of the procedure used to determine these thresholds, see Slotnick and Schacter (2004) or Garoff-Eaton et al. (2006). Spatial autocorrelation was modeled through convolution with a 4-mm FWHM Gaussian kernel in the current study.

In several analyses, to determine whether differential neural activity observed in one contrast was distinct from activity observed in another contrast, we used the masking function in SPM to exclude activity from the second contrast when analyzing the activity from the first. In these analyses, the same voxel and cluster extent thresholds described above (P < 0.01 and 40 voxels, respectively) were used for both the excluded contrast and the contrast of interest. Voxel coordinates are reported in Talairach space (Talairach and Tournoux 1988) and refer to the most significant voxel proximal to the center of each active region.

Event-related timecourses were extracted from active clusters of interest in order to characterize the nature of the neural activity in those regions (for details, see Slotnick and Schacter 2004). Using custom software written in MATLAB, linear trends were extracted from the timecourses. Statistics were performed on the mean event-related activity occurring 4–8 sec after stimulus onset unless otherwise noted, and activity occurring 0–4 sec before stimulus onset was used to correct for baseline activity. Given that these timecourses and the contrast effects analyses characterizing the neural activity in clusters defined by the random-effects analyses, one-tailed t-tests were used to compare timecourses associated with different event types within those active clusters. To compute the joint probability of observing a specific pattern of activity (i.e., the same relationship between timecourses associated with two event types) across multiple regions of interest, we used Fisher’s technique (Fisher 1973), as described by Slotnick and Schacter (2006).

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