

The Role of the Angular Gyrus in Semantic Cognition – A Synthesis of Five Functional Neuroimaging Studies

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The role of the angular gyrus in semantic cognition – A synthesis of five functional neuroimaging studies

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Abstract

Semantic knowledge is central to human cognition. The angular gyrus (AG) is widely considered a key brain region for semantic cognition. However, the role of the AG in semantic processing is controversial. Key controversies concern response polarity (activation vs. deactivation) and its relation to task difficulty, lateralization (left vs. right AG), and functional-anatomical subdivision (PGa vs. PGp subregions). Here, we combined the fMRI data of five studies on semantic processing ($n = 172$) and analyzed the response profiles from the same anatomical regions-of-interest for left and right PGa and PGp. We found that the AG was consistently deactivated during non-semantic conditions, whereas response polarity during semantic conditions was inconsistent. However, the AG consistently showed *relative* response differences between semantic and non-semantic conditions, and between different semantic conditions. A combined analysis across all studies revealed that AG responses could be best explained by *independent* effects of both task difficulty and semantic processing demand. Task difficulty effects were stronger in PGa than PGp, regardless of hemisphere. Semantic effects were stronger in left than right AG, regardless of subregion. These results suggest that the AG is independently engaged in both domain-general task-difficulty-related processes and domain-specific semantic processes. In semantic processing, we propose that left AG acts as a “multimodal convergence zone” that binds different semantic features associated with the same concept, enabling efficient access to task-relevant features.

Introduction

Semantic knowledge about objects, people and events in the world is crucial for core human cognitive abilities, such as object recognition and use, as well as language comprehension (Kiefer and Pulvermüller 2012; van Elk et al. 2014; Lambon Ralph 2014). The angular gyrus (AG) is widely considered a key brain region for semantic processing (for reviews, see Binder and Desai 2011; Seghier 2013; Price et al. 2015). This view is supported by meta-analyses of functional neuroimaging studies, which revealed consistent AG engagement for general semantic contrasts (e.g., words vs. pseudowords; Binder et al. 2009; Jackson 2021). Moreover, transcranial magnetic stimulation (TMS) studies indicate a causal role of left AG in general semantic processing (Sliwinska et al. 2015; Davey et al. 2015; Hartwigsen et al. 2016). It has been proposed that the AG acts as a cross-modal convergence zone that integrates semantic features related to various sensory-motor modalities (Binder 2016; Fernandino et al. 2016; Kuhnke et al. 2020b). This theory is corroborated by the AG's proximity to and connectivity with several sensory-motor cortices (Bonner et al. 2013; Binder and Fernandino 2015; Kuhnke et al. 2021).

However, the role of the AG in semantic processing is controversial. Key controversies concern response polarity and its relation to task difficulty, lateralization, and functional-anatomical subdivision. First, the AG often shows deactivation (rather than positive activation) during both semantic and non-semantic tasks, as compared to a resting baseline (Lambon Ralph et al. 2016; Humphreys et al. 2021). This response profile is in line with the AG's involvement in the default mode network (DMN), a set of brain regions that are functionally interconnected during rest and deactivated during attention-demanding tasks (Buckner et al. 2008; Raichle 2015). Notably, the amount of AG deactivation seems to be related to task difficulty (e.g., as measured by response times), where the AG shows less deactivation for easier conditions (Humphreys and Lambon Ralph 2017). This pattern has been observed for both semantic and non-semantic tasks, suggesting a domain-general, rather than semantics-specific role of the AG (Humphreys et al. 2015, 2021). Crucially, general semantic contrasts often compare easy vs. hard conditions (as in words vs. pseudowords). Therefore, it is unclear whether relative increases in AG activity for these contrasts indeed reflect semantic processing or merely domain-general task difficulty (Humphreys et al. 2021).

Second, it is unknown whether the left and right AG have the same or different functions in semantic cognition. In the meta-analysis of Binder et al. (2009), both left and right AG showed consistent engagement in semantic processing, albeit left AG showed a higher activation likelihood than right AG. In contrast, a more recent meta-analysis found exclusive recruitment of the left AG, but not right AG, across neuroimaging studies (Jackson 2021). Indeed, some individual studies reveal selective semantic effects in the left, but not right AG (Price et al. 2016; Kuhnke et al. 2020b), suggesting a potential functional lateralization.

However, the response profiles of left and right AG in semantic and non-semantic tasks have not been systematically compared.

Finally, it remains unclear whether the AG constitutes a functional unit or comprises multiple functional subdivisions. The AG can be anatomically subdivided into two distinct cytoarchitectonic subregions: an anterior subregion *PGa*, and a posterior subregion *PGp* (Caspers et al. 2006, 2008). As function is generally assumed to follow cytoarchitecture (Fedorenko and Kanwisher 2009), it seems likely that *PGa* and *PGp* correspond to two functionally distinct areas. Indeed, related to the first controversy, it is possible that the AG contains both a subregion engaged in semantic processing and a domain-general subregion that shows task-difficulty-related deactivation (Seghier et al. 2010). While functional subdivisions of the AG have been suggested previously (Seghier 2013; Noonan et al. 2013), it is unclear to what extent they correspond to the *PGa* vs. *PGp* distinction, and the response profiles of *PGa* and *PGp* have not been directly compared during semantic and non-semantic tasks.

To address these issues, we combined and re-analyzed the fMRI data of 5 different studies on semantic processing from our laboratory (Kuhnke et al. 2020b; Chapman and Hartwigsen 2021; Graessner et al. 2021; Turker et al. 2021; Martin et al. 2021). For each study, we investigated the response profiles of the same anatomical regions-of-interest (ROIs) for left and right *PGa* and *PGp*. Specifically, we asked whether each AG subregion shows significant activation or deactivation, as compared to the resting baseline, during semantic and non-semantic conditions. Moreover, we tested for relative activity differences between semantic and non-semantic conditions, as well as between different semantic conditions to probe the AG's sensitivity to semantic variables. In a combined linear-mixed-model analysis across all studies, we then investigated whether the response profile of each AG subregion can be best explained by task difficulty, semantic processing demand (i.e., whether the task involves semantic processing), or both.

The view that the AG is a domain-general region showing task-difficulty-related deactivation (Lambon Ralph et al. 2016; Humphreys et al. 2021) would predict that the AG is consistently deactivated during both semantic and non-semantic conditions, and that the level of AG activity can be explained by task difficulty alone. In contrast, the view that the AG is engaged in semantic processing (Seghier 2013; Binder and Fernandino 2015; Kuhnke et al. 2020b) would predict that AG responses cannot be explained by task difficulty alone, but it is crucial to consider semantic processing demand. Regarding laterality, we hypothesized that left AG might show stronger semantic effects than right AG, given that left AG showed more consistent engagement in meta-analyses of semantic processing (Binder et al. 2009; Jackson 2021). Regarding functional subdivision of the AG, some previous work would predict that *PGa* is involved in domain-general task-difficulty-related processes, whereas *PGp* is engaged

in semantic processing (Jefferies 2013; Noonan et al. 2013). Other research suggests that both PGa and PGp are involved in semantic processing, albeit with different semantic roles (Seghier 2013).

Materials and Methods

Studies

We combined the fMRI data of 5 different studies on semantic processing from our laboratory (Kuhnke et al. 2020b; Chapman and Hartwigsen 2021; Graessner et al. 2021; Turker et al. 2021; Martin et al. 2021). Table 1 presents an overview of all studies. Every study acquired blood oxygenation level dependent (BOLD) functional images in young and healthy human adults using a 3T MRI scanner. The total number of participants across all studies was 172. 36 participants participated in multiple studies, yielding 111 unique participants. This overlap was taken into account in combined analyses. Written informed consent was obtained from all participants. All studies were performed according to the guidelines of the Declaration of Helsinki and approved by the local ethics committee of the University of Leipzig. For further details on each study's experimental design, measurement procedure and processing, please see the individual publications.

For all subsequent analyses, each study's fMRI data were re-preprocessed to match voxel size (2.5 mm³) and smoothing kernel size (5 mm³ FWHM) in Montreal Neurological Institute (MNI) space. Moreover, we ensured that no study removed response time related activity from their data as a central question of the present study is whether AG responses can be explained by task difficulty.

ROI analyses

For each study, we extracted the response profiles from the same anatomical regions-of-interest (ROIs) for left and right PGa and PGp. Anatomical ROIs were defined as the maximum probability maps for these cytoarchitectonic areas (Caspers et al. 2006, 2008) provided by the *SPM Anatomy* toolbox (Eickhoff et al. 2005, 2006). For each experimental condition and participant, we estimated percent signal change (PSC) as compared to the resting baseline using the *MarsBaR* toolbox (Brett et al. 2002). For statistical inference, we first performed one-sample t-tests on each experimental condition to test for significant activation or deactivation from the resting baseline. Second, to investigate relative activity differences between experimental conditions, we ran repeated-measures ANOVAs, correcting for non-sphericity using the Huynh-Feldt method. Significant interactions were resolved via step-down ANOVAs and post-hoc paired t-tests. P-values were corrected for multiple comparisons using the Bonferroni-Holm method.

Table 1. Overview of all studies.

	Study	N	Age range	Tasks	Stimulus type	Stimulus modality	Stimulus conditions	Example trial
A	Chapman and Hartwigsen (2021)	41	21-35 years	taxonomic judgment; thematic judgment; scrambled judgment	pairs of line drawings (comprehension)	visual	taxonomically related, thematically related & unrelated images; <i>congruent & incongruent scrambled images</i>	taxonomic: moose & gorilla: taxonomically related?; thematic: monkey & banana: thematically related?; scrambled images: does left image mirror right?
B	Graessner et al. (2021)	33	19-34 years	explicit (meaningfulness judgment); implicit (lexical status comparison)	spoken 2-word phrases (comprehension)	auditory	meaningful, anomalous, pseudoword phrases; single words	explicit: "anxious horse" - meaningful?; implicit: "anxious gufel" - same lexical status?
C	Kuhnke et al. (2020b)	40	19-33 years	lexical decision; sound judgment; action judgment	written words (comprehension)	visual	words with a low or high association to sounds & actions; <i>pseudowords</i>	lexical decision: "Telephone" - word?; sound judgment: "Telephone" - strongly associated with sounds?; action judgment: "Telephone" - strongly associated with actions?
D	Martin et al. (2021)	30	21-34 years	overt semantic fluency; overt counting	written category names (production)	visual	easy & difficult semantic categories; <i>easy (forward) & difficult (backward) counting</i>	easy semantic categories: colors, body parts, animals; difficult semantic categories: flowers, metals, fish
E	Turker et al. (2021)	28	18-29 years	overt reading	written words (production)	visual	simple (2-syllable) & complex (4-syllable) words & pseudowords	read aloud: "Firma", "Bürgermeister", "Molte", "Perautsarung"

Semantic conditions are highlighted in bold font; non-semantic conditions are highlighted in italics.

Linear-mixed-model analysis across all studies

To examine effects of semantic processing and task difficulty on AG activation, we employed a linear-mixed-effects modeling approach across all studies. First, we used a goodness-of-fit comparison to determine the best-fitting model using the Akaike Information Criterion (AIC), where a model was considered meaningfully more informative than others if it decreased the AIC by at least two points (Burnham and Anderson 2004). To reduce overfitting, AIC takes into account model complexity by penalizing models with more parameters. Models predicted PSC across ROIs using various fixed and random effects. The optimal model for our data in terms of AIC was determined in a stepwise fashion, first determining the optimal random effects structure (i.e., individual random effects of task and condition, nested effects, etc.); next determining which measure(s) of task difficulty (response times or error rates) better predicted PSC; and finally determining which interactions of task difficulty, semantic processing demand, and ROI were optimal. In describing model selection, we name the best model and then describe the alternative models and their difference in AIC (Δ AIC).

Models were run using the *lme4* package (Bates et al. 2015) in *R* (version 4.1.1; R Core Team), and model AIC comparisons were performed using the *bbmle* package (Bolker 2020). Significance of effects from the optimal model and further models was derived using the Satterthwaite approximation for degrees of freedom from the *R* package *lmerTest* (Kuznetsova et al. 2017), and predicted plots of interactions were created using the *R* packages *sjPlot* (Lüdtke 2021) and *ggeffects* (Lüdtke 2018). Significant interactions from the optimal model were further investigated by specifying contrasts with the *R* package *hypr* (Rabe et al. 2020) in the manner suggested by Schad et al. (2020). In addition, versions of the optimal model were also run in each ROI individually to test effects of task difficulty and semantics in each ROI without assuming homogeneity of variance. These models were identical to the optimal model but excluded the ROI term.

Results

ROI analyses for each study

Study A: Chapman and Hartwigsen (2021)

In the study by Chapman and Hartwigsen (2021), 41 young and healthy adults had to decide whether pairs of object line drawings were taxonomically related (taxonomic task), or thematically related (thematic task). In both tasks, the drawings could be taxonomically related (e.g., *monkey* and *moose*), thematically related (e.g., *monkey* and *banana*), or unrelated (e.g., *monkey* and *telephone*). In a non-semantic control task, participants decided whether pairs of scrambled versions of images from the semantic tasks mirrored each other.

All AG-ROIs were deactivated in the non-semantic scrambled task and exhibited higher activity for semantic than non-semantic tasks. However, the four AG-ROIs showed distinct response profiles during the semantic tasks (ROI x TASK x CONDITION interaction) (Figure 1A; Tables S1-S3 for statistics). Left PGp was the only AG-ROI to show positive activation, which occurred for thematic pairs during thematic judgments. Both left AG subregions showed relatively higher activity for thematic pairs than taxonomic and unrelated pairs across semantic tasks.

Right PGa was deactivated for all conditions, with less deactivation for taxonomic and thematic than unrelated pairs across tasks. Finally, right PGp did not show activation differences from resting baseline in the semantic tasks. However, right PGp was relatively more engaged for thematic than taxonomic and unrelated pairs across tasks.

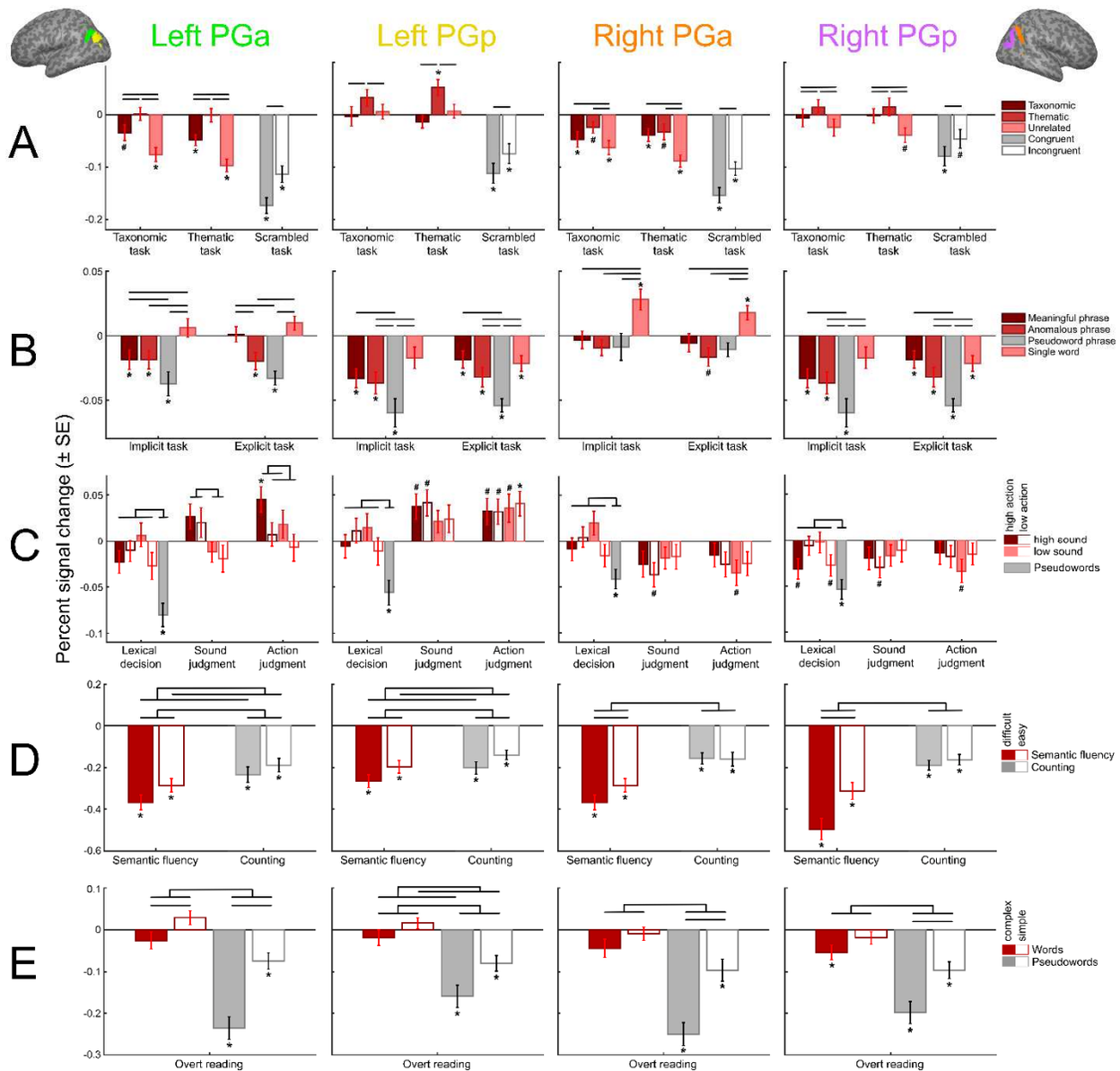


Figure 1. Response profiles of each AG subregion for every study. Mean percent signal change as compared to the resting baseline is plotted for each experimental condition (grouped by task). Error bars represent standard error of the mean. Semantic conditions are highlighted in red; non-semantic conditions are in gray. Black bars illustrate significant differences between experimental conditions. *: $p < 0.05$ (Bonferroni-Holm corrected); #: $p < 0.05$ (uncorrected).

Study B: Graessner et al. (2021)

In Graessner et al. (2021), 33 participants performed explicit (meaningfulness judgment) and implicit (lexical status judgment) semantic tasks on auditorily presented meaningful phrases (e.g., “fresh apple”), anomalous phrases (“awake apple”) and pseudoword phrases (“fresh gufel”), as well as single words (“apple”).

Left PGa showed a distinct response profile from the other AG-ROIs (ROI x TASK x CONDITION interaction) (Figure 1B; Tables S4-S6). In both tasks, left PGa was significantly deactivated for anomalous and pseudoword phrases. Pseudoword phrases elicited stronger deactivation than meaningful and anomalous phrases. However, selectively in the explicit task (i.e., not in the implicit task) and selectively in left PGa, meaningful phrases induced relatively higher activity than anomalous phrases (TASK x CONDITION interaction).

All other AG-ROIs showed a main effect of CONDITION. Left and right PGp were significantly deactivated for all phrases in both tasks. Pseudoword phrases induced stronger deactivation as compared to meaningful phrases in both regions, and as compared to anomalous phrases in left but not right PGp.

Right PGa was significantly activated for single words, but not for phrases, in both tasks. Single words produced higher activity than all phrase types (which did not significantly differ).

Study C: Kuhnke et al. (2020b)

In the study by Kuhnke et al. (2020b), 40 participants performed three different tasks—lexical decision, sound judgment, and action judgment—on written words with a high or low association to sounds and actions (e.g., “guitar” was a high sound–high action word). Pseudowords acted as a non-semantic control condition.

In the lexical decision task, none of the AG-ROIs showed significant (de-)activation for the four word types, but all regions demonstrated strong deactivation for pseudowords that differed significantly from activity for words (Figure 1C; Tables S7-S9).

In the semantic judgment tasks, the AG-ROIs exhibited distinct response profiles (ROI x TASK x SOUND and ROI x TASK x ACTION interactions). Left PGa showed relatively higher activity for high- than low-sound words during sound judgments. During action judgments, left PGa was more strongly engaged for high- than low-action words. High-sound high-action words elicited significant positive activation as compared to the resting baseline.

In contrast, left PGp showed a main effect of TASK, driven by higher positive activation for all word types during semantic judgments than lexical decisions. (Sound and action judgments did not significantly differ.)

Contrary to the left AG regions, right PGa and PGp showed no positive activation vs. rest in any condition, and no significant activity differences between semantic conditions.

Study D: Martin et al. (2021)

In the study by Martin et al. (2021), 30 healthy adults performed an overt semantic fluency task on easy categories (e.g., colors, animals) and difficult categories (e.g., metals, flowers), as well as an overt counting task with easy (forward) and difficult (backward) conditions.

All AG-ROIs were significantly deactivated during all conditions (Figure 1D; Tables S10-12). Moreover, all AG regions exhibited stronger deactivation during the semantic fluency than counting task (main effect of TASK).

However, left and right AG regions showed a striking difference in their relative responses (ROI x TASK x DIFFICULTY interaction). Left PGa and PGp were more deactivated for difficult than easy conditions across both tasks (main effect of DIFFICULTY). In contrast, right PGa and PGp showed a selective difficulty effect in the semantic fluency task, but not in the counting task (TASK x DIFFICULTY interaction).

Study E: Turker et al. (2021)

In Turker et al.'s (2021) study, 28 participants read aloud simple (two-syllable) and complex (four-syllable) words (e.g., "Firma", "Bürgermeister") and pseudowords (e.g., "Molte", "Perautsarung").

All AG-ROIs were significantly deactivated for both simple and complex pseudowords (Figure 1E; Tables S13-15). Words did not induce significant activation or deactivation, as compared to the resting baseline. As an exception, right PGp was deactivated for complex words.

However, left and right AG-ROIs exhibited distinct relative responses (ROI x WORD x COMPLEXITY interaction). Left PGa and PGp showed a complexity effect (complex vs. simple) for both words and pseudowords. In contrast, right PGa and PGp showed a selective complexity effect for pseudowords, but not for words.

Commonalities and differences between studies

In all studies, all AG-ROIs were consistently deactivated during non-semantic conditions. As the only exception, right PGa did not show a significant difference from resting baseline in study B (Graessner et al. 2021).

In contrast, response polarity during semantic conditions was inconsistent between ROIs, studies, and conditions. The same AG subregion could show both positive activation and deactivation in different studies, or even in different conditions of the same study. Notably, both

study A (Chapman and Hartwigsen 2021) and study C (Kuhnke et al. 2020b) found significant positive activation in left AG during explicit semantic judgment tasks.

Relative activity differences between conditions were more consistent. In all studies, all AG-ROIs showed significant activity differences between semantic and non-semantic conditions. This semantic effect was generally driven by relatively higher activity for semantic than non-semantic conditions.

Crucially, left AG not only showed relative activity differences between semantic and non-semantic conditions, but also between different semantic conditions. The same pattern was sometimes, but not always, observed in right AG (e.g., non-significant differences in studies C and E).

Finally, in addition to functional differences between left and right AG, we found distinct response profiles for the cytoarchitectonic subregions PGa and PGp. Out of all AG-ROIs, left PGa seemed to exhibit the strongest sensitivity to fine-grained semantic manipulations. For example, in study B (Graessner et al. 2021), left PGa was the only AG-ROI that showed relatively higher activity for meaningful phrases (e.g., “fresh apple”) than anomalous phrases (e.g., “awake apple”) during explicit meaningfulness judgments. In study C (Kuhnke et al. 2020b), left PGa was the only AG subregion that showed a selective response to task-relevant semantic features, that is, higher activation for high- than low-sound words during sound judgments and for high- than low-action words during action judgments.

Linear-mixed-model analysis across all studies

To investigate whether the response patterns of each AG subregion could be best explained by task difficulty, semantic processing demand (i.e., whether the task involves semantic processing), or both, we performed a linear-mixed-model analysis across all studies.

First, to determine the optimal random effects structure for our model, we tested a baseline null model predicting percent signal change (PSC) with only the mean signal and random intercepts for participants. The models being compared always included random intercepts for participants, but they differed in the task and condition random effects. Results showed that the optimal random effects structure included a random intercept for participant and a random intercept for condition nested within task (Table 2A).

Table 2. AIC comparisons to determine the optimal model predicting AG response.

Comparison	Model	Δ AIC	df	
(A) Random effects	PSC ~ 1 + (1 participant) + (1 task/condition)	0	5	*
	PSC ~ 1 + (1 participant) + (1 task)	5.5	4	
	PSC ~ 1 + (1 participant) + (1 condition)	463.5	4	
	PSC ~ 1 + (1 participant)	1610	3	
(B) zRT & zER	PSC ~ zRT + (1 participant) + (1 task/condition)	0	6	*
	PSC ~ 1 + (1 participant) + (1 task/condition)	42.2	5	
	PSC ~ zER + (1 participant) + (1 task/condition)	43.6	6	
(C) Interactions	PSC ~ zRT + semantics + ROI + zRT:ROI + semantics:ROI + (1 participant) + (1 task/condition)	0	16	*
	PSC ~ zRT + semantics + ROI + zRT:semantics + zRT:ROI + semantics:ROI + (1 participant) + (1 task/condition)	0.3	17	
	PSC ~ zRT + semantics + ROI + zRT:semantics + zRT:ROI + semantics:ROI + zRT:semantics:ROI + (1 participant) + (1 task/condition)	5.4	20	
	PSC ~ zRT + semantics + ROI + semantics:ROI + (1 participant) + (1 task/condition)	30.4	13	
	PSC ~ zRT + semantics + ROI + zRT:semantics + semantics:ROI + (1 participant) + (1 task/condition)	30.7	14	
	PSC ~ zRT + semantics + ROI + zRT:ROI + (1 participant) + (1 task/condition)	34.9	13	

PSC ~ zRT + semantics + ROI + zRT:semantics + zRT:ROI + (1 participant) + (1 task/condition)	35.2	14
PSC ~ zRT + semantics + ROI + (1 participant) + (1 task/condition)	66.4	10
PSC ~ zRT + semantics + ROI + zRT:semantics + (1 participant) + (1 task/condition)	66.8	11
PSC ~ zRT + (1 participant) + (1 task/condition)	160.3	6

* = winning model; PSC = percent signal change; zRT = standardized RT; zER = standardized error; AIC = Akaike Information Criterion.

Next, we determined whether standardized error rates (zER) and/or standardized response times (zRT) were good predictors of PSC. Models with a single fixed effect—zER or zRT—were compared to our null intercept model. Results showed that including zRT as a fixed effect improved model fit, but the model including zER was not better than the null model (Table 2B). Thus, zRT was included as a fixed effect in further models and zER was excluded.

Finally, we compared models with three fixed effects—zRT, semantics, and ROI—and all combinations of interactions between the three. The model with the highest explanatory value included a 2-way interaction of zRT x ROI as well as a 2-way interaction of semantics x ROI (Table 2C). The ANOVA table for this optimal model is shown in Table 3. Although the next best-fitting model was within 2 Δ AIC from the optimal model, suggesting no meaningful difference, the additional zRT x Semantics variable implies that the next best-fitting model may be overfit, and so we interpret results from the more parsimonious optimal model.

Table 3. ANOVA table for the optimal model of AG responses across studies.

Factor	SS	MSE	DF	Den. DF	F	p
zRT	0.37	0.37	1	4822.9	45.68	< 0.001
Sem	0.09	0.09	1	27.7	11.50	0.002
ROI	0.35	0.12	3	5187.4	14.67	< 0.001
zRT x ROI	0.29	0.10	3	5187.4	12.18	< 0.001
Sem x ROI	0.33	0.11	3	5187.4	13.67	< 0.001

SS = sum of squares; MSE = mean squared error; DF = degrees of freedom; Den = denominator; Sem = effect of semantics.

A plot of the predicted interaction between zRT and ROI based on the optimal model (Figure 2A) suggested that the effect of zRT was strongest in PGa and weaker in PGp. A plot of the predicted interaction between semantics and ROI (Figure 2B) suggested that the effect of semantics was stronger in left than right AG-ROIs. These interactions were investigated further by adding specific ROI contrasts to the model (full model results in Supplementary Table S16).

Specifically, the contrast model included contrasts of PGa vs. PGp and left vs. right ROIs. We found that zRT interacted with the PGa vs. PGp contrast (coeff = -0.022, $p < 0.001$) but not with the left vs. right contrast (coeff = -0.004, $p = 0.47$), providing evidence that the zRT x ROI contrast was due to a larger effect of zRT in PGa than PGp, regardless of hemisphere. Moreover, effects of semantics interacted with the left vs. right contrast (coeff = 0.075, $p < 0.001$), but not with the PGa vs. PGp contrast (coeff = 0.009, $p = 0.46$). This indicates that the semantics x ROI interaction was driven by a stronger effect of semantics in left than right AG, regardless of subregion.

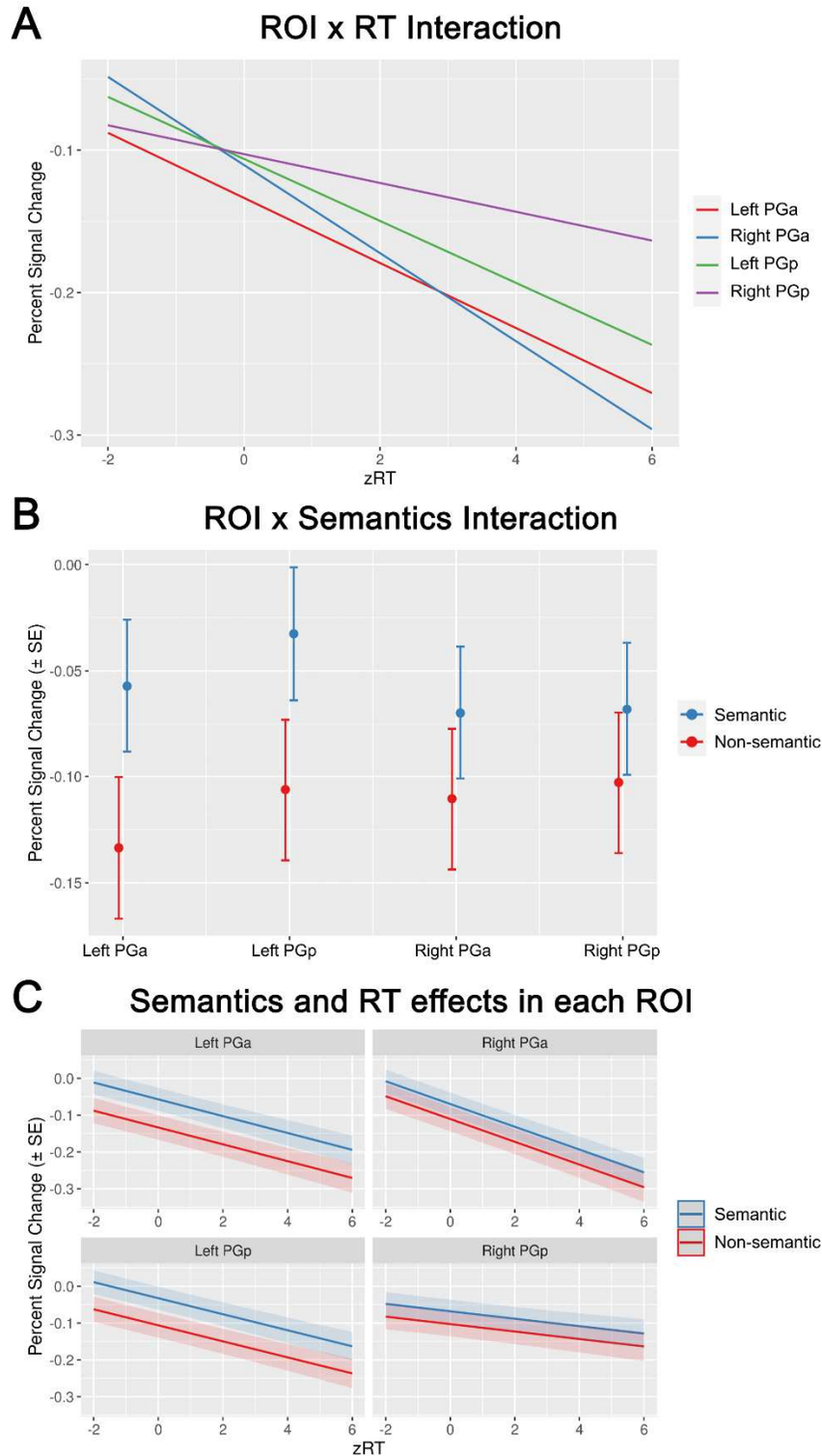


Figure 2. Results of the linear-mixed-model analysis. (A) Interaction of ROI and zRT on predicted percent signal change. (B) Interaction of ROI and semantics on predicted percent signal change. (C) Combined effects of semantics and zRT on predicted percent signal change in each ROI. Error bars and shaded areas represent standard error of the mean.

Finally, we examined the zRT and semantic effects in each ROI individually (Figure 2C; Supplemental Tables S17-S20). This step-down analysis allowed us to investigate the effects of task difficulty and semantics without assuming homogeneity of variance in each ROI. These models revealed that, in addition to the independent effects of difficulty and semantics across all AG ROIs, each individual ROI showed independent main effects of difficulty (zRT) (Left PGa coefficient = -0.013, $p = 0.015$; Left PGp coefficient = -0.024, $p < 0.001$; Right PGa coefficient = -0.027, $p < 0.001$; Right PGp coefficient = -0.021, $p < 0.001$) and semantics (Left PGa coefficient = 0.073, $p = 0.001$; Left PGp coefficient = 0.057, $p < 0.001$; Right PGa coefficient = 0.051, $p = 0.012$; Right PGp coefficient = 0.044, $p = 0.021$).

Overall, the mixed-model analysis across all studies revealed that AG responses are best explained by *independent* effects of task difficulty (as measured by zRT) and semantic processing demand. AG activity decreased with increasing task difficulty. This task performance effect was larger in PGa than PGp, regardless of hemisphere. Moreover, AG activity was relatively higher for semantic than non-semantic conditions. This semantic effect was larger in left than right AG, regardless of subregion. The model intercept is negative, indicating that the AG is deactivated on average. However, AG activity was sometimes positive, especially when the task was semantic (see Figure S1).

Discussion

This study tested the functional involvement of the angular gyrus (AG) in semantic cognition, focusing on three key issues: (1) response polarity (activation vs. deactivation) and its relation to task difficulty, (2) lateralization (left vs. right AG), and (3) functional-anatomical subdivision (PGa vs. PGp). To this end, we combined and re-analyzed the fMRI data of five studies on semantic processing from our laboratory (Kuhnke et al. 2020b; Chapman and Hartwigsen 2021; Graessner et al. 2021; Turker et al. 2021; Martin et al. 2021). For each study, we extracted the response profiles from the same anatomical regions-of-interest (ROIs) for left and right PGa and PGp.

We found that the AG was consistently deactivated during non-semantic conditions, as compared to the resting baseline. In contrast, response polarity was inconsistent during semantic conditions, involving both deactivation and activation in different studies, conditions and AG-ROIs. However, we consistently found *relative* response differences between semantic and non-semantic conditions, as well as between different semantic conditions. These effects were always observed in left AG, and often but not always in right AG.

A combined linear-mixed-model analysis across all studies revealed that these response profiles could be best explained by both task difficulty (as measured by response times; RT) and semantic processing demand *independently*. AG activity decreased with increasing task difficulty and was relatively higher for semantic than non-semantic conditions. Difficulty effects were stronger in PGa than PGp, irrespective of hemisphere. Semantic effects were stronger in left than right AG, regardless of subregion.

Theories of AG Function

Our findings support the view that the AG is engaged in semantic processing (Binder and Desai 2011; Seghier 2013; Kuhnke et al. 2020b), while they oppose the view that the AG is exclusively a domain-general region showing task-difficulty-related deactivation (Lambon Ralph et al. 2016; Humphreys et al. 2021). The domain-general view would have predicted that the AG is consistently deactivated during both semantic and non-semantic conditions, and any response differences between semantic and non-semantic conditions can be completely explained by task difficulty differences. In contrast, the semantics view would have predicted that AG responses cannot be explained by task difficulty alone, but it is crucial to consider semantic processing demand (i.e., whether the task involves semantic processing).

In line with the domain-general view, we found consistent deactivation of the AG during non-semantic conditions. Moreover, AG activity was related to task difficulty in both semantic and non-semantic conditions. However, response polarity during semantic conditions was inconsistent and involved positive activations (e.g., in the studies by Kuhnke et al. 2020b; Chapman and Hartwigsen 2021). Crucially, contrary to the domain-general view and in support of the semantics view, AG activity levels could not be explained task difficulty alone, but semantic processing demand proved essential: Models based on task difficulty alone were substantially outperformed by models that also included semantic processing demand. The optimal model included two-way interactions of ROI x RT and ROI x Semantics, indicating that effects of RT and semantics on activity levels in each AG-ROI were independent. These results strongly support the view that the AG is engaged in semantic processing.

Response Polarity and Semantic Processing in the AG

The inconsistency of AG response polarity during semantic conditions suggests that responses vs. rest are unreliable evidence to assess the AG's role in semantic processing, contrary to the arguments in some previous work (Humphreys et al. 2015, 2021). More generally, comparisons against rest are problematic as the resting baseline itself is not process-neutral. "Resting" can

involve mind wandering, autobiographical memory, as well as self-referential and introspective processes (Andreasen et al. 1995; Stark and Squire 2001; Morcom and Fletcher 2007; Andrews-Hanna 2012). It is particularly problematic that all these processes may involve the retrieval of semantic information (Binder et al. 1999, 2009). Thus, the AG might be “deactivated” during attention-demanding tasks as the semantic processing that occurs during rest is interrupted (Seghier 2013). In other words, AG deactivation may indeed reflect its involvement in semantic processing (Binder et al. 1999, 2009). Under this view, it is not surprising that AG activity in our semantic conditions often did not significantly differ from the resting baseline. A semantic region would only be predicted to show positive activation (above rest) when the task involves semantic processing to a *greater* extent than during the resting state. Indeed, we found positive activation for some semantic conditions (e.g., in Kuhnke et al. 2020b; Chapman and Hartwigsen 2021). These conditions have in common that they explicitly required the retrieval of individual semantic features.

Compared to absolute responses vs. rest, *relative* responses between different experimental conditions were much more consistent: The AG consistently showed differential activity between semantic and non-semantic conditions, and between different semantic conditions. This strongly suggests that the AG is sensitive to semantics. Moreover, these findings corroborate the view that relative responses between conditions, not absolute responses vs. rest, should constitute the main focus of neuroimaging studies on AG function (Stark and Squire 2001; Finn 2021).

Task Difficulty Effects in the AG

Several recent studies have claimed that semantic effects in the AG are likely to be an artifact of difficulty rather than semantic processing per se (Humphreys et al. 2015, 2021). That is, the authors point out that some typical semantic contrasts that reveal larger relative activation in the AG may be confounded with difficulty (e.g., words vs. pseudowords, concrete vs. abstract words) and that their effects in the AG are reversible (Pexman et al. 2007; Graves et al. 2017). Indeed, multiple studies have shown difficulty effects in the AG, such that more difficult conditions show increased deactivation compared to less difficult conditions (Hahn et al. 2007; Humphreys et al. 2015). The current study showed that, across five experiments—including three with a words vs. pseudowords contrast—task difficulty and semantic processing demand show *independent* effects in the bilateral AG. AG activity decreased with increasing task difficulty across semantic and non-semantic conditions. Independently, AG activity was relatively higher during semantic than non-semantic conditions. Notably, task difficulty effects and semantic effects were also

orthogonal in regional preference: Task difficulty effects were stronger in PGa than PGp, regardless of hemisphere. Semantic effects were stronger in left than right AG, regardless of subregion. Thus, our results are consistent with the claim that the AG indexes task difficulty, but they provide strong evidence against the claim that semantic effects in the AG are explained by difficulty. Indeed, our results showed semantic effects in the AG in all studies, regardless of task manipulation or input modality (Figure 1, red vs. gray), consistent with previous meta-analyses of semantic processing (Binder et al. 2009; Jackson 2021).

The independence of difficulty effects from semantic effects in the AG provides evidence that the AG is likely to have a domain-general role in addition to being sensitive to semantic processing. This view is supported by a previous fMRI study which revealed that the same regions of the default mode network (DMN), including AG, can show domain-general task difficulty effects (activation for pseudowords vs. words) and domain-specific semantic effects (decoding of high vs. low imageability words) (Mattheiss et al. 2018). Moreover, this view is in line with modern network-based views of cognitive neuroscience, suggesting that the function of a brain region depends on its interactions with other areas in a given task (Seghier 2013). Under this view, the same brain region can have multiple different functions by virtue of being connected to different regions in different tasks (Bassett and Sporns 2017). Indeed, the AG exhibits particularly flexible functional connectivity in different cognitive tasks (Chai et al. 2016; Kuhnke et al. 2021). The AG seems to be involved in *both* domain-general task-difficulty-related processes and domain-specific semantic processes. These are not mutually exclusive.

Lateralization

Left AG showed stronger semantic effects than right AG. Our combined analysis across all studies revealed that left AG exhibits larger activity differences between semantic and non-semantic conditions. In individual studies, left AG always showed activity differences between semantic and non-semantic conditions, as well as between different semantic conditions. Right AG also often showed these effects, but not always. For example, in study C (Kuhnke et al. 2020b), left but not right AG was engaged for sound and action features of word meaning. In study E (Turker et al. 2021), only left but not right AG was sensitive to word complexity. In study B (Graessner et al. 2021), right PGa did not distinguish pseudoword and real-word phrases.

These results are in line with a previous neuroimaging meta-analysis demonstrating a more consistent recruitment of left than right AG during semantic processing (Binder et al. 2009). However, they are inconsistent with meta-analyses suggesting exclusive recruitment of left but not right AG (Jackson 2021; Hodgson et al. 2021). Our findings suggest that right AG is also

sensitive to semantics, but plays a weaker role than left AG, at least under “normal” conditions in young and healthy human adults. In support of this view, Jung-Beeman (2005) summarized evidence that both the left and right hemispheres are engaged in semantic cognition; however, the right hemisphere seems to perform coarser computations than the left.

As a hypothesis for future work, we propose that the right AG might compensate when the left AG is perturbed or even damaged. Such potential mechanisms of adaptive plasticity could be investigated in future studies combining non-invasive brain stimulation (e.g., TMS) with a neuroimaging readout (e.g., fMRI) (Bergmann et al. 2016; Hartwigsen 2018; Hartwigsen and Volz 2021). Previous studies using combined TMS-fMRI have revealed adaptive changes in task-related activation (Binney and Lambon Ralph 2015; Hallam et al. 2016; Jung and Lambon Ralph 2016; Hartwigsen et al. 2017) and/or effective connectivity (Hallam et al. 2016; Jung and Lambon Ralph 2016; Hartwigsen et al. 2017) after TMS was applied before a semantic task. Some of these studies provide evidence for flexible redistribution in the semantic network, with a stronger, compensatory contribution of homologous right-hemispheric areas after perturbation of a key left-hemispheric semantic node (e.g., Binney and Lambon Ralph 2015; Jung and Lambon Ralph 2016). Analogously, future TMS-fMRI studies could investigate whether right AG can compensate when left AG is perturbed during semantic tasks. Such combined TMS-fMRI studies would also be of clinical relevance as they could indicate whether and how the semantic system can compensate for actual AG damage, e.g. following a stroke (Hartwigsen 2016, 2018).

Functional-Anatomical Subdivision

We observed distinct response profiles for the cytoarchitectonic subregions of the AG, *PGa* and *PGp*. Specifically, our combined analysis across all studies revealed that domain-general task difficulty effects were stronger in *PGa* than *PGp*, regardless of hemisphere.

These results partially support and partially refute previous functional-anatomical subdivisions of the AG. Noonan et al. (2013) performed a meta-analysis of functional neuroimaging studies on executive control during semantic processing. They found that left dorsal AG (~*PGa*) and adjacent intraparietal sulcus (IPS) showed increased activity for semantic tasks with a higher executive demand. In contrast, left ventral AG (~*PGp*) was engaged for semantic vs. phonological tasks, but insensitive to executive demands. Based on these results, Jefferies (2013) proposed that dorsal AG / IPS is part of the multiple demand network (MDN) involved in domain-general executive control (Duncan 2010), whereas ventral AG supports semantic integration. Similarly, Humphreys et al. (2021) argued that “dorsal AG (*PGa*)/lateral bank of intraparietal sulcus (IPS) serves a distinctive function to the ventral AG (central *PGp*)” because

“dorsal and ventral AG show opposing effects of task difficulty in semantic and visuospatial tasks; dorsal areas show a greater response when difficulty is increased, whereas ventral AG shows the inverse pattern, stronger deactivation for harder tasks” (p. 453). In line with these proposals, we found stronger task difficulty effects in PGa than PGp, suggesting a stronger contribution of PGa to domain-general task-difficulty-related processes. However, in contrast to these previous views, we found the same *negative* relationship between neural activity and task difficulty in all AG-ROIs: Activity decreased with increasing task difficulty.

This response pattern contradicts the expected response pattern of a control-related MDN region (i.e., increased activity for increased difficulty) and is more consistent with the response of a DMN region (cf. Noonan et al. 2013). Previous reports of control-related activation in PGa may have reflected activation “spillover” from nearby MDN regions, such as IPS (Duncan 2010; Whitney et al. 2012). This is especially plausible in a coordinate-based meta-analysis (as in Noonan et al. 2013), which involves smoothing activation peaks in standard space using Gaussian kernels (Eickhoff et al. 2009). Indeed, a more recent meta-analysis of semantic control, which updated the Noonan et al. (2013) study with novel data and methodology, found no consistent AG engagement (Jackson 2021; also see Hodgson et al. 2021). Moreover, a high-resolution subject-specific parcellation study revealed that IPS is part of the core MDN, whereas the AG is anti-correlated with the MDN and more likely to belong to the DMN (Assem et al. 2020).

Crucially, in our study, all AG subregions—including bilateral PGa—also showed domain-specific semantic effects, independent of their domain-general task difficulty effects. These findings strongly suggest that bilateral PGa supports not only domain-general processes, but also domain-specific semantic processes (Mattheiss et al. 2018). Indeed, while the combined analysis across all studies indicated similar semantic effects (i.e., activity differences between semantic and non-semantic conditions) for PGa and PGp, left PGa seemed to exhibit the highest sensitivity to fine-grained semantic manipulations in individual studies. For example, in the study by Graessner et al. (2021), left PGa was the only AG region that showed an activity difference between meaningful phrases (e.g., “fresh apple”) and anomalous phrases (e.g., “awake apple”) when this subtle semantic difference was task-relevant. In Kuhnke et al. (2020b), left PGa was the only AG subregion that selectively responded to action and sound features of word meaning when these were task-relevant. These results suggest that left PGa might be the AG subregion that is most relevant for semantic cognition.

This view of the left PGa is supported by Seghier (2013) who subdivided the left AG into anterior-dorsal and ventral subregions based on four functional neuroimaging studies (Sharp et al. 2009; Nelson et al. 2010; Seghier et al. 2010; Price and Ansari 2011). The anterior-dorsal

subregion shows a remarkably close and consistent correspondence with area PGa (see Figure 4 in Seghier 2013). Moreover, in all four studies, this area was highly sensitive to semantic variables. The other (more ventral) subregions were more variable across studies, both in location and in relation to semantic processing. Taken together, these previous findings and our current results support the view that left PGa constitutes a functional unit that can be functionally distinguished from its neighbors (e.g., PGp, IPS) and that is engaged in semantic cognition.

The Role of the AG in Semantics

A common view holds that the AG acts as a cross-modal convergence zone or “hub” that binds and integrates semantic features related to various sensory-motor modalities (Damasio 1989; Mesulam 1998; Binder and Desai 2011; Seghier 2013). This view is supported by the AG’s location at the junction between several sensory-motor processing streams (e.g., somatomotor, auditory, visual; Seghier 2013; Binder and Fernandino 2015; Margulies et al. 2016). Moreover, the AG shows extensive structural (Hagmann et al. 2008; Bonner and Price 2013) and functional (Tomasi and Volkow 2011; Kuhnke et al. 2021) connectivity with various sensory-motor cortices. Crucially, functional neuroimaging studies indicate that AG activity increases with the amount of semantic information that can be extracted from a given input (Binder 2016). At the level of individual concepts, AG activity is modulated by thematic associations (more/stronger > fewer/weaker associations; (Bar and Aminoff 2003), concreteness (concrete > abstract concepts; Binder et al. 2005), frequency (more > less frequent concepts; Graves et al. 2010), and familiarity (familiar > unfamiliar concepts; Woodard et al. 2007). Beyond the single-concept level, the AG is sensitive to compositionality at the phrase level (semantically related > unrelated word pairs; Price et al. 2015b; Schell et al. 2017), sentence level (semantically meaningful sentences > semantically anomalous sentences or word lists; Humphries et al. 2001; Oleser et al. 2007; Pallier et al. 2011), and narrative level (coherent stories > isolated sentences; Xu et al. 2005; Ferstl et al. 2008; Martín-Loeches et al. 2008). For instance, Humphries et al. (2006) demonstrated that left AG exhibits the weakest response to a list of unrelated words (“a ball the a the spilled librarian in sign through fire”), a stronger response to grammatical but meaningless phrases (“the freeway on a pie watched a house and a window”), an even stronger response to a list of related words (“on vacation a lost a and bag wallet a man the”), and by far the strongest response to semantically meaningful and grammatical sentences (“the man on a vacation lost a bag and a wallet”). Our current results are generally in line with these previous findings. For example, study A (Chapman and Hartwigsen 2021) shows increased AG activity for thematic associations between concepts. Study B (Graessner et al. 2021) demonstrates that the AG is

sensitive to phrase compositionality. Taken together, these results suggest that the AG integrates different semantic features into a coherent conceptual representation—be that a representation of a single object, or a more complex scene or story.

However, the role of a cross-modal hub of the semantic system is classically associated with the anterior temporal lobes (ATL) (Patterson et al. 2007; Lambon Ralph et al. 2016). In support of this view, evidence from semantic dementia (Patterson et al. 2007; Jefferies 2013), functional neuroimaging (Visser et al. 2010; Rice et al. 2015), and TMS (Pobric et al. 2010a, b) indicates a crucial role of the ATL in semantic processing across virtually all types of concepts.¹ Given the presumptive role of the ATL as a cross-modal semantic hub, proposing a similar role for the AG may seem redundant. If the ATL already acts as a cross-modal hub, what could be the function of the AG?

We propose that the AG constitutes a “multimodal” hub, whereas the ATL is an “amodal” hub. As an “amodal” hub, the ATL integrates semantic features into highly abstract representations that do not retain modality-specific information. Such an amodal representation seems necessary to explain the emergence of coherent semantic categories, which can combine superficially distinct entities (e.g., *pear* and *pineapple*) and distinguish superficially similar entities (e.g., *pear* and *lightbulb*) (Lambon Ralph et al. 2010). As a “multimodal” hub, on the other hand, the AG binds different semantic features associated with the same concept, while retaining modality-specific information. During online processing, AG could thereby enable efficient access to task-relevant semantic features (Kuhnke et al. 2020b, 2021).

Similar proposals have been put forth previously. For instance, Seghier (2013) argued that while integration and amodality have been associated with the ATL, the AG might support “first-order” integration that provides direct access to conceptual representations. Similarly, Reilly et al. (2016) proposed the AG to constitute a “low-order hub” engaged in multimodal feature binding, whereas the ATL acts as a “high-order” hub performing symbolic transformations on the bound features. During these non-linear transformations, modality-specific information is lost.

The multimodal–amodal hub theory is supported by several studies. Fernandino et al. (2016) found that AG activity during concreteness judgements correlated with the strength of sensory-motor associations for all modalities tested (action, sound, shape, color, motion). In contrast, ATL activity did not correlate with individual sensory-motor associations. In line with

¹ Although the broad term “anterior temporal lobe (ATL)” is often used to describe the semantic hub region, the precise anatomical location of the ATL hub is quite variable across studies, sometimes referring to the temporal pole, lateral portions of anterior superior, middle and inferior temporal gyri, or the anterior fusiform gyrus. Some work suggests that the most crucial cross-modal hub area is located in the anterior fusiform (Binney et al. 2010; Mion et al. 2010; Lambon Ralph et al. 2016).

these results, Kuhnke et al. (2020b) demonstrated that left AG responds to both sound and action features of concepts when these are task-relevant. Again, the ATL did not show modality-specific effects. However, the lateral ATL was engaged for abstract semantic information (i.e., words vs. pseudowords). In a follow-up study (Kuhnke et al. 2021), left AG was functionally coupled with auditory brain regions during sound feature retrieval, and with somatomotor regions during action feature retrieval. This suggests that left AG guides the retrieval of task-relevant semantic features via flexible coupling with different sensory-motor cortices. In contrast, the lateral ATL interacted with other high-level cross-modal areas, but not sensory-motor regions. Finally, TMS over left AG can selectively disrupt the retrieval of individual task-relevant semantic features (Kuhnke et al. 2020a; also see Pobric et al. 2010a; Ishibashi et al. 2011). In contrast, TMS over lateral ATL typically impairs semantic processing for all types of concepts (Pobric et al. 2010a, b).

Overall, these data suggest that AG and ATL play distinct, complementary roles during semantic processing. ATL acts as an “amodal” hub that represents an abstract similarity structure transcending individual modalities (Patterson and Lambon Ralph 2016). In contrast, AG acts as a “multimodal” hub that binds different semantic features of the same concept, enabling efficient access to task-relevant features (Seghier 2013; Reilly et al. 2016; Kuhnke et al. 2020b, 2021).

Conclusion

In conclusion, we found that AG responses are best explained by independent effects of task difficulty and semantic processing demand (Figure 3). Task difficulty effects were stronger in PGa than PGp, regardless of hemisphere. Semantic effects were stronger in left than right AG, regardless of subregion. These results indicate that the AG is involved in both domain-general task-difficulty-related processes and domain-specific semantic processes. In semantic processing, we propose that left AG acts as a “multimodal” hub which binds different semantic features associated with the same concept, enabling efficient retrieval of task-relevant features. While the right AG seems to play a weaker role under “normal” conditions in young and healthy adults, it is also sensitive to semantics and might compensate when the left AG is damaged.

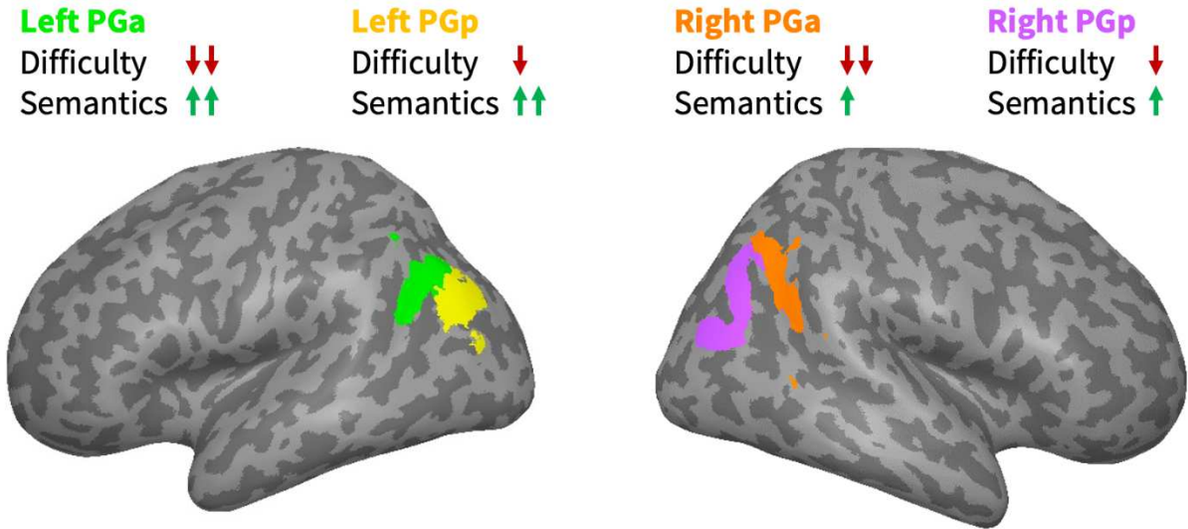


Figure 3. Overview of key findings. In all AG regions, neural activity depends on both task difficulty and semantic processing demand independently. Activity decreases with increasing difficulty, and is relatively higher for semantic than non-semantic conditions. Difficulty effects are stronger in PGa than PGp, regardless of hemisphere. Semantic effects are stronger in left than right AG, regardless of subregion.

Statements & Declarations

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Competing Interests

The authors declare no competing interests.

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Author Contributions

Philipp Kuhnke: Conceptualization, Investigation, Data curation, Formal analysis, Methodology, Writing—original draft, Writing—review and editing;

Curtiss A. Chapman: Conceptualization, Investigation, Data curation, Formal analysis, Methodology, Writing—original draft, Writing—review and editing;

Vincent K.M. Cheung: Conceptualization, Formal analysis, Methodology, Writing—review and editing;

Sabrina Turker: Investigation, Data curation;

Astrid Graessner: Investigation, Data curation;

Sandra Martin: Investigation, Data curation;

Kathleen A. Williams: Resources, Methodology;

Gesa Hartwigsen: Conceptualization, Funding acquisition, Supervision, Project administration, Writing—review and editing.

Data and Code Availability

Participant- and condition-specific behavioral data and AG response profiles, as well as analysis code for the ROI analyses and mixed models are openly available via the Open Science Framework (OSF): https://osf.io/ah5et/?view_only=54eb4120a476485ab0c72477213e9963

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