Categorical perception of race is mediated by distributed patterns of activity in the brain

Lily Tsoi¹, Yune S. Lee², Liane Young¹

¹ Department of Psychology, Boston College
² Department of Speech and Hearing Science, The Ohio State University

Contact Information:

Lily Tsoi
140 Commonwealth Avenue
McGuinn 300
Chestnut Hill, MA 02467
Email: lily.tsoi@bc.edu
Abstract

Race is an abstract social category that is often associated with perceptual cues. We used functional magnetic resonance imaging (fMRI) to investigate the cognitive and neural processes that support categorical perception of race: the warping of perceptual information (i.e., facial features related to race) into discrete racial groups. Participants viewed morphed faces along 10-step White-Black continua in the scanner and had their categorical boundaries measured outside the scanner. Despite the fact that participants generally perceived the faces in a non-linear (i.e., categorical) manner, changes in overall activity within regions for visual processing and top-down processing of visual stimuli linearly tracked with changes in facial features associated with racial ambiguity. Interestingly, the warping of face stimuli along a continuum into discrete race categories appears to be mediated not by overall activity within specific regions in the brain but by distributed patterns of activity in regions associated with the attention network. Additionally, ROI (region-of-interest)-based multivariate pattern analyses revealed contributions of face-processing regions (i.e., right FFA) and social cognition (i.e., left temporoparietal junction) in categorical race perception. Together, these findings point to different types of information afforded by activation-based and pattern-based analyses regarding processes involved in race perception.

Keywords: MVPA, race perception, social categorization

Words: 4994
1. Introduction

Each United States census, since the first one in 1790, has contained at least one question about racial identity. However, the ways in which questions have been posed and answers have been coded have changed dramatically over time (Pew Research Center, 2015). For example, in 1980 and 1990, if a respondent marked more than one race category (e.g., White; Black; American Indian, Eskimo, and Aleut; Asian and Pacific Islander), the Census Bureau re-categorized that person to a single race, typically the race of the respondent's mother. By contrast, in the year 2000, the U.S. Census Bureau introduced the option to be classified as more than one race, acknowledging the ever-increasing number of multiracial individuals in the United States. Indeed, race categorization relies not only on perceptual cues (e.g., skin color, facial structure) but also on shifting conceptual and cultural cues (e.g., changes in criteria used to label people by race; greater public acceptance and higher rates of interracial unions). The current study investigated how race is represented in the human brain, taking the contrast between Black and White faces as a case study.

The present research capitalized on a specific psychological finding: faces along a race continuum—including racially ambiguous faces—are typically categorized as a single race instead of both races in proportion to the position of the face along the continuum (Levin & Angelone, 2002). The phenomenon by which people perceive features varying along a continuum as discrete categories is known as categorical perception. Prior research on categorical perception has focused mainly on sensory and perceptual features in the auditory domain with regard to speech perception (Bidelman et al., 2013; Chang et al., 2010; Desai et al., 2008; Lee et al., 2012; Liberman et al., 1957; Myers & Swan, 2012;
Prather et al., 2009; Raizada & Poldrack, 2007) and music (Klein & Zatorre, 2015; Klein & Zatorre, 2011; Lee et al., 2011), as well as in the visual domain, with regard to color (Franklin et al., 2008; Özgen & Davies, 2002; Roberson et al., 2008), shapes and/or patterns (Goldstone, 1994; Newell & Bülthoff, 2002; Notman et al., 2005), and faces (Beale & Keil, 1995; Etcoff & Magee, 1992; Rotshtein et al., 2004). While many of these studies on categorical perception have relied on linearly varying a single specific perceptual feature (e.g., voice onset time, luminance), the current study focused on race categorization based on holistic facial features. Building on prior work on categorical perception of facial features in different forms (e.g., face identity, facial expression), as discussed below, the current study used face morphs, constructed by simultaneously manipulating a suite of perceptual facial features (e.g., skin shade, brow ridge, cheeks, and eye sockets).

Prior work has shown that categorical perception of complex visual features such as face identity and facial expression is supported by brain regions further along the visual processing stream and higher-level cognitive regions. For example, the right fusiform gyrus and fusiform face area (FFA) appears sensitive to perceived changes in face identity (e.g., from Marilyn Monroe to Margaret Thatcher) (Rotshtein et al., 2004). Similarly, a study using an adaptation paradigm shows sensitivity of the FFA and posterior superior temporal sulcus (pSTS) to perceived changes in identity (i.e., different individuals) as well as expression (i.e., angry, afraid, disgusted, happy) (Fox et al., 2009). Regions outside of this core face network shows further specificity: the middle STS shows release from adaptation when participants perceive a change in expression but not identity, whereas the precuneus shows release from adaptation for the opposite change. Relatedly, categorical perception of gender from faces appears to be represented by the orbitofrontal cortex (Freeman et al.,
2010). Overall, this body of work suggests that regions further along the visual processing stream and/or higher-level brain regions outside of the core face network that appear to support categorical perception of facial features like face identity and facial expression may also support categorical perception of race.

To examine regions that mediate categorical race perception, we used multivariate searchlight analyses, an approach that has been successfully used for discriminating neural patterns that map morphed stimuli into discrete categories (Lee et al., 2012). This pattern-based approach, unlike the conventional activation-based approach, allows us to identify subtle differences across conditions that share many physical features (Kriegeskorte, Goebel, & Bandettini, 2006).

Along with examining categorical race perception at the whole-brain level, we used functional ROI-based MVPA to examine whether categorical perception of race is also mediated by the FFA—an important region for categorical perception of facial features discussed above—as well as cortical regions involved in theory of mind (ToM), including bilateral temporoparietal junction (TPJ), precuneus, and medial prefrontal cortex (mPFC) (Fletcher et al., 1995; Gallagher et al., 2000; Gobbini et al., 2007; Saxe & Kanwisher, 2003). Behavioral research has revealed that people infer other people’s internal states (e.g., traits, mental states) rapidly and reflexively from perceptual information, including facial appearance (for reviews, see Olivola et al., 2014; Todorov et al., 2015). Indeed, neuroimaging work has revealed that regions in the ToM network are recruited when viewing natural scenes containing people (Wagner et al., 2011). Moreover, detecting animacy—a potential cue to socially relevant information—in moving shapes is sufficient to elicit activity in ToM regions (Wheatley et al., 2007).
In short, the present study investigates the cognitive and neural processes supporting categorical race perception, examining possible contributions of late visual processing (e.g., faces) and higher-level cognitive processing (e.g., social cognition).

2. Methods

2.1. Participants

Twenty-nine right-handed participants between the ages of 18 and 40 (mean ± standard deviation = 26.5 ± 6.46; 13 females) were recruited from the Boston community. All participants were native English speakers, had normal or corrected-to-normal vision, and reported having no history of psychiatric or neurological disorders. One participant fell asleep early in the scan session and was removed from the study. The remaining 28 participants consisted of 14 Caucasians (7 females), 12 African-Americans (4 female), 1 Asian-American (female), and 1 male of unspecified race. Participants gave written informed consent and were paid $25/hour for their participation. The study was approved by the Boston College Institutional Review Board.

2.2. Stimuli and Procedures

FaceGen Modeller 3.5 (Singular Inversions) was used to generate 3D faces that were morphed by race (from white to black). The European to African morphing slider was used, which contained 41 steps: faces every four steps along the slider were used as stimuli, starting with the third step and ending with the third-to-last step. The race morphing control combined 95 different features, such as skin shade (dark/ light), brow ridge (e.g., high/low), cheeks (e.g., round/gaunt), and eye sockets (dark / light); because the various
feature controls were correlated, adjusting the race morphing slider affected all of these features.

Ten face morphs varying along a continuum from White to Black were produced for each gender (Figure 1a). Each face was presented at a 20.05° yaw angle. An elderly version of each face was created as an oddball stimulus; there were 20 oddball stimuli in total. Face images (faces on a black background) and fixation crosses were centered on a gray background (Figure 1b). During fMRI scanning, these images were projected onto a screen (1024 x 768 pixel resolution) at the end of the magnet bore with an InFocus IN5542 projector. Each face image horizontally subtended a visual angle of approximately 11°.

Participants were instructed to indicate the appearance of any oddball stimuli (i.e., elderly-looking faces) via button box press with their right hand. There were 10 runs; each run lasted 4.2 minutes and consisted of 40 trials: 38 typical trials and 2 oddball trials. In a typical trial, the same face morph was presented three times (each for 0.5 s) with a blank screen (0.1 s) between face morphs. In an oddball trial, the same face morph was presented two times, and an elderly-looking version of the face morph was presented once. The temporal position of the oddball was random (i.e., first, second, or third face in the trial) but roughly matched: across 20 oddball trials, the average participant was presented with an oddball 6.83 times on the first position, 7.08 times on the second, and 5.92 times on the third. Due to technical issues, we were able to collect only 22 out of 28 participants’ oddball responses in the scanner. A fixation cross (4.5 s) was presented between all trials. Two sets of randomized stimulus presentation order sequences were created using the sequence optimizer optseq2 (Dale, 1999), and the order sequence was counterbalanced across
participants. The experiment was controlled by MATLAB 2008b and the Psychophysics Toolbox extensions (Brainard, 1997; Pelli, 1997) on a MacBook Pro.

![Stimuli for the fMRI experiment and behavioral results.](image)

**Figure 1.** Stimuli for the fMRI experiment and behavioral results. (a) Two sets of 10 faces morphed along a White-Black continuum: one male (top), one female (bottom). (b) Typical trial: the same face is presented three times (depicted in figure). Oddball trial: the same face is presented two times and the oddball (an elderly-looking version of the face) randomly appears in one of the three positions (not depicted). Participants are instructed to make a button-box response upon seeing an oddball. (c) Perceptions of faces along White-Black continua from 24 participants. Top: the average psychometric curve. The y-axis shows the percentage of responses labeled as “black” for each face token (i.e. morph-level) in the continuum (1=White; 10=Black). Error bars indicate SEM. Bottom: Participant count by categorical boundary.
2.3. Functional localizer tasks

Participants completed two functional localizer tasks. One functional localizer task (Epstein & Kanwisher, 1998) was used to define bilateral fusiform face area. A second localizer task (Dodell-Feder et al., 2011) was used to functionally define regions involved in ToM or social cognition more generally. Information about the two tasks can be found in Supplementary Materials.

2.4. Post-scan behavioral tasks

After the fMRI experiment, participants completed a behavioral task outside the scanner that measured their subjective categorical boundary for the face morphs during the experimental task in the scanner. Each face was presented ten times, and participants were instructed to indicate whether they viewed the face as white or black by pressing one of two buttons. The boundary was defined as the 50% crossover point between viewing the face as white or black on each participant’s psychometric curve. This boundary was used to define binary classes (white vs. black) for subsequent labeling of each participant’s neural data. Stimuli were presented using PsychoPy (Peirce, 2007) on a MacBook Pro. Finally, for exploratory purposes, a subset of participants ($N = 14$) completed a race Implicit Association Test (IAT), a measure of implicit racial bias.

2.5. fMRI data acquisition

Twenty of the participants were scanned in a Siemens 3T Tim Trio MRI scanner at the Center for Brain Science (Harvard University). Because the Center for Brain Science switched to a 3T Siemens Prisma scanner and because we wanted to continue scanning with the same protocols and the same type of scanner, we scanned the final nine
participants in a Siemens 3T Tim Trio MRI scanner at the Athinoula A. Martinos Imaging Center (Massachusetts Institute of Technology).

Participants were scanned using a 12-channel head coil. Thirty-six axial slices (3-mm isotropic voxels, 0.54-mm gap) were acquired using the following gradient-echo planar imaging (EPI) sequence parameters: repetition time (TR) = 2000 ms; echo time (TE) = 30 ms; flip angle (FA) = 90°; field of view (FOV): 216 x 216; interleaved acquisition. Anatomical data were collected with T1-weighted multi-echo magnetization prepared rapid acquisition gradient echo image sequences using the following parameters: TR = 2530 ms; TE = 1.64 ms; FA = 7°; 1-mm isotropic voxels; 0.5 mm gap between slices; FOV = 256 x 256.

2.6. fMRI data analyses

Data preprocessing and analyses were performed using SPM8 (http://www.fil.ion.ucl.ac.uk/spm) and custom software with MATLAB 2012a. All images were slice-time corrected, realigned to the first EPI of the first run, and spatially normalized into Montreal Neurological Institute (MNI) standard stereotactic space (EPI template) with preserved original voxel size.

2.6.1. Activation-based analyses. Neural responses were modeled in an event-related design using a general linear model (GLM), with conditions modeled as boxcar functions convolved with a canonical hemodynamic response function (HRF). The GLM included the six motion parameters as nuisance regressors. Trials during which an oddball appeared were removed from all analyses.

We performed activation-based analyses testing the parametric effect of racial ambiguity. The first analysis examined regions whose activity increased as faces became
less ambiguous and more racially defined. Predictor values were calculated by assigning low values (indicating racial ambiguity) to the middle face tokens (i.e., Face Tokens 5 and 6) and assigning increasingly higher values to more racially unambiguous face tokens; predictor values ranged from 1 (racially ambiguous) to 5 (racially defined). The second analysis examined regions whose activity increased as faces became more racially ambiguous. Predictor values were calculated by assigning higher values to the middle face tokens and lower values to face tokens closer to the ends of the continua.

We also examined regions that were recruited more for white vs. black faces and vice versa, in which faces were labeled as ‘white’ or ‘black’ using participants’ subjective categorical boundaries.

For each analysis, contrast images were submitted to random effects analyses, and the output contrasts were thresholded using a voxel-wise threshold of $p < .001$ (uncorrected), after which we corrected for multiple comparisons ($p < .05$) across the whole brain based on cluster extent and Gaussian random field theory (Friston et al., 1994; Worsley et al., 1992). Anatomical labels were retrieved using SPM Anatomy toolbox (v 2.2) (Eickhoff et al., 2005).

2.6.1 Searchlight analysis. For the searchlight analysis, fMRI time courses for all voxels were extracted from unsmoothed images and high-pass filtered with a 128 s cutoff. The signals were also mean-centered to normalize intensity differences among runs. Trials during which an oddball appeared were removed from all analyses. The preprocessed time courses were mapped to each face token (labeled 1 to 10; see Figure 1a) based on a GLM framework: first, a regressor for each face token per run was constructed by convolving the onset of each trial with the canonical HRF (hemodynamic response function). Next, the
mean height of the regressor was calculated. Lastly, if the height of the regressor at each time point was greater than the mean height, this time point was assigned to a particular face token. Time points were labeled as belonging to ‘black’ and ‘white’ classes based on participants’ subjective categorical boundaries.

We moved a 3-voxel radius sphere throughout the brain, centering on each voxel. In each searchlight sphere, a binary classification (white or black) was performed using a Gaussian Naïve Bayes classifier (Pereira et al., 2009; Raizada & Lee, 2013). For validating the classification, we used leave-one-run-out procedure, wherein data from one of the runs were reserved for testing, and remaining data were used for training (a total of 10 cross-validations). The output searchlight images for all participants were used to perform a group-level analysis. The group maps were thresholded using a voxel-wise threshold of $p < .001$ (uncorrected), after which we corrected for multiple comparisons ($p < .05$) across the whole brain based on cluster extent and Gaussian random field theory (Friston et al., 1994; Worsley et al., 1992). Anatomical labels were retrieved using SPM Anatomy toolbox (v 2.2) (Eickhoff et al., 2005).

2.6.2. ROI-based MVPA. We examined the spatial pattern of neural activity for white and black faces within regions involved in face and race processing (i.e., bilateral FFA) and regions involved in social cognition (i.e., right and left TPJ, precuneus, and dmPFC). Analyses were performed in MNI space. To define the FFA, beta values were estimated in each voxel for blocks corresponding to each of four types of stimuli (scenes, faces, objects, and scrambled objects) using a boxcar function. A contrast map was obtained for [faces > objects] in every participant. The FFA was defined as all voxels in a 9-mm radius of the peak voxel that passed threshold in the contrast image ($p < 0.001$, uncorrected, extent
cluster size > 10). To define regions involved in social cognition, beta values were estimated in each voxel for stories describing mental states (e.g., belief) or physical representations (e.g., photo). A contrast map was obtained for [belief > photo] in every participant. ROIs were defined as all voxels in a 9-mm radius of the peak voxel that passed threshold in the contrast image (p < 0.001, uncorrected, extent cluster size > 10). Information about the number of participants whose ROIs we were able to define and the MNI coordinates for those ROIs can be found in Table S1 (Supplementary Material).

The classification procedure of ROI-based MVPA was identical to that used for searchlight. Leave-one-run-out cross-validation was used for all analyses. An accuracy score averaged across training/testing set combinations was computed for each ROI and every individual.

3. Results

3.1.1 Behavioral results

Participants’ responses in the post-scan behavioral task confirmed that participants overall perceived a sharp categorical shift for faces near the middle of the continuum (Figure 1c). Four participants perceived the continua in a non-categorical manner; because we were interested in categorical race perception, these participants’ data were removed from all further analyses. The analyses below therefore reflect the data of the remaining 24 participants. Analyses examining the relationship between people’s responses in the post-scan task and implicit race bias revealed no meaningful relationship (see Supplementary Materials).
Additionally, we assessed participants’ performance on the oddball-detection task as a measure of participants’ attentiveness. Due to a coding error, we were able to consistently collect responses to oddballs on only the first and second positions within the trial. On average, participants responded to 5.44 ($SD = 2.64$) out of 6.83 oddballs and 5.50 ($SD = 2.64$) out of 7.08 oddballs in the first position and second position, respectively. Overall, these results suggest that participants were attending to the task.

3.1.2. fMRI results

We examined whether any regions showed a parametric effect of racial ambiguity (Figure 2; Table 1). Analyses revealed that activity in the following regions increased with decreases in racial ambiguity (or increases in racial unambiguity): the left lingual gyrus, left fusiform gyrus, left inferior frontal gyrus, and right posterior medial frontal cortex. In contrast, activity in bilateral mid orbital gyrus increased with increases in racial ambiguity.

![Figure 2. Parametric effects of racial ambiguity. Regions whose activity increased as faces became less racially ambiguous and more racially defined are shown in red (left lingual gyrus, left fusiform gyrus, and right posterior medial frontal cortex not depicted). Regions whose activity increased activity as faces became less racially defined and more racially ambiguous are shown in yellow (bilateral mid orbital gyrus). Images were cluster-level corrected (FWE) at $p < 0.05.$](image-url)
Table 1. Regions involved in the parametric effect of racial ambiguity

<table>
<thead>
<tr>
<th>Region name</th>
<th>MNI coordinates</th>
<th>t value</th>
<th># of voxels</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>x</td>
<td>y</td>
<td>z</td>
</tr>
<tr>
<td><strong>Regions whose activity increased as faces became more racially defined</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>L lingual gyrus</strong></td>
<td>-27</td>
<td>-94</td>
<td>-14</td>
</tr>
<tr>
<td>L lingual gyrus</td>
<td>-33</td>
<td>-88</td>
<td>-14</td>
</tr>
<tr>
<td>L fusiform gyrus</td>
<td>-33</td>
<td>-67</td>
<td>-14</td>
</tr>
<tr>
<td>L fusiform gyrus</td>
<td>-33</td>
<td>-55</td>
<td>-20</td>
</tr>
<tr>
<td>L cerebellum</td>
<td>-33</td>
<td>-73</td>
<td>-20</td>
</tr>
<tr>
<td><strong>L inferior frontal gyrus</strong></td>
<td>-48</td>
<td>17</td>
<td>22</td>
</tr>
<tr>
<td>L inferior frontal gyrus</td>
<td>-42</td>
<td>11</td>
<td>28</td>
</tr>
<tr>
<td>L inferior frontal gyrus</td>
<td>-45</td>
<td>29</td>
<td>19</td>
</tr>
<tr>
<td><strong>R posterior medial frontal cortex</strong></td>
<td>3</td>
<td>23</td>
<td>49</td>
</tr>
<tr>
<td><strong>Regions whose activity increased as faces became more racially ambiguous</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>L mid orbital gyrus</strong></td>
<td>-6</td>
<td>38</td>
<td>-11</td>
</tr>
<tr>
<td>R mid orbital gyrus</td>
<td>3</td>
<td>44</td>
<td>-8</td>
</tr>
<tr>
<td>Anterior cingulate cortex</td>
<td>0</td>
<td>50</td>
<td>-2</td>
</tr>
</tbody>
</table>

Note: Regions in bold are peak voxels; indented regions indicate sub-peak voxels. Regions listed here were cluster-level corrected at \( p < 0.05 \) (FWE).
Further analyses examining regions recruited more for white vs. black faces or black vs. white faces based on participants’ subjective categorical boundaries revealed no regions for either contrast. Together, these results suggest that although activation-based analyses reveal regions that track with physical changes leading to changes in racial ambiguity, they do not reveal regions that are overall recruited more for faces that participants subjectively perceive as white vs. black or vice versa. We examined whether categorical perception of race manifests instead as spatially distributed patterns of activity.

Searchlight analyses based on participants’ categorical boundaries revealed four regions: the right superior parietal lobule (with subpeaks in the right inferior parietal lobule), right middle frontal gyrus (with subpeaks in the right precentral gyrus), left inferior parietal lobule (with subpeaks in the left precentral gyrus), and left middle frontal gyrus (Figure 3; Table 2). Due to concerns that white and black participants would show different patterns of results, we compared the searchlight results for these two groups of participants; no significant differences were found for the two groups.
**Figure 3.** Group-level searchlight analysis based on participants’ subjective categorical boundaries reveal the left inferior parietal lobule, the left middle frontal gyrus, the right superior parietal lobule, and the right middle frontal gyrus (cluster-level corrected at $p < 0.05$, FWE).
Table 2. Searchlight results. Regions mediating categorical race perception based on participants’ subjective categorical boundaries (cluster-level corrected at \( p < 0.05 \), FWE)

<table>
<thead>
<tr>
<th>Region name</th>
<th>MNI coordinates</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>t value</th>
<th># of voxels</th>
</tr>
</thead>
<tbody>
<tr>
<td>R superior parietal lobule</td>
<td></td>
<td>36</td>
<td>-52</td>
<td>58</td>
<td>6.24</td>
<td>1009</td>
</tr>
<tr>
<td>R inferior parietal lobule</td>
<td></td>
<td>51</td>
<td>-37</td>
<td>55</td>
<td>5.94</td>
<td></td>
</tr>
<tr>
<td>R inferior parietal lobule</td>
<td></td>
<td>42</td>
<td>-43</td>
<td>55</td>
<td>5.83</td>
<td></td>
</tr>
<tr>
<td>R middle frontal gyrus</td>
<td></td>
<td>42</td>
<td>8</td>
<td>55</td>
<td>5.56</td>
<td>618</td>
</tr>
<tr>
<td>R precentral gyrus</td>
<td></td>
<td>51</td>
<td>8</td>
<td>49</td>
<td>5.30</td>
<td></td>
</tr>
<tr>
<td>L inferior parietal lobule</td>
<td></td>
<td>-27</td>
<td>-49</td>
<td>52</td>
<td>5.28</td>
<td>379</td>
</tr>
<tr>
<td>L precentral gyrus</td>
<td></td>
<td>-27</td>
<td>-4</td>
<td>49</td>
<td>4.95</td>
<td></td>
</tr>
<tr>
<td>L middle frontal gyrus</td>
<td></td>
<td>-42</td>
<td>20</td>
<td>46</td>
<td>4.83</td>
<td>73</td>
</tr>
<tr>
<td>L middle frontal gyrus</td>
<td></td>
<td>-42</td>
<td>11</td>
<td>52</td>
<td>4.52</td>
<td></td>
</tr>
<tr>
<td>L middle frontal gyrus</td>
<td></td>
<td>-36</td>
<td>26</td>
<td>46</td>
<td>4.27</td>
<td></td>
</tr>
</tbody>
</table>

Note: Regions in bold are peak voxels; indented regions indicate sub-peak voxels.
We also tested whether the spatial patterns of neural activity for white and black faces as defined subjectively are distinct from each other within (1) regions involved in face processing, i.e., bilateral fusiform face area (rFFA), and (2) regions involved in social cognition, i.e., right and left temporoparietal junction (rTPJ), precuneus, and dmPFC. We were able to define the left and right fusiform area, right temporoparietal junction, and precuneus in 21 out of 24 participants, left temporoparietal junction in 20 out of 24 participants, and dorsomedial prefrontal cortex in 15 out of 24 participants (Table S1). ROI-based MVPA showed that the spatial patterns of neural activity for white and black faces were distinguishable from each other in the left FFA and right FFA; classification accuracy was above chance (0.5) for both the left FFA (mean accuracy = 0.53, SD of accuracy = 0.04, \(t(20) = 4.40, p < 0.001\), one-tailed) and right FFA (mean accuracy = 0.52, SD of accuracy = 0.04, \(t(20) = 3.274, p = 0.002\), one-tailed). Classification accuracy was at chance level for the rTPJ, precuneus, and dmPFC (ps > 0.05) but above chance for lTPJ (\(M = 0.51, SD = 0.03, t(19) = 2.311, p = 0.017\)).

4. Discussion

The current study investigated the neural mechanisms supporting categorical perception of race. Despite the fact that participants generally perceived the faces in a non-linear (i.e., categorical) manner, changes in mean levels of activity within regions for visual processing and top-down processing of visual stimuli linearly tracked with changes in facial features associated with racial ambiguity. Interestingly, the warping of face stimuli along a continuum into discrete race categories appears to be mediated not by mean levels
of activity within specific regions in the brain but by distributed patterns of activity in the brain. Together, these findings point to different types of information afforded by activation-based and pattern-based analyses regarding processes involved in race perception.

Activation-based analyses suggest that increased activity in the left lingual gyrus, left fusiform gyrus, left IFG, and right posterior medial frontal cortex is linked to increases in racial unambiguity. Prior work has linked the lingual gyrus, a region along the ventral visual stream (Ungerleider & Haxby, 1994), to processing intermediate visual forms (e.g., curvature, radial and concentric patterns; (Gallant et al., 2000) and more visually complex images (Machielsen et al., 2000) as well as encoding visual memories (Bogousslavsky et al., 1987; Nenert et al., 2014; Rombouts et al., 1999; Ueno et al., 2007). There is also some evidence that the lingual gyrus, along with the fusiform gyrus, is important for object color processing (Chao & Martin, 1999; Hsu et al., 2012; Miceli et al., 2001; Simmons et al., 2007; Ueno et al., 2007b; Wang et al., 2013; but see Bogousslavsky et al., 1987; McKeefry & Zeki, 1997), though these two regions appear to contribute to color processing in different ways. According to some research, the lingual gyrus responds to any color stimuli regardless of task (e.g., as a sensory response), whereas the fusiform gyrus responds to color stimuli in tasks that require attention to color information (Beauchamp et al., 1999; Hsu et al., 2012; Simmons et al., 2007). In short, color processing may be important for processing increasingly racially defined faces, which converges with behavioral evidence showing that in children and adults, skin color plays a prominent role in race categorization (Dunham et al., 2015). Additionally, other work has shown involvement of the IFG and right medial frontal cortex in processing own-race vs. other-race faces, with these frontal regions
suggested to support top-down processing of visual stimuli like faces (Feng et al., 2011). Why top-down processing of visual stimuli would increase for more racially defined faces is unclear; indeed, prior work has proposed that activation of the left IFG is found when people have to select among competing sources of information to guide a response (Thompson-Schill et al., 1997). Racially defined faces, as compared to racially ambiguous faces, ought to make selection easier. One possible explanation could be that the problem of selection is not salient until people are faced with racially defined faces. Another explanation could be that top-down processing of more racially defined faces leads people to shift their attention to various features that disambiguate the race of a face.

Our results also indicate that increased activity in the mid orbital gyrus is linked to increased racial ambiguity. This finding aligns well with work showing that the orbitofrontal cortex (OFC) is recruited for decisions under uncertainty or ambiguity (Elliott et al., 2000; Hsu, 2005; Krain et al., 2006). Intriguingly, this region appears to be involved even when participants in the scanner were not making decisions that depended on assessing the racial ambiguity of faces.

Categorical race perception, on the other hand, is mediated not by overall activity but by distributed patterns of activity in regions typically involved in top-down control of visual attention (e.g., superior parietal lobule) (Corbetta et al., 2008; Parlatini et al., 2017) and gatekeeping between top-down cognitive control and bottom-up sensory-driven attention (e.g., middle frontal gyrus) (Japee et al., 2015). Some researchers have proposed that the frontoparietal network provides a ‘priority map’, integrating factors related to bottom-up and top-down attention (Bisley & Goldberg, 2010; Katsuki & Constantinidis, 2014). Indeed, a recent study has revealed frontoparietal contributions to working memory
by representing feature-specific information about relevant stimuli and by mediating top-down cognitive control (Ester et al., 2015). This frontoparietal network may be encoding stimuli features related to race and/or encoding race information based on top-down factors like prior knowledge (e.g., past experiences with people of different races).

Finally, we focused on specific regions of interest (ROIs) based on our a priori hypotheses regarding face-processing regions (i.e., FFA) and regions involved in social cognition (i.e., bilateral TPJ, precuneus). Research in social neuroscience has directly investigated how the FFA responds to faces that differ by race. Studies using conventional activation-based analyses have found preferential activation of the fusiform gyrus to own-race versus other-race faces (Feng et al., 2011; Golby et al., 2001; Lieberman et al., 2005; Natu et al., 2011); however, evidence points to the idea that the fusiform gyrus may not be responding to race specifically, but to motivationally relevant social categories (Van Bavel et al., 2008, 2011). Indeed, when race is made orthogonal to group membership, the fusiform gyrus responds similarly to White and Black faces but differentially to own-group versus other-group faces regardless of race (Van Bavel et al., 2008). Although overall activity in the fusiform gyrus may be driven by motivationally relevant social categories, recent work using MVPA reveals that the spatial patterns of activity for Black and White faces within the fusiform gyrus and specifically the FFA are distinguishable from each other above chance level (Contreras et al., 2013; Ratner et al., 2013). This work suggests that race is in fact encoded in the fusiform gyrus. Evidence also suggests that the right FFA supports categorical perception of visual features. Indeed, prior work on the neural basis of categorical perception of visual features such as facial identity or expression showed that the right FFA is sensitive to perceived (categorical) changes but not physical changes in
facial identity and expression (Fox et al., 2009; Rotshtein et al., 2004). Our results show that the left and right FFA mediate categorical perception of race. This finding provides converging evidence for the role of the FFA in categorical perception of visually accessible features related to faces. Importantly, though, our findings also reveal that categorical race perception does not appear to be mediated by the entire ToM network. Instead, our findings support a specific role for the lTPJ. Prior work has implicated the lTPJ in processing perspective differences regardless of rational (goal-oriented) action (Aichhorn et al., 2009; Aichhorn et al., 2006; Perner et al., 2006). Future work may wish to explore how race categorization, particularly in the case of ambiguous face morphs, recruits the processing of perspective differences.

4.1. Limitations and caveats

While the focus of this paper is on categorical perception of race, we acknowledge that race can sometimes be perceived in a continuous manner (indeed, a small minority of our participants showed this pattern). Recent work in social psychology reveals that when people have the explicit option to use the Multiracial label for Black-White biracial individuals, the majority of people use the Multiracial label (Chen & Hamilton, 2012). However, the frequency with which it is used is less than the use of the label White for White individuals and Black for Black individuals. More importantly, the likelihood of using the Multiracial label for Black-White biracial individuals decreases under cognitive load or time pressure (Chen & Hamilton, 2012). Some researchers therefore suggest that, while people are able to categorize Black-White biracial individuals as Multiracial when making more deliberate or reflective categorizations, they nevertheless categorize biracial individuals as Black when making rapid or reflexive categorizations (Peery &
Bodenhausen, 2008). Our aim in the current work was to probe this rapid, reflexive categorization of race, given its contribution to the formation and expression of implicit racial attitudes and prejudice—the focus of abundant work in social psychology (Dovidio et al., 2002; Dovidio et al., 1997; Goff et al., 2008; Greenwald et al., 1998; Lai et al., 2014; McConnell & Leibold, 2001; Wittenbrink et al., 1997).

We also acknowledge that not all individuals engage in the same race categorization processes. Social psychological and sociological work has revealed categorization of racially ambiguous people to reflect the principle of hypodescent (i.e., categorization of mixed-race individuals to the minority group) (Davis, 1991). The general finding is that people tend to categorize Black-White biracial individuals as Black than as White (Halberstadt et al., 2011; Ho et al., 2011; Peery & Bodenhausen, 2008). However, not all people show this effect (indeed, our participants overall did not show this effect). For example, the extent to which a person categorizes racially ambiguous faces as Black may depend on whether that person is liberal or conservative (Krosch et al., 2013). Other work has revealed that White, but not Black, participants tend to transfer negative implicit attitudes about a Black individual to a Black-White biracial individual or another Black individual but not to a White individual (Chen & Ratliff, 2015). Future work investigating the cognitive processes supporting categorical race perception should increase our understanding of attitudes toward racially mixed individuals.

4.3 Conclusion

Researchers have consistently focused on select areas when studying the neural basis of race processing: the visual cortex, fusiform gyrus, FFA, and the amygdala. In the current study on categorical race perception, we extend and complement the previous
evidence by showing that categorical perception is mediated by not only regions along the late visual processing stream but also higher-level cortical regions that fall outside these typical regions of interest. Our use of the whole brain searchlight approach combined with ROI-based MVPA allows us to investigate the entire cortical network involving both perceptual, attentional, and conceptual processing systems, affording greater insight into the neural and cognitive correlates of categorical perception of race.

Acknowledgments

We thank the staff at the Center for Brain Science at Harvard and the Martinos Center at MIT for their support and members of the Boston College Morality Lab for useful discussions. This work was supported by the Alfred P. Sloan Foundation (L.Y.; grant number BR2012) and a National Science Foundation Graduate Research Fellowship (L.T.; grant number 1258923).
References


CATEGORICAL RACE PERCEPTION


Supplementary Material

Functional localizer tasks

To define fusiform face area:

One functional localizer task (Epstein & Kanwisher, 1998) was used to define bilateral fusiform face area. Participants performed a one-back task in which they were required to press a button whenever two identical stimuli appeared in a row. Participants viewed 16 blocks of color photographs, four blocks for each of four stimulus types: scenes, faces, objects, and scrambled objects. Each block consisted of 16 trials, two of which contained a repeated image; different images were used across blocks. Block order was pseudorandomized. Images were presented for 600 ms with a 400 ms interstimulus interval. Participants completed two runs of this task; each run lasted 5.6 minutes.

To define regions involved in theory of mind:

A second localizer task (Dodell-Feder, Koster-Hale, Bedny, & Saxe, 2011) was used to functionally define regions involved in ToM or social cognition more generally. The task consisted of 20 visually presented stories matched in linguistic complexity, with 10 stories in each of two conditions: (1) stories requiring the inference of another person’s false beliefs and (2) stories requiring the inference of outdated (i.e., false) physical representations (e.g., an outdated photograph; for all stimuli, see http://saxelab.mit.edu/superloc.php). Each story was presented on the screen for 10 s, followed by a true/false question about the story (4 s). Participants completed two runs of this task; each run lasted 4.5 minutes.
**Table S1.** Localizer results. Mean values are provided for peak coordinates, # of voxels, and t values.

<table>
<thead>
<tr>
<th>Region of interest</th>
<th># of people (out of 24)</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th># of voxels</th>
<th>t value</th>
</tr>
</thead>
<tbody>
<tr>
<td>rFFA</td>
<td>21</td>
<td>42</td>
<td>-54</td>
<td>-20</td>
<td>37</td>
<td>6.72</td>
</tr>
<tr>
<td>lFFA</td>
<td>21</td>
<td>-40</td>
<td>-64</td>
<td>-17</td>
<td>31</td>
<td>6.39</td>
</tr>
<tr>
<td>rTPJ</td>
<td>21</td>
<td>53</td>
<td>-55</td>
<td>21</td>
<td>80</td>
<td>8.04</td>
</tr>
<tr>
<td>lTPJ</td>
<td>20</td>
<td>-51</td>
<td>-57</td>
<td>26</td>
<td>71</td>
<td>6.57</td>
</tr>
<tr>
<td>precuneus</td>
<td>21</td>
<td>-1</td>
<td>-57</td>
<td>38</td>
<td>80</td>
<td>7.28</td>
</tr>
<tr>
<td>dmPFC</td>
<td>15</td>
<td>4</td>
<td>54</td>
<td>30</td>
<td>47</td>
<td>5.55</td>
</tr>
</tbody>
</table>
Relationship between implicit race bias and responses on the race categorization post-scan task

We fit a sigmoid function to each participant's psychometric curve

\[ y = \frac{1}{1 + e^{-\frac{(x-a)}{b}}} \]

and determined the intercept \((a)\) and slope \((b)\) for each participant. Neither the slope (indicating the abruptness with which participants switched from consistently labeling faces along a continuum as one category to another category) nor the intercept (point along the continuum corresponding to a 'black' label 50% of the time) correlated with implicit race bias as measured with the IAT (Figure S1).
Figure S1. Slope (top) and intercept (bottom) from participants’ psychometric curves by implicit race bias as measured by a race Implicit Association Test (IAT). Slope refers to the abruptness with which participants switched from consistently labeling faces along the continuum as one category (‘white’) to another category (‘black’). Intercept refers to the point along the continuum corresponding to a ‘black’ label 50% of the time. Neither correlations were significant.