Predictive coding across the left fronto-temporal hierarchy during language comprehension

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We used magnetoencephalography (MEG) and event-related potentials (ERPs) to track the time-course and localization of evoked activity produced by expected, unexpected plausible, and implausible words during incremental language comprehension. We suggest that the full pattern of results can be explained within a hierarchical predictive coding framework in which increased evoked activity reflects the activation of residual information that was not already represented at a given level of the fronto-temporal hierarchy ("error" activity). Between 300 and 500 ms, the three conditions produced progressively larger responses within left temporal cortex (lexico-semantic activation of residual information that was not already represented at a given level of the fronto-temporal hierarchy ("error" activity). Decrease in these responses between 500 and 1000 ms indicates that the posterior part of the left temporal cortex (orthographic prediction error) is suppressed. Between 200 and 500 ms, the mid- and anterior temporal cortices are activated in parallel, particularly over the left posterior fusiform, posterior inferior frontal, and anterior lateral temporal cortices (feedback activity that produced top-down error), whereas highly implausible inputs activated left inferior frontal cortex, posterior fusiform (unsuppressed orthographic prediction error/reprocessing), and medial temporal cortex (possibly supporting new learning). Therefore, predictive coding may provide a unifying theory that links language comprehension to other domains of cognition.

Key words: ERP; hierarchy; MEG; prediction; probabilistic.

Introduction

One of the most amazing feats of human cognition is our ability to comprehend language by transforming streams of linguistic input into a high-level representation of real-world events (an "event model"; Radwan and Zacks 2011). Language comprehension can be understood as a process of "probabilistic inference": The use of prior knowledge, encoded within an internal generative model, to infer the underlying high-level representation that best "explains" the bottom-up input (Kuperberg and Jaeger 2016). According to an influential theory of brain function, probabilistic inference is approximated using an optimization algorithm known as "predictive coding" (Mumford 1992; Rao and Ballard 1999; Friston 2005; Clark 2013; Spratling 2017). Predictive coding claims that higher levels of the cortical hierarchy, representing information over longer spatiotemporal scales, continually generate top-down predictions of activity at lower cortical levels, which represents information over shorter spatiotemporal scales. As new bottom-up information becomes available to lower levels, top-down predictions attempt to "suppress" the activity produced by these inputs. Any residual (unexplained) information encoded in the input, i.e. "prediction error," is then used to update higher-level representations, allowing them to produce more accurate predictions. Inference is complete when, over multiple iterations of the algorithm, prediction error is minimized across all levels of the cortical hierarchy.

In the language domain, studies of speech perception (Blank and Davis 2016; Sohoglu and Davis 2020) and visual word-recognition (Price and Devlin 2011; Heilbron et al. 2020b) have shown that predictable inputs produce less neural activity than unpredictable inputs in regions that encode low-level phonological and orthographic information. This has been taken as evidence for the suppression of lower-level prediction error by accurate top-down predictions. An important open question is whether the principles of hierarchical predictive coding can also account for the neuroanatomical localization and timing of evoked neural activity produced during higher-level language comprehension, which requires us to infer information represented at multiple time-scales as new linguistic inputs unfold in real time.

300–500 ms

MEG studies have shown that single words, presented without any prior context, produce robust event-related (evoked) responses: Increases in time-locked neural activity, relative to baseline. These evoked responses begin in early perceptual regions, and are followed by subsequent peaks as activity rapidly flows from posterior to anterior regions of ventral and lateral temporal cortices, and then to inferior frontal and orbitofrontal cortices. Between 300 and 500 ms, the mid- and anterior temporal and inferior frontal cortices are activated in parallel, particularly over the left hemisphere (e.g. Dale et al. 2000; Dhond et al., 2001; Haigren et al. 2002; Marinkovic et al. 2003). This feedforward sweep of activity propagates to the scalp surface, manifesting in both EEG and MEG as the N400 waveform, which is thought to reflect the initial influence of the incoming stimulus on the current state of semantic memory (Kutas and Federmeier 2011).
A large body of studies using event-related potentials (ERPs) has established that an N400 is triggered by each meaningful content word in a sentence (Van Petten & Kutas 1990, 1991; Payne et al. 2015). During sentence comprehension, the amplitude of the N400 is highly sensitive to the lexical predictability of an incoming word in relation to its prior context (Kutas and Hillyard 1984; DeLong et al. 2005; Federmeier et al. 2007), with expected words evoking a smaller N400 than unexpected but plausible words (e.g. “For the sleepover, you should bring a blanket and a pillow/game”). There is also evidence for an effect of contextual plausibility on the N400, over and above the effect of lexical predictability (Kuperberg et al. 2020; Nieuwland et al. 2020), with unexpected plausible words evoking a smaller N400 than implausible words (e.g. “For the sleepover, you should bring a blanket and a game/sneeze”). However, it remains unclear which regions in the fronto-temporal hierarchy contribute to the effects of lexical predictability and contextual plausibility. Functional magnetic resonance imaging (fMRI) studies have had difficulty addressing this question because the sluggish hemodynamic response is largely insensitive to transient, feedforward neural activity (Furey et al. 2006), like that reflected by the N400 (Lau et al. 2008; Geukes et al. 2013; Lau et al. 2013, 2016). Although previous MEG and intracranial EEG studies have shown that the effects of sentence context on the N400 can localize to both temporal and inferior frontal cortices (e.g. Helenium et al. 1998; Halgren et al. 2002; Maess et al. 2006; Ibara et al. 2007; Heilbron et al. 2020a), none of these previous studies have separately manipulated lexical predictability and contextual plausibility. Determining which levels of the fronto-temporal hierarchy are modulated by these two factors is important because it can help distinguish between different neurobiological frameworks of language comprehension, including predictive coding.

One possibility is that the effects of lexical predictability and contextual plausibility on the N400 both localize to temporal and inferior frontal regions, with graded increases in activity across expected, unexpected plausible, and implausible words. This would be consistent with a recent computational model (Rabovsky et al. 2018) that interprets the N400 as an implicit update in a single hidden layer that maps directly from lexical inputs to an event model. Assuming that this single state is distributed across both frontal and temporal cortices, the more expected the new input, the smaller the update, and the smaller the evoked response it should produce across these two regions. We will refer to this as a “distributed state” account (see Fig. 1A for summary).

A second possibility is that, in plausible sentences, the N400 effect of lexical predictability localizes to regions of the left temporal cortex that support lexical processing (i.e. expected < unexpected = implausible), whereas, in unpredictable sentences, the N400 effect of contextual plausibility localizes to the inferior frontal cortex (i.e. expected = unexpected < implausible). This account is motivated by theories that draw a sharp distinction between lexico-semantic access (Lau et al. 2008, 2009) and event-level integration (Brothers et al. 2015; Nieuwland et al. 2020). Unlike the “distributed state” account, it assumes a hierarchical architecture in which semantic information associated with individual words (lexico-semantic information) is encoded at a shorter time-scale at lower levels of the fronto-temporal hierarchy (temporal cortex; Price 2012), whereas a full representation of the prior context is encoded and maintained over a longer time span within higher-level regions (e.g. the prefrontal cortex; Lerner et al. 2011). According to this framework, in plausible sentences, it should be easier to access lower-level lexico-semantic representations of expected than unexpected plausible words because prior lexico-semantic predictions would have already pre-activated these representations within the left temporal cortex (see Wang et al. 2018). This would be consistent with MEG and intracranial studies showing that the semantic priming effect localizes to regions within the left temporal cortex (Nobre and McCarthy 1995; Lau et al. 2013; Lau and Nguyen 2015), and that this facilitation effect is enhanced when top-down prediction is encouraged (Lau et al. 2016). In contrast, according to this account, the effect of contextual implausibility is not driven by differences in prior lexico-semantic prediction. Instead, inferring an implausible event leads to additional difficulties in integrating it with real-world knowledge, with this integration being mediated by the left inferior frontal cortex (Hagoort et al. 2004). We will refer to this as the “prediction-integration” account (see Fig. 1B for summary).

A third possibility is that the effects of lexical predictability and contextual plausibility on the N400 localize only to temporal regions (expected < unexpected plausible < implausible). This account also assumes a hierarchical architecture, with the N400 reflecting activity at the lower lexico-semantic level. In contrast to the “prediction-integration” account, however, it assumes that comprehenders not only predict semantic features associated with specific lexical items (lexico-semantic predictions), but also semantic features that are associated with whole semantic categories (e.g. <animate> vs. <inanimate>). When new input becomes available, these semantic-level predictions will also serve to facilitate the retrieval/access of unexpected plausible (vs. implausible) lexico-semantic representations. Therefore, on this account, all contextual effects on incoming words—both lexical predictability and contextual plausibility—are hypothesized to “bottom-out” at the lower lexico-semantic level of representation (MacDonald et al. 1994; Smith and Levy 2013). Updates of higher-level states, which are maintained over a longer time-scale, are assumed not to influence the N400, although the precise reasons for this are not usually specified. We will refer to this account as the “lexico-semantic facilitation” account (see Fig. 1C for summary).

Finally, hierarchical predictive coding predicts a fourth possible pattern. Graded N400 modulation across the three conditions within temporal regions, but nongraded modulation within the inferior frontal cortex (see Fig 1D). Similar to the “prediction-integration” and the “lexico-semantic” facilitation accounts, predictive coding assumes a hierarchical architecture. Here, we assume that, during language comprehension, lexico-semantic information associated with individual words is encoded at a shorter time-scale within the temporal cortex, and that a higher-level event model is represented over a longer time-scale within the prefrontal cortex.

Within the temporal cortex, predictive coding posits that the prior context generates predictions that pre-activate semantic features, both of expected individual words and whole semantic categories, similar to the “lexico-semantic facilitation” account. Therefore, predictive coding also predicts progressively larger N400 responses across the three conditions within the temporal cortex (expected < unexpected plausible < implausible). However, instead of directly attributing these evoked effects to changes induced by the input in lexico-semantic states, predictive coding attributes them to differences in the magnitude of lexico-semantic prediction error—activity produced by a distinct population of lexico-semantic “error units” that cannot be suppressed/explained by top-down predictions from the higher-level event model (see Fig 2, left, for computational simulations see Nour Eddine 2021; Nour Eddine et al. in press).
Within the inferior frontal cortex, predictive coding attributes differences in amplitude of the N400 to differences in the magnitude of prediction error produced at the higher level of the event model, i.e. activity within higher-level error units that is not suppressed by predictions based on real-world knowledge (see Fig. 2, right). Within this framework, there should be no difference in the magnitude of higher-level prediction error produced by expected and unexpected plausible inputs because, in both cases, the event model is plausible (i.e. consistent with real-world knowledge), and so any higher-level prediction error/evoked response is suppressed. However, in the case of implausible inputs, higher-level prediction error is no longer suppressed (see Rao and Ballard 1999). Therefore, predictive coding predicts an enhanced N400 response in the inferior frontal cortex only to implausible inputs (expected = unexpected plausible < implausible).

### 600–1,000 ms

Beyond the N400 time-window, previous ERP studies have shown that unpredicted inputs can also produce additional evoked responses between 600 and 1,000 ms, particularly in rich, schema-constraining contexts. This later-stage activity manifests at the scalp surface as a set of positive-going waveforms between 600 and 1,000 ms, with different scalp distributions depending on whether the interpretation is plausible or implausible (Van Petten and Luka 2012; DeLong et al. 2014; Brothers et al. 2020; Kuperberg et al. 2020). Unexpected words that yield plausible interpretations produce a late frontally-distributed positivity (Federmeier et al. 2007; Brothers et al. 2020; Kuperberg et al. 2020), whereas anomalous inputs that yield highly implausible interpretations produce a late posteriorly-distributed positivity, also known as the semantic P600 (Kuperberg et al. 2003; Kuperberg 2007; van de Meerendonk et al. 2009; Brothers et al. 2020; Kuperberg et al. 2020).

Various theoretical proposals have been offered to explain these late ERP responses. For example, it has been proposed that the late frontal positivity reflects lexical inhibition (Ness and Meltzer-Asscher 2018), or a large late shift of the event model (Brothers et al. 2020; Kuperberg et al. 2020), whereas the late posterior positivity/P600 has been linked to reprocessing of the perceptual input that is triggered by linguistic conflict (van de Meerendonk et al. 2009; Brothers et al. 2020; Kuperberg et al. 2020; Brothers et al. 2021). However, we know very little about the neuroanatomical sources of this later-stage evoked activity. Although fMRI is more sensitive to later-stage feedback activity than earlier stimulus-driven feedforward transient responses (Furey et al. 2006; like the N400), the sluggish hemodynamic response also captures much later activity that extends beyond 1,000 ms, and that is likely to reflect processing of subsequent words and offline reflections about the meaning of a sentence as a whole. Even though several MEG and intracranial studies have explored the effects of semantic context during sentence comprehension, they have generally failed to report effects beyond the N400 time-window.

Hierarchical predictive coding provides a biologically-motivated computational framework for understanding these late evoked responses. Within a nonstationary hierarchical generative framework (cf. Qian et al. 2012; Gershman et al. 2014), new information that is inconsistent with the comprehender’s prior event model will induce attempts to retrieve new schema from long-term memory (Franklin et al. 2020; Kuperberg 2021). In the case of an unexpected plausible input, this retrieval will be successful, resulting in the generation of new top-down predictions. Predictive coding posits that, as these new predictions flow down the cortical hierarchy between 600 and 1,000 ms, they will produce top-down error, i.e. they will activate error units that carry residual top-down information that is not already encoded.
This study

Here, we used MEG and ERPs to track the time-course and localization of evoked neural activity, produced in the first 1,000 ms following word onset, during incremental language comprehension. Participants read predictable multi-sentence discourse contexts, e.g. “The lifeguards received a report of sharks right near the beach. Their immediate concern was to prevent any incidents in the sea. Hence, they cautioned the...” Critical words in the final sentence were (i) expected (e.g. “swimmers”), (ii) unexpected but plausible (e.g. “trainees”), or (iii) highly implausible (e.g. “drawers”).3 To examine the full time-course and neuroanatomical localization of the evoked responses produced by each condition, we carried out a distributed source-localization analysis of the MEG data, which is relatively undistorted by conductivities of the skull and scalp. To test our a priori hypotheses, we carried out planned comparisons between each pair of conditions across a left-lateralized search region where we expected effects to be maximal (particularly the effects of lexical predictability, see Federmeier 2022), and within three 200 ms time-windows of interest (300–500 ms, corresponding to the N400 time-window, and 600–800 ms and 800–1,000 ms, corresponding to the first and second halves of the time-window associated with the late positivity ERP effects). We also report activity at every 100 ms across both hemispheres (left hemisphere findings are shown in the Results section, and right hemisphere results are shown in Supplementary Materials).

Fig. 2. Between 300 and 500 ms (N400): Proposed predictive coding across the left fronto-temporal hierarchy. In rich, constraining discourse contexts, top-down predictions, based on stored real-world and linguistic knowledge, are propagated down the cortical hierarchy (faint blue diagonal arrows) before new bottom-up input becomes available. Lexico-semantic level (pink): Within regions of the left temporal cortex that support lexico-semantic processing, the lexico-semantic representation of the incoming word is inferred throughout the N400 time-window within “state units” (ST_i−1 → ST_i). Within a distinct set of “error units”, residual lexico-semantic information that is not suppressed (cannot be explained) by the top-down predictions (PE = max(0, ST−Pr), simplified in the figure as PE = ST−Pr), produces lexico-semantic prediction error, which manifests as the evoked N400 response. Lexico-semantic prediction error, and evoked activity within left temporal cortex is therefore smallest to expected words (e.g. “swimmers,” not shown), larger to unexpected plausible words (e.g. “trainees,” left), where some semantic features (e.g. <animate>) were predicted, and largest to implausible words (e.g. “drawers,” right) where no semantic features were predicted. Event model (green): Throughout the N400 time-window, lexico-semantic prediction error passes up to left inferior frontal cortex (red arrows), where it induces updates of the higher-level event model. Unexpected plausible (left): These updates yield a plausible high-level event model that is congruent with predictions about the possibility/plausibility of real-world events. Therefore, higher-level prediction error is suppressed, and the evoked response within left inferior frontal cortex is no larger than to expected inputs. In addition, the newly-updated event model (ST_i) now generates correct top-down predictions (blue curly arrow) that “switch off”/suppress lower-level lexico-semantic prediction error, leading to the reduction of the evoked response within left temporal cortex at the end of the N400 time-window. Implausible right): Updates of the event model yield a highly implausible interpretation (e.g. “lifeguards cautioned drawers”) that cannot be explained by real-world knowledge predictions, resulting in a larger higher-level prediction error at the level of the event model, and an enhanced evoked response within left inferior frontal cortex (red arrow). Moreover, because it is more difficult to converge on an implausible interpretation, top-down lexico-semantic predictions will be less accurate and less likely to suppress lower-level lexico-semantic prediction error (blue dotted curly arrow), further enhancing and prolonging the evoked response within left temporal cortex.

within prior states (see Rao and Ballard 1997, 1999), both at the level of the event model and at the lexico-semantic level. This framework therefore predicts that, relative to expected inputs, unexpected plausible inputs will produce late evoked effects both within inferior frontal and temporal cortices (see Fig. 3, left).

In contrast, in the case of a highly implausible input that yields an anomalous interpretation (e.g. “...they cautioned the “drawers”), the brain is unable to retrieve new schema, and it will therefore continue to generate incorrect top-down predictions. This will result in a failure to switch off prediction error at still lower levels of the cortical hierarchy that encode lower-level linguistic representations. This account therefore predicts that highly implausible words will produce an enhanced later evoked response within lower-level regions such as the posterior fusiform cortex, which supports sublexical orthographic processing (Price and Devlin 2011; Heilbron et al. 2020b), i.e. orthographic reprocessing (see Fig. 3, right).
Finally, in addition to collecting MEG data, we also collected EEG data simultaneously. This was important because MEG and EEG do not capture precisely the same underlying signal (Ahlfors et al. 2010; see Supplementary Materials for further discussion). Therefore, by collecting EEG data using the same stimuli and in the same participants, we were able to replicate previous ERP findings (Kuperberg et al. 2020). A full description is provided there as well as in Supplementary Materials. Briefly, the discourse contexts of each scenario were constraining (average cloze probability of the most probable word: 68%), as quantified in a cloze norming study that was carried out in participants who were recruited through Amazon Mechanical Turk. These contextual constraints came from the entirety of the discourse context—the first two sentences plus the first few words of the third sentence before the critical word. In all scenarios, these first few words of the third sentence included an adjunct phrase of 1–4 words, followed by a pronominal subject that referred back to the noun with the highest cloze probability for that context. To create the unexpected plausible scenarios, each context was paired with a noun of zero (or very low) cloze probability, but that was still plausible in relation to this context. To create the implausible scenarios, each context was paired with a noun that violated the animacy constraints of the preceding verb (which constrained either for an animate or an inanimate noun).

The stimuli were based on three of the conditions used in a previous ERP study (Kuperberg et al. 2020). A full description is provided there as well as in Supplementary Materials. Briefly, the discourse contexts of each scenario were constraining (average cloze probability of the most probable word: 68%), as quantified in a cloze norming study that was carried out in participants who were recruited through Amazon Mechanical Turk. These contextual constraints came from the entirety of the discourse context—the first two sentences plus the first few words of the third sentence before the critical word. In all scenarios, these first few words of the third sentence included an adjunct phrase of 1–4 words, followed by a pronominal subject that referred back to the first two sentences, a verb and a determiner.

To create the expected scenarios, each context was paired with the noun with the highest cloze probability for that context. To create the unexpected plausible scenarios, each context was paired with a noun of zero (or very low) cloze probability, but that was still plausible in relation to this context. To create the implausible scenarios, each context was paired with a noun that violated the animacy constraints of the preceding verb, see Table 1. In all scenarios, the critical noun was followed by three additional words to complete the sentence.
Table 1. Example of each experimental condition, together with stimuli characteristics.

Prior discourse-constraining context: “The lifeguards received a report of sharks right near the beach. Their immediate concern was to prevent any incidents in the sea. Hence, they cautioned the...”

<table>
<thead>
<tr>
<th>Scenario type</th>
<th>Critical word</th>
<th>Constraint(b)</th>
<th>Cloze(b)</th>
<th>SSV(c)</th>
<th>Length</th>
<th>Freq.</th>
<th>OLD</th>
<th>Conc.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expected</td>
<td>swimmers...</td>
<td>69%</td>
<td>69%</td>
<td>0.18</td>
<td>5.69</td>
<td>1.53</td>
<td>1.93</td>
<td>4.30</td>
</tr>
<tr>
<td>Unexpected</td>
<td>trainers</td>
<td>(14%)</td>
<td>0.1%</td>
<td>(0.01)</td>
<td>7.46</td>
<td>0.61</td>
<td>2.61</td>
<td>4.15</td>
</tr>
<tr>
<td>plausible</td>
<td></td>
<td>(14%)</td>
<td>(0.5%)</td>
<td>(0.06)</td>
<td>(2.22)</td>
<td>(0.88)</td>
<td>(0.86)</td>
<td></td>
</tr>
<tr>
<td>Implausible</td>
<td>drawers...</td>
<td>66%</td>
<td>0%</td>
<td>0.01</td>
<td>7.11</td>
<td>0.81</td>
<td>2.47</td>
<td>4.21</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(16%)</td>
<td>(0%)</td>
<td>(0.05)</td>
<td>(2.04)</td>
<td>(0.85)</td>
<td>(0.81)</td>
<td></td>
</tr>
</tbody>
</table>

The critical words (underlined here although not in the experiment itself) were followed by three additional words, indicated here with three dots. Means are shown with the standard deviations in parentheses. Length: Number of letters. The lexical constraint of each discourse context was calculated by identifying the most common completion across participants who saw that context in a cloze norming study (see Supplementary Materials), and tallying the proportion of participants who provided this completion. Cloze probabilities of critical words were calculated based on the percentage of respondents in the cloze norming study who provided the critical noun. SSV: Semantic similarity values, quantifying the semantic relatedness between the critical words and the “bag of words” within the prior contexts, based on latent semantic analysis. Freq.: Log frequency values of critical words, retrieved from the English Lexicon Project. OLD: Orthographic Levenshtein Distance values of critical words, retrieved from the English Lexicon Project. Conc.: Concreteness ratings of critical words (Brysbaert et al. 2014).

When constructing these materials, we attempted to minimize the possibility of bottom-up priming between words in the prior context and the critical word. First, the critical noun and the preceding verb were never strongly associated, as confirmed in a separate cloze norming study (described in Kuperberg et al. 2020, see pages 17–18). In that cloze norming study, participants provided continuations for very short sentence contexts constructed around the verb (proper name + verb + determiner, e.g. “Jane cautioned the...”). Critical nouns were provided only rarely as a completion (mean cloze = 6%, standard deviations, SD = 7%). This meant that, in the expected condition of the present study, the critical word was separated from any semantically related words in the prior discourse by a sentence boundary (see Van Petten et al. 1997 for evidence that purely associative priming between individual pairs of words is short-lived, and likely plays a minimal role in driving discourse contextual effects on the N400).

Second, we carefully matched the unexpected plausible and implausible conditions on the degree of semantic relatedness between the critical word and the full “bag of words” in the prior context, operationalized using semantic similarity values, extracted using latent semantic analysis (Landauer and Dumais 1997), see Table 1 (t(1024) = 0.09, P = 0.93, d = 0.01).

Each list contained 25 expected, 25 unexpected plausible, and 50 implausible scenarios. This ensured that each participant saw an equal proportion of plausible and implausible scenarios. Conditions were counterbalanced across lists such that, across all participants, (i) the same 25 discourse contexts appeared in all three conditions, but no individual saw the same context more than once, and (ii) the same 25 critical words appeared in the 25 unexpected plausible scenarios and 25 of the implausible scenarios, but no individual saw this word more than once. We made the a priori decision to include all 50 implausible scenarios in our main analyses in order to maximize power to source-localized effects (see below). As a byproduct of counterbalancing the same high constraint contexts across the three conditions, the critical words in the expected scenarios had fewer letters, smaller orthographic neighborhoods, and were more frequent than in the other two conditions (all ts > 5, Ps < 0.001, ds > 0.54), see Supplementary Materials for an additional analysis that addresses potential concerns regarding these differences. However, all these lexical features were matched between the unexpected plausible and the implausible conditions (all ts < 1.74, Ps > 0.08, ds < 0.16), see Table 1.

As discussed in Supplementary Materials, each list additionally included 100 scenarios with less constraining contexts (again, 50% plausible and 50% implausible). The 50 low constraint implausible scenarios served as fillers, and the 50 low constraint plausible scenarios served as a fourth experimental condition. In the present study, we focused on the three high constraint conditions, which were perfectly matched prior to the onset of the critical word. Analyses that directly contrast the expected, high constraint plausible, and low constraint plausible words will be reported in a separate manuscript.

Overall procedure

Participants took part in two separate experimental sessions: one in which we simultaneously collected MEG/EEG data, and one in which we collected structural and functional MRI data. The present manuscript focuses on the EEG and MEG datasets. Below, we describe the acquisition and analysis of the MEG/EEG data, as well as the structural MRI data, which was used to constrain MEG source-localization. A report of the fMRI dataset, together with detailed comparisons with the EEG/MEG dataset, will appear in a separate manuscript.

Participants

Thirty-three participants took part, but one MEG/EEG dataset was excluded because of technical problems. Here, we report the results of 32 MEG/EEG datasets (16 females, mean age: 23.4; range: 18–35). All participants were native speakers of English (with no other language exposure before the age of 5), were right-handed, and had normal or corrected-to-normal vision. All were screened to exclude past or present psychiatric and neurological disorders, and none were taking medication affecting the Central Nervous System. The study was approved by the Mass. General Brigham Institutional Review Board (IRB), and written informed consent was obtained from all participants.

Stimuli presentation and task

Stimuli were presented using PsychoPy 1.83 software and projected on to a screen in white Arial font (size: 0.1 of the screen height) on a black background. On each trial, the first two sentences appeared in full (each for 3,900 ms, 100 ms interstimulus interval, ISI), followed by a fixation (a white “+” character), which was presented for 550 ms, followed by a 100 ms ISI. Then the third
sentence was presented word by word (each word for 450 ms, 100 ms ISI).

Participants’ task was to judge whether or not the scenario “made sense” by pressing one of two buttons after seeing a ‘?’ which appeared after each scenario (1,400 ms with a 100 ms ISI). Response fingers were counterbalanced across participants. This task encouraged active coherence monitoring during online comprehension and was intended to ensure that comprehenders detected the anomalies, which is necessary to produce a late posterior positivity/P600 response (Sanford et al. 2011). In addition, following 24/200 trials (distributed semi-randomly), participants answered a “YES/NO” comprehension question that appeared on the screen for 1,900 ms (100 ms ISI). This encouraged participants to comprehend the scenarios as a whole, rather than focusing on only the third sentence in which the anomalies appeared.

Following each trial, a blank screen was presented with a variable duration that ranged from 100 to 500 ms. This was then followed by a green fixation (++++) for a duration of 900 ms, followed by an ISI of 100 ms. These green fixations were used to estimate the noise covariance for the MEG source-localization (see below). To ensure precise time-locking of stimuli, we used frame-based timing, which synced stimulus presentation to the frame refresh rate of the monitor (for example, a 450 ms word presentation would be displayed for exactly 27 frames on our 60 Hz monitor).

Stimuli were presented in 8 blocks, each with 25 scenarios. Blocks were presented in random order in each participant. Participants took part in a short practice session before the formal experiment to gain familiarity with the stimulus presentation and tasks.

**Data acquisition**

**MEG and EEG data acquisition**

Participants sat inside a magnetically shielded room (IMEDCO AG, Switzerland). The MEG data were acquired with a Neuromag VectorView system (Elekta-Neuromag Oy, Finland) with 306 sensors—102 triplets, with each triplet comprising two orthogonal planar gradiometers and one magnetometer. The EEG data were acquired at the same time using a 70-channel MEG-compatible scalp electrode system (BrainProducts, München), and referenced to an electrode placed on the left mastoid. An electrode was also placed on the right mastoid and a ground electrode was placed on the left collabone. Electrooculogram (EOG) data were collected with bipolar recordings: Vertical EOG electrodes were placed above and below the left eye, and horizontal EOG electrodes were placed on the outer canthus of each eye. ECG data were also collected with bipolar recordings: electrocardiogram (ECG) electrodes were placed a few centimeters under the left and right collabones. Impedances were kept at <20 kΩ at all scalp sites, at <10 kΩ at mastoid sites, and at <30 kΩ at EOG and ECG sites. Both MEG and EEG data were acquired with an online band-pass filter of 0.03–300 Hz and were continuously sampled at 1,000 Hz.

To record the head position relative to the MEG sensor array for later co-registration of the MEG and MRI coordinate frames, the locations of three fiduciary points (nasion and two auricular), four head position indicator coils, all EEG electrodes, and at least 100 additional points, were digitized using a 3Space Fastrak Polhemus digitizer, integrated with the Vectorview system.

**Structural MRI data acquisition**

In order to create participant-specific head model for MEG source-localization, we acquired structural MRIs using a 3T Siemens Trio scanner with a 32-channel head coil for all participants.

T1-weighted high-resolution structural images were obtained using the following parameters: 1-mm isotropic multi-echo magnetization-prepared rapid gradient-echo, MP-RAGE, time to repetition (TR): 2.53 s; flip angle: 7°; and 4 echoes with TE: 1.69, 3.55, 5.41, and 7.27 ms.

**ERP analysis**

 stringent procedures were applied to ensure the accuracy of the subsequent subtractions. The data analyses included an independent component analysis (ICA) to remove ICA components associated with eye-movement (one component on average was removed per participant). Finally, we visualized the artifact-free trials, and removed any additional trials with residual artifact. On average, 6.5% of trials were removed from each condition (equally distributed across the 3 conditions: F(2,62) = 2.31, P = 0.08, n² = 0.086), yielding, on average, 23 trials in the expected and unexpected plausible conditions, and 46 trials in the implausible condition. Finally, the data of bad channels were interpolated using spherical spline interpolation (Perrin et al. 1989). In each participant, at each site, we then calculated ERPs, time-locked to the onset of critical words, in each of the conditions, applying a −100 ms pre-stimulus baseline.

We next averaged these voltages across all time points and electrode sites within each of three spatiotemporal regions of interest (ROIs) that were selected, prior to capture the N400, the late frontal positivity and the late posterior positivity/P600 ERP components (Kuperberg et al. 2020). The N400 was operationalized as the average voltage across 10 electrode sites within a central region (Cz, C1, C2, C3, C4, CPz, CP1, CP2, CP3, and CP4), averaged across all sampling points between 300 and 500 ms; the late frontal positivity was operationalized as the average voltage across 8 electrode sites within a prefrontal region (FPz, Fp1, Fp2, Fp3, Fp4, AFz, AF3, and AF4), averaged across sampling points between 600 and 1,000 ms; the late posterior positivity/P600 was operationalized as the average voltage across 11 electrode sites within a posterior region (Pz, P1, P2, P3, P4, P0z, PO3, PO4, Oz, O1, and O2), averaged across sampling points between 600 and 1,000 ms. We carried out planned statistical comparisons between each pair of conditions. We also report post-hoc ERP analyses across an earlier 200–300-ms time-window in Supplementary Materials.

**MEG analysis**

**MEG preprocessing, individual averaging, and sensor-level visualization**

MEG data were analyzed using version 2.7.4 of the minimum norm estimate (MNE) software package in Python (Gramfort et al. 2014). In each participant, in each run, MEG sensors with excessive noise were visually identified and removed from further analysis. This resulted in the removal of 7 (on average) of the 306 MEG sensors. Signal-space projection (SSP) correction was used to correct for ECG artifact. Trials with eye-movement and blink artifacts were automatically removed (Gramfort et al. 2014). Then, after applying a band-pass filter at 0.1–30 Hz, we segmented epochs from −100 to 1,000 ms, relative to the onset of the critical words.
We removed epochs with additional artifact, as assessed using a peak-to-peak detection algorithm (the pre-specified cutoff for the maximal amplitude range was 4 × 10⁻¹² T/m for the gradiometer sensors and 4 × 10⁻¹² T for the magnetometer sensors). On average, 6.7% trials in each condition were removed (equally distributed across the 3 conditions: F(2,62) = 0.87, P = 0.42, η² = 0.027), yielding, on average, 21 artifact-free trials in the expected and unexpected plausible conditions, and 42 artifact-free trials in the implausible condition.

In each participant, in each block, at each magnetometer sensor and at each of the two gradiometers at each site, we calculated event-related fields (ERFs), time-locked to the onset of critical words in each of the three conditions, applying a −100-msec pre-stimulus baseline. We averaged the ERFs across blocks in sensor space, interpolating any bad sensors using spherical spline interpolation (Perrin et al. 1989). We created gradiometer and magnetometer sensor maps to visualize the topographic distribution of ERFs across the scalp. In creating the gradiometer maps, we used the root mean square of the ERFs produced by the two gradiometers at each site.

**MEG source-localization in individual participants**

Each participant’s cortical surface was first reconstructed from their structural T1 MP-RAGE image using the FreeSurfer software package developed at the Martinos Center, Charlestown, MA (http://surfer.nmr.mgh.harvard.edu). We used MNE-Python (Gramfort et al. 2014) to estimate the sources of the ERFs evoked by critical words in each of the 3 conditions, on each participant’s reconstructed cortical surface using minimum-norm estimation (MNE, Hämäläinen and Sarvas 1989).

In order to calculate the inverse operator in each participant—the transformation that estimates the underlying neuroanatomical sources for a given spatial distribution of activity in sensor space—we first needed to construct a noise-covariance matrix of each participant’s MEG sensor-level data, as well as a forward model in each participant (the model that predicts the pattern of sensor activity that would be produced by all dipoles within the source space).

To construct the noise-covariance matrix in each participant, we used 650 ms of MEG sensor-level data recorded during the presentation of the green inter-trial fixations (we used an epoch from 100 to 750 ms, which cut-off MEG data measured at the onset and offset of these fixations in order to avoid onset and offset evoked responses). We concatenated these fixations across blocks. To construct the forward model in each participant, we needed to (i) define the source space—the location, number, and spacing of dipoles, (ii) create a Boundary Element Model (BEM), which describes the geometry of the head and the conductivities of the different tissues, and (iii) specify the MEG-MRI coordinate transformation—the location of MEG sensors in relation to the head surface.

The source space was defined on the white matter surface of each participant’s reconstructed MRI and constituted 4,098 vertices per hemisphere, with three orthogonally-oriented dipoles at each vertex (two tangential and one perpendicular to the cortical surface). We defined these vertices using a grid that decimated the surface into meshes, with a spacing of 4.9 mm between adjacent locations (spacing: “oct6”). We created a single compartment BEM by first stripping the outer non-brain tissue (skull and scalp) from the pial surface using the watershed algorithm in FreeSurfer, and then applying a single conductivity parameter to all brain tissue bounded by the inner skull. We specified the location of the MEG sensors in relation to the head surface by manually aligning the fiducial points and 3D digitizer (Polhemus) data with the scalp surface triangulation created in FreeSurfer, using the mne_analyze tool (Gramfort et al. 2014).

We then calculated the inverse operator in each participant, setting two additional constraints. First, we set a loose constraint on the relative weighting of tangential and perpendicular dipole orientations within the source space (loose = 0.2). Second, we set a constraint on the relative weighting of superficial and deep neuroanatomical sources (depth = 0.8) in order to increase the likelihood that the minimum-norm estimates would detect deep sources.

We then applied each participant’s inverse operator to the ERFs of all magnetometer and gradiometer sensors calculated within each block. We estimated activity at the dipoles that were orientated perpendicular to the cortical surface at each vertex (pick_ori = “normal”). Each of these perpendicular dipoles had both a positive and a negative value, which indicated whether the currents were outgoing or ingoing respectively. We chose to retain the two polarities of each estimated dipole for further analyses for two reasons. First, this approach allowed us to include all trials in each of our 3 conditions, thereby maximizing power without inflating our estimate of noise in the conditions with more trials (if we had chosen to simply estimate the magnitude of each dipole by squaring the positive and negative values to yield positively-signed estimates, we would have artificially inflated the noise estimates in the implausible conditions, which had twice as many trials as the expected and the unexpected plausible conditions). Second, by retaining this polarity information, we were able to determine whether any statistical differences between conditions were driven by differences in the magnitude and/or differences in the polarity of the dipoles evoked in each condition (see Supplementary Materials for further discussion).

Then, for each condition in each block, we computed noise-normalized dynamic Statistical Parametric Maps (dSPMs; Dale et al. 2000) on each participant’s cortical surface at each time point, and averaged these values across blocks within each participant. Finally, the source estimates for each participant were morphed on the FreeSurfer average brain, “fsaverage” (Fischl et al. 1999), for group averaging and statistical analysis.

**Statistical analysis of MEG source-level evoked activity**

To statistically analyze the source-localized evoked MEG responses, we carried out pairwise t-tests between each pair of conditions on the signed estimated dSPM values at each sampling point from 300 to 1,000 ms after critical word onset. Pairwise t-tests were calculated at each vertex in source space, over a large left-lateralized search region where we expected effects to be maximal (particularly the effects of lexical predictability, see Federmaner 2022). This search area included left lateral temporal cortex, left ventral temporal cortex, left medial temporal cortex, left lateral parietal cortex, left lateral frontal cortex, and left medial frontal cortex, and was defined on the Desikan-Killiany Atlas (Desikan et al. 2006; Supplementary Fig. 1A, see online supplementary material for a color version of this figure).

We used permutation-based cluster-mass procedures (Maris and Oostenveld 2007) to correct for multiple comparisons in time and space. First, at each vertex and at each time point, any data points that exceeded a pre-set uncorrected significance threshold of 1% (i.e. P ≤ 0.01) were -log10 transformed, and the rest were zeroed. The use of unsigned -log-transformed F-values allowed us to account for effects with the opposite polarities. A single neuroanatomical source that is located on one side of a sulcus...
can appear on the cortical surface as adjacent dipoles of opposite polarity (outgoing and ingoing currents) because of signal bleeding to the other side of the sulcus (Hamalainen et al. 1993; see Figs 6 and 7). The use of unsigned P-values therefore ensured that adjacent effects of opposite signs were treated as a single underlying source in the statistical analyses.

Second, to minimize multiple comparisons over time, we averaged the transformed P-values at each vertex within three a priori time-windows of interest: 300–500 ms, corresponding to the N400 time-window, and 600–800 ms and 800–1,000 ms, corresponding to the first and second halves of the timewindow associated with late positive ERP effects. Then, to minimize multiple comparisons over space, these transformed P-values were further averaged across vertices within 140 spatial patches of approximately equal size (Khan et al. 2018), shown in Supplementary Fig 1B. This resulted in 140 separate test statistics in each time-window of interest, i.e. 140 spatiotemporally-clustered test statistics. This approach ensured that data points within the same time-window and spatial patch (i.e. each spatiotemporal cluster) were treated as originating from a single underlying source. We should note that, although this approach increases statistical power, it also constrains our statistical inference to the spatial resolution of each patch and to the temporal resolution of our a priori time-windows.

Finally, we computed a null distribution for our spatiotemporally-clustered test statistics, by carrying out exactly the same procedure as that described above, but this time randomly assigning condition labels within each participant, with 10,000 permutations. For each randomization, we took the largest value across all spatial patches as our cluster-mass statistic. To test our hypotheses, we compared each of the observed spatiotemporally-clustered test statistics against this null distribution. If an observed cluster-level statistic fell within the highest 5.0% of the distribution, we considered it to be significant. Note that this nonparametric cluster-based approach is robust to any differences in signal-to-noise resulting from different numbers of trials in the expected, unexpected plausible, and implausible conditions.

When displaying the results, we show both the uncorrected and cluster-corrected results on the cortical surface. We projected the averaged -log10 transformed uncorrected P-values (P < 0.05) within each time-window at each vertex on to the “taverage” brain (Fischl et al. 1999). We then grouped together all spatial patches that reached cluster-level significance into the neuroanatomical regions that are shown in Supplementary Fig 1A and listed in Supplementary Table 1 (defined using the Desikan-Killiany Atlas; Desikan et al. 2006). If one or more patches within a neuroanatomical region reached cluster-level significance, we circled the region in red.

We also carried out several additional analyses, which are reported in Supplementary Materials: (i) an analysis of a subset of the MEG data to address potential concerns regarding differences between the expected and implausible conditions in the number of trials per condition, and in the lexical properties of the critical words, (ii) an analysis to determine whether there were effects in an earlier 200–300-ms time-window, and (iii) an analysis of an analogous search region within the right hemisphere.

Finally, in Supplementary Materials, we report an exploratory cross-ROI multivariate decoding analysis, which was intended to provide preliminary multivariate data to clarify or support certain points in our interpretation of the evoked effects, as discussed in the main manuscript.

### Results

#### Behavioral

Participants correctly judged the plausibility of 89.42% scenarios (SD: 8.83%) and answered 80.12% (SD: 11.50%) of the comprehension questions correctly, suggesting that they were engaged in comprehension (see Supplementary Materials).

#### ERP

The ERP results (Fig 4 and Table 2) replicate previous findings (Kuperberg et al. 2020). Between 300 and 500 ms, the N400 amplitude increased across the three conditions. Between 600 and 1,000 ms, the unexpected plausible words produced a larger late frontal positivity than both the expected and implausible continuations, whereas the implausible words produced a larger late posterior positivity/P600 than both the expected and unexpected plausible continuations.

#### MEG

To facilitate a qualitative comparison between MEG sensor-level results and the scalp-recorded ERP results, we show the sensor-level MEG results in Fig 5 (we note, however, that MEG statistical analyses were carried out in source space). Between 300 and 500 ms, the sensor-level MEG findings showed a similar graded increase in evoked activity, see Fig 5A (the evoked response to the implausible continuations was larger in MEG than in ERP, see Supplementary Materials for discussion). Between 600 and 1,000 ms, the unexpected plausible and implausible words produced larger responses than the expected words, and the topographic sensor maps revealed distinct patterns for each effect, see Fig 5B.

#### Source-localized MEG: evoked effects

300–500 ms

As shown in Fig 6, between 300 and 500 ms, expected, unexpected plausible, and implausible words produced graded increases in activity within multiple regions of left lateral, ventral, and medial temporal cortices. In medial temporal cortices, the effects were also driven by a dipole going in the opposite direction to the expected words. In addition, the implausible words produced a larger response than both the expected and unexpected plausible words within left inferior frontal cortex, as well as a larger response than the expected words within anterior cingulate cortex.

600–1,000 ms

As shown in Fig 7, by 500 ms, the activity produced by the unexpected plausible words within left temporal cortex had diminished. Between 600 and 1,000 ms, however, these continuations produced a response within left inferior frontal cortex, and re-activated the left middle temporal cortex, with a dipole going in the opposite direction to that produced in the N400 time-window.

The evoked activity produced by the implausible words was quite different. By 500 ms, the response produced by these continuations within the left inferior frontal cortex had diminished, but the activity produced within left temporal and posterior fusiform cortices continued into the 600–1,000-ms time-window. In the latter half of this time-window, the implausible words also re-activated the left inferior frontal cortex, with a dipole going in the opposite direction to that produced between 300 and 500 ms. Finally, throughout the 600 and 100-ms window, the implausible words also produced a large dipole within left medial temporal cortex, again with the opposite polarity to that produced in the N400 window.
**Fig. 4.** ERP results. A) Grand-averaged ERP waveforms elicited by critical words in each of the three conditions, shown at three representative electrode sites: Cz, FPz, and Pz. Expected: solid black line; unexpected plausible: solid red line; implausible: dashed blue line. Negative voltage is plotted upwards. Dotted boxes are used to indicate the time-windows corresponding to the N400 (300–500 ms), the late frontal positivity (600–1,000 ms), and the late posterior positivity/P600 (600–1,000 ms) ERP components. B) Voltage maps show the topographic distributions of the ERP effects produced by contrasting expected, unexpected plausible, and implausible critical words between 300 and 500 ms (left panel) and between 600 and 1,000 ms (right panel). Note that the N400 effects and the late positivity effects are shown at different voltage scales to better illustrate the scalp distribution of each effect.

**Table 2.** ERP statistical results.

<table>
<thead>
<tr>
<th>Spatiotemporal region</th>
<th>Contrast</th>
<th>t-Value</th>
<th>df (31)</th>
<th>P-value</th>
<th>Effect size (d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N400 (Central region, 300–500 ms)</td>
<td>Unexpected &gt; Expected</td>
<td>-5.31</td>
<td>&lt; 0.001</td>
<td>0.94</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Implausible &gt; Expected</td>
<td>-7.72</td>
<td>&lt; 0.001</td>
<td>1.36</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Implausible &gt; Unexpected</td>
<td>-4.63</td>
<td>0.001</td>
<td>0.82</td>
<td></td>
</tr>
<tr>
<td>Late frontal positivity (Prefrontal region,</td>
<td>Unexpected &gt; Expected</td>
<td>3.03</td>
<td>0.005</td>
<td>0.53</td>
<td></td>
</tr>
<tr>
<td>600–1,000 ms)</td>
<td>Implausible = Expected</td>
<td>1.51</td>
<td>0.14</td>
<td>0.24</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Unexpected &gt; Implausible</td>
<td>2.52</td>
<td>0.018</td>
<td>0.42</td>
<td></td>
</tr>
<tr>
<td>Late posterior positivity/P600 (Posterior</td>
<td>Unexpected = Expected</td>
<td>1.91</td>
<td>&lt; 0.001</td>
<td>1.12</td>
<td></td>
</tr>
<tr>
<td>region, 600–1,000 ms)</td>
<td>Implausible &gt; Expected</td>
<td>5.75</td>
<td>&lt; 0.001</td>
<td>0.99</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Implausible &gt; Unexpected</td>
<td>5.75</td>
<td>&lt; 0.001</td>
<td>0.99</td>
<td></td>
</tr>
</tbody>
</table>

Pair-wise statistical comparisons showed that, within left inferior frontal cortex, there was a significant difference in comparing both the unexpected plausible and the implausible words with the expected words (600–800 ms), with both effects being driven by dipoles going in opposite directions in the two conditions. A direct comparison between the unexpected plausible and implausible conditions, however, revealed no differences within left inferior frontal cortex. Within left lateral temporal cortex, there were significant differences in comparing the unexpected plausible words with both the expected (800–1,000 ms) and implausible words (600–800 ms). Both these effects were driven by dipoles going in the opposite direction in each of the two conditions. In posterior fusiform cortex there was a significant difference in comparing the implausible words with both the expected (600–800 ms) and unexpected plausible words (600–800 ms and 800–1,000 ms). Similarly, within medial temporal cortex, the response produced by the implausible words differed from that produced by both other conditions (600–800 ms and 800–1,000 ms).

**Discussion**

We used MEG and EEG to track the spatiotemporal dynamics of evoked activity produced by expected, unexpected plausible, and implausible words during language comprehension. At the scalp surface, our ERP findings replicate previous studies by showing that, between 300 and 500 ms, the three conditions produced progressively larger N400 responses (Kuperberg et al. 2020; Nieuwland et al. 2020), and that, between 600 and 1,000 ms,
A 300–500 ms. The N400 effect of lexical predictability localizes to the left temporal cortex. In our ERP recordings, the N400 was smaller to the expected than to the unexpected plausible words, replicating many previous ERP studies (e.g. Kutas and Hillyard 1984; DeLong et al. 2005; Federmeier et al. 2007). MEG source-localization showed that this effect of lexical predictability localized to multiple regions within the left temporal cortex.

B 600–1,000 ms. By simultaneously collecting MEG data and source-localizing the evoked response in both time-windows, we were able to show, for the first time where in the brain these effects are produced.

Several previous MEG (e.g. Helenius et al. 1998; Halgren et al. 2002; Maess et al. 2006; Ihara et al. 2007) and intracranial (McCarthy et al. 1995) studies have reported effects of sentence context on the N400 within temporal and/or inferior frontal cortices. However, most of these previous studies have directly contrasted expected and implausible words, without dissociating the effects of lexical predictability and contextual plausibility. It has therefore been difficult to directly link activity produced at lower and higher levels of the fronto-temporal hierarchy to processing at different levels of representation (mapping between word-forms, semantic features, and real-world knowledge).

In addition, no previous MEG or intracranial study has source-localized the evoked effects of semantic context beyond the N400 time-window. By determining precisely when and how evoked activity across the fronto-temporal hierarchy is modulated following the onset of incoming words in both time-windows, our findings shed new light on the cognitive architecture and neurobiology of online language comprehension.

**300–500 ms**

**The N400 effect of lexical predictability localizes to the left temporal cortex**

In our ERP recordings, the N400 was smaller to the expected than to the unexpected plausible words, replicating many previous ERP studies (e.g. Kutas and Hillyard 1984; DeLong et al. 2005; Federmeier et al. 2007). MEG source-localization showed that this effect of lexical predictability localized to multiple regions within the left temporal cortex.

**unexpected plausible** and **implausible** continuations produced two spatially-distinct late positivities (Van Petten and Luka 2012; DeLong et al. 2014; Brothers et al. 2020; Kuperberg et al. 2020). By simultaneously collecting MEG data and source-localizing the evoked response in both time-windows, we were able to show, for the first time where in the brain these effects are produced.

Several previous MEG (e.g. Helenius et al. 1998; Halgren et al. 2002; Maess et al. 2006; Ihara et al. 2007) and intracranial (McCarthy et al. 1995) studies have reported effects of sentence context on the N400 within temporal and/or inferior frontal cortices. However, most of these previous studies have directly contrasted expected and implausible words, without dissociating the effects of lexical predictability and contextual plausibility. It has therefore been difficult to directly link activity produced at lower and higher levels of the fronto-temporal hierarchy to processing at different levels of representation (mapping between word-forms, semantic features, and real-world knowledge).

In addition, no previous MEG or intracranial study has source-localized the evoked effects of semantic context beyond the N400 time-window. By determining precisely when and how evoked activity across the fronto-temporal hierarchy is modulated following the onset of incoming words in both time-windows, our findings shed new light on the cognitive architecture and neurobiology of online language comprehension.
The N400 effect of contextual plausibility localizes to both temporal and inferior frontal cortices

In ERPs, we also observed a smaller N400 to unexpected plausible words, relative to implausible words, again replicating previous findings (Kuperberg et al. 2020; Nieuwland et al. 2020). We note that this effect of contextual plausibility on the scalp-recorded N400 was much larger in MEG than in ERP, which has important implications for the functional interpretation of the N400 in the presence of an overlapping late posterior positivity/P600.
Fig. 7. MEG source-level activity produced by the expected, unexpected plausible and implausible critical words in the late 600–1,000 ms time-window. A) Signed dynamic statistical parametric maps (dSPMs) produced by expected (top) unexpected plausible (middle), and implausible (bottom) critical words are shown at 100-ms intervals from 500 until 1,000 ms. All dSPMs are displayed on the FreeSurfer average surface, “fsaverage” (Fischl et al. 1999), thresholded at 0.15, with red indicating outgoing currents (positive dSPM values) and blue indicating ingoing currents (negative dSPM values). Ingoing and outgoing currents that are directly adjacent to one another are interpreted as reflecting a single underlying dipole (neuroanatomical source). This is because, if an underlying dipole/source is situated on one side of a sulcus, the signal can bleed into the other side, leading to the appearance of an adjacent dipole in the opposite direction (Hämäläinen et al. 1993). The full dynamics of these source activations (at all sampling points) are shown as videos in Supplementary Materials. B) Statistical maps contrasting unexpected plausible and expected words (top), implausible and expected words (middle), and implausible and unexpected plausible words (bottom) are shown between 600 and 800 ms (left) and between 800 and 1,000 ms (right). Averaged -log10 transformed uncorrected P-values (P < 0.05) at each vertex are shown on the “fsaverage” surface. We grouped together all spatial patches that reached cluster-level significance into the neuroanatomical regions that are shown in Supplementary Fig. 1A and listed in Supplementary Table 1 (defined using the Desikan-Killiany Atlas; Desikan et al. 2006). If one or more patches within a neuroanatomical region reached
Supplementary Materials

MacDonald et al. 1994
Mamashli et al. 2019
Kuperberg et al. 2020
Lyu et al. 2019
Marslen-Wilson et al. 1988
Marslen-Wilson 1987
Altmann and Steedman 2011
Brouwer et al. 2017
Baggio and Hagoort 2011
Tanenhaus and Trueswell 1995

Within the left temporal cortex, the effect of contextual plausibility broadly localized to the same regions as the effects of lexical predictability, described previously. Supplementary analyses also revealed some plausibility effects in homologous regions of the right temporal lobe (lateral and medial, see Supplementary Materials). Again, we interpret these smaller evoked responses within the temporal cortex as reflecting facilitated processing at the lexico-semantic level. However, in this case, instead of resulting from the pre-activation of a specific upcoming word, this facilitation resulted from the pre-activation of distributed features associated with a broad semantic category (i.e., animacy-based features; Wang et al. 2020). Thus, lexico-semantic processing was easier for the plausible unexpected inputs, where some semantic features were pre-activated, than for the implausible inputs where no semantic features were pre-activated (Paczyński and Kuperberg 2011, 2012; Kuperberg et al. 2020).

In the prefrontal cortex, the effect of contextual plausibility localized to the inferior portion of the left frontal and orbitofrontal cortices, as well as to their right hemisphere homologues (see Supplementary Materials). We interpret these smaller inferior frontal evoked responses to the unexpected plausible vs. the implausible continuations as reflecting the relative ease of mapping plausible event representations onto longer-term real-world knowledge.

Architecture implications: Towards a predictive coding framework

In the Introduction, we laid out four general frameworks of language comprehension. Although each of the first three frameworks can explain a subset of our findings in the N400 time-window, they cannot explain the full set of results across both temporal and inferior frontal cortices. For instance, both the "distributed state" and the "lexico-semantic facilitation" accounts can explain the graded N400 modulation across the three conditions within the temporal cortex. However, the "distributed state" account would have predicted similar graded effects within the inferior frontal cortex (Fig. 1A), whereas the "lexico-semantic facilitation" account would have predicted no prefrontal N400 modulation at all (Fig. 1B). The "prediction-integration" account can explain why the N400 in the inferior frontal cortex was selectively enhanced to implausible continuations. However, this framework would have incorrectly predicted nongraded N400 modulation across the three conditions within the temporal cortex, with an attenuation of the N400 only to the expected continuations (Fig. 1C).

We argue that the full pattern of evoked activity produced between 300 and 500 ms across the fronto-temporal hierarchy can be explained by the computational principles of hierarchical predictive coding (Fig. 1D). We emphasize that this predictive coding framework shares several important features with the other three neurobiological models. Like these other frameworks, it assumes that language comprehension is both interactive and incremental (cf. Marslen-Wilson 1987; Altmann and Steedman 1988; Marslen-Wilson et al. 1988; MacDonald et al. 1994; Tanenhaus and Trueswell 1995), with the prior context influencing the initial feedforward sweep of evoked activity produced by incoming words between 300 and 500 ms (the N400 response). Also similar to these other models, it assumes that, in the 300 and 500-ms time-window, information is continually transferred across temporal and inferior frontal cortices in both predictable and unpredictable sentences (see Baggio and Hagoort 2011, and see Lyu et al. 2019; Mamashli et al. 2019; Liu et al. 2020 for recent empirical evidence). Finally, similar to both the "prediction-integration" and the "lexico-semantic facilitation" accounts, predictive coding posits a hierarchical organization of representations across the fronto-temporal hierarchy. During discourse comprehension, we assume that comprehenders incrementally built a higher-level event model that was maintained over a relatively long time-scale within the prefrontal cortex, and that interacted both with real-world knowledge, represented over a still longer time-scale, as well as with lower-level lexico-semantic representations, encoded within the temporal cortex at a shorter time-scale.

From a computational perspective, however, there is a key difference between predictive coding and other frameworks. Although prior computational models of the N400 have assumed that evoked activity reflects changes in the state of neural activity at a given level of the cortical hierarchy (e.g. Brouwer et al. 2017; Rabovsky et al. 2018), predictive coding posits that the magnitude of the evoked response reflects the magnitude of prediction error, i.e. activity produced within a distinct set of "error units". At each level of the cortical hierarchy, these error units only encode information that is not suppressed (or "explained") by predictions produced by state units at the cortical level above (Rao and Ballard 1999, Friston 2005). Thus, predictive coding provides an intuitive biological explanation for the origin of the N400 neural response: Specifically, new unpredicted inputs trigger increased firing and postsynaptic potentials within these error units, which, in turn, results in a larger evoked N400 response.

Within the temporal cortex, predictive coding attributes the graded increases in N400 amplitude across the three conditions (expected < unexpected plausible < implausible) to graded increases in the magnitude of prediction error at the lexico-semantic level. In the case of expected continuations, lexico-semantic prediction error is fully suppressed by prior top-down predictions of the semantic features of specific upcoming words; in the case of unexpected plausible continuations, lexico-semantic prediction error is partially suppressed by prior top-down predictions of animacy-linked semantic features (see Wang et al. 2020), and, in the case of implausible continuations, lexico-semantic prediction error is not suppressed at all.

Within the inferior frontal cortex, predictive coding attributes the nongraded modulation of the N400 across the three conditions (expected = unexpected plausible < implausible) to differences in magnitude of higher-level prediction error produced at the level of the event model. According to this framework, there was no difference in the higher-level prediction error/evoked N400 activity to the expected and unexpected plausible words in inferior frontal cortex because, in both these conditions, information encoded in the plausible higher-level event model was explained/suppressed by predictions from still longer-term real-world knowledge. Despite the absence of an inferior frontal N400 cluster-level significance, we indicate the region using a red circle. The unexpected plausible versus expected contrast revealed significant clusters within left middle temporal and inferior frontal cortices (driven by dipoles going in opposite directions in the two conditions). The implausible versus expected contrast revealed significant clusters within left posterior (occipitotemporal) fusiform cortex (driven by a dipole to the implausible words), left inferior frontal cortex (driven by dipoles going in opposite directions in the two conditions), and within the medial temporal cortex (driven by a dipole to the implausible words). See also Supplementary Materials for analyses over the right hemisphere (Supplementary Figs. 2–4).
effect for this contrast, unpredicted information was still shared between the left temporal and inferior frontal cortices in this time-window, as evidenced by a small but significant cross-ROI decoding effect (see Supplementary Materials). In contrast, when the implausible continuations produced updates of the higher-level event model, these could not be explained/suppressed by predictions based on real-world knowledge, giving rise to a higher-level prediction error and an enhanced evoked response within this region (see Fig 2, right). This explanation follows directly from Rao and Ballard’s discussion in their original description of hierarchical predictive coding in the visual system (Rao and Ballard 1999); So long as newly encoded unpredicted input is consistent with the more general statistics of natural environmental inputs, it should not result in a large increase in activity within higher-level regions because any higher-level prediction error is continually suppressed. However, when “the statistics differ in certain drastic ways from natural statistics” (Rao and Ballard 1999, p. 84), then this will lead to a larger response within higher-level regions of the cortical hierarchy.

Finally, predictive coding also posits that an inability to converge on a plausible interpretation in the prefrontal cortex should result in less accurate top-down predictions, which will fail to fully “switch off” lower-level lexico-speech prediction error within the temporal cortex within the N400 time-window (see Fig 2, right). This failure to suppress lexico-speech prediction error may have also contributed to the enhanced N400 response produced by the implausible (relative to the unexpected plausible) words within the temporal cortex, as well as a prolongation of this effect into the later time-window.

600–1,000 ms

Replicating previous ERP findings, unexpected plausible and implausible words produced two distinct late positivities between 600 and 1,000 ms: A late frontal positivity to the unexpected plausible words (Federmann et al. 2007; Van Petten and Luka 2012; DeLong et al. 2014; Brothers et al. 2020; Kuperberg et al. 2020), and a late posterior positivity/P600 to the highly implausible continuations (Kuperberg 2007; van de Meerendonk et al. 2009; Van Petten and Luka 2012; DeLong et al. 2014; Brothers et al. 2020; Kuperberg et al. 2020; Brothers et al. 2021). Once again, our source-localized MEG results constrain our understanding of the functional significance of this late-stage evoked activity.

Late evoked effects to unexpected plausible words within left inferior frontal and middle temporal cortices

It has been proposed that the left frontal positivity evoked by unexpected plausible words reflects a successful high-level shift in the event model, together with feedback to the lower lexico-speech level (Brothers et al. 2015, 2020; Kuperberg et al. 2020). Our MEG findings are consistent with this account. Between 600 and 1,000 ms, unexpected plausible, relative to expected, continuations produced late evoked effects within both left inferior frontal and left middle temporal cortices.

Hierarchical predictive coding offers a computational-level explanation for this late-stage activity. Within this framework, these late evoked responses are attributed to top-down error—that is, activity produced by new top-down predictions at a given level of the hierarchy that cannot be explained by its prior state (Rao and Ballard 1997, 1999). More specifically, according to this framework, if an unpredicted incoming word leads the brain to retrieve new schema-relevant information from long-term memory (cf. Franklin et al. 2020, e.g. infer what trainees might be doing in a beach scenario), then this will result in the generation of new schema-relevant predictions (Kuperberg 2021) that are propagated down the cortical hierarchy. When these new top-down predictions activate the left inferior frontal cortex, they will produce top-down error at the level of the event model, explaining the larger late evoked response to the unexpected plausible continuations within this region, and when they reach the left temporal cortex they will produce top-down error at the lexico-semantic level, explaining the larger late evoked response within the left middle temporal cortex (see Fig 3, left). In the visual system, it has been proposed that, following an initial bottom-up sweep of activity, this type of feedback re-activation ensures that lower-level regions encode information that is consistent with global gestalt representations that are encoded in higher cortical areas (Lee and Mumford 2003).

Consistent with this interpretation, the late evoked effect within the left middle temporal cortex was driven by a dipole with the opposite polarity to the dipole produced within the left temporal cortex in the earlier N400 time-window. Although the precise significance of this dipole reversal is unclear (see Supplementary Materials for further discussion), it provides evidence that this later evoked effect (top-down lexico-semantic error) was functionally distinct from the earlier stimulus-driven evoked N400 effects observed between 300 and 500 ms (bottom-up lexico-semantic prediction error). This dipole reversal also provides evidence that this late evoked activity does not simply reflect a response to the subsequent word. Finally, an exploratory multivariate analysis revealed a small but significant above-chance cross-ROI decoding effect between left frontal and left temporal regions, suggesting that the unexpected information was indeed shared between these two brain regions within this late time-window (see Supplementary Materials).

Late evoked effects to highly implausible words within posterior fusiform, inferior frontal, and medial temporal cortices

In the present study, the implausible words were not simply implausible—they were also anomalous (e.g. “cautioned the ‘drawers’”); that is, they conflicted with the state of the hierarchical generative model as a whole. This conflict may explain why, relative to expected words, the anomalous activated the anterior cingulate cortex in the earlier N400 time-window (Botvinick 2007; see also Ide et al. 2013).

It has been proposed that the late posterior positivity/P600 produced by highly implausible continuations between 600 and 1,000 ms is linked to a conflict-driven reprocessing at lower levels of linguistic representation (van de Meerendonk et al. 2009; Brothers et al. 2020; Kuperberg et al. 2020; Brothers et al. 2021). Consistent with this theory, the implausible words produced a robust late evoked effect within the posterior (occipitotemporal) fusiform cortex—the so-called “visual word-form area”—that supports orthographic processing (Price and Devlin 2011; Helligen et al. 2020b; see Supplementary Materials for additional discussion on the relationship between the ERP and MEG evoked effects in this late time-window).

Predictive coding again provides a mechanistic account of this lower-level orthographic reprocessing (see Fig 3, right). Within this framework, the late evoked activity within the posterior fusiform cortex is attributed to the production of prediction error at a still lower orthographic level of representation (Price and Devlin 2011) because higher cortical levels failed to generate accurate top-down predictions that would have otherwise suppressed this low-level error within this late time-window. Specifically, after inferring an anomalous event (e.g. “lifeguards cautioned drawers”) between 300 and 500 ms, it was not possible to shift the event model by retrieving new
stored schemas from long-term memory. Therefore, between 600 and 1,000 ms, incorrect predictions based on the prior context and real-world knowledge would have continued to be propagated down the cortical hierarchy. Within the left middle temporal cortex, these top-down lexico-semantic predictions (e.g., “swimmers”/<animate>) would have been incompatible with the lexico-semantic information that was inferred from the bottom-up input (e.g., “drawers”/<inanimate>), resulting in a destabilization of the lexico-semantic state. As a result of this destabilization, the left middle temporal cortex would have produced noisy, inaccurate orthographic predictions. Upon reaching the posterior fusiform cortex, these predictions would have failed to suppress orthographic prediction error, leading to the enhanced evoked response within this region (orthographic reprocessing).

Predictive coding can also explain why, in this late time-window, the anomalous continuations re-activated the left inferior frontal cortex, with a dipole of the opposite polarity to that produced by these continuations in the N400 time-window. Within this framework, this late inferior frontal evoked response reflected top-down error produced the level of the event model when the inaccurate predictions, based on real-world knowledge (<lifeguards cautioned swimmers>), failed to explain the implausible event (<lifeguards cautioned draw>) that had been inferred within the N400 time-window. Indeed, an exploratory cross-ROI decoding analysis revealed no evidence of shared information between left inferior frontal and lateral temporal/fusiform regions within this later time-window (see Supplementary Materials).

Finally, the implausible continuations also produced a dipole within the medial temporal cortex throughout the 600–1,000 ms time-window, again with the opposite polarity to that produced in the N400 time-window. We speculate that this medial temporal activity supported new learning/adaptation, which was triggered by the failure of the current generative model to explain the input—that is, to minimize error across the cortical hierarchy (see further below).

Open questions

We have argued that the full time-course of evoked activity produced within temporal and inferior frontal cortices in response to expected, unexpected plausible, and implausible words, can be understood within a hierarchical predictive coding framework of language comprehension. This interpretation leaves many open questions that will be important to address in future studies.

First, it will be important for future studies to parametrically manipulate lexical predictability and plausibility in order to determine precisely how these factors modulate evoked activity within left temporal and inferior frontal cortices at both early and later stages of processing. It will also be important to understand whether and how evoked activity in both time-windows is modulated by the lexical and discourse constraint of the prior context. For example, according to predictive coding, between 300 and 500 ms, evoked activity produced by unexpected plausible words within left temporal cortex should not be modulated by prior contextual constraint, consistent with scalp-recorded N400 findings (Kutas and Hillyard 1984; Federmeier et al. 2007; Kuperberg et al. 2020).

Second, consistent with a predictive coding architecture, we have assumed that, at each level of the cortical hierarchy, the amplitude of the evoked response largely reflects the magnitude of prediction error—activity within populations of error units that cannot be explained/suppressed by the level above (Friston 2005). A key claim of predictive coding is that these error units are computationally distinct from state units, which encode representational information, regardless of its predictability. As such, this framework predicts that, even if new information does not produce an increased evoked response, we should still be able to decode this information using multivariate methods, which are sensitive to representational information, regardless of its magnitude. It will therefore be important for future studies to test this hypothesis directly by using a combination of univariate and multivariate methods. For example, future studies should follow up our preliminary finding, reported in Supplementary Materials, that, within the left inferior frontal cortex, despite failing to produce a larger N400 response (higher-level prediction error) between 300 and 500 ms, it was still possible to decode new unexpected plausible information within this region in this same time-window. Similarly, future studies should test the hypothesis that, within the left temporal cortex, despite producing a very small evoked N400 response, it should be possible to decode expected information within the N400 time-window (see Kok et al. 2012; Bell et al. 2016 for evidence of dissociations between univariate and multivariate activity to expected inputs in low-level visual perception).

A third and related set of questions concerns the nature of information flow across the cortical hierarchy. Like most other neurobiological frameworks of language comprehension, predictive coding assumes that, in plausible sentences, both unexpected and expected information are shared across multiple regions of the cortical hierarchy between 300 and 500 ms (e.g., Baggio and Hagoort 2011; Lyu et al. 2019; Mamashli et al. 2019; Liu et al. 2020). What remains unclear is precisely how this information is transmitted between regions. For example, some researchers have hypothesized that different frequency bands of oscillatory neural activity carry top-down predictions (e.g. slow beta/alpha) and bottom-up prediction error (e.g. fast gamma; Arnal and Giraud 2012; Bastos et al. 2015; Lewis and Bastiaansen 2015; Bastos et al. 2020). It will be important for future studies to test this hypothesis more explicitly.

Finally, an important set of open questions concerns the relationship between evoked neural activity and longer-term learning, particularly given the robust effects observed in the medial temporal cortex in both the 300–500 and 600–1,000-ms time-windows. In the present study, we focused on the role of evoked activity (prediction error) in relation to inference, i.e. the process of online language comprehension. However, prediction error is also thought to play a critical role in learning in both linguistic (Elman 1990; Dell and Chang 2014) and nonlinguistic (Rescorla 1988) domains. Recent computational models, using standard connectionist architectures, have shown that N400 prediction errors can, in principle, drive incremental learning (see Rabovsky and McRae 2014, Rabovsky et al. 2018; Fitz and Chang 2019; see also footnote 2 in Introduction). Predictive coding offers a biologically plausible algorithm for instantiating this type of longer-term learning (Rao and Ballard 1999; Whittington and Bogacz 2019; see Nour Eddine et al. in press for discussion).

In addition to this relatively slow cortical adaptation, which may be linked to the N400, it is also possible that linguistic anomalies may trigger a more rapid form of learning that is often discussed in the P600 literature (e.g. Coulson et al. 1998; Hanulikova et al. 2012). Notably, in the present study, semantic anomalies produced a distinct reverse-dipole effect in the 600–1,000 ms time-window within medial temporal cortices (see McClelland et al. 1995; O’Reilly and Rudy 2001 for more general discussion of distinct modes of medial temporal function in relation to
slower vs. faster learning). In future studies, it will be important to determine whether and how evoked activity in the medial temporal lobe, in both early and late time-windows, influences different types of longer-term learning.

Conclusions

Of course, no single study can provide definitive evidence for any single model of language comprehension. As we have noted, several of our individual findings, including the graded increase in N400 activity within the temporal cortex, and reprocessing of anomalies within posterior fusiform cortex, are also consistent with other psycholinguistic or neurobiological models. Here, we have interpreted the full pattern of findings within a single computational framework—predictive coding—which has been proposed as a unifying theory of brain function, in multiple domains of perception and cognition (Clark 2013), including lower-level aspects of language processing (Price and Devlin 2011; Blank and Davis 2016; Sohoglu and Davis 2020). Our findings suggest that the computational principles of predictive coding may also explain the time-course of evoked activity produced across the fronto-temporal network that supports higher-level language comprehension.

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Supplementary material

Supplementary material is available at Cerebral Cortex online. Conflict of interest statement: None declared.

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