

# Self-regulation via neural simulation

Michael Gilead<sup>a,b,1</sup>, Chelsea Boccagno<sup>a</sup>, Melanie Silverman<sup>a</sup>, Ran R. Hassin<sup>c</sup>, Jochen Weber<sup>a</sup>, and Kevin N. Ochsner<sup>a,1</sup>

<sup>a</sup>Department of Psychology, Columbia University, New York, NY 10027; <sup>b</sup>Department of Psychology, Ben-Gurion University of the Negev, Be'er Sheva 8410501, Israel; and <sup>c</sup>Department of Psychology, The Hebrew University of Jerusalem, Jerusalem 9190501, Israel

Edited by Marcia K. Johnson, Yale University, New Haven, CT, and approved July 11, 2016 (received for review January 24, 2016)

**Can taking the perspective of other people modify our own affective responses to stimuli? To address this question, we examined the neurobiological mechanisms supporting the ability to take another person's perspective and thereby emotionally experience the world as they would. We measured participants' neural activity as they attempted to predict the emotional responses of two individuals that differed in terms of their proneness to experience negative affect. Results showed that behavioral and neural signatures of negative affect (amygdala activity and a distributed multivoxel pattern reflecting affective negativity) simulated the presumed affective state of the target person. Furthermore, the anterior medial prefrontal cortex (mPFC)—a region implicated in mental state inference—exhibited a perspective-dependent pattern of connectivity with the amygdala, and the multivoxel pattern of activity within the mPFC differentiated between the two targets. We discuss the implications of these findings for research on perspective-taking and self-regulation.**

perspective-taking | emotion regulation | mPFC | simulation | amygdala

**T**he ability to respond adaptively in the face of emotionally challenging situations is essential to mental and physical health. So much so, in fact, that emotion dysregulation is a core feature of virtually every form of psychopathology. Given this, it isn't surprising that the last decade has seen enormous growth in behavioral and brain research asking how we can effectively regulate our emotions. Although this work has made many important advances (1, 2), it has focused almost entirely on cognitive regulatory strategies that involve controlling attention to and/or rethinking the meaning of stimuli and events. As such, this work has completely overlooked the way in which social cognitive processes can be used to regulate our emotions.

The use of social cognition to regulate emotion was suggested by classic works in social psychology (3), which noted that by simulating others' perspective on the world we could shape our own experience and behavior. It is exemplified by "(Stanislavski) method actors" who understand a role by attempting to generate within themselves the presumed thoughts and feelings of a character, thereby allowing themselves to go beyond the written words in the script and respond as their character would (4). It is also present in everyday life when we seek guidance with respect to emotional dilemmas by asking ourselves how a friend, family member, mentor or religious figure (e.g., "What would Jesus do?") would respond in that situation.

In the current research we asked whether and how taking the perspective of other people can modify our own affective responses to stimuli. For example, by thinking of how someone more brave than ourselves would respond to a situation, we might down-regulate negative emotions, decrease aggression, and calm frazzled nerves. Alternatively, by thinking of how someone more sensitive and anxious would respond to the situation, we might enhance vigilance and increase reactivity to threatening situations.

To address these possibilities, we conducted a neuroimaging experiment investigating whether seeing the world through the eyes of a "tough" vs. a "sensitive" person can up-regulate or down-regulate affective responding, respectively. Furthermore, we sought to delineate the neural mechanisms by which such perspective-dependent regulatory consequences transpire.

Although no prior work has addressed these questions, per se, the literatures on emotion regulation (1, 5–11) and perspective-taking (12–18) can be integrated to generate testable hypotheses. On one hand, research on emotion regulation has shown that activity in lateral prefrontal cortex (i.e., dorsolateral prefrontal cortex and ventrolateral prefrontal cortex) and middle medial prefrontal cortex (i.e., presupplementary motor area, anterior ventral midcingulate cortex, and anterior dorsal midcingulate cortex) (19) supports the use of cognitive strategies to modulate activity in (largely) subcortical systems for triggering affective responses, such as the amygdala, thereby altering individuals' emotional responses (2). On the other hand, research on perspective-taking has shown that drawing inferences about the mental states of others (also known as "mentalizing")—as would be involved in simulating their perspective on an event—is supported by a network of regions centered on the anterior medial frontal cortex, specifically, the pregenual anterior cingulate cortex (pgACC) and the dorsomedial prefrontal cortex (dmPFC) (13, 19, 20).

Based on this literature, we formulated two hypotheses. First, we predicted that by taking the perspective of a target person, an individual could change behavioral and brain markers of affective responding, thereby providing evidence that one is emotionally experiencing the world the way the target would. Second, we predicted that these regulatory effects would be supported not by lateral prefrontal regions implicated in attentional and cognitive control, but rather, by dorsomedial prefrontal regions involved in perspective-taking. Put another way, we predicted that perspective-taking related activity in the anterior mPFC would regulate activity in neural systems for affective responding.

To test these hypotheses, we collected whole-brain fMRI data while participants attempted to predict the affective responses of other individuals. Before scanning, participants were presented with descriptions of two people, who they were led to believe had

## Significance

**As Harper Lee tells us in *To Kill a Mockingbird*, "You never really understand a person until you consider things from his point of view, until you climb in his skin and walk around in it." Classic theories in social psychology argue that this purported process of social simulation provides the foundations for self-regulation. In light of this, we investigated the neural processes whereby humans may regulate their affective responses to an event by simulating the way others would respond to it. Our results suggest that during perspective-taking, behavioral and neural signatures of negative affect indeed mimic the presumed affective state of others. Furthermore, the anterior medial prefrontal cortex—a region implicated in mental state inference—may orchestrate this affective simulation process.**

Author contributions: M.G., M.S., R.R.H., J.W., and K.N.O. designed research; M.G., C.B., and M.S. performed research; M.G. and J.W. analyzed data; and M.G. and K.N.O. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

<sup>1</sup>To whom correspondence may be addressed. Email: mgilead@bgu.ac.il or ochsner@psych.columbia.edu.

This article contains supporting information online at [www.pnas.org/lookup/suppl/doi:10.1073/pnas.1600159113/-DCSupplemental](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1600159113/-DCSupplemental).

previously participated in the experiment. These descriptions suggested that one person was likely to be emotionally sensitive and squeamish, whereas the other was likely to be rugged and tough. Next, participants viewed neutral and negative affect-inducing images and evaluated the images from either their own or the tough or sensitive targets' perspective.

We examined the effect of perspective-taking on multiple behavioral and brain markers of affective responding, including reports of the target's predicted affective reactions to stimuli, activation in the amygdala (which is the brain region most strongly associated with detecting, encoding, and promoting responses to affectively relevant and especially potentially threatening stimuli) (21, 22), and finally, a recently identified picture-induced negative emotion signature (PINES) (23). PINES is a distributed, whole-brain multivoxel activation pattern developed using machine learning techniques that can reliably predict levels of negative affect elicited by aversive images. Because this signature is not affected by general arousal and is not reducible to activity in the amygdala, it provides a neural marker of negative affect independent of participants' own self-reports. We predicted that both neural measures of negative affective responding (amygdala and PINES) would simulate the presumed affective state of the target person; namely, negative affect-related activity would be up- vs. down-regulated for the sensitive (vs. tough) perspective.

To address the prefrontal systems that might support perspective-taking and regulate affective responding, we used a combination of connectivity and multivoxel pattern analyses to identify a brain region whose activity was associated with amygdala up-regulation when adopting the sensitive perspective and/or down-regulation for the tough perspective—and whose distributed pattern of activity provided evidence that it differentially represented the two perspectives. As noted, we predicted this region to be located in the anterior mPFC.

## Results

### Does Perspective-Taking Modulate Affective Processing?

**Behavioral ratings.** A manipulation check showed that participants reported more negative affect in response to negative than to neutral images [ $F(1,23) = 572.56$ ,  $P < 0.001$ ]. We conducted an ANOVA to see whether the perspective manipulation indeed altered participants predicted affective response. The results showed a significant interaction [ $F(1,23) = 202.08$ ,  $P < 0.001$ ], such that affect ratings were lower when participants viewed negative images from the perspective of the tough (mean = 2.564, SD = 0.118) vs. the sensitive target [mean = 3.793, SD = 0.103;  $t(23) = 12.60$ ,  $P < 0.001$ ]. There was no significant difference in ratings for neutral images from the perspective of the tough (mean = 1.071, SD = 0.021) and sensitive (mean = 1.117, SD = 0.028) targets [ $t(23) = 1.58$ ,  $P = 0.126$ ; Fig. 1]. There were also no significant differences in response latencies for the sensitive (mean = 921.57, SD = 129.19)

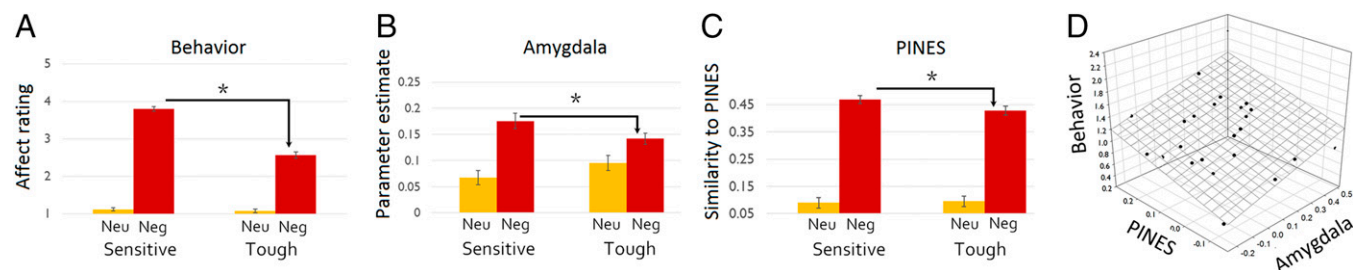
and tough (mean = 941.90, SD = 120.97) perspectives [ $t(23) = 0.8$ ; not significant].

For negative images, affect ratings from the self perspective (mean = 3.140, SD = 0.496) were higher than those for the tough target [ $t(23) = 5.08$ ,  $P < 0.001$ ] and lower than those for the sensitive target [ $t(23) = 8.10$ ,  $P < 0.001$ ]. For neutral images, affect ratings from the self perspective (mean = 1.058, SD = 0.104) did not differ from the tough perspective [ $t(23) = 0.48$ , not significant] and were lower than those for the sensitive perspective [ $t(23) = 2.63$ ,  $P = 0.015$ ].

Based on participants' affect ratings for the tough, sensitive, and self perspectives, we calculated for each participant a measure of "similarity to sensitive/tough target" that indexed the extent to which affect ratings from the self perspective were more similar to one target or the other. This measure, alongside with other neural and self-report measures of self-other similarity, indicated that, overall, participants did not identify more with one perspective or another and that the level of self-other similarity did not modulate our key measures (see *SI Experimental Procedures*, Fig. S1, and Table S1 for details of these analyses).

**Amygdala analysis.** As a first step in examining whether perspective-taking modulates affective processing, we defined the right and left amygdala as anatomical regions of interest based on the Harvard-Oxford probabilistic atlas (using voxels with a 50% or higher probability of being labeled as the amygdala) and extracted parameter estimates for the six conditions (negative/neutral × sensitive/tough/self). As predicted, in the left amygdala, when participants observed the images from their own perspective, activation was higher for negative (mean = 0.177, SD = 0.167) than for neutral (mean = 0.086, SD = 0.148) images [ $t(23) = 2.77$ ,  $P = 0.005$ ]; likewise, in the right amygdala, activation was higher for negative (mean = 0.149, SD = 0.149) than for neutral (mean = 0.087, SD = 0.112) images [ $t(23) = 2.15$ ,  $P = 0.020$ ].

After establishing that amygdala activity is responsive to the presentation of aversive images when viewing them from one's own perspective, we asked whether the amygdala was modulated when taking a tough or sensitive perspective. To do so, we conducted a  $2 \times 2$  ANOVA with perspective (sensitive/tough) and valence (negative/neutral) as within-participant factors. As predicted, the results showed an interaction of perspective and valence in both the right [ $F(1,23) = 6.77$ ,  $P = 0.007$ , partial  $\eta^2 = 0.227$ ] and the left amygdala [ $F(1,23) = 2.96$ ,  $P = 0.049$ , partial  $\eta^2 = 0.114$ ]. In the right hemisphere, amygdala activation was lower when viewing negative images from the perspective of the tough (mean = 0.141, SD = 0.098) vs. the sensitive target (mean = 0.175, SD = 0.120) [ $t(23) = 2.02$ ,  $P = 0.027$ ]; there was no significant difference in activation for neutral images from the perspective of the tough (mean = 0.095, SD = 0.092) and sensitive (mean = 0.067, SD = 0.089) targets [ $t(23) = 1.38$ ,  $P = 0.180$ ]. In the left hemisphere, there was a marginally significant effect wherein



**Fig. 1.** (A) Behavioral ratings of negative affect in response to negative images were higher for sensitive (vs. tough) targets. (B) right amygdala response to negative images was higher when adopting the sensitive (vs. the tough) perspective. (C) When participants adopted the sensitive (vs. tough) perspective, their neural response to negative images reflected higher levels of negative affect, measured as the level of similarity to the PINES pattern. Error bars denote within-participant SEs. (D) Participants who exhibited a greater difference in amygdala activity and PINES expression for the tough vs. sensitive target subsequently estimated greater differences in predicted negative affect for these targets.

amygdala activation was lower when viewing negative images from the perspective of the tough (mean = 0.175, SD = 0.130) vs. sensitive target (mean = 0.200, SD = 0.129) [ $t(23) = 1.45$ ,  $P = 0.079$ ]. There was no significant difference in activation for neutral images from the perspective of the tough (mean = 0.100, SD = 0.117) and sensitive (mean = 0.078, SD = 0.105) targets [ $t(23) = 1.04$ ,  $P = 0.306$ ]. For both negative and neutral images, the self perspective did not differ from the sensitive perspective in either the right or left amygdala ( $P > 0.27$ ); likewise, the self perspective did not differ from the tough perspective in either the right or left amygdala ( $P > 0.64$ )—suggesting that the perspective  $\times$  valence interaction was not driven solely by either the tough or sensitive perspective.

Although the amygdala was more active for negative vs. neutral images when viewed from the self's perspective, it could be argued that different subregions of the amygdala may be differentially engaged under the self and other conditions. To address this concern, we conducted a whole-brain search based on the self negative  $>$  self neutral contrast. This contrast yielded significant activation across several brain regions, including the left and right amygdala, which we then masked with anatomically defined amygdala regions based on the Harvard-Oxford probabilistic atlas. The interaction of perspective and valence remained significant in the right amygdala cluster [47 voxels, peak coordinate,  $x = 18$ ,  $y = -3$ ,  $z = -18$ ;  $F(1,23) = 5.25$ ,  $P = 0.015$ ]; however, the interaction in the left amygdala (102 voxels, peak MNI coordinate,  $x = -12$ ,  $y = -6$ ,  $z = -18$ ) did not attain significance [ $F(1,23) = 1.25$ ,  $P = 0.136$ ]. In light of this, we limited our subsequent analyses to the right amygdala cluster.

**PINES analysis.** Another concern is that, although amygdala activation is strongly associated with the processing of negatively valenced stimuli, it is sometimes activated when processing positive stimuli (24), which may reflect a more general role for the amygdala in detecting and encoding of goal-relevant stimuli (25–27). These findings suggest that the amygdala's role in negative affect be indirect, which complicates attempts to rely on its activation as a neural marker of negative affective responses.

In light of this, we sought to strengthen our claim that emotional perspective-taking modulates negative affective processing by using a recently identified PINES (23). The PINES is a whole brain activation pattern developed using machine learning techniques that can reliably predict self-reported emotional responses to aversive images. As noted, prior work (23) has shown that this signature is not affected by general arousal, and is not reducible to patterns of activity in the amygdala. Thus, it provides an independently validated neural marker of experienced affective negativity.

We first validated the PINES method in the current dataset by showing that the PINES expression score was significantly higher when observing negative (mean = 0.490, SD = 0.192) vs. neutral (mean = 0.069, SD = 0.152) images from the perspective of the self [ $t(23) = 11.24$ ,  $P < 0.001$ ]. Furthermore, regardless of perspective, the PINES score was higher for negative (mean = 0.462, SD = 0.175) vs. neutral (mean = 0.094, SD = 0.156) images viewing conditions [ $F(1,23) = 349.18$ ,  $P < 0.001$ ]. Having established that the PINES pattern differentiates images as a function of their negativity in our dataset, we investigated the effect of perspective (sensitive/tough) and valence (negative/neutral) on the degree of affective negativity, as gauged by the PINES expression score. As predicted, there was an interaction between perspective and valence, mirroring the effect in the amygdala [ $F(1,23) = 7.72$ ,  $P = 0.005$ ]. The PINES expression score was lower when viewing negative images from the perspective of the tough (mean = 0.428, SD = 0.173) vs. sensitive target (mean = 0.468, SD = 0.188) [ $t(23) = 1.83$ ,  $P = 0.039$ ]; there was no significant difference in expression for neutral images from the perspective of the tough (mean = 0.095, SD = 0.092) vs. the sensitive (mean = 0.067, SD = 0.089) targets [ $t(23) = 1.28$ ,  $P = 0.210$ ]. The PINES calculation was done on a trial-based model that we used for multivariate

analyses [PINES, multivoxel pattern analysis (MVPA), and pattern similarity analysis]. The results of the analysis are identical when using an aggregated-trial model. Valence  $\times$  perspective interaction:  $F(1,23) = 7.72$ ,  $P = 0.005$  (see *SI Experimental Procedures* for further details concerning PINES method).

**Brain-behavior correlation.** Having established that the different measures of affective response are each impacted by the perspective taken, an important next question was whether and how they are related. In particular, it is important to demonstrate that one or both of the neural measures—amygdala activity and/or PINES score—predict self-reports of negative affective experience, as such correlations would support the idea that the neural regions supporting simulation of the tough vs. sensitive target's emotions lead to changes in a reports of affective experience.

To address this issue, we calculated for each participant a measure of each type of effect on a measure of affective response (i.e., the behavioral effect, PINES effect, and amygdala effect), as the difference between negative and neutral conditions for the sensitive vs. tough perspectives [i.e., sensitive (negative-neutral) – tough (negative-neutral)]. As predicted, the results showed that participants who exhibited a greater difference in amygdala activity for the tough vs. sensitive target subsequently exhibited a greater difference in their behavioral evaluations of the affective states of these targets ( $r = 0.38$ ,  $P = 0.033$ ). Likewise, participants who exhibited a greater difference in PINES scores for the tough vs. sensitive target subsequently exhibited a greater difference in their behavioral evaluations of the affective states of these targets ( $r = 0.39$ ,  $P = 0.014$ ). Interestingly, there was no correlation between the PINES effect and amygdala effect ( $r = 0.01$ ), as would be expected based on prior work establishing the PINES that suggested they could be independent predictors of negative affect (23). A multiple regression with both the PINES effect and amygdala effect as predictors and the behavioral effect as the dependent variable showed a significant effect for the PINES ( $b = 1.193$ , SE = 0.562, 95% CI: 0.023, 2.364,  $P = 0.046$ ) and a marginally significant effect for the amygdala ( $b = 0.742$ , SE = 0.363, 95% CI: –0.103, 1.499,  $P = 0.053$ ;  $R^2 = 0.295$ ). Thus, our results suggest that each of the two patterns made a unique contribution to changing reports of affective responding.

### What Are the Neural Systems That Support the Perspective-Based Modulation of Affective Processing?

**Psychophysiological interaction.** To identify regions that may play a key role in the perspective-taking-based regulation of amygdala activity, we conducted a psychophysiological interaction (PPI) analysis (28). This analysis was done by creating regressors for each of the experimental conditions, the amygdala time series, and interaction terms for the amygdala time series and the experimental conditions. The difference of the relevant PPI-term regression coefficient, i.e., [(amygdala time series)  $\times$  (sensitive negative)]  $>$  [(amygdala time series)  $\times$  (tough negative)] was then subjected to a second-level random effects analysis, which also included a between-participants covariate coding for the average difference in affect rating across conditions.

This analytical strategy allowed us to identify regions that during sensitive negative trials were more positively correlated with right amygdala activation and/or during tough negative trials we more negatively correlated with amygdala activity and exhibited this pattern more so for participants that displayed greater perspective-related modulation of affective response (i.e., greater behavioral effect). The resulting analysis yielded a cluster of 203 voxels in the anterior mPFC (specifically, pgACC and dmPFC; peak MNI coordinate,  $x = -9$ ,  $y = 54$ ,  $z = 15$ ; Fig. 2), which survived the  $P < 0.05$ , whole-brain corrected significance threshold determined by AlphaSim. Masking out this anterior mPFC cluster did not alter the results of the PINES analysis.

In other words, participants who showed the greatest perspective-dependent modulation of affective experience also showed the





modulated through an interaction with a brain system that subserves such model-based, conceptual capacities.

As noted earlier, our results suggest that this system involves the anterior mPFC. This region is widely implicated in conceptual thought in general (33–35) and social cognition in particular (17, 36). To give one example, recent work shows that multivoxel patterns of activity in the anterior mPFC can be used to predict which one of two individuals a participant is thinking about (20). The current research dovetails and builds on this prior work by showing that anterior mPFC doesn't just support inferences about others states and traits but supports simulation of their perspective on world, thereby changing the way that we appraise the affective significance of events and subsequently respond to them.

**Implications for Models of the Self-Regulation of Emotion.** An important implication of the current findings is the suggestion that perspective-taking could have emotion regulatory benefits. In the current study, participants did not have the explicit goal of up- or down-regulating their emotions, and yet, merely trying to understand the emotions of tough vs. sensitive others modulated the activity in a brain system involved in the generation of negative affect. Thus, our research suggests that the attempt to “walk in the shoes” of an emotionally resilient individual may cause people to feel less unpleasant in the face of adversity.

Accordingly, it may be possible to harness the type of emotional perspective-taking studied here as an emotion regulation strategy, aimed at helping individuals cope with emotional distress. Extant research within the field of emotion regulation has shown that people can effectively down-regulate negative affect by using top-down cognitive control (2). However, a limitation of many cognitive emotion regulation strategies is that they depend on attentional, linguistic, and working memory systems supported by lateral prefrontal regions. Lateral prefrontal regions are not fully developed until late adolescence (37) and can be disrupted under severe stress (38). Thus, the finding that perspective-based regulation of the amygdala relies on anterior medial rather than lateral prefrontal regions may suggest a new pathway for effective emotion regulation.

Specifically, a simulation-based emotion regulation strategy may be important in populations for which strategies dependent on lateral PFC may be problematic because lateral frontal functionality is compromised or yet to develop (39). For example, future studies could investigate whether young children may especially benefit from being taught how to regulate their emotions using simulative pretend play (“imagine that you are a big boy/girl”).

More broadly, the current findings highlight that there may be a plurality of computations and neural pathways by which emotion-regulatory consequences can occur. In this way, the current findings contribute to our growing understanding of the complexity of neural interactions that subserve important behavioral outcomes. Hopefully, future research extending the findings described herein could shed further light on strategies that support adaptive socioemotional functioning.

## Experimental Procedures

**Participants.** Twenty-four right-handed participants (12 females; average age, 20.5 y; SD = 2.577; range 18–28 y) participated in the experiment for monetary compensation. All were native-level English speakers, all had normal or corrected vision, and none had a history of neurological or psychiatric disorders. Sample size was determined a priori, based on previous neuroimaging studies showing regulation-related modulation of amygdala activity (2). Three additional participants were excluded from the final analysis (one for missing data and two for failing to comply with task instructions, as evident by deviation of more than 3 SDs from the mean affect rating in at least one task condition). Participants gave written consent before taking part in the experiment. The study was approved by the Institutional Review Board of Columbia University.

## Materials.

**Target description questionnaires.** The descriptions of the tough and sensitive targets were given in the form of printed questionnaires that were ostensibly

filled out by two previous participants. At the top of each questionnaire, a name appeared in hand-written text. Both names were matched to each participants' sex. The questionnaire contained demographic details (e.g., place of birth) and responses to personal questions (e.g., music preferences, hobbies). The key differences between the two types of targets arose from the way each one had supposedly responded to particular questions. In actuality, the answers had been pretested to elicit perceptions that one target was tough and the other sensitive. For example, the tough character worked as an EMT and enjoyed action and horror movies and loud music. By contrast, the sensitive character worked as a graphic designer and liked classical music and romantic comedies. Furthermore, in one of the free response items the tough target described him/herself as being relatively resilient and the sensitive character described him/herself as being relatively sensitive. These characteristics were embedded within more mundane details to bolster the believability of the experiment.

**Affective stimuli.** Fifty-four negative images (mean normative valence = 2.76, mean normative arousal = 5.91, on a 1–9 scale) and 54 neutral images (mean normative valence = 5.32, mean normative arousal = 3.15) were taken from the International Affective Picture System (40). Both negative and neutral images were divided to three lists, matched for arousal and valence. An additional set of six similarly valenced and arousing negative images were used during training.

## Behavioral Procedure.

**Prescanning.** After providing consent, participants were asked to fill out a questionnaire describing various demographic and personal details about themselves. They were told that in the experiment they will be asked to predict the emotions of previous participants and that we need their answers to the personal details questionnaire to use them for the next participant. In actuality, this questionnaire was only administered to bolster the believability of the experiment, and it was not subsequently used. Immediately after filling out the questionnaire, participants were given the “character description” questionnaires, which were in the same format as the one they filled out. They were asked to read the answers of each previous participant carefully and form an impression of them in their mind.

Participants then were instructed on the task they would perform inside the scanner. They were told that they will be presented with images and that each image will be preceded either by a cue with the name of the participant whose perspective they should take or by a cue asking them to take their own perspective. Each image would be followed with a screen asking them to rate the affective response (either of themselves or the target individual) the image elicits. They were then told that they should rate the images based upon the perspective they were cued with, and that these answers would be compared with the previous participants' actual ratings. We told participants that trials wherein they gave the rating from their own perspective would be used for the next participants (in actuality, self-perspective trials were used to identify the neural substrates of spontaneous emotional response). Participants' goal was to predict the previous participants' responses as accurately as possible. To increase the incentive to do so, participants were told that if they were in the top 10% of participants in terms of accuracy, they will receive a \$100 bonus (in actuality, the bonus criteria was based on scanner movement). Participants then performed a short training on the task that involved completing sample trials guided by the experimenter.

Finally, as a pretask manipulation check, participants were asked to recall the answers for each of the two previous participants' questionnaires. Whenever participants made a mistake, the questions were repeated later on until participants arrived at 100% recall accuracy.

**Scanner task.** The task consisted of 108 trials (18 negative images and 18 neutral images for each of the three perspectives) that were divided into three functional runs. Each run contained 36 trials (6 negative and 6 neutral for each of the three perspectives) and lasted 10 min and 48 s.

Stimuli were presented using E-Prime 2.0 (Psychology Software Tools). Each experimental trial began with the presentation a cue with the name of the participant whose perspective they should take, or a cue asking them to take their own perspective, shown for 2 s. After a jittered fixation period (1–5 s), participants viewed the affective image for 6 s. The image was replaced by a screen that appeared for 3 s, asking them to rate the affective reaction to the image from the perspective they were asked to adopt (1 = neutral, 5 = very bad). The trial concluded with a second jittered fixation period (3–9 s). Stimuli were displayed in random order and the assignment of images to the three perspective conditions was counterbalanced across participants.

**Postscan.** At the end of the study, participants completed standardized questionnaires assessing individual differences in affective responding [Beck depression inventory (41) and state-trait anxiety inventory (42)] and perspective-taking [interpersonal reactivity index (43)]. None of these

individual-difference measures were significantly correlated with our dependent variables of interest (PINES scores, amygdala activity, affect ratings) nor did they moderate the effect of perspective (or the interaction of perspective and valence) on these dependent variables. In light of this, they are not discussed in results section.

**ACKNOWLEDGMENTS.** M.G. thanks M. Perkas, L. Chang, B. Dore, C. Helion, J. Shu, B. Denny, and R. Martin for assistance. M.G. is supported by fellowships from the Fulbright and Rothschild foundations. Completion of the manuscript was supported by National Institute on Aging Grant AG043463, National Institute of Child Health and Human Development Grant HD069178, and National Institute of Mental Health Grant MH090964 (to K.N.O.).

- Ochsner KN, Silvers JA, Buhle JT (2012) Functional imaging studies of emotion regulation: A synthetic review and evolving model of the cognitive control of emotion. *Ann N Y Acad Sci* 1251:E1–E24.
- Buhle JT, et al. (2014) Cognitive reappraisal of emotion: A meta-analysis of human neuroimaging studies. *Cereb Cortex* 24(11):2981–2990.
- Mead GH (1934) *Mind, Self, and Society: From the Standpoint of a Social Behaviorist (Works of George Herbert Mead, Vol. 1)* (Univ of Chicago Press, Chicago).
- Bandelj N (2003) How method actors create character roles. *Sociol Forum* 18(3): 387–416.
- Gross JJ (1998) Antecedent- and response-focused emotion regulation: Divergent consequences for experience, expression, and physiology. *J Pers Soc Psychol* 74(1): 224–237.
- McRae K, et al. (2010) The neural bases of distraction and reappraisal. *J Cogn Neurosci*, 22(2):248–262.
- Ochsner KN, Bunge SA, Gross JJ, Gabrieli JDE (2002) Rethinking feelings: An fMRI study of the cognitive regulation of emotion. *J Cogn Neurosci* 14(8):1215–1229.
- Ochsner KN, Gross JJ (2008) Cognitive emotion regulation: Insights from social cognitive and affective neuroscience. *Curr Dir Psychol Sci* 17(2):153–158.
- Wager TD, Davidson ML, Hughes BL, Lindquist MA, Ochsner KN (2008) Prefrontal-subcortical pathways mediating successful emotion regulation. *Neuron* 59(6):1037–1050.
- Kross E, Davidson M, Weber J, Ochsner K (2009) Coping with emotions past: The neural bases of regulating affect associated with negative autobiographical memories. *Biol Psychiatry* 65(5):361–366.
- Diekhof EK, Geier K, Falkai P, Gruber O (2011) Fear is only as deep as the mind allows: A coordinate-based meta-analysis of neuroimaging studies on the regulation of negative affect. *Neuroimage* 58(1):275–285.
- Frith CD, Frith U (2006) The neural basis of mentalizing. *Neuron* 50(4):531–534.
- Mitchell JP, Banaji MR, Macrae CN (2005) The link between social cognition and self-referential thought in the medial prefrontal cortex. *J Cogn Neurosci* 17(8):1306–1315.
- Mitchell JP, Macrae CN, Banaji MR (2006) Dissociable medial prefrontal contributions to judgments of similar and dissimilar others. *Neuron* 50(4):655–663.
- Tamir DI, Mitchell JP (2010) Neural correlates of anchoring-and-adjustment during mentalizing. *Proc Natl Acad Sci USA* 107(24):10827–10832.
- Saxe R, Kanwisher N (2003) People thinking about thinking people. The role of the temporo-parietal junction in “theory of mind”. *Neuroimage* 19(4):1835–1842.
- Skerry AE, Saxe R (2014) A common neural code for perceived and inferred emotion. *J Neurosci* 34(48):15997–16008.
- Spunt RP, et al. (2015) Amygdala lesions do not compromise the cortical network for false-belief reasoning. *Proc Natl Acad Sci USA* 112(15):4827–4832.
- de la Vega A, Chang LJ, Banich MT, Wager TD, Yarkoni T (2016) Large-scale meta-analysis of human medial frontal cortex reveals tripartite functional organization. *J Neurosci* 36(24):6553–6562.
- Hassabis D, et al. (2014) Imagine all the people: How the brain creates and uses personality models to predict behavior. *Cereb Cortex* 24(8):1979–1987.
- Phan KL, Wager T, Taylor SF, Liberzon I (2002) Functional neuroanatomy of emotion: A meta-analysis of emotion activation studies in PET and fMRI. *Neuroimage* 16(2): 331–348.
- Stillman PE, Van Bavel JJ, Cunningham WA (2015) Valence asymmetries in the human amygdala: Task relevance modulates amygdala responses to positive more than negative affective cues. *J Cogn Neurosci* 27(4):842–851.
- Chang LJ, Gianaros PJ, Manuck SB, Krishnan A, Wager TD (2015) A Sensitive and Specific Neural Signature for Picture-Induced Negative Affect. *PLoS Biol* 13(6): e1002180.
- Breiter HC, et al. (1996) Response and habituation of the human amygdala during visual processing of facial expression. *Neuron* 17(5):875–887.
- Cunningham WA, Brosch T (2012) Motivational salience: Amygdala tuning from traits, needs, values, and goals. *Curr Dir Psychol Sci* 21(1):54–59.
- Lindquist KA, Wager TD, Kober H, Bliss-Moreau E, Barrett LF (2012) The brain basis of emotion: A meta-analytic review. *Behav Brain Sci* 35(3):121–143.
- Whalen PJ, Phelps EA (2009) *The Human Amygdala* (Guilford Press, New York).
- Friston KJ, et al. (1997) Psychophysiological and modulatory interactions in neuroimaging. *Neuroimage* 6(3):218–229.
- Goldman AI (2005) Imitation, mind reading, and simulation. *Perspective on Imitation, from Neuroscience to Social Science*, eds Hurley S and Chater N (MIT Press, Cambridge, MA) Vol 2, pp. 79–93.
- Epley N, Keysar B, Van Boven L, Gilovich T (2004) Perspective taking as egocentric anchoring and adjustment. *J Pers Soc Psychol* 87(3):327–339.
- Gordon R (1986) Folk psychology as simulation. *Mind Lang* 1(2):158–171.
- Ryle G (1949) *The Concept of Mind* (Hutchinson, New York).
- Fairhall SL, Caramazza A (2013) Brain regions that represent amodal conceptual knowledge. *J Neurosci* 33(25):10552–10558.
- Binder JR, Desai RH, Graves WW, Conant LL (2009) Where is the semantic system? A critical review and meta-analysis of 120 functional neuroimaging studies. *Cereb Cortex* 19(12):2767–2796.
- Honey CJ, Thompson CR, Lerner Y, Hasson U (2012) Not lost in translation: Neural responses shared across languages. *J Neurosci* 32(44):15277–15283.
- Denny BT, Kober H, Wager TD, Ochsner KN (2012) A meta-analysis of functional neuroimaging studies of self- and other judgments reveals a spatial gradient for mentalizing in medial prefrontal cortex. *J Cogn Neurosci* 24(8):1742–1752.
- Gogtay N, et al. (2004) Dynamic mapping of human cortical development during childhood through early adulthood. *Proc Natl Acad Sci USA* 101(21):8174–8179.
- Raio CM, Oederer TA, Palazzolo L, Shurick AA, Phelps EA (2013) Cognitive emotion regulation fails the stress test. *Proc Natl Acad Sci USA* 110(37):15139–15144.
- Coccaro EF, Sripada CS, Yanowitch RN, Phan KL (2011) Corticolimbic function in impulsive aggressive behavior. *Biol Psychiatry* 69(12):1153–1159.
- Lang PJ, Bradley MM, Cuthbert BN (2008) *International Affective Picture System (IAPS): Affective Ratings of Pictures and Instruction Manual* (University of Florida, Gainesville, FL), Tech Rep A-8.
- Beck AT, Steer RA, Garbin MG (1988) Psychometric properties of the Beck depression inventory: 25 years of evaluation. *Clin Psychol Rev* 8(1):77–100.
- Marteau TM, Bekker H (1992) The development of a six-item short-form of the state scale of the Spielberger State-Trait Anxiety Inventory (STAI). *Br J Clin Psychol* 31(Pt 3): 301–306.
- Davis MH (1983) Measuring individual-differences in empathy: Evidence for a multi-dimensional approach. *J Pers Soc Psychol* 44(1):113–126.