

Understanding particularized and generalized conversational implicatures: Is theory-of-mind necessary?

Wangshu Feng ^{a,b}, Hongbo Yu ^c, Xiaolin Zhou ^{b,d,e,f,*}

^a Research Institute of Foreign Languages, Beijing Foreign Studies University, Beijing 100089, China

^b School of Psychological and Cognitive Sciences, Peking University, Beijing 100871, China

^c Department of Psychological and Brain Sciences, University of California Santa Barbara, Santa Barbara, CA 93106-9660, USA

^d Institute of Linguistics, Shanghai International Studies University, Shanghai 200083, China

^e Key Laboratory of Behavior and Mental Health, Peking University, Beijing 100871, China

^f PKU-IDG/McGovern Institute for Brain Research, Peking University, Beijing 100871, China



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ABSTRACT

A speaker's intended meaning can be inferred from an utterance with or without reference to its context for particularized implicature (PI) and/or generalized implicature (GI). Although previous studies have separately revealed the neural correlates of PI and GI comprehension, it remains controversial whether they share theory-of-mind (ToM) related inferential processes. Here we address this issue using functional MRI (fMRI) and transcranial direct current stimulation (tDCS). Participants listened to single-turn dialogues where the reply was indirect with either PI or GI or was direct for control conditions (i.e., PIC and GIC). Results showed that PI and GI comprehension shared the multivariate fMRI patterns of language processing; in contrast, the ToM-related pattern was only elicited by PI comprehension, either at the whole-brain level or within dorsal medial pre-frontal cortex (dmPFC). Moreover, stimulating right TPJ exclusively affected PI comprehension. These findings suggest that understanding PI, but not GI, requires ToM-related inferential processes.

1. Introduction

Imagine that Pat asks the hotel's front-desk clerk about where his friend went. The clerk responds by saying: "Some of guests are already leaving". In this conversation, the listener needs not only to decode the context-invariant "sentence meaning", but also to infer the implicated meaning (conversational implicature) beyond the literal expression (Grice, 1989; Hagoort & Levinson, 2014; Noveck & Reboul, 2008), which can be further classified into particularized conversational implicature (PI) and generalized conversational implicature (GI) (Grice, 1975). Here the utterance can convey both a GI, which is normally carried by the usage of a certain linguistic expression (e.g., *some of*) in the utterance and can be achieved without referring to the context of utterance, and a PI, which is intimately associated with the specific context of the utterance. Specifically, the clerk's use of the term "*some of*" warrants a GI: *Some but not all of* guests are already leaving, because the clerk used a weak scalar term "*some of*" on a scale, instead of a stronger one (e.g., *all*). Thus, GI is independent of the particulars of its context. In contrast, in the above example, the indirect reply may convey

a PI, "perhaps your friend has already left". Yet, if Pat is asking for the time, the same utterance can be interpreted as "it must be late".

In linguistic pragmatics, it is an ongoing debate, with three most influential theories, about whether interpreting GI and PI involve distinct or identical cognitive processes. Default Theory focuses on the important feature that GI is carried by the usage of certain sub-sentential locutions or structures of utterances, instead of by the particulars of the context of utterance (Chierchia, 2004; Horn, 2004; Levinson, 2000). Thus, according to this account, GI is computed by an automatic and effortless system that supports default inferences, whereas PI is computed by a separate one that supports context-sensitive inferences. In contrast, Relevance Theory holds that both types of implicature are recovered in comprehension by a single pragmatic system (Carston, 2004; Sperber & Wilson, 1986). According to this theory, understanding the speaker's meaning of an utterance is a process of searching for an optimally relevant interpretation under the particular context of the utterance, with the constraint of spending as little processing effort as possible. Once the interlocutor gets an interpretation crossing the relevance threshold, he or she will take it as what the speaker wanted to

* Corresponding author at: Institute of Linguistics, Shanghai International Studies University, Shanghai 200083, China.

E-mail address: xz104@pku.edu.cn (X. Zhou).

convey (Sperber & Wilson, 1986). That is to say, both PI and GI are derived from the same cognitive processes, which take contextual considerations into account from the beginning. Finally, Semantic Minimalism offers a more balanced view, which draws a distinction between semantic and pragmatic processing according to whether the recovery of the content can rely on computational operations alone (Borg, 2004; Cappelen & Lepore, 2005). Although both types of implicature require information beyond the strictly semantic information at hand, GI does not constitute full pragmatic content like PI, since it can be generated only by knowing the fact that the word “some” usually contains the meaning of “some but not all” in daily conversations. Thus, GI comprehension does not recruit the holistic, general pragmatic system that is recruited to generate PI, but involves a more limited system which runs on the basis of statistical facts about what speakers have communicated in past experience (Borg, 2009). In other words, PI is derived from fully context-based inferences, while GI is derived from constraint-based inferences.

Comparing the neurocognitive mechanisms underlying PI and GI comprehension would allow us to choose between these theoretical approaches. Prior neuroimaging studies have separately investigated the neural processes of comprehending PI and GI. On the one hand, studies adopting a reading or listening comprehension task showed that the neural substrates of PI comprehension can be divided into two subsystems (Hagoort & Levinson, 2014; Hagoort, 2013): a core language network responsible for filling in the semantic gap between the literal meaning of an utterance and its context (Ferstl & von Cramon, 2001; Siebörger, Ferstl, & von Cramon, 2007), and a theory-of-mind (ToM) network, which is commonly invoked by processes of inferring mental states of other individuals. The ToM network typically consists of medial prefrontal cortex (mPFC), bilateral TPJ, precuneus, and bilateral anterior superior temporal sulcus (Mar 2011; Van Overwalle & Baetens, 2009), and among these regions, dorsal mPFC (dmPFC) and right TPJ are likely to be the core regions supporting ToM processes (Schurz, Radua, Aichhorn, Richlan, & Perner, 2014). Studies on indirect reply used natural conversations as stimulus materials, in which the reply utterance was served as a direct or indirect reply to its preceding question (Bašnáková, Weber, Petersson, van Berkum, & Hagoort, 2014; Feng et al., 2017; Jang et al., 2013; Shibata, Abe, Itoh, Shimada, & Ueda, 2011; Tettamanti et al., 2017; van Ackeren, Smaragdi, & Rue-schemeyer, 2016). By comparing indirect reply to direct reply, these studies identified a set of brain regions that are linked to PI comprehension, including left (and right) inferior frontal gyrus (IFG), right (and left) middle temporal gyrus (MTG), mPFC, right (and left) TPJ, and precuneus.

On the other hand, neuroimaging studies of GI adopted a picture-sentence verification paradigm (Shetreet, Chierchia, & Gaab, 2014; Zhan, Jiang, Politzer-Ahles, & Zhou, 2017), comparing experimental conditions of mismatched generalized implicature (e.g., *some rabbits have keys*, following a cartoon in which all rabbits have keys), matched generalized implicature (e.g., *some rabbits have keys*, following a cartoon in which two of five rabbits have keys), no-implicature control (e.g., *every rabbit has keys*, following a cartoon in which all rabbits have keys). Participants were presented a cartoon and a sentence, and were required to decide if the sentence matched the picture. Shetreet et al. (2014) found that the mismatched and matched GI conditions commonly activated left IFG relative to a control condition. By comparing the mismatch and match GI conditions, they further found that GI mismatch activated additionally mPFC/anterior cingulate cortex and left middle frontal gyrus (MFG). The authors speculated that GI processing is possibly associated with semantic processing (IFG) and high-order cognitive functions (mPFC), like conflict control or ToM. Using similar constructions, Zhan and colleagues (2017) found that both GI mismatch and semantic mismatch activated bilateral ventral IFG, whereas GI mismatch uniquely activated left dorsal IFG and basal ganglia, relative to semantic mismatch. The activation in basal ganglia, together with IFG, suggests that the processing of GI mismatch may involve executive

functions beyond semantic unification.

Although these two lines of research on PI and GI have made remarkable achievements, we still have little direct evidence for the relationship between PI and GI processing for the following reasons. First, for the studies on GI, the picture-sentence verification paradigm provides a temporary linguistic context in which a GI recovered from the sentence is inconsistent with its paired picture (in the mismatch condition). Due to the fact that implicatures could be potentially cancelled by linguistic or extra-linguistic cues (Eckardt, 2007; Grice, 1975), it is difficult to know to what extent the neurocognitive mechanism revealed in this paradigm truly reflects the system underlying GI comprehension in normal conversations. Second, these two lines of research used different experimental paradigms and different participants, making it difficult to compare the findings across PI and GI processing. Third, previous neuroimaging research used mostly univariate data analysis approaches and showed common activations in IFG and mPFC for PI and GI processing. However, such overlapping brain activity does not necessarily imply shared neural representations and cognitive processes. Thus, to what extent comprehension of PI and GI share the same neurocognitive processes is still an open question.

Here we aim to identify both the shared and distinct neurocognitive processes underlying PI and GI comprehension by comparing these two types of conversational implicature in the same experiment. In particular, we aim to investigate whether ToM processing is necessary for interpreting both PI and GI. To this end, we adopted a listening comprehension task in which participants were required to interpret the speaker’s meaning of an utterance that warrants either a PI or a GI. We also asked the same participants to perform a ToM task. We would first identify brain regions and neural representations associated with PI and GI processing by conducting univariate analysis and multivariate pattern analysis (MVPA). Specifically, we would train PI, GI, and ToM classifiers to examine to what extent PI, GI and ToM share neural representations with each other. Then we tested the potential causal relationship between the neural activity of a ToM-related region (right TPJ) and PI and GI comprehension by using high definition transcranial direct current stimulation (HD-tDCS) over right TPJ. Overall, our design and methodology allow a direct comparison between PI and GI comprehension in the same experimental setup and a direct examination of whether ToM is part of the neurocognitive mechanisms underlying implicature understanding.

In the listening comprehension task, we presented participants with single-turn dialogue scenes, each of which contained a cover story, a yes/no question, and a reply. For the critical conditions, the reply served as an indirect answer to the question; understanding its meaning either relied on knowledge of the context in PI condition, or did not in GI condition. For their respective controls (i.e., PIC and GIC), essentially the same sentence was used, but as a direct reply to the preceding question (see Table 1 for examples). Given that intentionally interpreting speaker’s meaning of an utterance is a prerequisite for generating conversational implicature (Bach, 2006), we asked participants to make judgment as to what the reply utterance intended to convey. In addition, participants were asked to perform a false belief task (Dodell-Feder, Koster-Hale, Bedny, & Saxe, 2011). We chose the false belief task to assess ToM for reasons that it is the most classic task to reflect the development of ToM in children and it is also one of the most frequently used task to reflect the neural correlates of ToM.

Grounding on previous research, we predicted that PI processing would recruit the core language network, consisting of bilateral IFG and MTG, and ToM network, consisting of mPFC, precuneus and bilateral TPJ, while GI processing would also involve activations in IFG and mPFC. Importantly, the data would allow us to test the three possible predictions for the relationship between PI and GI processing. Default Theory predicts that PI and GI processing involve separated neural representations, thus we could identify two different neural patterns: one pattern responds to PI generation but not GI, and the other responds to GI generation but not PI. Relevance Theory predicts that PI and GI

Table 1

Examples of the dialogue scenarios in the four experimental conditions, translated into English.

Condition	Cover Story	Dialogue
PI	In a movie city, a director is going to finish off the shoot of her first literary film. The following is the dialogue between the director and her friend.	<p>Q: Will my film be successful at the box office? 我的电影会收获高票房吗?</p> <p>A: It is hard for audiences to really enjoy a literary film. 观众们很难真正欣赏文艺片。</p>
PIC		<p>Q: Do audiences like literary films? 观众们会喜欢文艺片吗?</p> <p>A: It is hard for audiences to really enjoy a literary film. 观众们很难真正欣赏文艺片。</p>
GI	After completing his performance, the supporting actor is removing makeup in the backstage of the theater. The following is the dialogue between the actor and the director.	<p>Q: Did everyone like our performance? 每个人都喜欢我的表演吗?</p> <p>A: Some of the audiences enjoyed your performance. 有的观众欣赏你的表演。</p>
GIC		<p>Q: Did everyone like our performance? 每个人都喜欢我的表演吗?</p> <p>A: Not all of the audiences enjoyed your performance. 不是所有观众欣赏你的表演。</p>

processing share the same or similar neural representation; thus we could hardly distinguish the fMRI multivariate patterns of PI and GI processing. Semantic Minimalism predicts that PI and GI processing share similar language processing systems, but distinguish in inferential processing. Accordingly, we could identify a neural pattern of language processing that responds to both PI and GI generation, and an inference-related pattern that specifically responds to PI processing.

2. fMRI experiment

2.1. Methods

2.1.1. Participants

Twenty-nine university students were recruited for the fMRI experiment. One participant was excluded from data analysis on the basis of binary judgment accuracy (three SDs lower than group average), leaving 28 participants for data analysis (14 females; mean age 21.5, SD = 1.9). All participants were right-handed Chinese native speakers with normal or corrected-to-normal vision. None of them suffered from neurological, psychiatric, or hearing disorders. This study was approved by the Ethics Committee of the School of Psychological and Cognitive Sciences at Peking University. Written informed consents were obtained from all the participants.

2.1.2. Design and materials

We used single-turn dialogue scenarios as stimulus materials. Each dialogue scenario was comprised of three parts - a cover story, a yes/no question, and an indirect or direct reply to the preceding question (Table 1, see *Supplementary Materials* for pretests). In the critical conditions (i.e., PI and GI), the reply was indirectly related to the question. For the control of PI condition, namely PIC, we used the same sentence as a direct reply to the preceding question. For the control of GI condition, namely GIC, we replaced the weak scalar term (e.g., *some of*) in the reply utterance of GI condition with its implicated meaning (e.g., *not all*), and thus the modified utterance served as a direct reply to the question. Various pairs of scalar items were included in GI pairs to minimize the repetition of certain lexical items, such as, “*some of* (*Youde/Youxie* in

Chinese) vs. *all* (*Suoyou/Quanbu*)”, “*sometimes* (*Youshi/YouDeShihou/Youshihou*) vs. *always* (*Zongshi/Zong*)”, “*sometimes* vs. *often* (*Jingchang/Changchang/Shichang/Chang*)”, “*occasionally* (*Ouer/Otuyou*) vs. *often*”, “*many times* vs. *all the time/everyday*”, “*may* (*Keneng/Yexu*) vs. *must* (*Yiding/Kending*)”, “*want/try* (*Xiang/Dasuan/Nuli/Changshi*) to do something vs. *succeed in doing something*”, “*strive* (*Zhengqu*) to do something vs. *promise* (*Baozheng*) to do something”, and “*a little adv.* vs. *very adv.*”. For each scenario, the question was strongly expected to receive a “yes” or a “no” answer and the reply gave a definite answer. Within each pair of scenarios, both direct and indirect replies were equivalent in giving a definite answer (“yes” or “no”) to the preceding questions. For the PI pairs, half of the replies answered “yes” to the questions while the other half answered “no”. However, for the GI pairs, all replies would give negative answers to the questions, rendering interpreting the scalar implicature of a weak term (i.e., the stronger term is not true) necessary for understanding the speaker’s meaning of the reply. For example, in Table 1, the utterance “*Some of the audiences enjoyed your performance*” triggered a “no” answer to the question “*Did everyone like our performance*”. In this case, to understand the reply, listeners need to know that the usage of “*some of*” warrants a GI “*some but not all*”. But, in the case that the same utterance gives a “yes” answer to the question “*Did anyone like our performance*”, it is unnecessary for listeners to notice that “*some of*” has the meaning of “*not all*”.

Apart from the scenarios in the four conditions, we created filler scenarios, which were similar to the critical scenarios in form and content. For each filler scenario, the question included a stronger term. Among these filler scenarios, 20 replies with strong terms were direct answers to the preceding questions, while the other 20 replies with weak terms were indirect. We added these fillers to balance the yes/no judgment of the scenarios, and to balance the yes/no response to replies with strong/weak terms (“*all*”, “*some*”, “*always*”, “*sometimes*” etc.), which made the materials more diversified and prevented the participants from formulating a certain response strategy.

To simulate natural conversations in daily life, all parts of dialogue scenarios were presented aurally. Fourteen Chinese native speakers were recruited to record specified parts of materials. One female and one male speaker were responsible for recording the cover stories, while six other female and six other male speakers recorded the single turn dialogues. For a particular scenario, the dialogue always occurred between a female and a male speaker. Each auditory stimulus was recorded in a sound-proof booth with a microphone (RODE NT1-A), digitized at 11.0 kHz sampling rate in a 16-bit format, and equated for the maximum sound intensity.

2.1.3. Procedures

For fMRI scanning, participants first performed a listening comprehension task. This task was separated into two sessions, each lasting about twenty minutes. All scenarios were divided into four experimental lists based on a Latin-square design, with each list further separated into two sessions. Each list consisted of 120 scenarios, including 20 scenarios for each experimental condition (i.e., PI, PIC, GI, and GIC) and 40 fillers. Scenarios in each list were sorted pseudorandomly, such that 1) no more than three scenarios in a certain experimental condition showed up consecutively; and 2) no more than four scenarios requiring an identical response showed up consecutively. In each trial, participants experienced the following events. First, a fixation cross was presented in the middle of the screen and remained for a jittered duration ranging from 1.5 to 5.5 s, before a blank screen lasting 0.1 s. Next, participants clearly heard the cover story, the question and the reply sequentially; at the meantime, only a fixation point was shown on the screen. We set up a fixed interval of 1 s after the presentation of the cover story, as well as a jittered interval ranging from 0.5 to 1.5 s between the presentation of the question and the reply. Finally, two option characters (“yes” on the left and “no” on the right) were presented and remained on the screen for 3 s immediately after the presentation of the reply utterance. Participants had to make a forced binary judgment as accurately and

quickly as possible as to whether the latter speaker really intended to answer “yes” or “no” to the question. The judgment was indicated by a button press with the index or middle finger of the participants’ right hand. Reaction time (RT) was measured as the latency of his/her response to the presentation of “yes” and “no” choices.

After the listening comprehension task, participants also completed a ToM task in the scanner. Stimulus materials of this task were obtained from the SaxeLab website (<http://saxelab.mit.edu/localizers>; credit David Dodell-Feder, Nicholas Dufour, and Rebecca Saxe), containing 10 “false belief” and 10 control stories. We first translated these stories and its corresponding statements into Chinese. Then an English-Chinese bilingual, with English as his native language, translated the Chinese version back to English. This English translation and the original version were almost identical, indicating that the Chinese version was consistent with what the English version intended to convey. For each trial, a story was visually shown for 12 s, followed by a statement about the preceding story for 4 s. Each participant made a binary judgment as to whether the statement was True or False according to the story. A fixation interval of 12 s was presented between the trials.

Prior to fMRI scanning, all participants received written instructions concerning how to complete the tasks and performed a short practice for each task. After scanning, each participant completed a Chinese version of Autism Spectrum Quotient (AQ) questionnaire which is intended to measure individuals’ social skills (Baron-Cohen, Wheelwright, Skinner, Martin, & Clubley, 2001). The subscale scores of this questionnaire reflect the degree of autistic-like social and communication difficulties; that is, the higher the score, the poorer the social or communication skills.

2.1.4. Data acquisition and preprocessing

Functional images were gathered on a research-dedicated 3-Tesla MRI scanner (GE MR750, General Electric, Fairfield, Connecticut), with a T2*-weighted echo-planar imaging sequence. Each volume contained 35 transversal slices, with repetition time/echo time/flip angle = 2000 ms/30 ms/90°, slice thickness/inter-slice gap = 4 mm/0.75 mm, field of view = 192 × 192 mm², resolution within slice = 64 × 64, and voxel size = 3.0 × 3.0 × 4.0 mm³. Slices of each volume were acquired in an interleaved order. Head movements were minimized using pillows and cushions within the head coil.

The fMRI data preprocessing was conducted using SPM8 (Wellcome Centre of Human Neuroimaging, London; <https://www.fil.ion.ucl.ac.uk/spm/>). The first five volumes in each session were excluded from data analysis to allow the MR stabilization. Images were time sliced and realigned to the sixth volume to correct for head-motion artifacts. We used a high-pass temporal filter (cutoff period = 128 s) to remove low-frequency drifts in fMRI time series. We spatially normalized all functional images into the standard Montreal neurological institute (MNI) space by matching gray matter, white matter, and cerebrospinal fluid (Ashburner & Friston, 2005) and resampled to 3 × 3 × 3 mm³ voxel. On this basis, the normalized data was smoothed using a 6-mm full-width half-maximum Gaussian kernel. No participants’ head movements exceeded 3 mm.

2.1.5. Univariate analysis

Whole-brain analyses were conducted using the generalized linear model (GLM) of SPM8 firstly at the participant level and secondly at the group level. For each session, all regressors were constructed as a boxcar function convolved with the canonical hemodynamic response function (HRF).

For the listening comprehension task, we defined nine/ten regressors in the GLM at the participant-level to model the following events: the auditory presentation of the cover story, the question and the reply, and the participants’ response. More specifically, the reply presentation was separately modeled by six/seven regressors, corresponding to four critical conditions (i.e., PI, PIC, GI, and GIC) and two types of fillers, as well as the misunderstood replies if the participant response was incorrect. The presentation of the cover story and the question, and the

participants’ response were modeled by three regressors of no interest, respectively. Six rigid body parameters calculated from the realignment procedure were additionally included to correct for head-motion artifacts. The onset and duration of each regressor were defined as the actual onset and duration of each auditory stimulus. The simple main effect was examined in each experimental condition to identify brain regions significantly activated for each condition. For the group level analysis, a flexible factorial repeated-measures ANOVA was conducted on the participant-level contrast images of each experimental condition. At the group level, we used a cortical mask to exclude the cerebellum and conducted further analyses within this mask. We defined two contrasts for the two types of conversational implicatures, respectively, comparing the PI and GI conditions to their corresponding controls.

For the ToM task, the participant-level models were created by using a GLM with the false belief and control conditions as regressors of interest. The duration of each regressor contained the duration of the story reading (12 s) and the True/False judgment (4 s). At the group level, the two contrast maps corresponding to the two conditions from each participant were fed into a flexible factorial design. We defined one contrast comparing the false belief condition to the control.

Conjunction Analysis. To explore regions that were activated in interpreting both types of conversational implicatures, we further performed an SPM ‘conjunction null’ analysis (Nichols, Brett, Andersson, Wager, & Poline, 2005) with (PI > PIC) ∩ (GI > GIC) (Friston, Holmes, Price, Büchel, & Worsley, 1999).

Parametric Analysis. To further reveal the functions of dmPFC during the comprehension of PI and GI, we conducted group-level parametric analyses using small volume correction within a dmPFC region-of-interest (ROI) to explore whether the dmPFC activation in PI/GI processing correlated with individual differences in social skills. The dmPFC ROI was defined by the co-activation of the contrasts PI > PIC and GI > GIC in the conjunction analysis at a relatively liberal threshold of voxel-level $p < 0.01$ uncorrected (1038 voxels in total). At the group-level, we used the measure of social skills (a subscale of AQ questionnaire) as a between-participant covariate and activations in the contrasts PI > PIC and GI > GIC recorded from the participant-level analyses as dependent variables, constructing two regression models, respectively. We next defined a sphere of 6-mm radius centered on the group peak coordinates identified by the parametric analysis (MNI coordinates: [9, 32, 49]; see *Results 3.3*), and extracted the parameter estimates from this sphere in the contrast map PI > PIC and GI > GIC, respectively. Pearson correlation coefficients were computed between the scores of social skills and the dmPFC activation in the contrasts PI > PIC and GI > GIC, respectively. We then performed a statistical comparison of correlation to formally test whether the correlation coefficients were significantly different through Fisher’s Z-transform method and Zou’s confidence interval (CI) method (Zou, 2007). Both methods were performed using the cocor 1.1–3 R package (<http://comparingcorrelations.org/>; Diedenhofen & Musch, 2015).

Psychophysiological interaction (PPI) analysis. Given that dmPFC was found to be involved in generating both PI and GI (see *Results 2.2.2*), our interest lied in whether the functional interplay between dmPFC and other regions was modulated by the type of conversational implicature. For this purpose, we conducted a PPI analysis (Friston et al., 1997) with dmPFC revealed in the abovementioned conjunction analysis as the seed region, and calculated a PPI map corresponding to the contrast between PI and GI. The regression model contained three regressors and six head motion parameters. The first regressor, called physiological regressor, was the fMRI signals from a 6 mm-radius sphere centered on the group peak coordinates in the co-activated dmPFC (MNI coordinates: [-9, 38, 43]; see Table S1 in *Supplementary Materials*); the second, called psychological regressor, was the design vector (PI vs. GI); the third was calculated as the interaction between the physiological and psychological regressor.

All results were thresholded at $p < 0.001$ uncorrected at voxel-level and $q < 0.05$ family-wise error (FWE) corrected for multiple

comparisons at cluster-level (whole-brain or within the dmPFC ROI using small-volume-correction; Chen, Jimura, White, Maddox, & Poldrack, 2015).

2.1.6. Multivariate pattern analysis

To identify the distributed neural representations of PI and GI processing, we used linear support vector machines (SVMs) to train multivariate fMRI pattern classifiers for PI and GI, respectively. We implemented the SVMs using Spider toolbox (<https://people.kyb.tuebingen.mpg.de/spider>). We trained three classifiers on individual contrast maps to discriminate PI from PIC, GI from GIC, and PI from GI. For illustration purposes, we carried out bootstrap tests to assess the significance of voxel classifier weights. We performed SVMs on 10,000 bootstrap samples (with replacement). In each voxel, two-tailed, uncorrected *p*-value was computed according to the distribution of classifier weights. For the whole-brain analysis, the weight maps were thresholded at *p* < 0.001 uncorrected (cluster size >10) to illustrate clusters that contributed most reliably to the classification (c.f., Wager et al., 2013). In classification and further similarity analysis, we used all the voxels in the training data. We performed a force-choice test with a leave-one-participant-out cross-validation method (cf., Chang, Giannaros, Manuck, Krishnan, & Wager, 2015; Woo et al., 2014) to calculate the classification accuracies of the SVM classifiers for PI vs. PIC and GI vs. GIC. The classifier trained to discriminate between PI and PIC (i.e., PI classifier) and the classifier trained to discriminate between GI and GIC (i.e., GI classifier) represented the neural patterns that were modifiable by PI and GI (Woo et al., 2014; Woo, Chang, Lindquist, & Wager, 2017). On the one hand, if PI and GI comprehension shared neural representations, then the PI classifier should accurately discriminate GI from GIC, and the GI classifier should accurately discriminate PI from PIC. On the other hand, if the cross-validated accuracy for classifier trained to discriminate between PI and GI was significant, there might be distinct cognitive processes between PI and GI comprehension.

Next, we used the Neurosynth Image Decoder (<http://neurosynth.org/decode>; Yarkoni, Poldrack, Nichols, Van Essen, & Wager, 2011) to quantify the neural representation similarity between our pattern classifiers and reverse-inference maps obtained from previous studies (i.e., thousands of published neuroimaging studies included in the Neurosynth database at Jan 2017). The Pearson correlation coefficients (*r*) between the unthresholded weight map of PI/GI classifier and the reverse inference z-map of each of the 2911 terms in the Neurosynth database was calculated to indicate pattern similarity. Here, we focused on pattern correlations between PI/GI comprehension and 15 core concepts in psychology and psycholinguistics: *attention, memory, knowledge, cognitive control, decision, emotion, reasoning, intention, theory of mind, language, orthographic, phonological, lexical, syntactic, and semantic*.

Furthermore, we investigated to what extent language and ToM processing could be involved in PI and GI comprehension. Independently defined language and ToM prototypical brain patterns by the term “language” and “theory mind” in the Neurosynth database, were used to discriminate PI and GI from their respective controls. With a leave-one-participant-out cross-validation scheme, we computed the classification accuracies of the language and ToM pattern classifier for PI vs. PIC, GI vs. GIC, and PI vs. GI.

With the same procedure above, we also trained a ToM classifier to discriminate the false belief and control conditions in the ToM task both on the whole-brain and within dmPFC ROI. For the ROI analysis, predefined voxels (the number of voxels = 1038) from the conjunction analysis illustrated above were selected as training and testing data. We calculated the classification accuracies of the ToM classifiers both on the whole-brain and within dmPFC ROI for PI vs. PIC, GI vs. GIC, and PI vs. GI.

2.2. Results

2.2.1. Behavioral results

For fMRI scanning, a 2 (scenario pair: PI pair vs. GI pair) × 2

(implicature: critical vs. control) repeated-measures ANOVA for participants’ task accuracy revealed a significant interaction, $F(1,27) = 8.20$, $p = 0.008$, $\eta_p^2 = 0.23$ (see Table 2). Tests for simple effects indicated that for the GI pairs, accuracies were lower in the GI condition than in its corresponding control condition, $p < 0.001$; this effect was not significant for the PI pairs, $p = 0.08$. Trials with incorrect response or no response within the time limit (3 s) were excluded from the following behavioral and fMRI analyses.

A 2×2 repeated-measures ANOVA for participants’ RTs revealed a significant interaction, $F(1,27) = 23.69$, $p < 0.001$, $\eta_p^2 = 0.47$. Tests for simple effects indicated that for the PI pairs, RTs were longer in the PI condition than in its corresponding control condition, $p < 0.001$; this effect was smaller for the GI pairs, $p = 0.004$. To deal with the possible speed-accuracy tradeoff in PI and GI conditions, we calculated the inverse efficiency score in each condition, which consisted of the average RT of correct trials divided by accuracy (Townsend & Ashby, 1978, see Table 2). An ANOVA for inverse efficiency scores showed also a significant interaction, $F(1,27) = 8.95$, $p = 0.006$, $\eta_p^2 = 0.25$: the inverse efficiency scores were larger in the PI condition than in its control; this effect was smaller for the GI pair. These findings indicated that understanding utterances with conversational implicature involves more complex pragmatic inferential processes, relative to utterances without conversational implicature, and that understanding PI seemed to be more difficult than understanding GI.

In addition, after the experiment, all fMRI participants read each scenario again and rated how indirectly the reply was related to the preceding question on a 7-point visual analog scale, ranging from “the most direct” to “the most indirect”. For this after-experiment indirectness rating, a 2×2 repeated-measure ANOVA for rating scores showed a significant interaction, $F(1,27) = 52.41$, $p < 0.001$, $\eta_p^2 = 0.66$. Tests for simple effects showed that for the PI pairs, the replies were more indirect in the PI condition than in its corresponding control condition, $p < 0.001$; this effect was smaller for the GI pairs, $p < 0.001$. These results suggested that the replies with conversational implicatures were considered to be more indirect than ones without such implicatures.

2.2.2. Whole-brain univariate analysis

To identify neural correlates of PI and GI comprehension, we examined, respectively, the contrasts PI > PIC and GI > GIC at the whole-brain level. The contrast PI > PIC (Fig. 1A and Table S1 in Supplementary materials) revealed activations in bilateral IFG, MTG, TPJ, mPFC (extending posteriorly to pre-SMA), precuneus (extending to post cingulum cortex), and bilateral MFG. The contrast GI > GIC (Fig. 1B and Table S1) revealed activations in bilateral IFG, left MTG, and mPFC/pre-SMA. Note that, after masking out the activations of the contrast GI > GIC at a voxel-level threshold *p* < 0.01 uncorrected, the contrast PI > PIC showed activations in bilateral anterior temporal lobe, bilateral TPJ, middle mPFC, and precuneus (Fig. 1D, in blue); after masking out the activations of the contrast PI > PIC, the contrast GI > GIC showed activation in pre-SMA (Fig. 1D, in orange).

A whole-brain conjunction analysis of the contrasts PI > PIC and GI > GIC revealed clusters of activation in bilateral IFG, left MTG, and dmPFC (extending to pre-SMA), as shown in Fig. 1C and Table S1. These results indicated that the comprehension of PI and GI may involve both overlapping and distinct neural correlates.

Table 2

Mean accuracy, RT, inverse efficiency score, and degree of indirectness, and standard deviation (in parenthesis) for each condition.

Measurement	PI	PIC	GI	GIC
Accuracy (%)	93.8 (5.2)	95.9 (4.9)	89.1 (7.5)	97.1 (4.0)
RT (ms)	852 (275)	586 (235)	669 (251)	577 (249)
Inverse Efficiency (RT/ Acc)	917 (320)	616 (256)	756 (299)	595 (260)
Indirectness	4.82 (0.94)	2.10 (0.53)	3.40 (1.04)	2.04 (0.77)

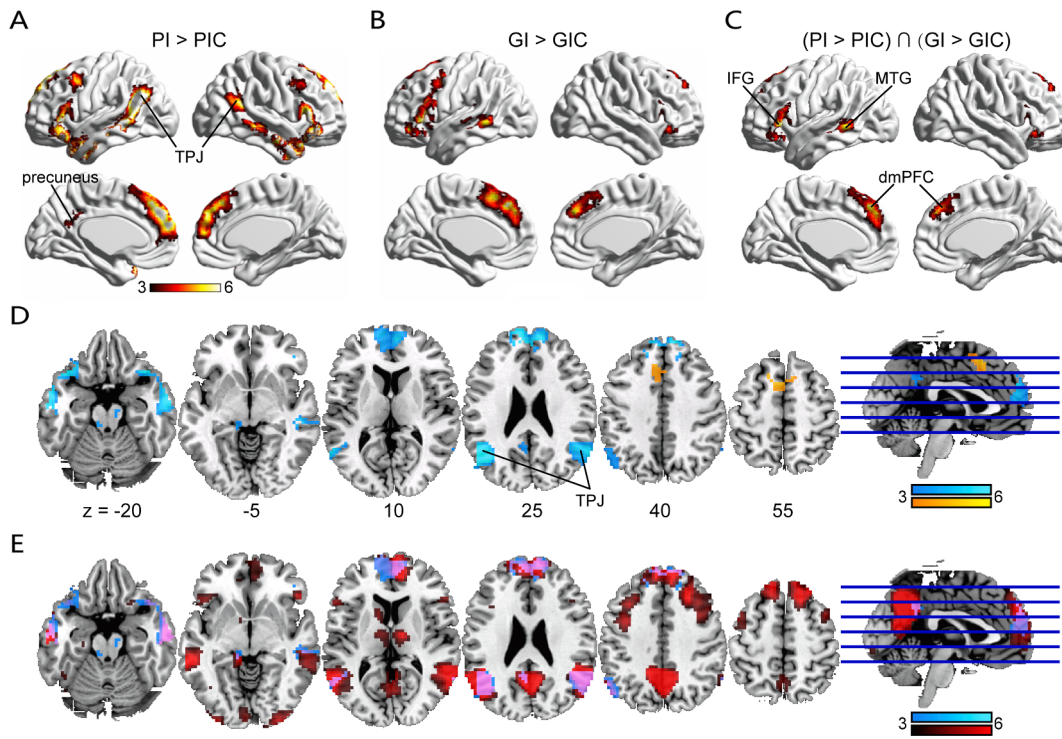


Fig. 1. Results of the whole-brain univariate analyses. The activations were revealed by the contrast PI > PIC (A), the contrast GI > GIC (B), and the conjunction of these two contrasts (C). (D) shows PI-specific activations (shown in blue) and GI-specific activations (shown in orange). (E) shows PI-specific activations (shown in blue), and the activations were revealed by the false belief > control contrast (shown in red). Pink clusters are the overlapping areas of the above two contrasts.

To identify brain regions activated by ToM processing, we examined the false belief > control contrast at the whole-brain level. This contrast evoked clusters of activation in bilateral TPJ extending inferiorly to anterior temporal gyrus, mPFC, precuneus extending to post cingulum cortex, bilateral IFG and MFG. These results are highly consistent with the ToM network identified in previous studies (Dodell-Feder et al., 2011; Lee & McCarthy, 2016). As shown in Fig. 1E, PI-specific activations (in blue) were almost completely embedded in ToM processing network identified in this study (in red).

2.2.3. Whole-brain multivariate pattern analysis

To test the hypothesis that PI and GI processing have shared neural representations, we first trained and tested multivariate patterns at the whole-brain level. Multivariate fMRI pattern classifier trained to dissociate PI vs. PIC could discriminate PI from its control with 96% accuracy (95% confident interval (CI): 90–100%, $p < 0.001$). When this classifier was applied to discriminate GI and its control, an accuracy approaching 100% (95% CI: 100–100%, $p < 0.001$) was obtained. Similarly, the classifier trained to dissociate GI vs. GIC could discriminate GI condition from its control with 96% accuracy (95% CI: 90–100%, $p < 0.001$), and could be generalized to discriminate PI vs. PIC with an accuracy of 96% (95% CI: 90–100%, $p < 0.001$). These findings provided evidence for the existence of functionally shared neural representations for PI and GI. In addition, we found that the classifier trained to dissociate PI vs. GI could discriminate PI condition from GI condition with 96% accuracy (95% CI: 91–100%, $p < 0.001$). Although such between-item comparison is informal, this finding may offer the possibility of distinction between these two processes.

Fig. 2A displays the thresholded whole-brain weight maps of the classifiers that discriminate PI (vs. PIC) and GI (vs. GIC), respectively (bootstrap tests with 10,000 iterations, a threshold of $p < 0.001$ uncorrected for illustration purpose only). PI vs. PIC was predicted by activations in bilateral IFG, left anterior temporal lobe, right anterior MTG, bilateral TPJ and mPFC, while GI vs. GIC was predicted by increased activity in bilateral IFG, left posterior MTG and mPFC.

As a quantitative method, the neural similarity analyses with Neurosynth found that the PI classifier was positively correlated with prototypical brain patterns associated with language or ToM processing (with the terms *language*, *semantic*, *theory mind*, *intention*), while the GI classifier was only correlated with prototypical brain patterns associated with language processing (*language*, *semantic*; Fig. 2B).

We further examined the extent to which PI and GI engage language and ToM processing, by showing how activation patterns of these two processes classify neural representations of PI and GI. “Language” and “ToM” prototypical brain patterns (Fig. 2D), defined by the meta-analytic database (term “language” and “theory mind” respectively), were used to discriminate PI and GI from their own controls (see Fig. 2C). The “Language” pattern performed significantly above chance in discriminating both PI vs. PIC (82%, 95% CI: 68–93%, $p < 0.001$) and GI vs. GIC (79%, 95% CI: 64–91%, $p = 0.004$), but performed at chance level in discriminating PI vs. GI (64%, 95% CI: 50–78%, $p = 0.18$), suggesting that PI and GI comprehension engaged essentially the same neural pattern of language processing. In contrast, the “ToM” pattern could discriminate both PI vs. PIC (93%, 95% CI: 84–100%, $p < 0.001$) and PI vs. GI (89%, 95% CI: 79–97%, $p < 0.001$), but performed at chance level in discriminating GI vs. GIC (61%, 95% CI: 45–76%, $p = 0.34$), indicating that the ToM-related inferential processes were possibly unique to PI comprehension. Note that use of the “ToM” pattern defined by the localizer task yielded the same results of abovementioned classification: discriminating PI vs. PIC with 96% accuracy (95% CI: 89–100%; $p < 0.001$) and PI vs. GI with 97% accuracy (95% CI: 89–100%, $p < 0.001$), but at chance level in discriminating PI vs. GI (68%, 95% CI: 53–83%, $p = 0.09$).

2.2.4. dmPFC functions differentially for PI and GI

The whole-brain MVPA showed that the difference between PI and GI processing was in the neural representations related to ToM and intention consideration. However, the univariate analysis demonstrated that dmPFC, which is considered to be one of the core regions of ToM network, was involved in both PI and GI processing. Within the co-

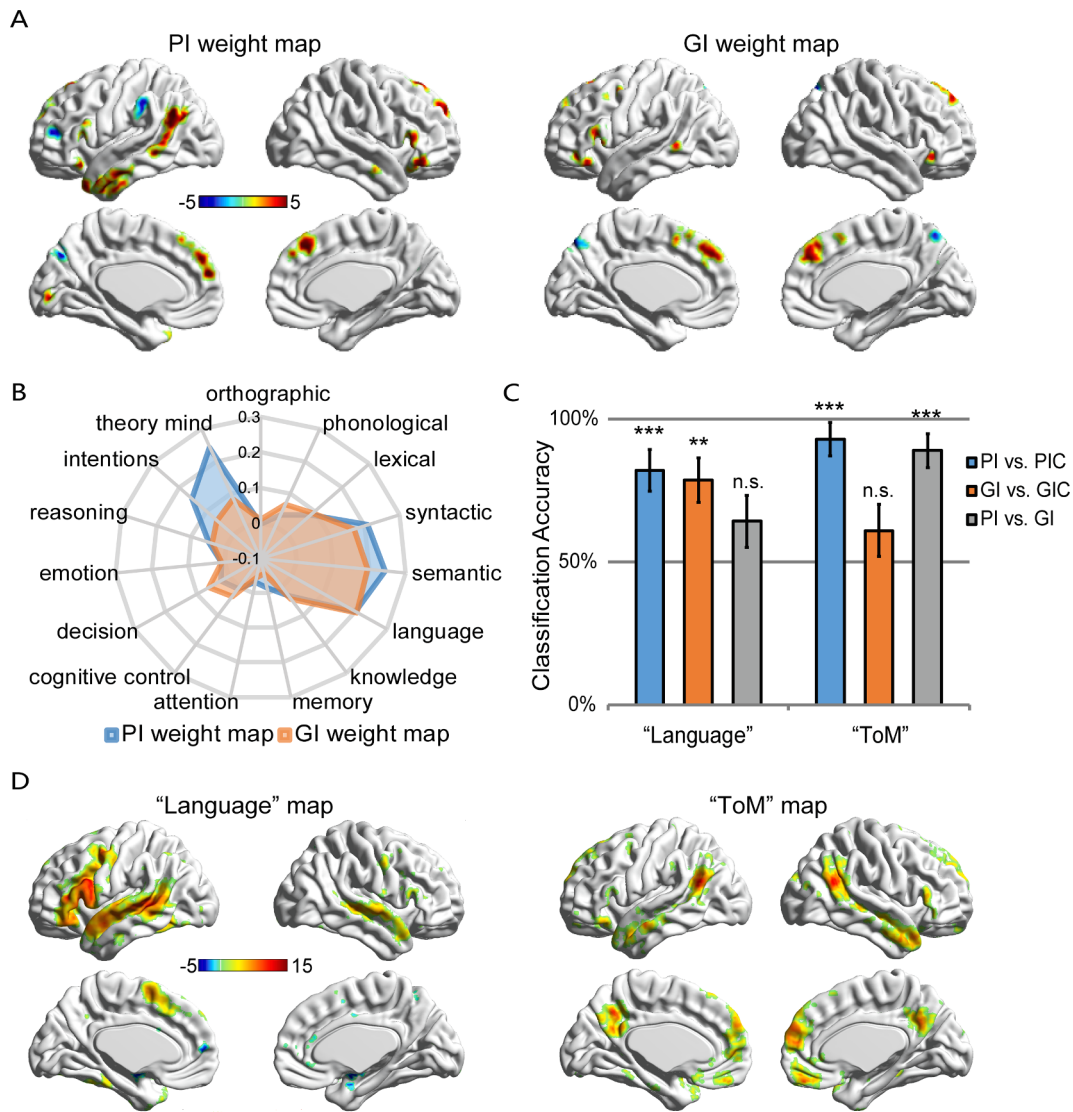


Fig. 2. Results of the whole-brain MVPA. (A) The whole-brain weight maps show voxels whose activity reliably classify PI vs. PIC conditions (i.e., PI weight map) or GI vs. GIC condition (i.e., GI weight map). Positive (warm color) and negative (cool color) weights indicate that more PI/GI processing was predicted by increased and reduced activity, respectively. (B) shows the results of neural similarity analysis using Neurosynth Image Decoder. (C) shows the accuracy of the "Language" map (left three bars) and the ToM map (right three bars) classifying PI vs. PIC, GI vs. GIC, and PI vs. PIC. Error bars represent SEs. ** $p < 0.01$, *** $p < 0.001$, n.s. not significant. (D) shows the prototypic language and ToM maps derived from Neurosynth database.

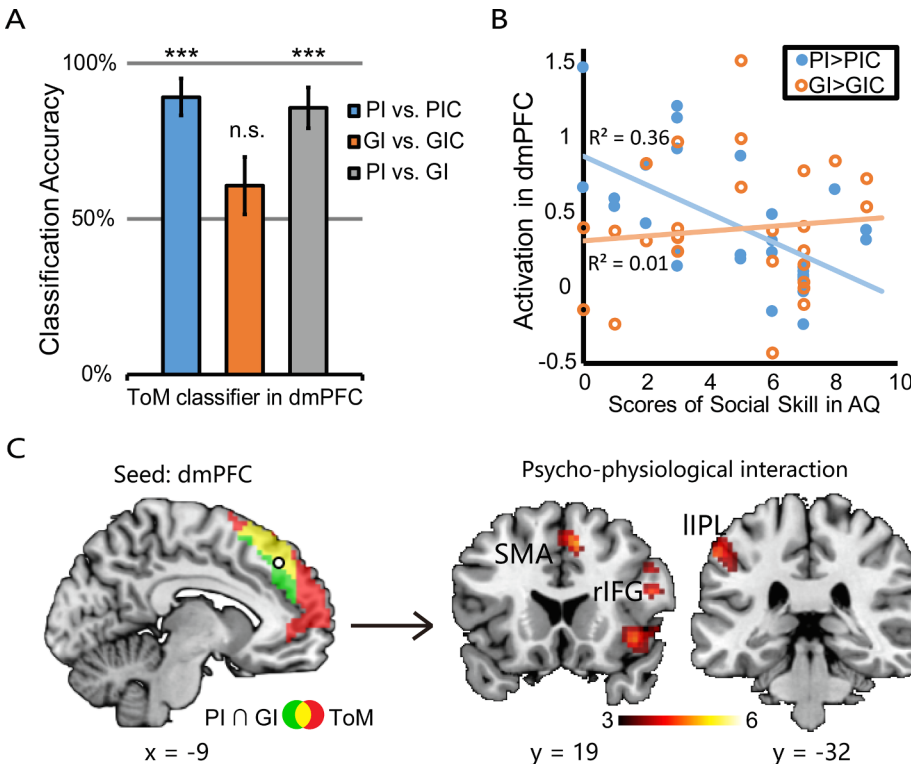
activated dmPFC, 58.6% voxels were also significantly activated by ToM task (see Fig. 3C). Given these seemingly contradictory findings, we further investigated whether dmPFC played an identical role in PI and GI processing.

We first hypothesized that if a "ToM" neural classifier within dmPFC could discriminate PI vs. PIC, but not GI vs. GIC, then it is reasonable for us to believe that PI and GI employed distinct neural representations in dmPFC. To test this hypothesis, we trained a "ToM" multivariate pattern within *a priori* dmPFC ROI to discriminate the false belief condition and its control in the ToM task. This dmPFC ROI was obtained from the univariate conjunction analysis of the contrast PI > PIC and GI > GIC. The cross-validation test showed that this "ToM" classifier could discriminate the false belief condition from its control with 100% accuracy (95% CI: 100–100%, $p < 0.001$). When applied to discriminate the four experimental conditions (Fig. 3A), this "ToM" classifier performed significantly above chance in discriminating both PI vs. PIC (89%, 95% CI: 79–97%, $p < 0.001$) and PI vs. GI (86%, 95% CI: 73–96%, $p < 0.001$). However, this classifier performed at chance level in discriminating GI vs. GIC (61%, 95% CI: 45–76%, $p = 0.34$), consistent

with the whole-brain MVPA classification. These findings provided support to the hypothesis that interpreting PI and GI has distinct neural representations within dmPFC. Specifically, the representation of PI, but not GI, may involve a ToM-related inferential component.

Secondly, we carried out univariate parametric analyses for activation in dmPFC ROI. We added the participants' social skills (as measured by AQ questionnaire; see *Supplementary Materials* for details) as group-level covariates for the PI > PIC and GI > GIC contrasts in two separate models. As shown in Fig. 3B, the magnitude of activation in dmPFC (peak coordinates: [9, 32, 49]; cluster size = 12; $p_{\text{FWE}} = 0.041$, small-volume corrected) negatively correlated with the social skills scores during PI processing ($r = -0.60$, $p = 0.001$), but not during GI processing ($r = 0.10$, $p = 0.61$). A direct comparison confirmed that the two correlation coefficients differed significantly, $z = -3.22$, $p = 0.001$, with 95% CI being [-1.05, -0.29]. These findings indicated that individuals' social skills modulated dmPFC activation during PI processing, but had no effect on GI processing.

Finally, we conducted a PPI analysis by using the *a priori* dmPFC (peak coordinates: [-9, 38, 43]) as seed region. We found that dmPFC



showed significantly stronger functional interplay with several brain regions, including precentral gyrus, left inferior parietal lobe (IPL), right IFG pars opercularis and pars orbitalis (extending to right anterior insula), and pre-SMA during PI processing, relatively to GI processing (Fig. 3C and Table S2).

2.3. Discussion

In the current experiment, we investigated the neural representations of PI and GI comprehension. Results from both univariate and multivariate fMRI data analyses consistently demonstrate that comprehension of PI and GI share a language processing component but differ in that PI but not GI comprehension further relies on ToM-related inferential processes.

In the effect of PI comprehension (comprising indirect replies with PI vs. direct replies), there were activations in bilateral IFG, MTG, mPFC (extending to pre-SMA), TPJ, precuneus, and MFG, which essentially replicated previous findings on PI comprehension (Bašnáková, Weber, Petersson, van Berkum, & Hagoort, 2014; Feng et al., 2017; Shetreet, Chierchia, & Gaab, 2014; Shibata, Abe, Itoh, Shimada, & Umeda, 2011; van Ackeren, Smaragdi, & Rueschemeyer, 2016). IFG and MTG, as core regions of language network, have been implicated in recovering literal content (Ferstl & von Cramon, 2001; Xu, Kemeny, Park, Frattali, & Braun, 2005), while mPFC, TPJ and precuneus constitute a “ToM network” that is typical for tasks involving higher-order, ToM-related inferential processes (Koster-Hale & Saxe, 2013; Van Overwalle & Baetens, 2009). Moreover, instead of creating a pragmatically mismatch context (e.g., the sentence verification or picture-sentence verification paradigm), the current listening comprehension task revealed the neural substrates of GI processing beyond the scope of previous studies. By contrasting indirect replies with GI against direct replies, we showed that interpreting GI also more reliably activates bilateral IFG, left MTG, and mPFC (extending to pre-SMA) than understanding direct replies. Thus, PI and GI processing may engage common neural substrates from the perspective of overlapping fMRI activations.

The brain regions in the “ToM network” can support multiple

Fig. 3. Results of the ROI analyses. (A) shows the cross-validated accuracy of the ToM map within the dmPFC ROI classifying PI vs. PIC, GI vs. GIC, and PI vs. GI. (B) shows the results of ROI-based parametric analyses. The parameter estimates corresponding to four experimental conditions were extracted from the dmPFC (based on the parametric analysis). (C) shows the overlapping area (shown in yellow) within dmPFC between the ToM network (shown in red) and the co-activation of PI and GI (shown in green), and the results of PPI analysis.

cognitive functions other than the classic ToM-related processes (i.e., inferring the mental states of other people, such as false belief reasoning). For example, some studies proposed that the dmPFC activation observed in discourse comprehension may reflect some general functions shared by ToM and discourse comprehension (Mason & Just, 2011); TPJ region supports cognitive control or attention (Carter & Huettel, 2013; Lee & McCarthy, 2016); and ToM network supports working memory of social information (Meyer, Spunt, Berkman, Taylor, & Lieberman, 2012). Therefore, the activation of “ToM” network does not necessarily imply the involvement of the typical ToM processing. To avoid the informal reverse inference of a cognitive process from activation in a certain brain region or system on the basis of a biased literature review (Aguirre, 2003; Poldrack, 2006), we performed more fine-grained analyses, combining the MVPA approach with independent neural representations drawn from large-scale meta-analysis, to clarify whether the overlapping activations arise from the same or different neural representations (Peelen & Downing, 2007) and to identify what cognitive processes were engaged among all likelihood (Poldrack, 2011). On the one hand, our results of whole-brain multivariate pattern decoding provide considerable evidence for the argument that the PI and GI comprehension engage the same neural representation of language processing, which may well be recruited by constructing and maintaining a coherent representation of utterances in the discourse (Menenti, Petersson, Scheeringa, & Hagoort, 2009; Rapp, Mutschler, & Erb, 2012). On the other hand, our results demonstrate that the neural representation engaged in performing ToM-like inferential processes is merely observable during PI comprehension, not during GI comprehension. Combined with the results of univariate analysis, these findings suggest that the comprehender’s ToM-related network is selectively recruited to infer speaker’s aims and intentions by recovering the meaning bound up with specific context.

The dmPFC was involved in both PI and GI comprehension. In the ROI analyses, however, we found that there were differences in the common activation of dmPFC during interpreting PI and GI. First, ROI-based MVPA showed that PI processing activated a ToM-related fMRI pattern within dmPFC, but GI processing did not. It means that

interpreting GI engages only weakly ToM-like inferential processing at best. Second, activation in dmPFC strongly correlated with individuals' social skills during PI processing, but not during GI processing. Third, dmPFC showed significantly stronger functional connectivity with SMA, premotor cortex, right IFG and left IPL during PI processing, relatively to GI processing. The latter pattern of frontal and parietal activity is associated with domain-general cognitive/executive control (Duncan, 2010; Ye & Zhou, 2009a, 2009b). Given that PI comprehension is generally more difficult than GI comprehension, it is reasonable to predict that PI may require additional cognitive processing to monitor and resolve the conflicts between sentential representations in discourse. Thus, the increased functional connectivity may reflect how the cognitive control system was involved in pragmatic inference during PI comprehension. Thus, a related idea is that this region is engaged in strategic inferential processing to establish the relation between utterances in discourse (Ferstl, Neumann, Bogler, & von Cramon, 2008; Ferstl & von Cramon, 2002; Kuperberg, Lakshmanan, Caplan, & Holcomb, 2006). The activation in dmPFC during GI comprehension could reflect a more general and encapsulated inferential process (Ferstl & von Cramon, 2001, 2002; Mason & Just, 2011), such as the one underlying the logical reasoning of specific terms (e.g., *some = not all*).

Nevertheless, our findings are fully congruent with the idea that dmPFC contains multiple, different neural populations that encode distinct mental states. The dmPFC is involved in a variety of high-order cognitive functions. Although dmPFC is one of the central regions in ToM processing (Van Overwalle, 2009), dmPFC is also recruited by ToM-unrelated inductive reasoning (Ferstl & von Cramon, 2002; Siebörger et al., 2007). We suggest that the dmPFC activity in the current study is probably more related to the activation of social information and situational context during generating PI, compared with GI. Specifically, in understanding indirect replies with PI, dmPFC is co-activated with other ToM-related regions, including TPJ and precuneus, and the dmPFC activity supports ToM-like inferential processing in order to infer the current mental state of the speaker in a particular context.

3. Brain stimulation (HD-tDCS) experiments

Given that ToM-related inferential processes may play a critical role in generating PI, but not GI, we performed two independent brain stimulation experiments, using HD-tDCS to test the causal role of a ToM-related brain region (right TPJ) in processing the two types of conversational implicature. In our fMRI experiment, right TPJ was specifically activated during interpreting PI, but not during interpreting GI. This region is generally considered as a critical brain region of the ToM network (Krall et al., 2015; Lee & McCarthy, 2016; Mar 2011; Saxe & Powell, 2006), responsible for extracting and integrating social information from the bulk of information (Carter & Huettel, 2013; Schaafsma, Pfaff, Spunt, & Adolphs, 2015). Moreover, previous studies have revealed that the anodal brain stimulation to right TPJ could improve ToM-related processing in social interaction (Santiesteban, Banissy, Catmur, & Bird, 2012; Sowden, Wright, Banissy, Catmur, & Bird, 2015), and the cathodal stimulation to right TPJ could reduce such function (Leloup, Miletich, Andriet, Vandermeeren, & Samson, 2016; Young, Campodon, Hauser, Pascual-Leone, & Saxe, 2010). Therefore, we selected right TPJ region to deliver tDCS.

3.1. Methods

3.1.1. Participants

Sixty-seven (37 females; mean age = 21.3, SD = 2.4, range 18–28 years) and eighty-eight (56 females; mean age = 20.7, SD = 2.0) university students, who did not take part in either the pretests or the fMRI experiment, participated in one anodal and one cathodal tDCS experiments, respectively. For the anodal experiment, a sub-group of the participants ($n = 34$, 22 females) received anodal tDCS over right TPJ,

whereas the other sub-group ($n = 33$, 15 females) received sham stimulation over the same area. For the cathodal experiment, 46 participants (26 females) received cathodal tDCS over right TPJ, whereas 42 participants (30 females) received sham stimulation over the same area. Five additional participants were excluded from the anodal experiment and seven from the cathodal experiment, due to incomplete data collection or their poor task performance (three SDs longer than average in reaction times or lower in task accuracy).

All the participants were right-handed Chinese native speakers with normal or corrected-to normal vision. None of them suffered from neurological, psychiatric, or hearing disorders. This study was approved by the Ethics Committee of the School of Psychological and Cognitive Sciences at Peking University, and written informed consents were obtained from all the participants.

3.1.2. Procedure

Two independent tDCS experiments were completed. Both experiments were double-blind; that is, neither the participants nor the experimenter who gave instructions to the participants was aware of the assigned type of brain stimulation. HD-tDCS was delivered using a multichannel stimulation adapter (SoterixMedical, 4 × 1-C3) connected to the constant current stimulator (SoterixMedical, Model 1300-A). Five Ag-AgCl sintered ring electrodes were embedded in an EEG cap and connected to the scalp with electrode gel. To deliver stimulation over right TPJ, one central electrode was placed on CP6, and four return electrodes surrounding it were placed over C4, T8, P8, and P4, following previous tDCS studies (Santiesteban et al., 2012; Price, Peelle, Bonner, Grossman, & Hamilton, 2016; Sowden et al., 2015). For active anodal/cathodal stimulation, the direct current climbed to 1.5 mA over 30 s and maintained constant for 20 min before ramping down. Stimulation started 5 min ahead of the listening comprehension task and covered the whole course of the task. For sham stimulation, the participants only received a 30-seconds ramp-up and a 30-seconds ramp-down stimulation at the beginning of the experiment.

Participants performed a listening comprehension task and a ToM task in turn. The procedure of the listening comprehension task and the ToM task in the tDCS experiments was the same as that in the fMRI experiment, with an exception that in the tDCS experiments, the listening comprehension task included 12 trials for each experimental condition and 24 filler trials. For both the listening comprehension task and the ToM task, we acquired the performance accuracy and RT for each trial.

3.1.3. Data analysis

We conducted repeated-measures ANOVAs for the accuracy and RT of the listening comprehension task in the two tDCS experiments. We also analyzed the accuracy of the ToM task to quantify individuals' task performance, and to further verify the effects of brain stimulation over right TPJ. The reason why we did not use RT of the ToM task as a dependent variable was that there was no enough trials with correct response to analyze in some cases, especially when participants who received cathodal brain stimulation over right TPJ interpreted false belief stories. To further determine the role of ToM processing in comprehending PI, we conducted mediation analyses with the INDIRECT macro for SPSS (<http://www.afhayes.com>) with 20,000 bootstrap iterations (Preacher & Hayes, 2008). Two separate mediation models, for the anodal and cathodal experiments respectively, were tested with the brain stimulation type as the independent variable, the RT difference for the contrast between PI and PIC as the dependent variable, and the performance accuracy difference between false belief and control conditions as the mediator.

3.2. Results

As a manipulation check, we first analyzed behavioral data of the ToM task. We conducted two separate 2 (tDCS type: anodal/cathodal vs.

sham) \times 2 (inference type: belief vs. control) repeated measures ANOVAs on participants' task accuracy. For the anodal experiment (Fig. 4A left panel), a marginally significant interaction between the two factors was revealed, $F(1, 65) = 3.48, p = 0.067, \eta_p^2 = 0.05$. Simple effect analysis revealed that for the sham group, the accuracy rate was lower in false belief condition ($70.6 \pm 2.7\%$) than in the control condition ($81.2 \pm 2.2\% ; p < 0.001, \eta_p^2 = 0.21$); for the anodal group, there was no significant difference in accuracy between false belief condition ($80.0 \pm 2.6\%$) and control condition ($83.8 \pm 2.1\% ; p = 0.14, \eta_p^2 = 0.03$). For the cathodal experiment (Fig. 4A right panel), the analysis also showed a marginally significant interaction, $F(1, 86) = 3.81, p = 0.054, \eta_p^2 = 0.04$. Simple effect analysis revealed that for the sham group, the accuracy rate was lower in false belief condition ($71.7 \pm 2.7\%$) than in control condition ($81.7 \pm 1.9\% ; p = 0.001, \eta_p^2 = 0.12$). This effect was larger for the cathodal group (false belief, $65.4 \pm 2.6\%$ vs. control, $83.3 \pm 1.8\% ; p < 0.001, \eta_p^2 = 0.33$). These findings confirmed that enhancing or disrupting right TPJ functions through tDCS facilitates or hinders ToM-related inferential processes.

We then analyzed behavioral data in the listening comprehension task. For each experimental condition, participants correctly responded to more than 95% of all trials. For the anodal experiment (Fig. 4B left panel), a 2 (tDCS type: anodal vs. sham) \times 2 (scenario pair: PI pair vs. GI pair) \times 2 (implicature: critical condition vs. control condition) repeated measures ANOVA on participants' RTs revealed a significant three-way interaction between tDCS type, scenario pair and implicature, $F(1, 65) = 4.30, p = 0.042, \eta_p^2 = 0.06$. Separate ANOVAs on the tDCS effect were carried out for the PI and GI scenario pairs, respectively. For the PI pair, there was a significant interaction between tDCS type and implicature, $F(1, 65) = 4.12, p = 0.046, \eta_p^2 = 0.06$. Tests for simple effects showed that for the sham group, the RTs were longer in the PI condition (765 ± 49 ms) than in the PIC condition (583 ± 40 ms; $p < 0.001, \eta_p^2 = 0.41$), while this effect was much larger for the anodal group (PI, 827 ± 48 ms vs. PIC, 566 ± 40 ms; $p < 0.001, \eta_p^2 = 0.59$), suggesting that the anodal stimulation over right TPJ causally slowed down responses to the indirect replies with PI. For the GI pair, there was neither a main effect of tDCS type, nor an interaction between tDCS type and implicature ($F_s < 1$),

indicating that the anodal brain stimulation over right TPJ could not affect GI comprehension.

The same pattern of results was obtained in the cathodal experiment (Fig. 4B right panel). The ANOVA on RT showed a significant three-way interaction, $F(1, 86) = 4.28, p = 0.042, \eta_p^2 = 0.05$. Separate ANOVAs on the tDCS effect were carried out for the PI and GI scenario pairs. For the PI pair, there was a significant interaction between tDCS type and implicature, $F(1, 86) = 4.97, p = 0.028, \eta_p^2 = 0.06$. Tests for simple effects showed that for the sham group, the RT was longer in the PI condition (690 ± 34 ms) than in the PIC condition (514 ± 27 ms; $p < 0.001, \eta_p^2 = 0.33$), and this effect was much larger for the cathodal group (PI, 793 ± 33 ms vs. PIC, 534 ± 26 ms; $p < 0.001, \eta_p^2 = 0.54$), indicating that the cathodal stimulation over right TPJ causally showed down responses to the indirect replies with PI. For the GI pair, there was neither a main effect of tDCS type, nor an interaction between tDCS type and implicature ($F_s < 1.5$), indicating that the cathodal brain stimulation over right TPJ could not affect GI comprehension.

To further explore the relationship between brain stimulation over right TPJ and behavioral performance on PI, we examined the indirect pathway from tDCS stimulation via ToM ability (the accuracy difference between false belief and control conditions) to PI comprehension. Results showed that the association between brain stimulation over right TPJ and PI comprehension could be mediated by ToM ability, for both anodal (the indirect effect estimate \pm SE = 22.97 ± 15.77 , 95% CI = [0.59, 65.25]) and cathodal (16.84 ± 13.19 , 95% CI = [0.41, 57.03]) experiments (Fig. 4C). Similar analyses could not be conducted for GI comprehension, as the brain stimulation over right TPJ exhibited no effect on it.

3.3. Discussion

Previous studies have consistently showed that the brain stimulation over right TPJ could causally affect ToM processing (Leloup et al., 2016; Santiesteban et al., 2012; Sowden et al., 2015; Young et al., 2010). Here, to further clarify the functions of ToM network in PI and GI comprehension by distinguishing its causal roles, we selected right TPJ region to

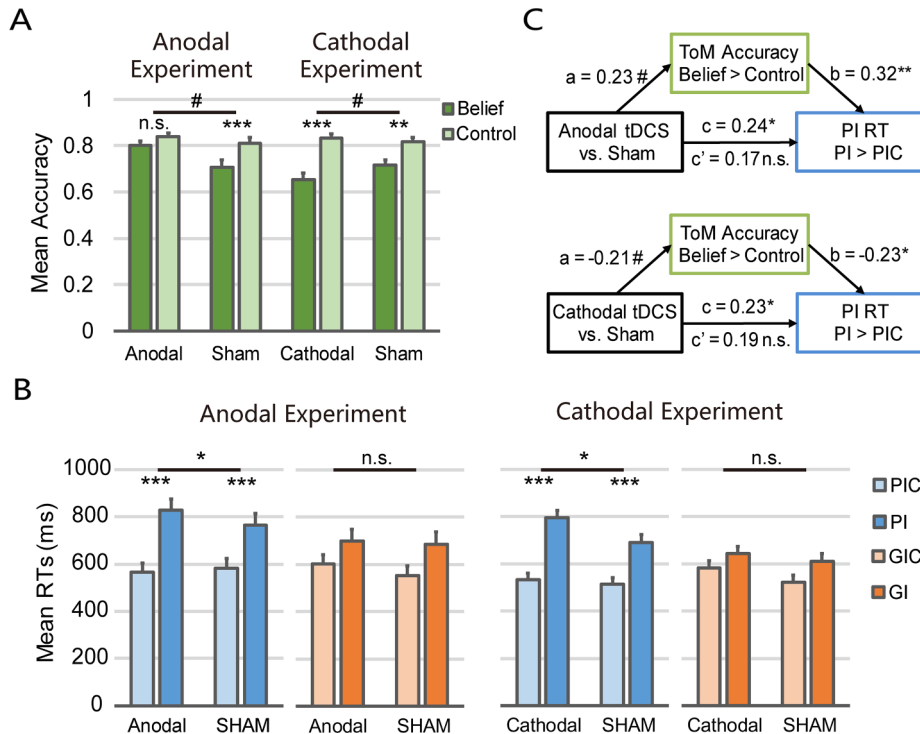


Fig. 4. tDCS results for the ToM task (A) and the listening comprehension task (B). (C) The indirect pathway from the brain stimulation over right TPJ, via ToM ability, to PI comprehension. Error bars represent between-subject SEs. # $p < 0.07$, * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, n.s. not significant.

deliver tDCS.

First of all, results of the ToM task verified the validity of tDCS manipulation by showing that enhancing or disrupting right TPJ functions through tDCS did facilitate or hinder ToM-related inferential processes. More importantly, both anodal and cathodal stimulation causally engendered slower responses to the indirect replies with PI, and the individual's ToM ability mediates the influence of tDCS on PI comprehension. But, neither anodal nor cathodal stimulation over right TPJ impacted responses to the indirect replies with GI. According to previous studies, ToM ability is tightly related to pragmatic language processing (Cummings, 2017), such as irony comprehension (Martin & McDonald, 2004; Monetta, Grindrod, & Pell, 2009), proverb comprehension (Brüne & Bodensteiner, 2005), and the interpretation of indirect speech (Cuerva et al., 2001; Muller et al., 2010). In interpreting an utterance, a comprehender is always to infer and identify the speaker's intentions in a certain linguistic expression. When the speaker's meaning of an utterance relies highly on particular context (as in PI condition), the complexity of such inferential processing increases. Hence during PI processing, the comprehender's communicative-pragmatic performance would be sensitive to his/her ToM ability in discerning the speaker's current intentions. Right TPJ, as a core region of ToM network, is selectively necessary for individuals' PI processing.

Surprisingly, similar to cathodal stimulation, anodal stimulation over right TPJ disrupted PI processing, whereas it did improve the individuals' theory-of-mind ability. This finding is incongruent with our prediction, although it does not invalidate our conclusion that changing the neural activity in right TPJ would affect PI processing but not GI processing. To this finding, one possible explanation is that the individuals' increased ToM ability by anodal stimulation may go well beyond what might be needed to interpret PI in the stimuli and may lead her/him to overthink the speaker's intention behind the indirect replies with PI (Talbert, 2017). Given that individuals do not know the formulaic answer of the speaker's meaning on many occasions in daily life, they will be inclined to consider more possible interpretations of the speaker's meaning when they have adequate resources of ToM processing. This might imply that the relationship between ToM ability and efficiency of interpreting PI is inverse U-shaped, rather than simply linear. In other words, when individuals' ToM ability is either too high or too low, she/he would have difficulties in understanding conversational implicatures as quickly as possible. This can also explain why some previous research failed to find a stable linear relationship between ToM ability and communicative-pragmatic language skills in healthy populations (e.g., Brüne & Bodensteiner, 2005; Gavilán & García-Albea, 2011; Piovan, Gava, & Campeol, 2016).

Another potential explanation is that listeners may be able to comprehend PI by flexibly switching between a language-dominated strategy and a ToM-dominated strategy depending on which strategy is the most efficient in a given context. Stimulating one of these systems will disrupt the cognitive flexibility. For example, consider the example shown in Table 1. To successfully understand this conversation, one may use two different strategies. The language-dominated strategy mainly relies on the successful retrieval of the critical semantic information embedded in the local and global contexts. Here the critical information is that the film mentioned in this conversation is a literary film. If this contextual information is successfully retrieved, one will know that the answer sentence equals to the sentence that "It is hard for audiences to really enjoy your film". In contrast, the ToM-dominated strategy requires limited contextual semantic information. Consider an extreme case that the participant remembered nothing about the cover story. To perform the PI task, one may use the knowledge that people tend to use an indirect way to express an unhappy message. By analyzing the mental states underlying the question using ToM, one would know that the "no" answer is the appropriate answer, so that the seemingly indirect or ambiguous answer is more likely to mean "no". Individuals may use these two strategies in a flexible way: the language-dominated strategy requires more memory load but can provide more accurate information;

the ToM-dominated strategy requires less contextual semantic information but involves more thinking about speaker's mental states. If anodal stimulation of right TPJ facilitates the ToM-dominated strategy, the participants might take more time to understand the PI and to decide whether the answer was the right one. Obviously, the current arguments are just speculations; the potential mechanistic links between ToM system and PI comprehension need further exploration, e.g. behavioral experiments with large sample sizes.

4. General discussion

In this study, the fMRI results demonstrate that comprehension of PI and GI shares a language processing component but differs on whether ToM-related inferential processes are also involved (PI comprehension) or not (GI comprehension). The results from the two tDCS experiments additionally provide causal evidence showing that stimulating a critical ToM region (right TPJ) can exclusively affect PI comprehension, via the modulation of the ToM ability. These findings have fundamental implications for the linguistic debates between Default Theory, Relevance Theory, and Semantic Minimalism concerning to what extent PI and GI processing share the same neurocognitive processes. PI and GI processing are neither identical as Relevance Theory predicts, nor completely distinct as Default Theory predicts. Our findings seem to fit well with Semantic Minimalism in that generating PI and GI share an identical core language system, responsible for enriching semantic content of the utterance; such content further feeds to two systems – a general pragmatic system for generating PI and a more limited system for generating GI (Borg, 2004; 2009).

One important finding of our study was that a language processing system is shared for PI and GI comprehension. This was supported by four pieces of evidence. First, PI and GI comprehension commonly activated the core language brain regions (bilateral IFG and left MTG), relative to their respective controls. Second, PI classifier that trained to discriminate between PI and PIC could discriminate GI from its control, while GI classifier could also discriminate PI from its control. Third, both PI and GI classifiers were positively correlated with language-related prototypical brain patterns defined by the Neurosynth meta-analytic database. Forth, the "Language" prototypical pattern could discriminate PI/GI from their respective controls, whereas it could not distinguish these two types of implicature. Taking together the above evidence, we demonstrated the shared language processing system between PI and GI comprehension. This conclusion is in accord with the Relevance Theory and the Semantic Minimalism that generating GI should at least partially recruited the same cognitive processes as generating PI (Borg, 2009; Sperber & Wilson, 1986). However, the Default Theory does not provide a theoretical space to accommodate the shared processing for PI and GI generation (Levinson, 2000).

Apart from bilateral IFG and left MTG, we revealed common activation in dmPFC for PI and GI comprehension. This set of common activations largely overlapped with the so-called general inference network (Mason & Just, 2011). In evaluating intentional and physical inferences Mason and Just (2011) observed a common set of activations that support both types of inference-making. The shared network, consisted of bilateral IFG, left STG, bilateral anterior temporal lobe, and mPFC, is responsible for successful inference during different task demands (Ferstl & von Cramon, 2001, 2002; Mason & Just, 2011). It is possible that PI and GI comprehension in the current study engaged such a general inference network, whose functioning might be heavily dependent upon the language network. This argument would support the idea of Semantic Minimalism that GI is a kind of abstraction from PI: individuals start by generating GI as PI, but later they learn to achieve this simply by knowing some common rules (Borg, 2009).

More importantly, the current study demonstrated that a crucial difference between the generation of PI and GI is the involvement of ToM processing. Firstly, the fMRI experiment showed that the ToM-related neural representation is only reflected in PI comprehension,

not in GI comprehension, whether at the whole-brain level or within the co-activated dmPFC region. Secondly, the tDCS experiments revealed that the brain stimulation over right TPJ could causally affect PI comprehension through its impacts upon the ToM ability, but it does not affect GI comprehension. These findings consistently indicated that the cognitive processes underlying PI and GI generation are distinct, supporting the intuitive distinction between PI and GI by Grice (1975). Thus, these findings are compatible with the accounts of either Default Theory or Semantic Minimalism. Overall, the evidence from this study suggests that compared to Default Theory and Relevance Theory, Semantic Minimalism provides more felicitous theoretical description of the cognitive processes underlying PI and GI generation and the relationship between these two types of implicatures.

Considering that we used the verbal false belief task to investigate the neural representation associated with ToM processing in the fMRI experiment and to measure the individuals' ToM ability in the tDCS experiments, one thing is noteworthy. In this ToM task, the false belief condition contains short discourses describing false beliefs, while the control condition contains discourses describing outdated photographs and maps (Dodell-Feder et al., 2011). Although this design roughly matched the domain-general inferences about outdated representations, the stimuli used in the target and the control conditions were not strictly matched in terms of linguistic variables. A recent study that matched some basic linguistic variables (such as length, sentence number, word frequency, and number of strokes per word) found that bilateral anterior superior temporal sulci and TPJ showed stronger activation in the false belief condition than the control condition from the beginning sentences of the stories, whereas the false-belief reasoning are supposed to occur only at the ending sentence of the false belief story (Lin et al., 2018). This finding indicates that these ToM-related brain activations may also reflect neurocognitive processes other than inferential processing, such as social concept retrieval. For the current study, analyses using both ToM map and language map from Neurosynth are free from the potential confounding between ToM and language processing in the ToM task. Moreover, we had reasons to believe that PI comprehension recruited ToM-like inference beyond social concept retrieval. First, in the current study, the contrast PI > PIC essentially revealed the full set of ToM network, instead of only regions linked to social concept retrieval. Second, dmPFC, which is unrelated to social concept retrieval, reflected a ToM neural pattern in understanding PI.

Finally, the role of the cognitive/executive control system in implicature comprehension is a concern. The cognitive control system, typically consisting of dmPFC, IFG, premotor cortex, and IPL, is considered to support adaptive behaviors, allowing individuals to deal with change and challenge. Previous studies indicated that the pragmatic difficulties following brain damage are due to domain-general cognitive/attentional control deficits (see Martin & McDonald, 2003). Thus, it is reasonable to predict that the cognitive control system plays a role in implicature comprehension. However, the current study did not provide strong evidence that the cognitive control system is directly engaged during either PI or GI generation. More specifically, the pattern similarity analysis with Neurosynth database did not reveal any significant correlation between PI/GI weight map and the prototypical brain patterns associated with "cognitive control" and "attention" (as shown in Fig. 2B). Nevertheless, we found that the ToM-related area (dmPFC) showed significantly stronger functional connectivity with the domain-general cognitive control network during PI comprehension, relative to GI comprehension. These findings suggest that the cognitive control system may be involved in implicature comprehension indirectly by regulating the dmPFC activity.

5. Conclusion

In this study, we identified both shared and distinct neurocognitive processes underlying PI and GI comprehension. By conducting both univariate analysis and MVPA of fMRI data, we demonstrate that PI and

GI processing engage a shared language processing component, whereas the PI but not GI comprehension requires neurocognitive processes associated with ToM and intention inference. Moreover, the ROI-based fMRI MVPA and functional connectivity results suggest that the computational processes in dmPFC may rely more on knowledge of situational or social information during PI processing, relatively to GI processing. Furthermore, tDCS results provide causal evidence showing that both anodal and cathodal tDCS to right TPJ results in slower PI comprehension, but neither of them impacts GI comprehension. Our findings not only provide a deeper insight into the neurocognitive mechanisms of understanding conversational implicature, but also have broader implications for reviewing linguistic distinctions between PI and GI from the neuroscientific perspective.

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