The neural dynamics of updating person impressions

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Person perception is a dynamic, evolving process. Because other people are an endless source of social information, people need to update their impressions of others based upon new information. We devised an fMRI study to identify brain regions involved in updating impressions. Participants saw faces paired with valenced behavioral information and were asked to form impressions of these individuals. Each face was seen five times in a row, each time with a different behavioral description. Critically, for half of the faces the behaviors were evaluatively consistent, while for the other half they were inconsistent. In line with prior work, dorsomedial prefrontal cortex (dmPFC) was associated with forming impressions of individuals based on behavioral information. More importantly, a whole-brain analysis revealed a network of other regions associated with updating impressions of individuals who exhibited evaluatively inconsistent behaviors, including rostrolateral PFC, superior temporal sulcus, right inferior parietal lobule and posterior cingulate cortex.

Keywords: impression formation; social cognition; person perception; dmPFC; IPFC; fMRI

INTRODUCTION

Human social interaction is as informationally rich as it is ubiquitous. As we spend countless hours engaging with other humans, we form impressions of the people around us—in large part, in an attempt to predict behavior. However, our fellow interaction partners are not always so consistent. As such, social interaction requires continuous, flexible updating of our initial impressions in light of new information.

An abundance of research has accumulated on the neural bases of ‘first impressions’. Much of this research has focused on initial appraisals of other people based on facial characteristics like attractiveness and perceived trustworthiness (for meta-analysis, see Mende-Siedlecki et al., 2011). First impressions based upon behavioral information have been extensively examined as well. This research has shown that our impressions of the people around us are powerfully influenced by the behaviors we come to associate with them (Todorov and Uleman, 2002; Bliss-Moreau et al., 2008; Todorov and Olson, 2008). Behavior-based impression formation can lead to automatic inferences regarding character traits (Todorov and Uleman, 2003), and further, can be generalized to similar-looking others (Verosky and Todorov, 2010). Typically, in such studies, people represented by faces paired with negative behavioral information are subsequently evaluated as being less trustworthy, and people paired with positive information are subsequently evaluated as being more trustworthy (Todorov and Olson, 2008).

Recent neuroimaging work has sought to identify brain regions crucial for forming impressions of others based upon behavioral information. The primary region associated with such tasks is the dorsomedial prefrontal cortex (dmPFC; Mitchell et al., 2004, 2005, 2006; Schiller et al., 2009; Baron et al., 2011; Cloutier et al., 2011a). Dovetailing with these results, the dmPFC has also been previously implicated in the spontaneous retrieval of affective person knowledge about faces previously learned in the context of behavioral information (Gobbini et al., 2004; Gobbini and Haxby, 2007; Todorov et al., 2007).

Regions such as the posterior cingulate cortex (PCC; Schiller et al., 2009; Freeman et al., 2010; Cloutier et al., 2011a), amygdala (Schiller et al., 2009; Baron et al., 2011), superior temporal sulcus (STS; Mitchell et al., 2005; Schiller et al., 2009; Freeman et al., 2010) and inferior frontal gyrus (IFG; Mitchell et al., 2005; Schiller et al., 2009; Baron et al., 2011; Freeman et al., 2010) have also been observed in conjunction with this type of impression formation task. However, while it is possible to speculate on a putative network of regions involved in impression formation, the preponderance of studies implicating the dmPFC in such tasks is undeniable.

Although there is a substantial body of research on first impressions, much less is known about how these impressions are updated. Impression formation is an ongoing process, and initial impressions must be updated on the basis of new, incoming information—which may be evaluatively inconsistent with previous impressions. Here, we explore a phenomenon we describe as impression updating—situations where new information learned about a target is evaluatively inconsistent with a previous impression, thus necessitating an update of that impression to account for the inconsistency.

Social psychology affords us a host of predictions regarding how person perception can be affected by such a turn of events (Reeder and Brewer, 1979; Fiske, 1980; Reeder and Spores, 1983; Skowronski and Carlston, 1987, 1989). Our impressions of other people may function as schemas that drive our expectations of their future behavior (Fiske and Linville, 1980). When we are faced with information that is inconsistent with a given schema, we are forced to reassess our impression to account for the new information (Srull and Wyer, 1989). However, despite previous behavioral work, neuroimaging investigations of impression updating have just begun. Some recent research has addressed the neural dynamics of how initial impressions are updated by behavioral information, in both electrophysiological (Rudoy and Paller, 2009) and neuroimaging contexts (Baron et al., 2011; Cloutier et al., 2011b; Ma et al., 2011). Baron and colleagues presented participants with untrustworthy-, trustworthy- and neutral-looking faces in the scanner, and in a subsequent phase, paired some of these faces with valenced behavioral information. Not only was the dmPFC more active during learning for faces paired with behaviors, but this activity correlated with a post-scan measure of learning, suggesting that in the context of this task, the dmPFC plays an important role in updating initial appearance-based impressions based upon behavioral information.
Especially relevant is a recent study by Ma and colleagues, in which participants read sets of behavioral descriptions that implied a specific trait about a particular individual. Critically, the last behavior was manipulated to be either consistent or inconsistent with that implied trait. Responses in the dmPFC were higher when this last behavior was trait-inconsistent, compared to when it was trait-consistent (Ma et al., 2011). Finally, another recent study by Cloutier and colleagues observed preferentially higher dmPFC activity when targets’ behaviors were incongruent with their social category (in this case, political affiliation), as opposed to when they were congruent (Cloutier et al., 2011b).

The current study focuses on evaluative impression updating over a long behavioral trajectory. To that aim, we presented participants with person targets who were paired with five descriptions of valenced behaviors (e.g. ‘Ron gave out toys at the children’s hospital during Christmas’), viewed consecutively. Half of the targets were paired with behavioral information that remained either consistently negative or consistently positive, thus requiring little demand for impression updating. The other half of the targets were paired with behavioral information that switched valence on the fourth trial. The desired effect is that the first three pieces of behavioral information create a strong expectation for that person to behave in a certain manner (for instance, acting like a good, law-abiding citizen)—an expectation that is subsequently violated on trials four and five, resulting in a high demand for impression updating.

We expected that participants would update their impressions of targets based upon new, inconsistent information. More importantly, consistent with other studies (Mitchell et al., 2004, 2005, 2006; Schiller et al., 2009), we expected that evaluative updating of impressions would recruit regions implicated in impression formation such as the dmPFC. Finally, based on recent studies (Cloutier et al., 2011b; Ma et al., 2011), we expected that in addition to these regions, evaluative updating would recruit regions involved in attention and cognitive control.

METHODS

Participants

Twenty-four (14 female) participants volunteered for the fMRI study and were paid $30 for their participation. They were between the ages of 18 and 45 years (mean = 25.3 years). All participants were right-handed, had normal or corrected-to-normal vision and reported no history of neurological illnesses or abnormalities. We acquired informed consent for participation approved by the Institutional Review Board for Human Subjects at Princeton University. All participants were fully debriefed at the completion of the experiment.

Face and behavior stimuli

Each participant saw a series of 50 faces taken from the book ‘Heads’ (Kayser, 1997), paired with positively and negatively valenced behaviors previously rated on goodness and kindness (Fuhrman et al., 1989). Each face was paired with five consecutively viewed behaviors, comprising one ‘target’. Targets were classified as either evaluatively consistent or inconsistent. Consistent targets consisted of a face paired with five behaviors of the same valence—either five straight positive behaviors (consistently positive) or five straight negative behaviors (consistently negative). Inconsistent targets consisted of a face paired with three behaviors of one valence, followed by two behaviors of the opposite valence—either three positive behaviors followed by two negative behaviors (positive-to-negative), or three negative behaviors followed by two positive behaviors (negative-to-positive). Additionally, participants sometimes saw control targets—faces presented alone on screen, without accompanying behaviors. All in all, participants encountered 50 total targets—10 targets corresponding to each of these five conditions.

Behaviors were combined together in groups of five such that each group within a given condition would be roughly equated on goodness and kindness. The average goodness and kindness ratings for each condition were as follows: consistently negative (M = 1.81, SD = 0.61), negative-to-positive (M = 4.79, SD = 3.15), consistently positive (M = 8.10, SD = 0.63), positive-to-negative (M = 4.83, SD = 3.20). Faces and behavior valences were counterbalanced between participants, such that each face was paired with each type of behavior group an equal number of times. Finally, each participant was given a unique, optimized target ordering, based upon a genetic algorithm (Wager and Nichols, 2003, http://wagerlab.colorado.edu/wiki/doku.php/help/ga/genetic_algorithm_for_fmri) to maximize statistical power. We note that while facial trustworthiness is not of interest in this study, the faces we used indeed varied on this dimension. That said, due to the counterbalancing of faces and behavior valences, any differences due to facial trustworthiness are assumed to be negligible.

Procedures

Participants were informed that they would be participating in a study on impression formation. They were told that they would be seeing a series of faces paired with behaviors, and that they would see multiple behaviors paired consecutively with each face. Participants were asked to form an impression of each target, altering that impression if necessary based on new information they learned as the task went along. Additionally, participants were told that picturing targets performing behaviors would likely aid in forming impressions. In scanner, they saw ten runs of face targets, each paired with five separate behaviors. Each run consisted of five face targets, one of each condition. Each run began with a 15 s presentation of a fixation cross. Each target was split into five face/bebehavior presentations. Faces and behaviors were presented together for 6 s. Next, a rating slide appeared for 4 s, during which the participant rated how trustworthy that individual was, based upon the information they had learned about him so far. Participants made their ratings with an MR-safe button box, on a scale ranging from 1 (very untrustworthy) to 4 (very trustworthy). Subsequently, a fixation cross appeared for 4 s. This series of events proceeded four more times per target (with the same face on the screen, paired with different behaviors each time). Following the fifth behavior, a new target appeared. All stimuli were projected onto a screen located at the rear of the bore of the magnet. Participants were able to view these stimuli via an angled mirror attached to the RF coil placed above their eyes.

Image acquisition

Blood oxygenation level-dependent (BOLD) signal was used as a measure of neural activation. Echo planar images (EPI) were acquired using a Siemens 3.0 Tesla Allegra head-dedicated scanner (Siemens, Erlangen, Germany) with a standard ‘bird-cage’ head coil (TR = 2000 ms, TE = 30 ms, flip angle = 80°, matrix size = 64 x 64). By using 32 interleaved 3-mm axial slices we were able to achieve near whole brain coverage. Prior to the primary data acquisition scan, a high-resolution anatomical image (TI-MPRAGE, TR = 2500 ms, TE = 4.3 ms, flip angle = 8°, matrix size = 256 x 256) was acquired for subsequent registration of functional activity to the participant’s anatomy and for spatially normalizing data across participants.

Image analysis

All fMRI data were analyzed with Analysis of Functional NeuroImages software (AFNI; Cox, 1996). The first four EPI images from each run were first realigned for motion correction. Each subject’s data were then co-registered to their own high-resolution MPRAGE image and spatially normalized to the MNI 152 template using a unique, optimized target ordering, based upon a genetic algorithm (Wager and Nichols, 2003, http://wagerlab.colorado.edu/wiki/doku.php/help/ga/genetic_algorithm_for_fmri) to maximize statistical power. We note that while facial trustworthiness is not of interest in this study, the faces we used indeed varied on this dimension. That said, due to the counterbalancing of faces and behavior valences, any differences due to facial trustworthiness are assumed to be negligible.
were discarded to allow the MR signal to reach steady-state equilibrium. Participants’ motion was corrected using a six-parameter 3D motion-correction algorithm following slice scan-time correction. Transient spikes were removed from the signal using the AFNI program 3dDespike. Subsequently, data were low-passed filtered with a frequency cut-off of 0.1 Hz following spatial smoothing with a 6-mm full width at half maximum (FWHM) Gaussian kernel. The signal was then normalized to percent signal change from the mean.

To identify regions that were more active when participants were forming impressions based on behaviors, we contrasted trials in which faces were paired with behaviors and trials in which faces were presented alone. This contrast yielded functional regions of interest (fROIs) involved in learning to associate behavioral information with faces, and by extension, forming behavior-based impressions of those person targets. We subsequently analyzed the parameter estimates in these fROIs as a function of the order of the behaviors (the first three vs the last two behaviors) and the evaluative consistency of the behaviors. Given the large number of fROIs yielded by the contrast of faces paired with behaviors and faces alone, the parametric map was thresholded at \( \alpha = 0.0001 \) (uncorrected). Furthermore, to select a minimum cluster size for corrected significance \( (P < 0.05) \), we performed a Monte Carlo simulation of null-hypothesis data, using the AlphaSim program included in the AFNI package. The Monte Carlo simulation indicated that a minimum cluster size of 8 voxels was appropriate.

To generate parameter estimates, we performed voxel-wise multiple regression on each participant’s preprocessed imaging data. Twenty-five regressors of interest (five 6000-ms trials per target \( \times 5 \) types of target) were convolved with a canonical hemodynamic response function and entered into our general linear model (GLM). Additionally, we included several regressors of no interest, including head motion estimates and time points representing rating slide presentations. Each participant’s parameter estimate maps were projected into Talairach space (Talairach and Tournoux, 1988) prior to performing any group-level analyses.

In addition to the fROI analyses, we performed a whole-brain analysis testing the interaction between trial number (last two trials \( vs \) first three trials) and evaluative consistency (consistent \( vs \) inconsistent). Finally, we performed separate whole-brain analyses contrasting the last two trials against the first three trials, in both consistent and inconsistent targets. Because we did not find reliable main effects of the valence of the behaviors and higher order interactions with this valence, we do not report analyses related to valence. However, we provide supplemental figures including the valence of the behaviors. All whole-brain analyses are reported using the same thresholding procedures as described above \( (P < 0.05 \text{ FDR-corrected}; \text{ voxel-wise threshold}, P < 0.005; \text{ minimum cluster-size threshold}, 31 \text{ voxels}) \).

**RESULTS**

**Behavioral results**

Because we were primarily interested in updating impressions, we focus on the changes in ratings in response to evaluatively inconsistent information. We computed separate averages across the first three and last two behaviors, isolating participants’ evaluations of our targets before and after the potential introduction of evaluatively inconsistent information. We further subtracted the ratings of control targets (faces presented without behavioral information) from the consistent and inconsistent targets’ ratings and recorded the absolute deviation from the control condition. These deviations provide a measure of the change in target evaluation. [See Supplementary Figure 1 for the means across all \( 5 \) (target type) \( \times 5 \) (trial number) conditions].

Participants updated their impressions of person targets based upon evaluatively inconsistent information. Specifically, the change in participants’ ratings from the first three to the last two behaviors was greater for inconsistent targets than consistent targets. A 2 (trial number: first three behaviors \( vs \) last two behaviors) \( \times 2 \) (consistency: consistent targets \( vs \) inconsistent targets) ANOVA revealed significant main effects of trial number \( [F(1,23) = 13.37, P < 0.001] \) and consistency \( [F(1,23) = 89.52, P < 0.001] \). Critically, we observed a significant interaction between trial number and consistency \( [F(1,23) = 69.92, P < 0.001] \), such that the absolute deviation in trustworthiness ratings from the first three to the last two behaviors was greater for inconsistent targets \( (M = 0.58, SE = 0.08) \) than for consistent targets \( (M = 0.29, SE = 0.04) \).

The mean response time across trials was 1119.41 ms \( (SE = 47.75) \). To test for potential differences in difficulty in processing information about consistent and inconsistent targets, we submitted the response times to a 2 (trial number: first three behaviors \( vs \) last two behaviors) \( \times 2 \) (consistency: consistent targets \( vs \) inconsistent targets) ANOVA. Neither main effect was significant, nor was the interaction between trial number and consistency. Nevertheless, we also tested for simple effects, and observed that the effect of trial number was not significant for either consistent \( [t(23) = 0.18, P = 0.858] \) or inconsistent targets \( [t(23) = -1.48, p = 0.153] \).

**fMRI results**

**Brain activity associated with impression formation**

We contrasted face-plus-behavior trials against face-alone trials. This method of localizing fROIs associated with forming impressions of person targets based on behavioral information is consistent with previous research (Schiller et al., 2009; Baron et al., 2011).

We observed 13 fROIs that responded more strongly to faces paired with behavioral information than to faces presented alone (Supplementary Table 1). We next tested which fROIs responded to the introduction of new behavioral information inconsistent with prior impressions, looking for a specific pattern of response, such that activity remained consistent or dropped from the first three trials (F3) to the last two trials (L2) for consistent and control targets, but increased for inconsistent targets.

The only fROI that produced this pattern of response was the dmPFC. As shown in Figure 1, activity increased in response to inconsistent information, but decreased when information was consistent. We performed a 3 (target type: inconsistent, consistent, control) \( \times 2 \) (trial number: first three trials \( vs \) last two trials) repeated-measures ANOVA on the \( \beta \) values extracted from this fROI, observing an interaction between consistency and trial number \( [F(2,46) = 5.45, P = 0.008, \eta^2 = 0.19] \). Splitting these analyses by target type, we observed that dmPFC signal rose from the first three trials to the last two trials for inconsistent targets \( [F(1,23) = 24.67, P < 0.001, \eta^2 = 0.52] \). Conversely, dmPFC signal change was not significant for consistent \( [F(1,23) = 1.21, P = 0.283, \eta^2 = 0.05] \) or control targets \( [F(1,23) = 0.934, P = 0.344, \eta^2 = 0.04] \). [See Supplementary Figure 2 for expanded analyses split by valence].

**Brain activity associated with updating impressions**

**Interaction analysis.** We sought to identify brain areas that showed a stronger L2 > F3 pattern for inconsistent targets than consistent targets, potentially reflecting their role in updating impressions based upon new, conflicting information. This interaction analysis showed that right IPL, left STS, PCC extending into the pulvinar, and bilateral rPFC were all significantly more active in the last two trials than the first three trials for inconsistent targets only (Table 1 and Figure 2). In addition, right STS showed a similar pattern, though this cluster did not surpass extent-based thresholding. Visualizations of signal change
but not to evaluatively consistent information, suggesting that it plays a role in updating of impressions. Within consistent targets, we observed no brain areas that were preferentially active during the last two trials, while bilateral fusiform gyrus, cuneus and right pulvinar were more active during the first three trials (Supplementary Table 2, Figure 3).

However, the L2 > F3 contrast within inconsistent targets yielded activity in dmPFC, PCC/precuneus, bilateral rIPFC, bilateral dIPFC, bilateral IPL, bilateral STS and left anterior insula (Supplementary Table 2, Figure 3). The reverse contrast, F3 > L2, yielded activity in bilateral fusiform, cerebellum, right lingual gyrus, and inferior occipital gyrus.

**DISCUSSION**

To explore the neural dynamics of updating person impressions, we presented participants with faces paired with behavioral descriptions that were either consistent or inconsistent in valence. As expected, forming impressions of these targets based upon behavioral information, compared to presentation of faces alone, activated a set of regions typically associated with similar impression formation tasks, including the dmPFC. Within this set of regions, only the dmPFC showed preferential activation to updating based on new, evaluatively inconsistent information, as opposed to updating based on information consistent with existing impressions. Additional whole-brain analyses pointed to a larger set of regions involved in updating of evaluative impressions, including bilateral rIPFC, bilateral STS, PCC and right IPL.

We also observed regions that did not respond differentially as a function of the evaluative consistency of the behaviors. Specifically, large portions of inferotemporal cortex, including the bilateral fusiform gyri, were less active for the last two trials than the first three trials for both consistent and inconsistent targets (Figure 3), most likely a result of habituation in response to the repeatedly-presented facial stimuli (Kanwisher and Yovel, 2006).

**The role of dmPFC in impression updating**

The results of the fROI analyses showed that the dmPFC was the only region that displayed enhanced responses to evaluatively inconsistent but not to evaluatively consistent information, suggesting that it plays an integral role in the evaluative updating of person impressions. This is consistent with previous conceptualizations of the dmPFC’s role in impression formation (Mitchell et al., 2004; 2005; 2006; Schiller et al., 2009; Baron et al., 2011; Ma et al., 2011). Surprisingly, the whole brain interaction analysis of evaluative consistency and order of behaviors only yielded sub-threshold dmPFC activity—a discrepancy most likely due to the low-power nature of our design. In fact, the simple contrast comparing the last two vs first three behaviors did yield a large dmPFC activation for inconsistent but not consistent targets (Figure 3).

Two recent studies have also linked the dmPFC to impression updating. Ma and colleagues observed increased dmPFC activity in response to targets that behaved in a manner inconsistent with specific traits they had been previously associated with (Ma et al., 2011). In addition, Cloutier and colleagues observed that the dmPFC also responded preferentially to instances where targets’ behaviors were inconsistent with their social category (e.g. a Democrat favoring small government). In the context of this recent research, the present study suggests that the dmPFC’s role in updating extends more broadly into instances of general evaluative inconsistency as well.

An alternative explanation of the increased dmPFC activity for inconsistent targets is that presenting inconsistent information on screen resulted in a less fluent reading experience. Hence, the increase in dmPFC activity is indicative of an increased difficulty associated with these targets. However, we observed no significant differences in response times across the last two trials between consistent and inconsistent targets, suggesting that our imaging results cannot be simply explained in terms of task difficulty.

**A functional network for updating impressions**

We now turn our attention to the other regions implicated in our analyses. How might the STS, IPL, rIPFC and PCC be acting in service of impression updating? The STS has been previously demonstrated to play an integral role in a variety of tasks associated broadly with social processing and social cognition (Hein and Knight, 2008). Neuroimaging research in the past decade has frequently implicated the STS in aspects of high-level person perception critical for social communication, for instance, biological motion (Allison et al., 2000; Vaina et al., 2001; Grossman and Blake, 2002; Pelphrey et al., 2003a; Puce and Perrett, 2003; Pelphrey et al., 2004a; Pelphrey et al., 2006) and facial expressions (static: Haxby et al., 2000; Hoffman and Haxby, 2000; Adolphs, 2002; LaBar et al., 2003; Calder and Young, 2005;
Meanwhile, the IPL has also been associated with a range of social cognitive functions, including gaze processing (Wicker et al., 1998; Pelphrey et al., 2003b; Pelphrey et al., 2004b; Calder et al., 2007), imitation (Iacoboni et al., 1999; Decety et al., 2002; Leslie et al., 2004), action perception in the service of understanding intentions (Gallese et al., 2004; Fogassi et al., 2005; Iacoboni et al., 2005; Montgomery and Haxby, 2008), self-other distinctions (Ruby and Decety, 2001; Ruby and Decety, 2003; Uddin et al., 2006) and shared representations (Keysers et al., 2004; Zaki et al., 2009).

Many of the functions listed above are inherently germane to impression updating. First and foremost, both the STS and IPL have been connected to aspects of face processing. The omnipresence of facial stimuli in our task certainly introduces a prevalent, if implicit demand to process facial features. Furthermore, as we told our participants that they should imagine targets performing the actions they were paired with, it is possibly not surprising that an area like the IPL, associated with action perception (especially social actions), should be implicated.

Of most relevance, a recent review of research on the social brain suggests that one function of the STS is to predict the behavior of social agents based on incoming information (Frith and Frith, 2010). Specifically, the authors offer evidence suggesting that activity in posterior STS increases when a social agent behaves in a manner that is inconsistent with prior expectancies. In previous research, this inconsistency has taken the form of unexpected shifts in gaze (Pelphrey et al., 2003b; Pelphrey et al., 2004a), as well as unexpected changes in actions (Saxe et al., 2004). In this sense, posterior STS activity in these tasks may be representing a social prediction error signal. Behrens and colleagues (2008) sought to directly test this possibility in a task in which participants made decisions based, in part, on a confederate’s advice. This advice was occasionally unexpectedly incorrect or correct, eliciting a prediction error correlating with an increase in posterior STS activity, a signal dissociable from reward-related non-social prediction error signals observed in the ventral striatum. The results of the present study are consistent with this framework. On trials when evaluatively inconsistent information was presented, our participants’ expectations were violated, and in turn, they were faced with the task of updating their impressions in order to better predict targets’ future actions.

Engell and Haxby, 2007; Ishai, 2008; dynamic: Ghazanfar et al., 2010; Said et al., 2010).

### Table 1: Regions showing significant differences in the interaction contrast of last two trials vs first three trials as a function of consistency

<table>
<thead>
<tr>
<th>Region</th>
<th>Lat</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>#Voxels</th>
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<tr>
<td>Interaction between L2 &gt; F3consistent and L2 &gt; F3consistent</td>
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<tr>
<td>Inferior parietal lobule</td>
<td>R</td>
<td>46.5</td>
<td>−64.5</td>
<td>47.5</td>
<td>317</td>
</tr>
<tr>
<td>PCC/pulvinar</td>
<td>−</td>
<td>1.5</td>
<td>−31.5</td>
<td>8.5</td>
<td>116</td>
</tr>
<tr>
<td>STS</td>
<td>L</td>
<td>−67.5</td>
<td>28.5</td>
<td>−2.5</td>
<td>86</td>
</tr>
<tr>
<td>Rostrolateral PFC</td>
<td>R</td>
<td>45.5</td>
<td>55.5</td>
<td>2.5</td>
<td>60</td>
</tr>
<tr>
<td>STS</td>
<td>L</td>
<td>−46.5</td>
<td>52.5</td>
<td>2.5</td>
<td>40</td>
</tr>
<tr>
<td>STS</td>
<td>R</td>
<td>64.5</td>
<td>−34.5</td>
<td>−9.5</td>
<td>28*</td>
</tr>
</tbody>
</table>

All clusters are significant at $P < 0.05$, after correction for multiple comparisons, unless indicated with an asterisk. $x$, $y$, $z$ coordinates reflect peak voxel location in Talairach coordinate system.

*Did not surpass cluster extent-thresholding ($k = 31$).
It seems likely that the STS and IPL are involved in processing specific to person targets in the context of this task. Conversely, the PCC and rlPFC are better suited to aid in more general, task-related processing during the updating impressions task.

While the PCC is typically associated with the default mode network (Gusnard and Raichle, 2001; Greicius et al., 2003; Buckner et al., 2008), it has also been implicated in a host of seemingly disparate processes, ranging from representation of subjective value (McCoy et al., 2005; Kable and Glimcher, 2007; Levy et al., 2010), to autobiographical memory retrieval (Maddock et al., 2001), to goal-directed cognition (Spreng et al., 2010). A recent reconceptualization of the PCC’s function attempts to reconcile these various functions within one parsimonious explanation, suggesting that the PCC is critical for adapting to changes in the environment (Pearson et al., 2011). This account of the PCC is extremely in step with the demands of the current experiment, wherein our participants had to identify relevant changes (i.e. behaviors inconsistent with existing impressions of person targets) and subsequently, adjust to those changes and act accordingly (i.e. update their impressions of person targets, as evidenced by changes in behavioral ratings).

The lateral PFC has also been linked to high-level cognitive processes, including maintaining abstract mental sets (Christoff et al., 2007), multitasking (Burgess et al., 2001; Burgess et al., 2003; Badre et al., 2004), and perhaps most importantly, the flexible exertion of cognitive control (Braver et al., 2003; Braver et al., 2009). Specifically, activity in the rostral portion of lateral PFC is associated with episodic control (Koechlin et al., 2003; Kouneiher et al., 2009)—in which a previously encountered cue modifies the perception or interpretation of present stimuli (Egner, 2009). In the context of the present study, this conceptualization of rlPFC’s role is particularly informative. The rlPFC activity in response to evaluatively inconsistent targets likely reflects the influence of previously learned information on participants’ evaluations of new information.

Limitations
Several low-level aspects of our design may be influencing our results. First and foremost, it is possible that the inclusion of trial-by-trial ratings is imposing an unnaturally high demand to update impressions upon our participants. While we concede that this is indeed a limitation of our approach, our intention was to collect a moment-to-moment measure of participants’ impressions. While we concede that this is indeed a limitation of our approach, our intention was to collect a moment-to-moment measure of participants’ impressions, so we could be absolutely certain that they showed behavioral evidence of updating. Future work could simply measure participants’ impressions only once following the presentation of all five behaviors.

Second, we employed a control condition (faces presented alone) in which we do not account for the reading that participants have to do in the consistent and inconsistent conditions. We chose to perform the faces-plus-behaviors vs faces alone contrast because it is consistent with previous related work (Schiller et al., 2009; Baron et al., 2011). More importantly, while this confound is unavoidable for our fROI analysis, our whole-brain analyses do not depend on this contrast.

Convergence with recent work
As discussed previously, recent studies involving trait-inconsistent updating (Ma et al., 2011) and category-inconsistent updating (Cloutier et al., 2011b) bear a great deal of relevance for the present investigation. Comparing between these three studies, we note interesting convergence in the neuroimaging results, even though they focus on different types of inconsistency. As Figure 4 shows, all three studies observed higher dmPFC, IPL, STS, PCC and IPFC activity when targets were behaviorally inconsistent, compared to when they were consistent.
Previous work has observed additional inconsistency-related activity in a more posterior region of mPFC (referred to as domain-general pmFC; Ma et al., 2011). One potential explanation for this divergence lies in the specific contrast with which Ma and colleagues obtained this result. While we chose to contrast the last two vs the first three trials in our behavior trajectories, they contrasted activity on only the critical fourth trial between target types (consistent vs inconsistent). In essence, the present analysis takes a more global perspective on the updating process as a whole, while Ma et al. (2011) isolated activity elicited at the precise moment when trait-inconsistent information was potentially presented. Running a similar analysis on our data yields activity in domain-general pmFC, as well (Supplementary Figure 4).

Taken together, these studies suggest that flexible updating of person impressions depends on the coordinated action of functional networks involved in social cognition and cognitive control. While this represents only a first step towards elucidating the neural dynamics underlying impression updating, a picture is beginning to come into focus, revealing a network of regions encompassing the dmPFC, IPL, STS, PCC and rIPFC, associated with this process.

**SUPPLEMENTARY DATA**
Supplementary data are available at SCAN online.

**Conflict of Interest**
None declared.

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