The speed of human social interaction perception

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Abstract

The ability to perceive others’ social interactions, here defined as the directed contingent actions between two or more people, is a fundamental part of human experience that develops early in infancy and is shared with other primates. However, the neural computations underlying this ability remain largely unknown. Is social interaction recognition a rapid feedforward process or a slower post-perceptual inference? Here we used magnetoencephalography (MEG) decoding to address this question. Subjects in the MEG viewed snapshots of visually matched real-world scenes containing a pair of people who were either engaged in a social interaction or acting independently. The presence versus absence of a social interaction could be read out from subjects’ MEG data spontaneously, even while subjects performed an orthogonal task. This readout generalized across different people and scenes, revealing abstract representations of social interactions in the human brain. These representations, however, did not come online until quite late, at 300 ms after image onset, well after feedforward visual processes. In a second experiment, we found that social interaction readout still occurred at this same late latency even when subjects performed an explicit task detecting social interactions. We further showed that MEG responses distinguished between different types of social interactions (mutual gaze vs joint attention) even later, around 500 ms after image onset. Taken together, these results suggest that the human brain spontaneously extracts information about others’ social interactions, but does so slowly, likely relying on iterative top-down computations.

1. Introduction

As fundamentally social primates, humans need to know who is doing what to whom, and why. Indeed, the ability to perceive and interpret social interactions between other agents is shared with other primates (Sliwa and Freiwald, 2017) and develops early in infancy (Hamlin et al., 2015a) and young children (Cowell Jason and Decety, 2015b). Further, social interaction representations are apparently computed in a specialized region of the posterior superior temporal sulcus (Isik et al., 2017; Walbrin et al., 2017). These findings underscore the importance of social interaction perception, but leave unanswered the question of how this information is extracted from visual input. In particular, is social interaction recognition a rapid feedforward process, akin to object recognition, or a slower post-perceptual inference?

Considerable evidence suggests that much of visual perception in primates, including face, scene, and “core” object recognition, is computed by rapid and largely feedforward pattern classification processes. First, these tasks in primates are well approximated by purely feedforward neural network models, not only in terms of accuracy but also in terms of the representations extracted (Khaligh-Razavi and Kriegeskorte, 2014; Radoslaw Martin Cichy, 2016; Yamins et al., 2014). Second, visual recognition in primates is fast, occurring within 200 ms of image onset, as expected of a largely feedforward process. These fast latencies have been demonstrated for face (Bentin et al., 1996; Dobs et al., 2016), scene (Cichy et al., 2016a; Greene and Hansen, 2018), and object (Carlson et al., 2013a; Isik et al., 2014; Yamins et al., 2014) recognition. In contrast, some visual information cannot be computed from bottom-up visual information alone. Object recognition under complex viewing conditions, such as occlusion takes longer (~300 ms), and cannot be performed with purely feedforward models (Rajaee et al., 2018; Tang et al., 2018, 2014). Generative models offer an attractive solution to these challenging vision problems (Wu et al., 2016; Yuille and Kersten, 2006). Rather than relying solely on bottom-up cues, these systems build models of objects and the world around them, and use these generated models as hypotheses to interpret incoming visual information.

Behavioral studies have suggested that the perception of social interactions shares some of the hallmarks of a classic visual pattern recognition problem: face recognition. First, people are better able to perceive social interactions when stimuli are presented upright rather

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of groups of individuals (Vestner et al., 2019). Others have tried to address this question with computational modeling. One study showed that different types of social interactions can be distinguished based on bottom-up visual cues (Blythe et al., 1999), but more recent work has suggested that top down or generative models are required to solve this problem (Ben-Yosef et al., 2017; Ullman et al., 2009). Importantly, all these modeling efforts focused on categorizing different types of social interactions (e.g., a hug versus a handshake), and it remains an open question whether feedforward computations are sufficient to simply detect social interactions (i.e., identify whether or not two agents are engaged in a social interaction).

Neurophysiologically, body-selective visual cortex responds more to bodies facing towards versus away from each other (Abassi and Papeo, 2019). This region may serve as a precursor to the pSTS “social interaction region”, which is selectively engaged in social interaction detection and categorization, independent of the number and body position of the agents (Elišk et al., 2017; Walbrin et al., 2017). It remains unknown, however, whether bottom-up information from relevant visual features, such as body pose, are sufficient to detect social interactions.

Neural dynamics are a useful tool to understand such computational questions. Understanding when different representations occur relative to each other can indicate how computations unfold over time and distinguish between different computational theories (e.g., fast feedforward processing versus slower recurrent computations, as described above). Here we used MEG decoding to ask whether social interactions can be detected via fast, feedforward processing. Using decoding methods, we ask whether the detection and categorization of a social interaction in a visual stimulus occurs on the rapid time scale of invariant object recognition (about 150 ms), as predicted from a feedforward pattern classification model, or more slowly, as expected if it requires a top-down inference. The perception of social interactions is a highly complex phenomenon that we operationalize in this study as follows. By social interactions we mean the relational process of two or more agents acting toward one another (for example looking at one another or “mutual gaze”) or a shared point of interest (for example looking at a common object or “joint attention”). We focused on these two gaze-based interactions because they are two fundamental types of social interactions that are easily depicted in images and among the earliest identified by infants (Beier and Spekela, 2012; Tomasello and Farrar, 1986). We use minimal yet naturalistic stimuli to depict these two cases of social interactions and contrast them to situations where the agents are acting independently (with separate objects of attention or “independent actions”) or where only one agent is directing their attention toward the other (“watch”). Perceiving these interactions is also a complex process that we break down here into two primary components: detection (whether or not two agents are involved in a social interaction) and categorization (which type of interaction the agents are involved in).

We find both the presence (detection) and type (categorization) of social interaction could be decoded from subjects’ MEG data, but this readout occurred quite late, at 300 ms and 500 ms for detection and categorization, respectively. In a second experiment, we showed that this readout did not occur earlier even when subjects performed an explicit social interaction detection task.

2. Material and methods

The below methods were pre-registered on the Open Science Framework platform: https://osf.io/3vnem/registrations. Any deviations from our pre-registration are noted as exploratory analyses.

2.1. Social interaction dataset

We created an image dataset depicting pairs of people interacting with each other or independently in different ways. There were five different conditions, shot across 12 scenes with 12 different actor pairs (60 images total, see Fig. 1A-B for example images). The five conditions differed in the way each pair of people were or were not interacting with each other. The different conditions include:

i. Mutual gaze – pair of actors is looking at each other. Social interaction.
ii. Joint attention – pair of actors looking at the same object. Social interaction.
iii. Independent actions 1 – two actors are engaged in separate independent actions. No social interaction.
iv. Independent actions 2 – two actors are engaged in separate independent actions (different actions from above). No social interaction.
v. Watch – one actor watches the other actor who is looking away. One-way interaction.

These five conditions were included to encompass our operational definition of a social interaction: two agents with actions directed towards each other (mutual gaze) or a shared point of interest (joint attention). We sought to distinguish these two relational actions from cases where the actors acted independently (independent actions) or a case where one agent is aware of the other and has “perceptual access” (watch), but the relation between the two agents is not present.

The images were shot in a way to minimize changes between conditions besides the objects of the actors’ actions (i.e., objects or other actors). Actors were spaced at roughly equal distances in all conditions and visual information like background and lighting were fixed across conditions. We further ensured that different visual features could not distinguish between scenes with versus without a social interaction using a deep convolutional neural network model (see ‘CNN model’ below).

2.2. Subjects

35 naïve subjects (19 for Experiment 1, and 16 different subjects for Experiment 2) between 18 and 45 years old with normal or corrected to normal vision participated in these experiments. Our experimental protocol was approved by the MIT Committee for the Use of Humans as Experimental Subjects. Three subjects were excluded from Experiment 1 based on a pre-defined behavioral exclusion criteria (<80% accuracy on behavioral task). Thus 16 subjects were included in each experiment. This sample size was based on prior MEG image decoding studies.

2.3. Experimental procedure

2.3.1. Experiment 1

In a first experiment, we asked whether and when social interaction information was spontaneously extracted by the brain. Subjects viewed the 60 images conveying different visually matched social and non-social scenes presented 30 times each in the MEG while they performed an orthogonal task. Subjects were instructed to fixate centrally and judge if the two people in each image were the same or different genders. The order of the 60 images was randomized within each block. The images were presented at 9 × 5 degrees of visual angle for 500 ms each with a central fixation cross. Each image was immediately followed by task instructions, and subjects responded yes or no. After each question a 200 ms fixation cross would appear before the next image. Task responses were self-paced, and the button order flipped halfway through the experiment to avoid motor confounds.

2.3.2. Experiment 2

In a second experiment, we sought to understand the role of task demands on the speed of social interaction information in the brain. The procedure was exactly the same as Experiment 1 except for three modifications. First, subjects viewed only 48 of the 60 images, excluding
“watch” condition. Second, to mitigate the effect of eye movements, images were presented smaller (5 × 2.8 degrees of visual angle) and for a shorter duration of 200 ms. While this does not remove the possibility of differential looking patterns between the conditions, it drastically reduces the need and time for subjects to make a saccade during each trial. Third, subjects performed an explicit social interaction task (“Are these two people engaged in a social interaction?”).

2.4. Eye tracking

We tracked the subjects’ left and right eye positions using an Eyelink 1000 eye tracker with a 9-point calibration. We were not able to achieve an accurate calibration for 4 subjects in Experiment 1 and 3 subjects in Experiment 2, so these subjects’ eye position data were excluded from the eye tracking analysis. To test whether there is stimulus-selective information was present in our eye tracking data, we performed the below
decoding procedures using the X,Y output of each eye as classifier features (see Decoding Methods for more details).

2.5. MEG acquisition and pre-processing

The MEG data were collected using an Elekta Neuromag Triux scanner with 306 sensors, 102 magnetometers and 204 planar gradiometers, with an online bandpass filter between 0.01 and 330 Hz. Subjects head position was continuously monitored throughout the experiment using five head position index (HPI) coils. First the signals were filtered using temporal Signal Separation and motion corrected (based on the position of the HPI coils) with Elekta Neuromag software. Next, Signal Space Projection (Tesche et al., 1995) was applied to correct for movement and sensor contamination. The MEG data were divided into epochs from −200 to 1000 ms, relative to video onset, with the mean baseline activity removed from each epoch. The signals were band-pass filtered from 0.1 to 100 Hz to remove external and irrelevant biological noise (Acunzo et al., 2012; Rousselet, 2012). The above preprocessing steps were all implemented using the Brainstorm software (Tadel et al., 2011).

2.6. MEG decoding

We analyzed the MEG data using the neural decoding toolbox for Matlab (Meyers, 2013). We averaged the data in each sensor into 10 ms non-overlapping bins, and trained and tested a new linear correlation coefficient classifier at each time point. We evaluated classification performance on independent, hold-out test data using 5-fold cross validation (CV) splits (training on 80% of the data and testing on the held out 20%) (Kriegeskorte et al., 2009). We performed feature selection using an ANOVA on only the training data (to avoid double dipping/ circularity) and selected the 25 sensors whose activity most significantly co-varied with the training labels. These selected sensors were fixed for testing. We repeated the entire decoding procedure at each time point 20 times and report the mean accuracy for each condition. See (Isik et al., 2018, 2014) for a more detailed description of the decoding methods and for more details on the decoding parameter selection.

The variables we decoded were:

1. 60-way image identity. Each image was repeated 30 times, and we divide the data into five cross validation splits with 6 trials per CV split. To increase signal to noise, we averaged the data from all 6 trials together.
2. Social interaction (mutual gaze and joint attention) vs. independent action images (2 non-interacting conditions per scenario).
3. Joint attention vs. mutual gaze.
4. Social interaction (mutual gaze) vs. watch.
5. Non-interacting images vs. watch.

In two additional exploratory analyses, we also decoded:

6. Mutual gaze images vs. non-interacting images.
7. Joint attention images vs. non-interacting images.

For tests 2–7, we ran the decoding in a manner that generalized across scenario. In particular, we trained our classifier on 10 scenarios and tested on the remaining two, held-out scenarios. For the generalization decoding, we averaged 30 trials together (note that in our pre-registration we stated we would average 24 trials together. This was an error as 24 is not divisible by the number of trials included in conditions 3–5 so would require us to exclude data from the decoding).

2.7. Statistical inference

We assessed decoding significance using non-parametric statistical tests that do not make assumptions about the underlying distribution of the data and accounts for its underlying noise structure (Pantazis et al., 2005). Specifically, we performed a sign permutation test that centers each subjects’ MEG data around chance and randomly multiplies it by +1 or −1. We repeated this procedure 1000 times to generate a null distribution. To correct for multiple comparisons, we used cluster correction in time with a cluster defining threshold of p < 0.05 and a corrected significance level of p < 0.05 (Cichy et al., 2016b; Mohsenzadeh et al., 2018).

2.8. CNN model

We ran our stimuli through a pre-trained feedforward deep neural network: VGG-16 trained on Imagenet (Simonyan and Zisserman, 2014). We asked if the output of each of the models’ five pooling layers could distinguish between images with vs. without a social interaction. First, to reduce the dimensionality of each layers’ output we performed PCA and selected the top 50 components from each layer (note our pilot data showed very similar results with 40–59 components). Within each layer, we then took the response to each image and, as with our MEG data, trained a linear classifier to distinguish between scenes with vs. without a social interaction on data from 10 of 12 scenes. We tested the linear classifier on data from the two held-out scenes. We repeated this procedure 20 times, holding out two random scenes each time.

2.9. Representational similarity analysis

We compared our MEG data to our behavioral data in Experiment 2 using representational similarity analysis. Extracting a simple MEG measure to correlate with behavioral data is not straightforward, and thus we used pairwise relationships. This representational similarity analysis (RSA) framework is a popular approach that has become standard for comparing neuroimaging and behavioral data that are in fundamentally different spaces (Kriegeskorte et al., 2008). To produce the MEG dissimilarity matrix, we followed a similar procedure to (Cichy et al., 2014). We first performed PCA on the MEG responses to the 60 images in our data to reduce the dimensionality of the data the number of components that explains 99.99% of original variance in data. We next calculated the dissimilarity between each pair of images based on their pairwise classification accuracy computed over those PCs. We repeated this at each time point to get a new dissimilarity matrix.

To produce the behavioral dissimilarity matrix, we used a behavioral metric that took into account both the subjects’ reaction time and response. Simply relying on their response would reduce behavior to a binary variable, giving us little dynamic range to detect correlations with the MEG data. Using just reaction time would not separate a very fast response for social interactions from a very fast response for independent actions (which should be represented as maximally different). We thus calculated a metric that scaled subjects’ responses by their reaction time: Response*(1-RT/max(RT)), where RT is reaction time, and the response is +1 for social interactions and −1 for independent actions. This scale places a very fast social interaction response and a very fast independent action response on opposite ends of its scale (responses ranged from +1 for the fastest social interaction responses to −1 for the fastest non-social interaction responses). Slower responses are represented between these two extremes as they are likely more behaviorally ambiguous (i.e., a very slow response for social interaction is considered to be a weaker behavioral response, and thus represented lower on our scale, than a very fast response for social interaction). For each subject, we calculated the average pairwise difference between each image pair to construct our behavioral dissimilarity matrix.

3. Results

3.1. Experiment 1

3.1.1. Late, spontaneous readout of social interactions

To identify MEG signals that contain information about the presence
of social interactions, sixteen naive subjects viewed visually matched images of different actor pairs in one of five different social or non-social conditions shot in one of 12 different scenes (60 total images, Fig. 1A–C). The five conditions were: 1) joint attention (two actors looking at the same object, a classic form of social interaction), 2) mutual gaze (two actors looking at each other, a different form of social interaction), 3) independent action 1 (two actors engaged in separate independent actions, i.e., no social interaction), 4) independent action 2 (a different instance of the two engaged in separate independent actions, no social interaction), 5) watch (one actor watching the other who is looking away, a one-way interaction or “perceptual access”). We defined social interactions broadly to include either joint attention or mutual gaze. The different conditions were well matched in terms of visual information, and a standard deep neural network model could not distinguish between scenes with versus without a social interaction in a manner that generalized across scenes (Fig. S1). In a post-MEG behavioral experiment, subjects rated the joint attention and mutual gaze images as significantly more social than the independent action images (Fig. 1D, p = 1.9 × 10^-12, two-sided t-test) and also slightly more interesting (Fig. 1E, p = 0.003). Mutual gaze images were rated as slightly more social than joint attention images (p = 0.017), but there was no difference in their interest rating (p = 0.41).

During MEG recording, each subject viewed each of the 60 images 30 times, randomized within block, while performing an orthogonal task. In particular, subjects were asked if the two actors were of the same or different gender. This task was balanced across actor pairs and scenes and the presence versus absence of social interactions. First, to replicate prior visual decoding studies and to ensure data quality, we asked whether we could decode the 60 individual images based on subjects’ MEG signals. These images included different scenes and actors, and hence differ in many visual properties. We trained a linear classifier on the response at each 10 ms time bin on 80% of the trials, and tested it on the remaining 20%. We found that we could significantly decode which image subjects viewed beginning at 60 ms after image onset (Fig. 2A). This time course of image decoding replicates several prior MEG decoding studies (Carlson et al., 2013b; Cichy et al., 2014; Isik et al., 2014), and presumably reflects primarily early visual processing. We next trained and tested a classifier at each training timepoint and each testing timepoint to generate a matrix of decoding accuracies across all train and test time points (King and Dehaene, 2014; Fig. S2A). Decoding accuracy was highest on the diagonal, when the classifier was trained and tested at the same time point, and was only significant during a narrow time window around the diagonal. This finding suggests that the neural signals are highly dynamic, in line with previously reported results of visual decoding (Cichy et al., 2014; Isik et al., 2014; Zhang et al., 2011).

We next asked the central question of this experiment: when do MEG signals encode information about whether a scene contains a social interaction (joint attention and mutual gaze conditions vs. independent action conditions)? To obtain abstract representations of social interactions, invariant to visual scene and actor information, we trained our classifier on data from subjects viewing 10 of the 12 scenarios and tested on the two held-out scenarios. This is a strong test of generalization, as the images within each scenario are much more visually similar to each other than they are to the other images in the dataset. We found that we could indeed read out the presence versus absence of a social interaction invariant to scene (Fig. 2B). This readout occurred relatively late however, beginning at 310 ms after stimulus onset. Applying the same temporal generalization approach as before, we found again a primarily diagonal decoding pattern, indicating transient social interaction representations (Figure S2B). These results suggest that humans spontaneously form abstract representations of social interactions, but this occurs later than the time scale of primarily feedforward processes as in the case of invariant object recognition.

3.1.2. Social interaction decoding cannot be explained by visual interest or eye movements

We next asked if other experimental factors could account for this social interaction decoding. First, subjects rated the social interaction images as slightly more visually interesting than the non-interacting images, but this was not uniformly true across image pairs. In an exploratory analysis (not included in our pre-registration), we took the half of the image pairs with the smallest difference in interest ratings between the social and non-social images. We found that although there was no longer a significant difference in subjects behavioral interest ratings (mean rating 2.9 ± 0.44 and 2.8 ± 0.35, p = 0.29), we could still decode scenes with vs. without a social interaction (Figure S3). Overall the times course of decoding looked very similar to that for all images (though the onset of significant decoding did not occur until 400 ms, likely due to lower power). Thus, differences in generic attention or interest are unlikely to account for our ability to decode the presence of social interactions.

We next asked if subjects’ eye movements varied systematically across interacting and non-interacting images. To do this, we followed the same decoding procedure as with the MEG data, but instead used subjects x,y eye-position as input to our classifier. We found that we could indeed decode the presence of social interactions based on
subjects’ eye position (Figure S4A). To address this alternative account of our findings, in a second experiment (see below) we presented the images at a smaller visual angle and for a shorter duration. While we were still able to decode scenes with vs. without a social interaction based on subjects’ MEG data, we could no longer do so based on eye position (Figure S4B).

3.1.3. Distinct representation of two-way social interactions

We next asked what information is driving our ability to decode social interactions. Is it sufficient for one agent to be aware of the other (“perceptual access”), or is an actual two-way social interaction necessary? To answer this question, we asked if we could decode one-way “watch” images (in which one agent sees the other but not vice versa) from two-way “mutual gaze” images. We found that we could decode watch versus mutual gaze at similar latency to social interaction read out (Fig. 3a, onset 330 ms), although this result was statistically less robust than decoding of the presence of a social interaction (only reaching significance for two brief time periods). Note that this decoding is based on half as much data as the social interaction detection analysis above, so the read out is necessarily noisier. Interestingly, we could not decode watch images from independent action images (Fig. 3b). These findings suggest that it is the presence versus absence of a two-way social interaction that is a distinct and spontaneously represented property of the image, not the mere presence of one-way perceptual access from one agent to the other.

3.1.4. Late readout of type of third-party social interaction

Finally, beyond simply detecting the presence of a social interaction, we asked whether the MEG signal contained information about the different types of social interactions in our dataset: joint attention vs. mutual gaze. Mutual gaze is perhaps the most perceptually obvious form of social interaction between two agents. But joint attention is also a fundamental form of social interaction that arises early in infancy (Scaife and Bruner, 1975) and may be critical in language learning (Tomasello and Farrar, 1986). Do perceivers spontaneously distinguish between these two forms of social interaction, and if so, when? We found that we could distinguish joint attention vs. mutual gaze in a manner that generalized across scenes, but only quite late, at 600 ms after image onset (Fig. 3c). This analysis makes use of only half the data as the analysis of social interaction detection and thus could fail to detect earlier discriminative information (but see Experiment 2/Fig. 6 for a replication of these results).

In sum, the results of Experiment 1 suggest that both the presence and type of social interactions are spontaneously represented in the brain, but this information comes online very late, well beyond the timescale of primarily feedforward processes.

3.2. Experiment 2

3.2.1. Social interaction perception is slow even during an explicit task

In a second experiment we asked if there were any conditions under which social interaction information could be extracted more quickly. In particular, can these neural computations occur faster if the subject is explicitly asked to behaviorally extract that information? This would be consistent with prior studies showing rapid visual readout that also used an explicit task (Thorpe et al., 1996). To test this hypothesis, we ran a second experiment with 16 additional naïve subjects who saw identical images in the MEG, but now instead of performing the same versus different gender task they performed an explicit social interaction task (i.e., does this image contain a social interaction?). We removed the “Watch” stimuli from this experiment because they are ambiguous in terms of whether they should or should not count as a social interaction, based on subject ratings in Experiment 1 (Fig. 1D). Further, since the latency for differentiating between social interactions and watch was the same as social interactions and independent actions (the main question of interest), we decided to leave this condition out to shorten experimental

Fig. 3. Decoding different types of social interactions from MEG signals in Experiment 1: Time series of (A) watch vs. social interaction (onset = 330 ms), (B) watch vs. independent actions (not significant), (C) mutual gaze vs. joint attention (onset = 600 ms). Error bars and statistical tests are the same as in Fig. 2.
time. The button order was flipped halfway through the study so that explicit motor responses could not account for our decoding results (see Methods). Subjects’ mean reaction time on the social interaction task was 1.0±0.25 s (mean±SD across subjects). As expected, we found no difference in the onset latency of image identity decoding (Fig. 4A).

Importantly, we found that even when subjects performed an explicit social interaction detection task, information about the presence vs. absence of a social interaction could again only be discriminated in MEG response at relatively late latencies after stimulus onset (290 ms, Fig. 4B). Could earlier social interaction signals exist that we did not have sufficient power to see in either experiment? To answer this question, in an exploratory analysis, we combined the data from Experiments 1 and 2 and re-ran our social interaction decoding analysis. Even with 32 subjects, the onset latency of social interaction detection did not change (Fig. 4C).

Next, we again asked if and when we could decode the type of social interaction (mutual gaze vs. joint attention) in Experiment 2, as we could in Experiment 1. While the latency moved slightly earlier, it was still quite late, with an onset of 530 ms after stimulus onset (Fig. 5A). In another exploratory analysis, we combined data across Experiments 1 and 2 and found that in all 32 subjects onset latency was 490 ms (Fig. 5B). The results of Experiment 2 serve as an internal replication, and confirm that even in the presence of task demands, social interaction perception is computed well beyond the timescale of visual pattern recognition.

Given the fact that mutual gaze images were rated as significantly more social than joint attention images (Fig. 1D), in an exploratory analysis we asked if there was any difference in each of their decoding timelines versus non-interacting images. We found that the two time courses looked very similar to each other and to our initial analysis where the two conditions were collapsed. In addition, the onset latency for both types of decoding was the same (320 ms) suggesting that social interaction detection occurs at this latency, no matter the type of social interaction depicted.

3.2.2. MEG-behavioral correlation

As with all decoding studies, it is important to investigate whether the neural information we extract is associated with perceptual judgements, or whether it is merely epiphenomenal (Grootswagers et al., 2018; Williams et al., 2007). One way to address this concern is to test whether readout performance is tied to subjects’ behavioral judgments and reaction times. Because we had these measures in our second experiment, we correlated each subject’s MEG and behavioral data using representational similarity analysis (RSA). Specifically, we computed the time-resolved dissimilarity matrix for our MEG data (48 × 48 pairwise image decoding accuracy) and a behavioral dissimilarity matrix (48 × 48 behavioral dissimilarity matrix), and correlated these measures within subject. We found that there was indeed a significant correlation between subjects’ behavioral responses and MEG data beginning 340 ms after stimulus onset (Fig. 6). These results suggest that the MEG signals we detected are behaviorally relevant to social interaction perception.

4. Discussion

Here we identified neural signals that contain information about the presence and type of third-party social interactions in a visual scene. These neural representations generalized across low-level visual features, and arose spontaneously, even when participants performed an orthogonal task. Crucially though, they arose relatively late compared to previously reported latencies for other types of visual pattern classification: 300 ms after stimulus onset for detection and 600 ms for categorization. These late latencies were found even when subjects performed an explicit social interaction detection task. Importantly, this neural readout was correlated with behavior, as measured through RSA. It is of course difficult to rule out earlier social interaction signals that we could not detect with MEG or with the size of our current sample, and there is no straightforward way to link multivariate decoding accuracies with standard effect size measures (Hebart and Baker, 2018). However, converging evidence from our two experiments suggest that the perception of social interactions in naturalistic real-world images is not based on purely feed-forward processing, but instead relies on slower,
presumably recurrent computations.

The latencies of social interaction detection and categorization (300 and 600 ms, respectively) are substantially later than those previously reported for different types of pattern recognition problems, such as invariant object recognition (Carlson et al., 2013a; Isik et al., 2014; Yamins et al., 2014). Prior work with ERPs (Thorpe et al., 1996) and physiology (Yamins et al., 2014) indicates that object recognition is quite fast, occurring between 100 and 200 ms after stimulus onset, even when the objects appear in complex backgrounds, like the natural images used in our study. Natural scene information has been shown to arise on a similarly fast time scale to object recognition (Cichy et al., 2016a; Greene and Hansen, 2018). While many psychological studies of social interaction detection have used minimalist displays (Papeo et al., 2017; Su et al., 2016; Vestner et al., 2019), it can be difficult to unconfound the presence of a social interaction from low-level visual properties in these stimuli. In a related study, we found decoding of facing (interacting) versus non-facing (independent) dyads occurred relatively quickly, however, we found the same effect when these stimuli were inverted. Further, we found that low-level image properties (pixels and early layers of a DNN) could distinguish between facing and non-facing dyads, even in a manner that generalized across agents and images (Isik et al., in prep).

It is worth noting that while the present study used static stimuli to aid in time locking, real-world social interactions are highly dynamic. Since much social information is conveyed via dynamic information, it is possible the use of static stimuli affected our onset latency. On the other hand, information conveyed in successive frames of a video would necessarily take longer to transmit and thus it seems unlikely that this would speed up the onset latencies identified above. Further, Experiment 2 showed that subjects were extremely good at detecting social interactions behaviorally even from brief presentations of static stimuli. The role of dynamics however remains an important open area to consider in future studies of social interaction perception.

In contrast to object and scene perception, there has been relatively little M/EEG decoding work on aspects of social perception. The N170 response (Bentin et al., 1996) is a strongly face-selective univariate response arising around 170 ms after image onset. However, recent decoding studies have shown that many aspects of face information are represented earlier than 170 ms. For example, age, gender and identity are all decodable around 100 ms (Dobs et al., 2018). Even emotion properties like expression (100 ms (Dima et al., 2018)) and valence and arousal (150 ms (Grootswagers et al., 2017)) have been shown to come online quickly. Beyond face properties, the emotional valence and self-relevance of communicative gestures can be decoded within 100 ms (Redcay and Carlson, 2015), and individual agents’ actions as early as 200 ms (Isik et al., 2018). Interestingly, like object and scene perception, these social dimensions all fall within the rough timescale of feedforward pattern recognition. The present study suggests that the detection of third-party social interactions occurs substantially later, and thus may be based on fundamentally different computations from these other visual and social recognition processes. A critical difference may be that, unlike face, emotion, action, and gesture recognition of individuals, social interaction recognition involves taking into account relational information between multiple agents.

Prior work has used ERPs to understand the dynamics of distinguishing different types of social interactions – intentional vs. unintentional harm in adults (Decety and Cacioppo, 2012; Hesse et al., 2016) and harm vs. help in infants (Jason M Cowell and Decety, 2015a, b) and children (Jason M Cowell and Decety, 2015a, b). These categorizations arise quite early, as early as 60 ms in adults and 100 ms in children, but also appear to have later components that are more behaviorally relevant in children, and line up with the timing of social interaction detection and categorization found in this study (Jason M Cowell and Decety, 2015a, b). In this study we do not see any early signals related to the different social interactions we tested. One possible explanation of this
difference is that our stimuli, unlike those previously used, are largely neutral in valence. It is possible that highly valenced stimuli are routed faster to relevant brain regions (although it should be noted that the above intentionality signals cannot be explained by valence). A comprehensive comparison of social interactions depicting difference valence and intentionality will be necessary to more thoroughly understand the neural dynamics of social interaction perception.

Typically, visual perception problems, such as object recognition, are broken down into two main components: detection and categorization (de la Rosa et al., 2011; Grill-Spector and Kanwisher, 2005; Marr and Nishihara, 1978), and we examined both those components here. In the rich space of social perception however, there are many other aspects to perception and understanding. Even social interaction categorization includes many more complex types of interactions and dimensions (such as valence, mentioned above) that we did not address here. Indeed, this study takes just a first step toward understanding the time course and computations underlying multi-agent social interaction perception. The simplified stimuli and paradigm represent only a small part of real-world social interaction perception, and many open questions remain. Beyond detecting simple, dyadic social interactions and distinguishing between different gaze events, what other social interaction information is spontaneously extracted by the brain? How do our brains code complex real-world complex social events such as a party or sporting event? And how fine-grained are the automatically-extracted representations of social interactions (positive vs. negative, different action categories, etc.)?

And perhaps most obviously, where in the brain do these social interaction representations originate? Our prior fMRI results indicate that a region in the right posterior superior temporal sulcus is selectively involved in the perception of social interactions (Isik et al., 2017). If this region does indeed underlie the decoding information we report here with MEG, does it receive input from purely from visual regions, or from higher level regions that code for information about individual social agents (Grossman et al., 2000; Puce et al., 1998; Saxe and Kanwisher, 2003) or the physical world around them (Fischer et al., 2016)? While detecting social stimuli and paradigm represent only a small part of real-world social interaction perception and many open questions remain. Beyond detecting simple, dyadic social interactions and distinguishing between different gaze events, what other social interaction information is spontaneously extracted by the brain? How do our brains code complex real-world complex social events such as a party or sporting event? And how fine-grained are the automatically-extracted representations of social interactions (positive vs. negative, different action categories, etc.)?

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