# Semantic object processing is modulated by prior scene context

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# **Abstract**

We typically encounter objects in a context, for example, a sofa in a living room or a car in the street, and this context influences how we recognize objects. Objects that are congruent with a scene context are recognised faster and more accurately than objects that are incongruent. Furthermore, objects that are incongruent with a scene elicit a stronger negativity of the N300/N400 EEG component compared to objects that are congruent with the scene. However, exactly how context modulates access to semantic object information is unknown. Here, we used a modelling-based approach with EEG to directly test how context influences the processing of semantic object information. Using representational similarity analysis, we first asked whether EEG patterns dissociated objects in congruent or incongruent scenes, finding that representational differences between the conditions emerged towards 300 ms. Next, we tested the relationship between EEG patterns and a semantic model based on property norms, revealing that the processing of semantic information for both conditions started around 150 ms, while after around 275 ms, semantic effects were stronger and lasted longer for objects in incongruent scenes compared to objects in congruent scenes. The timing of these effects overlapped with known N300/N400, suggesting previous congruency effects might be explained by differences in processing semantic object information. This suggests that scene contexts can provide a prior expectation about what kind of objects could appear, which might allow for more efficient semantic processing if the object is congruent with the scene, and extended semantic effects for incongruent objects.

Keywords: EEG; object recognition; context; semantics; RSA

## Introduction

In our daily lives, we easily recognise the objects around us. Yet, in certain situations we expect to see some objects more than others. For example, when walking down a street, we might expect to encounter a car but not an elephant. But if we visit a zoo, we would be much more likely to encounter an elephant in an enclosure than a car. In both scenarios, we recognise the object as a car and an elephant, however, the context in which we see these objects influences the way we perceive and respond to them. Objects that are congruent with their environment are recognised faster and more accurately than objects that are incongruent (Bar, 2004; Biederman et al., 1982; Davenport & Potter, 2004; Greene et al., 2015; Oliva & Torralba, 2007; Palmer, 1975). This is also reflected in neural processing, in that incongruent objects induce a later latency and a stronger negativity of the N300/N400 EEG components than congruent objects (e.g. Draschkow et al., 2018; Ganis & Kutas, 2003; Lauer et al., 2018; Lauer et al., 2020; Mudrik et al., 2010; Mudrik et al., 2014; Võ & Wolfe, 2013), which has been associated with the processing of semantic information (Kutas & Federmeier, 2011).

Much of what we do know about the semantic processing of visual objects comes from research where objects are presented isolated from the background or in a stream of unconnected events. This line of research indicates that in the first ~150 ms after the object appears, low- and middle-level object features are extracted, mostly in a feedforward fashion along the ventral visual stream (Cichy et al., 2016; DiCarlo et al., 2012; Lamme & Roelfsema, 2000). More complex visual features and semantic features are processed at later latencies, beginning after 150-200 ms, supported by recurrent dynamics within the ventral temporal lobes (Bankson et al., 2018; Chan et al., 2011; Clarke, 2019; Clarke et al., 2011, 2018; Kietzmann et al., 2019; Poch et al., 2015). In agreement with this object processing timeline, the effects of object-scene congruency on the N300/N400 EEG components occur at a similar time as semantic feature effects, which would allow for context to modulate object perception via semantic recurrent dynamics. However, the research so far does not answer the question of how semantic object information is accessed in these different situations, in terms of the timing of semantic activation. Three plausible scenarios are (1) that semantic object information is accessed faster for objects in congruent compared to incongruent environments, meaning that later processing of objects in incongruent environments leads to a N300/N400 congruency effect, (2) semantic access is initiated at the same time in both conditions, but continues for longer in the incongruent case, with the additional semantic activation related to congruency effects, or (3) that semantic access is initiated at the same time and for the same duration for both congruent and incongruent conditions, and differences in the magnitude of semantic access relate to congruency effects.

Here, we re-analysed EEG data by Draschkow and colleagues (2018) to test the hypothesis that neural effects of congruency on the N300/N400 components are driven by differences in accessing semantic object information, by combining computational models with Representational Similarity Analysis (RSA; Kriegeskorte et al., 2008) - a methodology that allows testing specific hypotheses about what object features contribute to neural signals during object processing (Bankson et al., 2018; Cichy et al., 2016; Clarke et al., 2018). For example, Clarke and colleagues (2018) demonstrated this approach using a model of semantics based on features from a property norming study (e.g. a car has the features 'has wheels', 'has a driver' and 'made of metal'), which was related to MEG signals. They reported the semantic model related to brain activity peaking around 250 ms after object onset - a latency similar to the onset of the N300/N400 component. This semantic

model, based on semantic features, provides not only the intriguing opportunity to directly test if context indeed modulates semantics, but also how it effects the temporal processing of objects in congruent and incongruent settings, adjudicating between the three scenarios we set out above. First, we used a simple congruency model that distinguishes between congruent and incongruent contexts, to establish when representational differences emerge that suggests a dissociation of processing between the conditions. Then we modelled the EEG data with a semantic model based on property norms that describes objects in terms of semantic features, to specifically test how congruency impacts the time course of processing semantic object features, and how this is different depending on contextual congruency between the object and the scene.

### Methods

We re-analysed EEG data reported by Draschkow and colleagues (2018). The data is freely available (https://github.com/DejanDraschkow/n3n4). Here we provide a short summary of the main aspects of the study design covering participants, procedure, EEG recording and pre-processing, as well as the specifications of our RSA analyses.

## Participants and procedure

Forty healthy participants viewed 152 scene images that were presented with either a semantically congruent or incongruent object (76 trials per condition). Each scene was paired with a congruent and an incongruent object, but participants saw each scene only once with either the congruent or the incongruent object (the conditions were counterbalanced across participants). At the beginning of each trial, a scene was presented for 500 ms, then a red dot appeared indicating the position where the object would appear. After 500 to 530 ms, the object was presented in the cued location of the scene and remained on the screen for 2000 ms (Fig 1, Fig 2A). The task was to report exact repetitions of scenes and objects (the repetition trials were excluded from subsequent analysis).

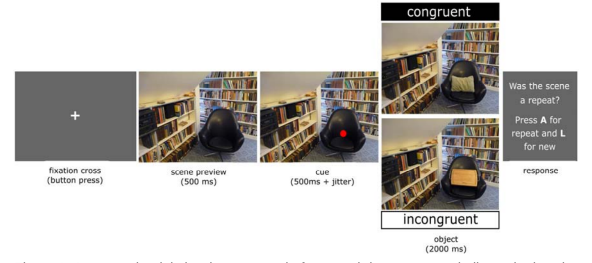


Figure 1. An example trial showing a scene before a red dot appears to indicate the location the object will appear. The object that appeared could either be congruent with the scene, in this example a cushion, or incongruent with the scene, in this example a chopping board. Each scene is only shown once to a participant, with either a congruent or incongruent object.

## EEG recording and pre-processing

EEG data was recorded with 64 active Ag/AgCl electrodes (Brain Products, GmhB), with a sampling rate of 1000 Hz. Data were down-sampled to 200 Hz, filtered between 0.1 Hz and 40 Hz, and eye and muscle artifacts were removed with independent component analysis. Epochs of 1100ms were created, from -200 ms to 900 ms centred around object onset, then baseline correction was applied from -200 ms to 0 ms. We started our analysis with the epoched data provided by Draschkow and colleagues (2018), however, we identified noisy trials by visual inspection, specifically those trials that contained high frequency noise or large amplitude signals beyond the range of normal activity. On average 2.9% of trials were removed (range 0-24%). All electrodes were included in the subsequent Representation Similarity Analysis (RSA).

## Representational Similarity Analysis

RSA was used to relate model-based congruency and semantic similarity between objects to the neural similarity based on EEG data.

#### Congruency model analysis

The consistency model contrasted objects presented in congruent and incongruent contexts. Subject-specific RDMs were constructed by first creating a vector to which we assigned a value of 1 for congruent scene-object trials and a value of 0 for incongruent scene-object trials, before calculating the Euclidean distance between each scene-object pair (Fig 2B).

From the EEG data, we extracted brain responses to each object for the time points from -200 ms to 900 ms in intervals of 5 ms, resulting in 221 time-points. Next, we calculated the correlation distance between each object pair at each time-point. This resulted in subject-specific RDMs at each time-point that summarised the neural similarity between the objects.

In the following step, we related the congruency model RDM with the brain response RDMs using Spearman correlation resulting in an RSA time-series per participant (Fig 2C). A random effects analysis assessed the model fit of the congruency model RDM and the brain RDMs at each time-point using a t-test against zero with an alpha of 0.01. To control for multiple comparisons across time we used a cluster-mass permutation test to assign p-values to clusters of significant tests (Maris & Oostenveld, 2007). For each permutation, the sign of RSA correlation time-series between the model and brain RDM was randomly flipped for each participant, before t-tests were performed on the permuted data, and the size of the largest cluster added to the permutation distribution. Finally, the cluster p-value for clusters in the original data were defined as proportion of the 10000 permutations (plus the observed cluster mass) that was greater than or equal to the observed cluster mass.

#### Semantic model analysis

The semantic model specified the semantic-feature similarity of object concepts based on a published set of property norms (Devereaux et al., 2014). The current version of the property norms is available from the Centre of Speech, Language, and the Brain (https://cslb.psychol.cam.ac.uk/propnorms). The property norms we used summarised how 826 different concepts related to 3026 different features (e.g. a zebra 'has stripes', 'eats grass' etc), allowing us to represent each concept by a collection of features that together define the concept (e.g. a zebra 'has legs, 'has stripes', but does not 'live in trees'). We matched the objects used by Draschkow and colleagues (2018) with concepts in the property norms. A matching concept was found for 118 out of 152 objects that were presented in a congruent context, and for 116 out of 152 objects that were presented in an incongruent context. Trials containing objects for which no match could be found were

excluded from further analysis. For the other trials, a semantic similarity space was defined by calculating the cosine distance between all possible pairs of objects, separately for objects that were presented in a congruent context and objects that were presented in an incongruent context, resulting in two semantic feature RDMs (Fig 2B).

The EEG data of each subject was separated for congruent and incongruent scene-object trials, before RDMs per time-point were calculated in the same way as for the congruency model, except that now two analyses were performed, one relating the semantic feature RDM to the brain RDMs for congruent trials, and one analysis relating the semantic feature RDM to the brain RDMs for the incongruent trials (Fig 2C). Significant differences between the RSA model fit for congruent and incongruent context conditions was additionally assessed with a cluster-based permutation test using paired sample t-tests.

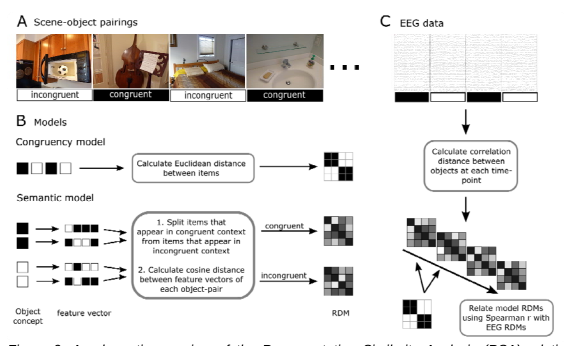


Figure 2. A schematic overview of the Representation Similarity Analysis (RSA) relating brain responses to the objects. (A) For each object an activation pattern was extracted (white represents congruent, and black represents incongruent trials). (B) Model RDMs. For the congruency model analysis all congruent object trials were assigned a value of 1 and all incongruent object trials a value of 0. Then the Euclidean distance between all object-pairs was calculated. The resulting RDM directly contrasts congruent and incongruent objects. For the semantic model analysis the congruent and incongruent object trials were analysed separately. Each object was assigned a corresponding concept feature vector. Then the cosine distance was calculated for all object-pairs, separately for consistent and inconsistent trials, resulting in RDMs that describe the similarity of objects based on semantics. (C) Relating models to brain signals. For each object the EEG response was extracted, and at each time-point the similarity between object pairs was calculated using correlation distance. Model RDMs were then related to EEG RDMs using Spearman correlation.

# Results

We combined computational modelling with RSA to test if congruency effects in N300/N400 EEG components were driven by semantic object information. First, we constructed a model of consistency to uncover when the processing of congruent and incongruent objects diverged. Then, using a model based on semantic features, we investigated the time-course of semantic processing of objects that were presented in either congruent or incongruent contexts.

#### Congruency model analysis

We first assessed whether neural patterns distinguished between objects presented in congruent and incongruent environments. RSA analysis of the EEG signals revealed that the congruency model distinguished between congruent and incongruent scene-object context trials, where we saw a significant relationship between the congruency model and EEG patterns from approximately 290 to 450 ms (cluster p = 0.022; Fig 3A, Table 1). The timing of this effect is in line with previous N300/N400 effects of congruency.

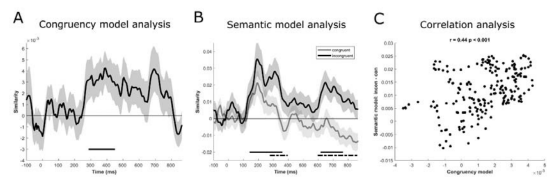


Figure 3. RSA results. (A) The consistency model fit shows the similarity based on Spearman correlation between the model RDM and the EEG RDMs at each time-point. Shaded area shows +- 1 standard error of the mean. The horizontal bar shows a statistically significant cluster. (B) The semantic model fit is shown separately for congruent (grey line) and incongruent (black line) conditions. The horizontal bars show statistically significant clusters for incongruent objects (black solid line) and the difference of semantic model fit between congruent and incongruent objects (black dotted line). (C) Correlational analysis relating the congruency model to the difference of semantic model fit between congruent and incongruent objects.

Table 1. EEG RSA effects

Model RDM	time-window	cluster-mass	cluster p-value
Congruency model	286-450 ms	74.26	p = .022
Semantic model: consistent	141-235 ms	46.11	p = .064
Semantic model: inconsistent	141-360 ms	132.25	p = .005
Semantic model: inconsistent	616-765 ms	80.78	p = .021
Semantic model: incon > con	276-395 ms	50.25	p = .049
Semantic model: incon > con	596-870 ms	169.26	p = .002

#### Semantic model analysis

While the congruency effect shows that object processing and scene information interact, it does not tell us about how semantic knowledge may be accessed differentially depending on the scene context. To address this, we constructed a semantic model based on semantic features that collectively describe each of the concepts. Two separate RDMs were created, one for objects that were presented in a congruent context and one for objects that were presented in an incongruent context. The model RDMs were then correlated with the corresponding brain RDMs, i.e. the congruent model RDM was related to brain activation RDMs to congruent objects and the incongruent model RDM to brain activation RDMs to incongruent objects.

For congruent objects, although the semantic model showed no statistically significant relationship to brain activity, it numerically performed well from around 140 ms to 235 ms (cluster p = 0.06). For incongruent objects, the semantic model fitted the brain responses significantly during two clusters, the first cluster including time-points around 140 ms to 360 ms (cluster p = 0.005) and a second cluster with time-points between around 620 to 765 ms (cluster p = 0.021; Fig 3B, Table 1). This might suggest that while semantic effects in both conditions seem to begin at similar times, approximately 150 ms after object onset, semantic effects for incongruent objects continue for a longer period of time. In order to test this, we compared the two conditions directly. The difference in model-fit between the two conditions was significant at two clusters, an early cluster including the time-points around 280 ms to 395 ms (cluster p = 0.049) and a second cluster including the time-points from approximately 600 ms to 870 ms (cluster p = 0.002; Fig 3B, Table 1). While cluster-based permutation testing does not allow precise estimation of effect on- and off-sets (Sassanhagen & Draschkow, 2019), qualitatively the time-windows of these clusters overlap with both our effects of the congruency model RDM, and known congruency effects in the N300/N400 (e.g. Draschkow et al., 2018; Ganis & Kutas, 2003; Lauer et al., 2018; Lauer et al., 2020; Mudrik et al., 2010; Mudrik et al., 2014; Võ & Wolfe, 2013), in addition to a regularly reported later effect coinciding with the P600 (e.g. Ganis & Kutas, 2003; Mudrik et al., 2010; Sauvé et al., 2017; Võ & Wolfe, 2013).

#### Correlation analysis

The model-based analysis revealed overlap between the congruency and the semantic model RDMs in the time-window from approximately 290 to 395 ms. Within this time-window, the congruency model successfully distinguished if an object was presented in congruent or incongruent context, and the semantic model displayed significantly better model fit with objects that were presented in an incongruent context compared to objects that were presented in a congruent context. In order to test if these two effects were related, we correlated the time courses of the congruency model fit with the time-course of the difference between the semantic model fit for the congruent and incongruent conditions. The results confirm a significant correlation (r = 0.44, p < 0.001; Fig 3C), demonstrating that the two analyses could be capturing the same temporal effect of congruency, which might suggest that effects of congruency are explained by differences in the processing of semantic object features.

#### Discussion

In the current study, we directly tested if scene context influenced object recognition through the modulation of processing semantic object information. We related the similarity

based on EEG activity in response to visual objects with both a model of congruency and a semantic model that was based on semantic object property norms. Both the congruency model and semantic model captured an effect of scene context on object processing in the time-window for which N300/N400 effects were previously reported (e.g. Draschkow et al., 2018; Ganis & Kutas, 2003; Lauer et al., 2018; Lauer et al., 2020; Mudrik et al., 2010; Mudrik et al., 2014; Võ & Wolfe, 2013). Additionally, the semantic model revealed a difference in processing of congruent and incongruent objects in a later time-window beyond ~ 600 ms, which has been reported in some previous studies (e.g. Ganis & Kutas, 2003; Mudrik et al., 2010; Sauvé et al., 2017. Võ & Wolfe, 2013). In these two time-windows, the semantic model displayed stronger fit for incongruent than for congruent objects, suggesting that the previously observed congruency effects were driven by the additional need for semantic processing of objects that were incongruent with their environment. This contrasts with alternative possibilities that semantic object information could have been accessed faster for objects in congruent compared to incongruent environments, or that semantic access was initiated at the same time and for the same duration for both congruent and incongruent conditions.

Our research is the first to employ a modelling-based approach to directly test the hypothesis that scene context influences object recognition by modulating the processing of semantic object information. The stronger model fit for incongruent objects beyond about 275 ms suggests that while both congruent and incongruent objects involve semantic processing beyond ~150 ms, semantic processes are extended in the incongruent condition. It may well be that this extended semantic processing for incongruent objects is what underpins the congruency effect, which seems to begin at a similar time to the divergence of semantic model fits across the two conditions.

Overall, our findings are in agreement with the framework that context generates expectations about objects we might encounter, and thereby affects the way objects are processed (Clarke, 2019). Our results, together with previous findings of congruency effects on N300/N400 EEG components, show stronger effects for objects that were unexpected compared to objects that were expected. This phenomenon is consistent with the predictive coding account (Friston, 2005) which states that the brain constructs prior expectations about upcoming sensory events based on experience, and generates an error response if the event does not match the expectation. In terms of scene-object congruency, exposure to a scene context could create a prediction about what objects are likely to appear in that scene. This is even more so here, as a fixation dot appeared prior to the object indicating the location the item would appear, thus limiting the range of likely object candidates. If the object is not congruent with the scene, and hence does not fit the prediction, it triggers a prediction error response causing delayed or enhancement of brain activity that is related to object processing, like we see for the N300/N400 EEG components (e.g. Draschkow et al., 2018; Ganis & Kutas, 2003; Lauer et al., 2018; Lauer et al., 2020; Mudrik et al., 2010; Mudrik et al., 2014; Võ & Wolfe, 2013). Similar effects of context on object recognition have been reported not only for scenes but also for other types of prior information like the presence of other objects (Auckland et al., 2007; Kovalenko et al., 2012; McPherson & Holcomb, 1999), and has been demonstrated in semantic priming studies (Renoult et al., 2012). This indicates that the semantic effects we see here reflect a more general mechanism that is not restricted to scenes. Taken together, having a prior expectation about object semantics might allow for more efficient semantic processing, with the effect that we see rapid and short semantic effects for congruent objects, and extended semantic effects for incongruent objects.

In their original work, Draschkow and colleagues (2018) demonstrated how congruency effects influencing the N300 and N400 components constitute highly related processes which allow the decoding of congruency across the two time-windows, finding significant

cross-decoding of congruency from about 200 ms after object onset. The consistency model analysis that we employed, likewise tested to distinguish between objects that were presented either in congruent or incongruent context. Our results highlight a similar time-window like the decoding analysis, thus confirming that a model-based approach is suitable to capture congruency effects in EEG data.

In addition to the congruency effects in the N300/N400 time-window, the semantic model analysis revealed a difference in processing of congruent and incongruent objects in a later time-window beyond ~600 ms. Effects in this time window were previously reported in similar studies, in which objects were embedded into scenes (e.g. Ganis & Kutas, 2003; Mudrik et al., 2010; Sauvé et al., 2017; Võ & Wolfe, 2013). However, the exact nature of these later effects remains unclear as they vary depending on task-demands (Ganis & Kutas, 2003; Võ & Wolfe, 2013; Sauvé et al., 2017). Võ and Wolfe (2013) reported a P600 component specifically in the context of mild syntactic violations of an object's position in scenes (object misplaced), but not for extreme syntactic violations (object in impossible position, for example in the air). One possible explanation for finding these effects in the current data is that objects were embedded in scenes, and this has likely induced a combination of semantic and mild syntactic violations in incongruent scene-object trials. For example, if a ball is embedded in a photo of a kitchen and is placed inside a microwave, then in addition to the semantic incongruency, a mild syntactic violation is also created. This combination of semantic and syntactic violations might explain RSA effects of semantic object processing in incongruent scenes during a similar time window to the previously reported P600 congruency effects.

In conclusion, our results revealed that while semantic processing begins around 150 ms after the object appears, the modulatory effect of the prior scene context starts around the onset of the N300 components, resulting in longer processing of objects that are incongruent with a scene compared to objects that are congruent. Additionally, we replicated effects in previously reported time-windows of the N300/N400 and P600 EEG components using a computational modelling approach. Importantly, our study highlights how object recognition processes are flexibly adapted based on prior information, in this case showing the dynamics associated with accessing semantic knowledge are modulated by the prior context.

# References

- Auckland, M. E., Cave, K. R., & Donnelly, N. (2007). Nontarget objects can influence perceptual processes during object recognition. *Psychonomic Bulletin and Review*, 14(2), 332–337. https://doi.org/10.3758/BF03194073
- Bankson, B. B., Hebart, M. N., Groen, I. I. A., & Baker, C. I. (2018). The temporal evolution of conceptual object representations revealed through models of behavior, semantics and deep neural networks. *NeuroImage*, 178(May), 172–182. https://doi.org/10.1016/j.neuroimage.2018.05.037
- Bar, M. (2004). Visual objects in context. *Nature Reviews Neuroscience*, *5*(8), 617–629. https://doi.org/10.1038/nrn1476
- Biederman, I., Mezzanotte, R. J., & Rabinowitz, J. C. (1982). Scene perception: Detecting and judging objects undergoing relational violations. *Cognitive Psychology*, *14*(2), 143–177. https://doi.org/10.1016/0010-0285(82)90007-X

- Chan, A. M., Baker, J. M., Eskandar, E., Schomer, D., Ulbert, I., Marinkovic, K., Cash, S. S., & Halgren, E. (2011). First-pass selectivity for semantic categories in human anteroventral temporal lobe. *Journal of Neuroscience*, *31*(49), 18119–18129. https://doi.org/10.1523/JNEUROSCI.3122-11.2011
- Cichy, R. M., Khosla, A., Pantazis, D., Torralba, A., & Oliva, A. (2016). Comparison of deep neural networks to spatio-temporal cortical dynamics of human visual object recognition reveals hierarchical correspondence. *Scientific Reports*, 6(January), 1–13. https://doi.org/10.1038/srep27755
- Clarke, A. (2019). Neural dynamics of visual and semantic object processing. In *Psychology of Learning and Motivation Advances in Research and Theory* (1st ed., Vol. 70). https://doi.org/10.1016/bs.plm.2019.03.002
- Clarke, A., Devereux, B. J., & Tyler, L. K. (2018). Oscillatory Dynamics of Perceptual to Conceptual Transformations in the Ventral Visual Pathway. *Journal of Cognitive Neuroscience*, 10. https://doi.org/10.1162/jocn
- Clarke, A., Taylor, K. I., & Tyler, L. K. (2011). The evolution of meaning: Spatio-temporal dynamics of visual object recognition. *Journal of Cognitive Neuroscience*, 23(8), 1887–1899. https://doi.org/10.1162/jocn.2010.21544
- Devereux, B. J., Tyler, L. K., Geertzen, J., & Randall, B. (2014). The Centre for Speech, Language and the Brain (CSLB) concept property norms. *Behavior Research Methods*, 46(4), 1119–1127. https://doi.org/10.3758/s13428-013-0420-4
- Davenport, J. L., & Potter, M. C. (2004). Scene consistency in object and background perception. *Psychological Science*, 15(8), 559–564. https://doi.org/10.1111/j.0956-7976.2004.00719.x
- DiCarlo, J. J., Zoccolan, D., & Rust, N. C. (2012). How does the brain solve visual object recognition? *Neuron*, 73(3), 415–434. https://doi.org/10.1016/j.neuron.2012.01.010
- Draschkow, D., Heikel, E., Võ, M. L. H., Fiebach, C. J., & Sassenhagen, J. (2018). No evidence from MVPA for different processes underlying the N300 and N400 incongruity effects in object-scene processing. *Neuropsychologia*, 120(July), 9–17. https://doi.org/10.1016/j.neuropsychologia.2018.09.016
- Friston, K. (2005). A theory of cortical responses. *Philosophical Transactions of the Royal Society B: Biological Sciences*, *360*(1456), 815–836. https://doi.org/10.1098/rstb.2005.1622
- Ganis, G., & Kutas, M. (2003). An electrophysiological study of scene effects on object identification. *Cognitive Brain Research*, *16*(2), 123–144. https://doi.org/10.1016/S0926-6410(02)00244-6
- Greene, M. R., Botros, A. P., Beck, D. M., & Fei-Fei, L. (2015). What you see is what you expect: rapid scene understanding benefits from prior experience. *Attention, Perception, and Psychophysics*, 77(4), 1239–1251. https://doi.org/10.3758/s13414-015-0859-8
- Kietzmann, T. C., Spoerer, C. J., Sörensen, L. K. A., Cichy, R. M., Hauk, O., & Kriegeskorte, N. (2019). Recurrence is required to capture the representational dynamics of the human visual system. *Proceedings of the National Academy of Sciences of the United States of America*, *116*(43), 21854–21863. https://doi.org/10.1073/pnas.1905544116

- Kovalenko, L. Y., Chaumon, M., & Busch, N. A. (2012). A pool of pairs of related objects (POPORO) for investigating visual semantic integration: Behavioral and electrophysiological validation. *Brain Topography*, 25(3), 272–284. https://doi.org/10.1007/s10548-011-0216-8
- Kriegeskorte, N. (2008). Representational similarity analysis connecting the branches of systems neuroscience. *Frontiers in Systems Neuroscience*, 2(November), 1–28. https://doi.org/10.3389/neuro.06.004.2008
- Kutas, M., & Federmeier, K. D. (2011). Thirty years and counting: Finding meaning in the N400 component of the event-related brain potential (ERP). *Annual Review of Psychology*, 62, 621–647. https://doi.org/10.1146/annurev.psych.093008.131123
- Lamme, V. A. F., & Roelfsema, P. R. (2000). The distinct modes of vision offered by feedforward and recurrent processing. *Trends Neurosci.*, 23, 571–579. https://doi.org/https://doi.org/10.1016/S0166-2236(00)01657-X
- Maris, E., & Oostenveld, R. (2007). Nonparametric statistical testing of EEG- and MEG data. Journal of Neuroscience Methods, 164(1), 177–190. https://doi.org/10.1016/j.jneumeth.2007.03.024
- McPherson, W. B., & Holcomb, P. J. (1999). An electrophysiological investigation of semantic priming with pictures of real objects. *Psychophysiology*, *36*(1), 53–65. https://doi.org/10.1017/S0048577299971196
- Mudrik, L., Lamy, D., & Deouell, L. Y. (2010). ERP evidence for context congruity effects during simultaneous object-scene processing. *Neuropsychologia*, *48*(2), 507–517. https://doi.org/10.1016/j.neuropsychologia.2009.10.011
- Mudrik, L., Shalgi, S., Lamy, D., & Deouell, L. Y. (2014). Synchronous contextual irregularities affect early scene processing: Replication and extension. Neuropsychologia, 56(1), 447–458. https://doi.org/10.1016/j.neuropsychologia.2014.02.020
- Lauer, T., Cornelissen, T. H. W., Draschkow, D., Willenbockel, V., & Võ, M. L. H. (2018). The role of scene summary statistics in object recognition. *Scientific Reports*, 8(1), 1–12. https://doi.org/10.1038/s41598-018-32991-1
- Lauer, T., Willenbockel, V., Maffongelli, L., & Võ, M. L. H. (2020). The influence of scene and object orientation on the scene consistency effect. *Behavioural Brain Research*, 394(July). https://doi.org/10.1016/j.bbr.2020.112812
- Oliva, A., & Torralba, A. (2007). The role of context in object recognition. *Trends in Cognitive Sciences*, 11(12), 520–527. https://doi.org/10.1016/j.tics.2007.09.009
- Palmer, S. E. (1975). The effects of contextual scenes on the identification of objects. *Memory & Cognition*, *3*(5), 519–526. https://doi.org/10.3758/BF03197524
- Poch, C., Garrido, M. I., Igoa, J. M., Belinchón, M., García-Morales, I., & Campo, P. (2015). Time-varying effective connectivity during visual object naming as a function of semantic demands. *Journal of Neuroscience*, *35*(23), 8768–8776. https://doi.org/10.1523/JNEUROSCI.4888-14.2015

- Renoult, L., Wang, X., Calcagno, V., Prévost, M., & Debruille, J. B. (2012). From N400 to N300: Variations in the timing of semantic processing with repetition. *NeuroImage*, 61(1), 206–215. https://doi.org/10.1016/j.neuroimage.2012.02.069
- Sassenhagen, J., & Draschkow, D. (2019). Cluster-based permutation tests of MEG/EEG data do not establish significance of effect latency or location. *Psychophysiology*, *56(6)*, e13335.
- Sauvé, G., Harmand, M., Vanni, L., & Brodeur, M. B. (2017). The probability of object–scene co-occurrence influences object identification processes. *Experimental Brain Research*, 235(7), 2167–2179. https://doi.org/10.1007/s00221-017-4955-y
- Võ, M. L. H., & Wolfe, J. M. (2013). Differential Electrophysiological Signatures of Semantic and Syntactic Scene Processing. *Psychological Science*, 24(9), 1816–1823. https://doi.org/10.1177/0956797613476955