

# Neural correlate of the construction of sentence meaning

Evelina Fedorenko<sup>a,b,1</sup>, Terri L. Scott<sup>c</sup>, Peter Brunner<sup>d,e</sup>, William G. Coon<sup>d,f</sup>, Brianna Pritchett<sup>g</sup>, Gerwin Schalk<sup>d,e,f</sup>, and Nancy Kanwisher<sup>g,1</sup>

<sup>a</sup>Department of Psychiatry, Harvard Medical School, Boston, MA 02115; <sup>b</sup>Department of Psychiatry, Massachusetts General Hospital, Boston, MA 02114; <sup>c</sup>Department of Speech, Language, and Hearing Sciences, Boston University, Boston, MA 02215; <sup>d</sup>National Center for Adaptive Neurotechnologies, Wadsworth Center, New York State Department of Health, Albany, NY 12208; <sup>e</sup>Department of Neurology, Albany Medical College, Albany, NY 12208; <sup>f</sup>Department of Biomedical Sciences, State University of New York at Albany, Albany, NY 12222; and <sup>g</sup>Brain & Cognitive Sciences Department, McGovern Institute for Brain Research, Massachusetts Institute of Technology, Cambridge, MA 02139

Contributed by Nancy Kanwisher, August 4, 2016 (sent for review February 17, 2016; reviewed by Daphne Bavelier and Ziv M. Williams)

The neural processes that underlie your ability to read and understand this sentence are unknown. Sentence comprehension occurs very rapidly, and can only be understood at a mechanistic level by discovering the precise sequence of underlying computational and neural events. However, we have no continuous and online neural measure of sentence processing with high spatial and temporal resolution. Here we report just such a measure: intracranial recordings from the surface of the human brain show that neural activity, indexed by  $\gamma$ -power, increases monotonically over the course of a sentence as people read it. This steady increase in activity is absent when people read and remember nonword-lists, despite the higher cognitive demand entailed, ruling out accounts in terms of generic attention, working memory, and cognitive load. Response increases are lower for sentence structure without meaning ("Jabberwocky" sentences) and word meaning without sentence structure (word-lists), showing that this effect is not explained by responses to syntax or word meaning alone. Instead, the full effect is found only for sentences, implicating compositional processes of sentence understanding, a striking and unique feature of human language not shared with animal communication systems. This work opens up new avenues for investigating the sequence of neural events that underlie the construction of linguistic meaning.

language | ECoG | compositionality | syntax | semantics

ow does a sequence of sounds emerging from one person's mouth create a complex meaning in another person's mind? Although we have long known where language is processed in the brain (1-3), we still know almost nothing about how neural circuits extract and represent the meaning of a sentence. A powerful method for addressing this question is intracranial recording of neural activity directly from the cortical surface in neurosurgery patients (i.e., electrocorticography or ECoG) (4, 5). Although opportunities for ECoG data collection are rare, determined by clinical—not scientific—priorities, they nonetheless offer an unparalleled combination of spatial and temporal resolution, and further provide direct measures of actual neural activity, rather than indirect measures via blood flow (as in PET, fMRI, and near infrared spectroscopy/optical imaging). ECoG data are particularly valuable for the study of uniquely human functions like language, where animal models are inadequate. Here we used ECoG to identify the neural events that occur online as the meaning of a sentence is extracted and represented.

Prior intracranial recording studies of language have largely focused on speech perception and production (e.g., refs. 6–11) and word-level processes (e.g., refs. 12–26). However, the most distinctive feature of human language is its compositionality: the ability to create and understand complex meanings from novel combinations of words structured into phrases and sentences (27). As a first step toward understanding the neural basis of sentence comprehension, we recorded intracranial responses while participants read sentences and three kinds of control stimuli. In each trial, a string of eight items (words or nonwords) were presented sequentially, enabling us to separately measure the neural response to each item individually; to encourage attention to all stimuli (whether meaningful or not),

participants decided at the end of each trial whether a probe word/ nonword appeared in the preceding string.

Our four stimulus conditions (materials adapted from ref. 28; examples are shown in the "Sample materials" table below) enabled us to orthogonally vary the presence of word meaning and sentence structure. Word-lists (W) included the same words as the sentences (S)

### Sample materials

Condition	Example			
Sentences	STEVE WAS LATE TO SCHOOL BECAUSE HE OVERSLEPT [probe: SCHOOL]			
	THE RED BALLOON ROSE UP INTO THE CLOUDS [probe: WENT]			
Word-lists	RAIN THE WORK BEHIND REACHED GREW KIDS OPENED [probe: GREW]			
	STOOD THE TIED CANDLE INTO SHED THE QUICKLY [probe: WALLET]			
Jabberwocky	THE GAR WAS SWARBING THE MUME FROM ATAR [probe: ATAR]			
	TOMAL HOTHED THE BLESPY NULO DURING THE VAYLANT [probe: FLORKY]			
Nonword-lists	PHREZ CRE EKED PICUSE EMTO PECH CRE ZEIGELY [probe: PHREZ]			
	PIV WUBA WOS PAFFING DEBON TRIENED LE KIF [probe: LOME]			

(scrambled across sentences), but lacked sentence structure. "Jabberwocky" (J) sentences were grammatical but largely meaningless, as they contained no real content words. Nonword-lists (N) contained neither sentence-level structure nor word meaning.

## **Significance**

How do circuits of neurons in your brain extract and hold the meaning of a sentence? To start to address this unanswered question, we measured neural activity from the surface of the human brain in patients being mapped out before neurosurgery, as they read sentences. In many electrodes, neural activity increased steadily over the course of the sentence, but the same was not found when participants read lists of words or pronounceable nonwords, or grammatical nonword strings ("Jabberwocky"). This build-up of neural activity appears to reflect neither word meaning nor syntax alone, but the representation of complex meanings.

Author contributions: E.F., G.S., and N.K. designed research; E.F., T.L.S., P.B., W.G.C., and G.S. performed research; E.F., T.L.S., P.B., B.P., G.S., and N.K. analyzed data; and E.F., T.L.S., G.S., and N.K. wrote the paper.

Reviewers: D.B., University of Geneva; and Z.M.W., Harvard Medical School.

The authors declare no conflict of interest.

<sup>1</sup>To whom correspondence may be addressed. Email: evelina.fedorenko@mgh.harvard. edu or ngk@mit.edu.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1612132113/-/DCSupplemental.

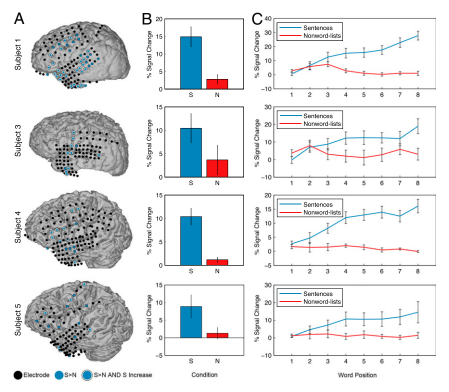


Fig. 1. (A) Cortical models of individual subjects showing all electrodes (black) (see Table 1 for numbers of electrodes), electrodes that show a significant sentences > nonword-lists (S > N) effect in odd-numbered runs (blue), and electrodes that show both a significant S > N effect and a monotonic increase across word positions in the sentence condition in odd-numbered runs, [i.e., our electrodes of interest, EOIs (blue, circled in white)]. (B) The  $\gamma$ -magnitude for sentences and nonword-lists averaged across word positions estimated in even-numbered runs (i.e., data independent from the data used to select the EOIs). (C) The  $\gamma$ -magnitude for sentences and nonword-lists in each of eight word positions in even-numbered runs (see SI Appendix, Part F for additional figures showing data not averaged within each word position). Error bars indicate SEMs over EOIs in both B and C.

Few prior studies have recorded intracranial responses during sentence comprehension. Noninvasive methods with high temporal resolution [event-related brain potential (ERPs) and magnetoencephalography] have revealed neural responses to specific types of events during sentence comprehension, such as syntactic or semantic violations (e.g., refs.29-36), but do not provide a measure of ongoing neural activity from focal cortical regions. Numerous prior studies with fMRI have identified cortical regions that respond strongly (e.g., refs. 28, 37-41) and selectively (42-44) during sentence comprehension (whether presented visually or auditorily) (28, 45), but fMRI lacks the temporal resolution to reveal the responses to individual words as a sentence is presented. Hence, little is known about how neural activity in specific cortical regions unfolds over the course of a sentence as the meaning of that sentence is extracted and represented.

Given the necessarily exploratory nature of this study, we took stringent measures to guard against the dangers of statistical

nonindependence and hidden degrees-of-freedom (46): all hypotheses, data analysis choices, and selection of specific electrodes were made based on analyses of only half the data (odd-numbered runs), before the other half of the data (even-numbered runs) were ever inspected.

Six epilepsy patients with subdurally implanted electrodes placed over left-hemisphere frontal, temporal, and parietal cortices were tested. We measured the time-course at each electrode of broadband  $\gamma$ -activity of the ECoG signal, which is closely related to spiking activity of neuronal populations directly underneath each recording electrode (47–50).

# Results

We first asked whether any electrodes produced a higher  $\gamma$ -response overall during the reading of sentences than nonword strings, as has been reported in numerous previous studies with fMRI (e.g., refs. 28, 51, 52). As expected, numerous electrodes in each subject, distributed across temporal and frontal regions,

Table 1. Numbers of total, analyzed, and language-responsive electrodes, as well as EOIs

Subject no.	Total electrodes	Analyzed electrodes	S > N Electrodes	S > N and S Increasing (i.e.,EOIs)
<b>S1</b>	120	117	45 (0.38*)	27 (0.23 <sup>†</sup> , 0.60 <sup>‡</sup> )
S3	112	84	11 (0.13)	9 (0.11, 0.82)
S4	134	124	15 (0.12)	9 (0.07, 0.60)
S5	98	87	15 (0.17)	6 (0.06, 0.33)

<sup>\*</sup>Proportion of S > N electrodes relative to analyzed electrodes.

 $<sup>^{\</sup>dagger}$ Proportion of S > N and S increasing electrodes relative to analyzed electrodes.

 $<sup>^{\</sup>ddagger}$ Proportion of S > N and S increasing electrodes relative to S > N electrodes.

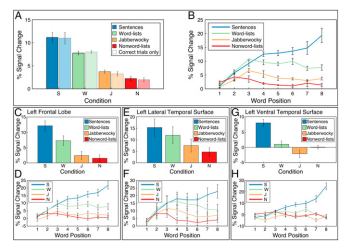


Fig. 2. (A) The γ-magnitude for all conditions (S, sentences; W, word-lists; J, Jabberwocky sentences; N, nonword-lists) averaged across word positions and subjects estimated in even-numbered runs. Darker bars show responses across all trials, lighter bars show responses across the subset of trials for which the memory-probe task was answered correctly. (B) The  $\gamma$ -magnitude for all conditions averaged across subjects in each of eight word positions in even-numbered runs. (C-H) The  $\gamma$ -magnitude for all conditions averaged across word positions (C, E, G) and in each of eight word positions (D, F, H) in even-numbered runs in three groups of EOIs: left frontal (C and D), left lateral temporal (E and F), and left ventral temporal (G and H). All four subjects had some EOIs in left frontal (total: 21 EOIs) and left lateral temporal (total: 20 EOIs) cortex, and two of the four had EOIs in left ventral temporal (total: 9 EOIs) cortex (see SI Appendix, Part G for numbers of EOIs in each group for each subject). Error bars indicate SEMs over subjects in all panels.

responded significantly more strongly to sentences than nonword strings (Fig. 1A and Table 1), when odd-run data only were analyzed (see SI Appendix, Part I for evidence that some electrodes show the opposite pattern, plausibly reflecting sensitivity to task

The most striking and novel finding from the analyses of oddrun data was that over half of the language-responsive electrodes further showed a monotonic increase of γ-power over the eight words in the sentence condition, while failing to increase for the control nonword-list condition. This monotonic increase appears to reflect a previously unreported marker of the neural construction of sentence meaning.

To rigorously test the significance of this finding, for each subject we selected electrodes of interest (EOIs) (Fig. 1A and Table 1) that showed (in odd-run data): (i) a significantly greater mean response to sentences than nonword-lists averaging across the eight word positions, and (ii) a monotonic increase over the eight positions in the sentence condition (Materials and Methods). Fifty-one such EOIs were identified across four subjects, distributed across the frontal and temporal cortex. These four subjects were used in all of the analyses reported in the text (Figs. 1–4, except Fig. 2 G and H). The remaining two subjects—with only two EOIs each-were excluded from the main analyses, but their results were qualitatively and quantitatively similar (SI Appendix, Part D). Response magnitudes of each EOI in each subject were then quantified from even-run data for each condition.

Indeed, the even-run data replicated the higher response to sentences than nonword-lists (P < 0.005) (Fig. 1B and Table 2). Critically, each subject also replicated the monotonic increase in γ-power for sentences (correlation between word position and response, P < 0.05), but not nonword-lists (Fig. 1C and SI Appendix, Part A; see also SI Appendix, Part B for similar results when using even-numbered runs for EOI selection). Reliable differences between the two conditions emerged between the second and fourth word position within the sequence (SI Appendix, Part A), suggesting that the ability to combine two or three words into a coherent representation was necessary for the sentence time-course to diverge significantly from that of the control, nonword-lists condition.

Sentences differ from nonword-lists in the presence of both word meaning and syntax. Is the observed response increase primarily driven by one of these factors? To find out, we measured responses to the two remaining conditions: word-lists (lexical meaning with little syntax) and Jabberwocky sentences (syntax with little meaning) (Sample Materials). Averaging across word positions, responses are highest for sentences, weaker for word-lists and Jabberwocky, and weakest for nonword-lists (Fig. 24 and Table 2), a pattern similar to the one observed previously in fMRI (28, 51). It is worth noting that unlike in fMRI, where the responses to word-lists and Jabberwocky are similar in magnitude (28), the ECoG response to word-lists is generally higher than the response to Jabberwocky (Fig. 24), and significantly so in many EOIs. In fact, across EOIs, each of the four subjects shows a significantly higher response to word-lists than Jabberwocky (P < 0.05) (Table 2). No EOI shows a

Table 2. Results from two-tailed paired samples t tests on the differences in PSC (with respect to baseline fixation) between pairs of experimental conditions

Subject	S–N	S-W	S–J	W-N	J–N	W–J
S1	$12.11 \pm 1.78$ $t(26) = 6.80$ $P < 10^{-6}$	$6.19 \pm 0.58$ $t(26) = 10.67$ $P < 10^{-10}$	$11.42 \pm 1.02$ $t(26) = 11.21$ $P < 10^{-10}$	$5.92 \pm 1.92$ t(26) = 3.08 $P < 10^{-2}$	$0.69 \pm 1.21$ t(26) = 0.57 P = 0.58	$5.23 \pm 0.92$ t(26) = 5.65 $P < 10^{-5}$
S3	$6.76 \pm 1.02$	$3.72 \pm 0.89$	$5.42 \pm 0.81$	$3.04 \pm 0.65$	$1.33 \pm 0.55$	$1.71 \pm 0.23$
	t(8) = 6.58	t(8) = 4.18	t(8) = 6.71	t(8) = 4.67	t(8) = 2.43	t(8) = 7.38
	$P < 10^{-3}$	$P < 10^{-2}$	$P < 10^{-3}$	$P < 10^{-2}$	P = 0.04	$P < 10^{-4}$
S4	$9.21 \pm 1.38$ t(8) = 6.66 $P < 10^{-3}$	$3.17 \pm 0.83$ t(8) = 3.80 $P < 10^{-2}$	$7.59 \pm 1.11$ t(8) = 6.83 $P < 10^{-3}$	$6.04 \pm 1.47$ $t(8) = 4.12$ $P < 10^{-2}$	$1.62 \pm 0.59$ t(8) = 2.75 P = 0.03	$4.42 \pm 1.03$ t(8) = 4.31 $P < 10^{-2}$
S5	$7.58 \pm 3.51$	$0.51 \pm 1.31$	$5.24 \pm 2.29$	$7.08 \pm 2.73$	$2.34 \pm 1.30$	$4.73 \pm 1.60$
	t(5) = 2.16	t(5) = 0.39	t(5) = 2.29	t(5) = 2.59	t(5) = 1.80	t(5) = 2.96
	P = 0.08	P = 0.72	P = 0.07	P = 0.05	P = 0.13	P = 0.03
Across subjects	$8.92 \pm 1.18$	$3.40 \pm 1.16$	$7.42 \pm 1.44$	$5.52 \pm 0.87$	$1.50 \pm 0.34$	$4.02 \pm 0.79$
	t(3) = 7.56	t(3) = 2.91	t(3) = 5.17	t(3) = 6.38	t(3) = 4.37	t(3) = 5.10
	$P < 10^{-2}$	P = 0.06	P = 0.01	$P < 10^{-2}$	P = 0.02	P = 0.01

Data are taken from even-numbered runs only and averaged across all word positions and EOIs. The first row of each cell denotes the average PSC difference with SEMs across EOIs. Significance levels are shaded with white: P < 0.01, midgray:  $0.01 \le P < 0.05$ , and dark-gray:  $P \ge 0.05$ .

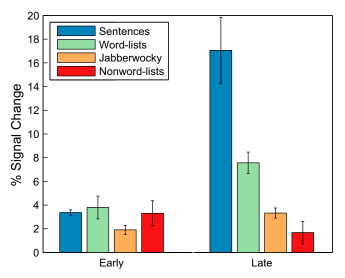


Fig. 3. High  $\gamma$ -magnitude during the early (first two words) and late (last two words) sentence positions estimated across EOIs in even-numbered runs for all conditions. Error bars indicate SEMs over subjects.

significant Jabberwocky > word-lists effect. This asymmetry is consistent with—though not necessarily directly related to—the more robust pattern information for lexical information than syntactic information revealed by multivoxel pattern analyses in fMRI (53).

Critically, the time-courses reveal that neither syntax nor lexical meanings on their own are sufficient to account for the full response increase in the sentence condition (Fig. 2B). This conclusion was supported by several two-way repeated-measures ANOVAs with two within-subjects factors: (i) condition and (ii) position (early = average of the first two words vs. late = average of the last two words). This approach allowed us to test whether the build-up effect in the sentence condition is significantly different from the patterns observed in the other conditions. Including all four conditions, we find a main effect of condition [F(3, 9) = 11.99,P = 0.005], a main effect of position [F(1, 3) = 70.08, P < 0.005], and a highly reliable condition by position interaction [F(3, 9)]18.18, P < 0.001]. Further, 2 × 2 ANOVAs comparing sentences against each of the other conditions revealed reliable interactions for sentences vs. nonword-lists [F(1, 3) = 22.12, P = 0.0182]and sentences vs. Jabberwocky [F(1, 3) = 23.23, P = 0.017], and a marginal interaction for sentences vs. word-lists [F(1, 3) = 9.551]P = 0.0537] (Fig. 3).

Although both the Jabberwocky and word-list conditions show an increase over the first three word positions, the response subsequently decreases, and correlations between word position and y were therefore not significant for either condition (Fig. 2B) and SI Appendix, Part A). The initial increase plausibly reflects participants' attempts to interpret the linguistic signal, a natural tendency given our general experience with language, where words combine to create complex meanings. However, upon realizing that no coherent interpretation is possible, these attempts are likely abandoned. Thus, the continuous response increase in the sentence condition apparently reflects neither syntactic processing nor word-level meaning alone, but the compositional construction of sentence meaning. Consistent with this high-level interpretation of the monotonic increase for sentences but not nonwords, the effect is also seen when the stimuli are presented auditorily rather than visually (SI Appendix, Part H).

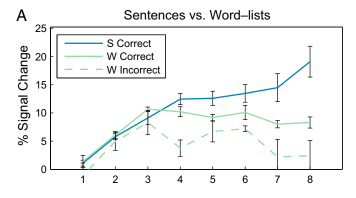
Examining the responses of individual EOIs across subjects suggests that conditions pattern similarly regardless of the EOI's anatomical location. To examine potential anatomical differences more systematically, we split our EOIs into three groups:

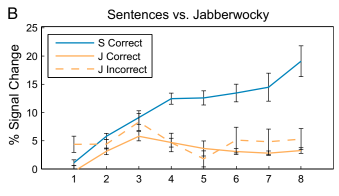
frontal lobe EOIs, EOIs located on the lateral temporal surface, and EOIs located on the ventral temporal surface (see SI Appendix, Part G for numbers of EOIs in each group). As shown in Fig. 2 C-F, the data patterns look remarkably similar for the frontal and lateral temporal EOIs. The left ventral temporal electrodes (Fig. 2 G and H) appear to show greater specificity, with strong build-up for sentences and none at all for the three other conditions. Although intriguing, only two of the four subjects contributed EOIs to this analysis, so it is not clear yet whether this apparent difference in functional response across anatomical locations reflects a general pattern. The distributed nature of the build-up effect contrasts with proposals in the neuroimaging literature that one particular region within the language network houses the core combinatorial/syntactic apparatus: for example, parts of the inferior frontal gyrus (e.g., refs. 54–56) or parts of the anterior temporal cortex (e.g., refs. 57–60). However, it accords with fMRI work that observes sensitivity to lexical and syntactic processing throughout the language network (e.g., refs. 28, 51, 53, 61, 62), and evidence that language regions form a highly integrated functional system (e.g., ref. 63).

### Discussion

Our most striking finding is that many language-responsive electrodes show a monotonic increase in  $\gamma$ -power over the course of the sentence as it is read or heard, and that this increase cannot be explained by the presence of either word meaning or sentence structure alone. What mental processes might underlie this increase in γ response? We can rule out several potential explanations linked to general cognitive factors. First, the build-up effect cannot be explained by general attention or arousal that is higher in the sentence condition than in the other three conditions. Performance on the memory-probe task provides a useful proxy for attention/arousal, and participants generally performed well on the memory-probe task across conditions (SI Appendix, Part C). Although accuracies were numerically higher in the sentence and word-list conditions than in the Jabberwocky and nonword-list conditions, performance was quite good even in the least-accurate (nonword-list) condition [>70%, not including subject 5 (S5), who misunderstood the instructions] (SI Appendix, Part C). These data accord with the subjective impression of performing this task: the nonword-list condition is the hardest, the Jabberwocky and word-list conditions intermediate, and the sentence condition is the easiest. This pattern goes in the opposite direction of that predicted by an attention/arousal/difficulty account. (It is worth noting that a number of electrodes do show a pattern of response indicative of sensitivity to difficulty/effort, with greater responses to nonword-list and Jabberwocky conditions than the sentence condition (SI Appendix, *Part I*), in line with prior fMRI findings (64).

Second, the build-up effect cannot be the result of an overall better performance on the memory-probe task in the sentence condition. One could, in principle, imagine that trials where the memory-probe task is answered correctly are the ones that show an increase over the course of the sentence. Because there are more correctly answered trials in the sentence condition, the build-up effect may emerge in the sentence condition, but not other conditions. This possibility is already unlikely, given the pattern of accuracies across conditions: for example, the wordlist condition does not show a significant build-up despite the fact that the accuracies for that condition are similar to those of the sentence condition. However, to test this possibility directly, we compared the time-courses of high  $\gamma$ -responses for correctly vs. incorrectly answered trials in each of the three conditions that failed to show a build-up effect (we could not perform this comparison for the sentence condition given that there were hardly any incorrectly answered trials). As Fig. 4 shows, the increase does not appear to depend on whether the trial was answered correctly: in the conditions other than the sentence condition, neither the correctly nor the incorrectly answered





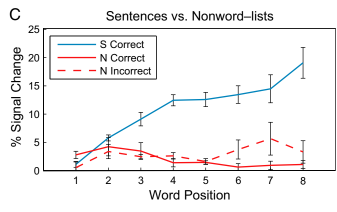


Fig. 4. Comparison of percent signal change (PSC) of correctly and incorrectly answered trials in the word-lists (A), Jabberwocky (B), and nonword-lists (C) conditions, averaged across subjects. For comparison, we include the average of the correctly answered sentence trials (there were too few subjects with incorrectly answered sentence trials to compute averages). Error bars indicate SEMs over subjects.

trials show an increase across positions. Fig. 2A further shows the overall magnitude for the correctly answered trials across the four conditions (lighter bars): the pattern of results looks almost identical to the analysis that includes all trials (darker bars).

Third, the build-up effect cannot be caused by the anticipation of the end of the trial because across conditions all trials were eight words/nonwords-long, thus the patterns should be similar across conditions. Furthermore, according to this hypothesis, one would predict a drop on the last word of the sentences—where the end of the trial is reached—which we do not observe (instead, the response keeps increasing). Similarly, the effect cannot be caused by the preparation to perform the memory-probe task, because the task is the same across conditions.

What about hypotheses that more specifically invoke language processing? Words that occur later in the sentence are more predictable (65-67), and hence easier to process. However, it is unclear why neural activity would increase for words that are easier to process, especially given that the N400 ERP component decreases for words in later sentence positions (67), as do reading times (68, 69). Another possibility is that the response increase reflects strengthening predictions about upcoming material. Evidence against this possibility is that the signal did not decrease substantially at points where a complete clause is formed, and thus no further material is necessary/predicted, or at the following word (SI Appendix, Part E). Thus, the most likely explanation of the response increase is that it reflects the increasing complexity of the evolving representation of the meaning of the sentence. In particular, because items must be held in memory for all conditions, and the steady increase in  $\gamma$  is greatest for sentences, that increase must reflect the representation of structured sentence meaning over and above the representation of individual lexical items.

It may be useful to link these results to classic spreadingactivation theories of lexico-semantic processing (e.g., refs. 70 and 71). In particular, words that occur in the context of phrases or sentences plausibly lead to more robust and longer-lasting representations because they receive reinforcement from related words, which tend to occur in close proximity in natural language (e.g., refs. 72 and 73). Thus, in sentences (compared with lists of unrelated words) the net amount of activation (i) should be higher overall and (ii) should increase as more words are incorporated into the evolving structure/meaning. Alternatively, these results may reflect the construction of a situation model, an abstract mental representation of a verbally described situation or event (e.g., refs. 74-77). Situation models are typically argued to be multidimensional, with the most common dimensions including protagonist, intentionality, causation, time, and space (e.g., ref. 76). Constructing a coherent situation model has been argued by many to be a signature of successful comprehension of a text (e.g., refs. 74, 75, 78-82). Although most empirical work on situation models has focused on discourse-level understanding, the concept also applies to smaller segments of connected text, like clauses and sentences.

To conclude, the increase in  $\gamma$ -response over the course of the sentence constitutes a neural correlate of the evolving representation of sentence-level meaning. This effect appears to be broadly distributed across the fronto-temporal language network rather than being localized to a particular brain region. The precise nature of this build-up effect remains to be determined, and we lay out several questions whose evaluation would help constrain the space of possible hypotheses.

First, is the build-up effect specific to language, or would other meaningful stimuli (e.g., movies) or structured meaningless stimuli (e.g., music) elicit it as well? The regions of the language network have been shown to respond to language stimuli in a highly selective manner (e.g., refs. 42-44), so it is unlikely that nonlinguistic stimuli would produce a similar build-up effect in these regions (although they might produce such an effect in other cortical regions if similar neural phenomena underlie the representation of complex structured representations in these other domains). However, this possibility remains to be evaluated. Second, is a syntactic frame required or would lists of semantically related words (e.g., cat-dog-pig) or words that could be combined to form a complex meaning (e.g., cookie-girl-eat) suffice? Third, what is the temporal scope of the effect? Clearly, the response cannot increase indefinitely; instead, it will plausibly eventually reach a plateau. The longer time-scale dynamics of the y build-up remain to be discovered. And fourth, what happens when an incoming word does not fit the structure or meaning constructed so far? From the ERP literature, we know that unexpected words lead to a larger N400 effect (29, 67). However, given that the build-up effect appears to reflect distinct mental computations, it would be important to characterize its behavior with respect to structurally and semantically unexpected words.

In summary, we report here a striking and robust phenomenon: a monotonic increase in neural activity over the course of a sentence as the subject processes it. This effect occurs for sentences, but not for word meanings or syntactic structure alone, ruling out most domain-general accounts and implicating in the effect a quintessential property of language: compositionality. Although these results certainly do not give us a full account of how sentence meaning is constructed and represented neurally, they provide an exciting new window into how that process unfolds over time, and a powerful method for understanding the sequence of neural events that underlie the extraction of complex linguistic meanings in future work.

## **Materials and Methods**

Participants. We recorded electrical activity from intracranial electrodes of six subjects (five female, aged 14-29 y) with intractable epilepsy who read sentences, lists of words, Jabberwocky sentences, and lists of nonwords. These subjects underwent temporary implantation of subdural electrode arrays at Albany Medical College to localize the epileptogenic zones and to delineate it from eloquent cortical areas before brain resection. All of the subjects gave informed written consent to participate in the study, which was approved by the Institutional Review Board of Albany Medical College. Two subjects were excluded from the main analyses because only two electrodes in each subject met our criteria for inclusion in this study. However, their data were qualitatively and quantitatively similar and are included in SI Appendix, Part D. One further subject was tested but excluded from all analyses because of difficulties in performing the task (i.e., pressing multiple keys, looking away from the screen) during the first five runs. After the first five runs, the subject required a long break during which a seizure occurred.

Materials and Procedure. In an event-related design, subjects read sentences, lists of words, Jabberwocky sentences, and lists of nonwords. The materials were adapted from ref. 28. Each event (trial) consisted of eight words/ nonwords, presented one at a time at the center of the screen. At the end of each sequence, a memory probe was presented (a word in the sentence and word-list conditions, and a nonword in the Jabberwocky and nonword-list conditions) and participants had to decide whether or not the probe had appeared in the preceding sequence by pressing one of two buttons. Two different presentation rates were used: S1, S5, and S6 viewed each word/ nonword for 450 ms (fast-timing), and S2, S3, and S4 viewed each word/ nonword for 700 ms (slow-timing). The presentation speed was determined before the experiment and was based on the participant's preferences. After the last word/nonword in the sequence, a fixation cross was presented for 250 ms, followed by the probe item (1.400-ms fast-timing, 1.900 ms slowtiming), and a postprobe fixation (250 ms). Behavioral responses were continually recorded. After each trial, a fixation cross was presented for a variable amount of time, semirandomly selected from a range of durations from 0 to 11,000 ms, to obtain a low-level baseline for neural activity.

Trials were grouped into runs to give participants short breaks throughout the experiment. In the fast-timing version of the experiment, each run included eight trials per condition and lasted 220 s, and in the slow-timing version, each run included six trials per condition and lasted 264 s. The total amount of intertrial fixation in each run was 44 s for the fast-timing version and 72 s for the slow-timing version. All subjects completed 10 runs of the experiment, for a total of 80 trials per condition in the fast-timing version and 60 trials per condition in the slow-timing version.

**Data Collection and Analysis.** The implanted electrode grids consisted of platinum-iridium electrodes that were 4 mm in diameter (2.3–3 mm exposed) and spaced with an interelectrode distance of 0.6 or 1 cm. The total numbers of implanted electrodes were 120, 128, 112, 134, 98, and 36 for the six subjects, respectively (Table 1). Electrodes were implanted on the left hemisphere for all subjects except S6, who had bilateral coverage (16 left hemisphere electrodes). Signals were digitized at 1,200 Hz. Recordings were

synchronized with stimulus presentation and stored using the BCI2000 software platform (83, 84). Upon visual inspection of the recordings, we removed reference electrodes, ground, and electrodes with high noise levels and interictal activity as revealed by independent analyses, which left 117, 84, 84, 124, 87, and 33 electrodes for the six subjects, respectively.

**Cortical Mapping.** We defined the brain anatomy of each subject using preoperative MRI scans, and the location of the electrodes using post-operative computed tomography (CT) imaging. We then created a 3D surface model of each subject's cortex from the MRI images, coregistered it with the location of the electrodes given by the CT images using Curry Software (Compumedics NeuroScan).

Extraction of ECoG Signal Envelope. ECoG recordings were first high-pass-filtered at a frequency of 0.5 Hz and spatially distributed noise common to all electrodes was removed using a common average reference spatial filter. Notch filters removed further noise at 60, 120, 180, and 240 Hz. An IIR bandpass filter was used to select high  $\gamma$ -frequencies (70–170 Hz) and the envelope of the ECoG signal (i.e., the magnitude of the analytic signal) in the high  $\gamma$ -band was computed by taking the absolute value of the Hilbert transform of the resulting signal. The signal envelopes were further low-pass-filtered at 100 Hz and down-sampled to 300 Hz to reduce noise.

## Selecting EOIs.

Step 1: Selection of language-responsive electrodes. Language-responsive electrodes were defined as electrodes in which the envelope of the high  $\gamma$ -signal is significantly higher for trials of the sentence condition than the nonwordlist condition. To do this, we first computed the mean of the signal envelope for each of the eight word positions (time-locked to the onset of each word/ nonword and averaging over the presentation window) in each trial for each condition in each electrode, using data from the odd-numbered runs only (see SI Appendix, Part F for sample EOIs without within-position averaging). We then computed the mean across the eight word positions in each trial for each condition in each electrode. Finally, we correlated the trial means with a vector of condition labels (sentences = 1, nonword-lists = -1). The resulting Spearman's  $\rho$  provided a benchmark against which to test the significance of any positive correlations. The condition labels vector was randomly reordered (via a permutation test without replacement) and a new Spearman's p was computed, and this process was repeated 1.000 times. The fraction of correlations from randomly assigned labels that produced a higher  $\rho$  than the benchmark correlation became our P value. Electrodes with  $P \le 0.01$  and a positive  $\rho$  were included in step 2.

Step 2: Selection of the subset of language-responsive electrodes exhibiting an increase across word positions in the sentence condition. These electrodes were defined as electrodes that exhibited a monotonic increase in high  $\gamma$ -signal over the course of a sentence. To do this, we computed the mean of the high  $\gamma$ -envelope for each word position in the sentence condition and then computed the Spearman's rank correlation coefficient, which measures statistical dependence without assuming linearity, between word position (one through eight) and mean signal magnitude at each position within each sentence trial, separately for each electrode. As in step 1, we again only used the data from odd-numbered runs. A two-tailed one-sample t test was performed and electrodes with mean correlations that were significantly different from zero ( $P \le 0.01$ ) were selected as our EOIs and were included in all of the analyses reported here.

ACKNOWLEDGMENTS. We thank Ted Gibson, Charles Jennings, Roger Levy, Kyle Mahowald, Steve Piantadosi, and Nathaniel Smith for providing helpful comments on this work; Eyal Dechter for help with setting up the experiment; Steve Piantadosi and Kyle Mahowald for providing some word-level features extracted from corpora; and Zuzanna Balewski and Ted Gibson for help with collecting norming data on Mechanical Turk. This research was supported by the NIH (Grants EB00856, EB006356, and EB018783), the US Army Research Office (Grants W911NF-08-1-0216, W911NF-12-1-0109, W911NF-14-1-0440), and Fondazione Neurone. E.F. was supported by National Institute of Child Health and Human Development Award HD-057522.

- Geschwind N (1970) The organization of language and the brain. Science 170(3961): 940–944.
- Binder JR, et al. (1997) Human brain language areas identified by functional magnetic resonance imaging. J Neurosci 17(1):353–362.
- 3. Bates E, et al. (2003) Voxel-based lesion-symptom mapping. Nat Neurosci 6(5):448–450.
- Jacobs J, Kahana MJ (2010) Direct brain recordings fuel advances in cognitive electrophysiology. Trends Cogn Sci 14(4):162–171.
- Mukamel R, Fried I (2012) Human intracranial recordings and cognitive neuroscience. *Annu Rev Psychol* 63:511–537.
- Chang EF, et al. (2010) Categorical speech representation in human superior temporal gyrus. Nat Neurosci 13(11):1428–1432.
- Bouchard KE, Mesgarani N, Johnson K, Chang EF (2013) Functional organization of human sensorimotor cortex for speech articulation. Nature 495(7441):327–332.
- Chang EF, Niziolek CA, Knight RT, Nagarajan SS, Houde JF (2013) Human cortical sensorimotor network underlying feedback control of vocal pitch. *Proc Natl Acad Sci* USA 110(7):2653–2658.
- Bouchard KE, Chang EF (2014) Control of spoken vowel acoustics and the influence of phonetic context in human speech sensorimotor cortex. J Neurosci 34(38):12662–12677.

- 10. Cogan GB, et al. (2014) Sensory-motor transformations for speech occur bilaterally. Nature 507(7490):94-98.
- 11. Leonard MK, Bouchard KE, Tang C, Chang EF (2015) Dynamic encoding of speech sequence probability in human temporal cortex. J Neurosci 35(18):7203-7214.
- 12. Hart J, Jr, et al. (1998) Temporal dynamics of verbal object comprehension. *Proc Natl* Acad Sci USA 95(11):6498-6503.
- 13. Tanji K, Suzuki K, Delorme A, Shamoto H, Nakasato N (2005) High-frequency gammaband activity in the basal temporal cortex during picture-naming and lexical-decision tasks | Neurosci 25(13):3287-3293
- 14. Mainy N, et al. (2008) Cortical dynamics of word recognition. Hum Brain Mapp 29(11): 1215-1230
- 15. Towle VL, et al. (2008) ECoG gamma activity during a language task: Differentiating expressive and receptive speech areas. Brain 131(Pt 8):2013–2027.
- 16. Sahin NT, Pinker S, Cash SS, Schomer D, Halgren E (2009) Sequential processing of lexical, grammatical, and phonological information within Broca's area. Science 326(5951):445-449.
- 17. Edwards E, et al. (2010) Spatiotemporal imaging of cortical activation during verb generation and picture naming. NeuroImage 50(1):291-301.
- 18. Kellis S, et al. (2010) Decoding spoken words using local field potentials recorded from the cortical surface. J Neural Eng 7(5):056007.
- 19. Thampratankul L, et al. (2010) Cortical gamma oscillations modulated by word association tasks: Intracranial recording. Epilepsy Behav 18(1-2):116-118.
- 20. Flinker A, Chang EF, Barbaro NM, Berger MS, Knight RT (2011) Sub-centimeter language organization in the human temporal lobe. Brain Lang 117(3):103-109.
- 21. Pei X, Barbour DL, Leuthardt EC, Schalk G (2011) Decoding vowels and consonants in spoken and imagined words using electrocorticographic signals in humans. J Neural Ena 8(4):046028.
- 22. Conner CR, Chen G, Pieters TA, Tandon N (2014) Category specific spatial dissociations of parallel processes underlying visual naming. Cereb Cortex 24(10):2741-2750.
- 23. Kojima K, et al. (2013) Gamma activity modulated by picture and auditory naming tasks: Intracranial recording in patients with focal epilepsy. Clin Neurophysiol 124(9): 1737-1744.
- 24. Trébuchon A, Démonet JF, Chauvel P, Liégeois-Chauvel C (2013) Ventral and dorsal pathways of speech perception: An intracerebral ERP study. Brain Lang 127(2):273-283.
- 25. Cibelli ES, Leonard MK, Johnson K, Chang EF (2015) The influence of lexical statistics
- on temporal lobe cortical dynamics during spoken word listening. Brain Lang 147:66-75. 26. Flinker A, et al. (2015) Redefining the role of Broca's area in speech. Proc Natl Acad Sci
- 27. von Humboldt W (1836) On language. On the Diversity of Human Language Construction and Its Influence on the Mental Development of the Human Species, ed Losonsky M (Cambridge Univ Press, New York).
- 28. Fedorenko E, Hsieh PJ, Nieto-Castañón A, Whitfield-Gabrieli S, Kanwisher N (2010) New method for fMRI investigations of language: Defining ROIs functionally in individual subjects. J Neurophysiol 104(2):1177-1194.
- 29. Kutas M, Hillyard SA (1980) Reading senseless sentences: Brain potentials reflect semantic incongruity. Science 207(4427):203-205.
- 30. Osterhout L, Holcomb PJ (1992) Event-related brain potentials elicited by syntactic anomaly. J Mem Lang 31(6):785-806.
- 31. Hagoort P, Brown C, Groothusen J (1993) The syntactic positive shift (SPS) as an ERP measure of syntactic processing. Lang Cogn Process 8(4):439–483.
- 32. Hagoort P (2008) The fractionation of spoken language understanding by measuring electrical and magnetic brain signals. Philos Trans R Soc Lond B Biol Sci 363(1493): 1055-1069.
- 33. Van Petten C, Luka BJ (2012) Prediction during language comprehension: Benefits, costs, and ERP components. Int J Psychophysiol 83(2):176-190.
- 34. Kaan E (2007) Event-related potentials and language processing: A brief overview. Lang Linguist Compass 1(6):571-591.
- 35. Kielar A, Panamsky L, Links KA, Meltzer JA (2015) Localization of electrophysiological responses to semantic and syntactic anomalies in language comprehension with MEG. Neurolmage 105:507-524.
- 36. Halgren E, et al. (2002) N400-like magnetoencephalography responses modulated by semantic context, word frequency, and lexical class in sentences. NeuroImage 17(3): 1101-1116
- 37. Kuperberg G, et al. (2003) Distinct patterns of neural modulation during the processing of conceptual and syntactic anomalies. J Cogn Neurosci 15(2):272-293.
- 38. Bavelier D, et al. (1998) Hemispheric specialization for English and ASL: Left invariance-right variability. NeuroReport 9(7):1537-1542.
- 39. Robertson DA, et al. (2000) Functional neuroanatomy of the cognitive process of mapping during discourse comprehension. Psychol Sci 11(3):255–260.
- 40. Noppeney U, Price CJ (2004) An FMRI study of syntactic adaptation. J Cogn Neurosci 16(4):702-713.
- 41. Snijders TM, et al. (2009) Retrieval and unification of syntactic structure in sentence comprehension: An FMRI study using word-category ambiguity. Cereb Cortex 19(7):
- 42. Fedorenko E, Behr MK, Kanwisher N (2011) Functional specificity for high-level linguistic processing in the human brain. Proc Natl Acad Sci USA 108(39):16428-16433.
- 43. Monti MM, Parsons LM, Osherson DN (2009) The boundaries of language and thought in deductive inference. Proc Natl Acad Sci USA 106(30):12554-12559
- 44. Monti MM, Parsons LM, Osherson DN (2012) Thought beyond language: Neural dissociation of algebra and natural language. Psychol Sci 23(8):914–922.
- 45. Braze D. et al. (2011) Unification of sentence processing via ear and eye: An fMRI study. Cortex 47(4):416-431.
- 46. Kriegeskorte N. Simmons WK, Bellgowan PS, Baker CI (2009) Circular analysis in systems neuroscience: The dangers of double dipping. Nat Neurosci 12(5):535-540.

- 47. Manning JR, Jacobs J, Fried I, Kahana MJ (2009) Broadband shifts in local field potential power spectra are correlated with single-neuron spiking in humans. J Neurosci 29(43):13613-13620.
- 48. Miller KJ, Sorensen LB, Ojemann JG, den Nijs M (2009) Power-law scaling in the brain surface electric potential. PLoS Comput Biol 5(12):e1000609.
- 49. Ray S, Maunsell JH (2011) Different origins of gamma rhythm and high-gamma activity in macaque visual cortex. PLoS Biol 9(4):e1000610.
- 50. Whittingstall K, Logothetis NK (2009) Frequency-band coupling in surface EEG reflects spiking activity in monkey visual cortex. Neuron 64(2):281–289.
- 51. Bedny M, Pascual-Leone A, Dodell-Feder D, Fedorenko E, Saxe R (2011) Language processing in the occipital cortex of congenitally blind adults. Proc Natl Acad Sci USA 108(11):4429-4434
- 52. Pallier C, Devauchelle AD, Dehaene S (2011) Cortical representation of the constituent structure of sentences. Proc Natl Acad Sci USA 108(6):2522-2527.
- Fedorenko E, Nieto-Castañon A, Kanwisher N (2012) Lexical and syntactic representations in the brain: An fMRI investigation with multi-voxel pattern analyses. Neuropsychologia 50(4):499-513.
- 54. Friederici AD, Bahlmann J, Heim S, Schubotz RI, Anwander A (2006) The brain differentiates human and non-human grammars: Functional localization and structural connectivity. Proc Natl Acad Sci USA 103(7):2458-2463.
- 55. Grodzinsky Y, Friederici AD (2006) Neuroimaging of syntax and syntactic processing. Curr Opin Neurobiol 16(2):240-246.
- 56. Hagoort P (2005) On Broca, brain, and binding: A new framework. Trends Cogn Sci 9(9):416-423
- 57. Vandenberghe R. Nobre AC. Price CJ (2002) The response of left temporal cortex to sentences. J Cogn Neurosci 14(4):550-560.
- 58. Brennan J. et al. (2012) Syntactic structure building in the anterior temporal lobe during natural story listening. Brain Lang 120(2):163-173.
- 59. Bemis DK, Pylkkänen L (2013) Basic linguistic composition recruits the left anterior temporal lobe and left angular gyrus during both listening and reading. Cereb Cortex 23(8):1859-1873.
- Baron SG, Osherson D (2011) Evidence for conceptual combination in the left anterior temporal lobe. NeuroImage 55(4):1847-1852.
- 61. Blank I, Balewski Z, Mahowald K, Fedorenko E (2016) Syntactic processing is distributed across the language system. NeuroImage 127:307-323.
- 62. Bautista A, Wilson SM (2016) Neural responses to grammatically and lexically degraded speech. Lang Cogn Neurosci 31(4):567-574.
- 63. Blank I, Kanwisher N, Fedorenko E (2014) A functional dissociation between language and multiple-demand systems revealed in patterns of BOLD signal fluctuations. J Neurophysiol 112(5):1105-1118.
- 64. Fedorenko E, Duncan J, Kanwisher N (2013) Broad domain generality in focal regions of frontal and parietal cortex. Proc Natl Acad Sci USA 110(41):16616-16621.
- 65. Marslen-Wilson W, Tyler LK (1975) Processing structure of sentence perception. Nature 257(5529):784-786.
- 66. Levy R (2008) Expectation-based syntactic comprehension. Cognition 106(3):1126-1177.
- 67. Kutas M, Federmeier KD (2011) Thirty years and counting: finding meaning in the N400 component of the event-related brain potential (ERP). Annu Rev Psychol 62:621-647.
- 68. Graesser AC (2013) Prose Comprehension Beyond the Word (Springer Science & Business Media, Medford, MA).
- 69. Smith NJ, Levy R (2013) The effect of word predictability on reading time is logarithmic. Cognition 128(3):302-319.
- 70. Quillian MR (1969) The teachable language comprehender: A simulation program and theory of language. Commun ACM 12(8):459–476
- 71. Collins AM, Loftus EF (1975) A spreading-activation theory of semantic processing. Psychol Rev 82(6):407-428.
- 72. Lund K, Burgess C (1996) Producing high-dimensional semantic spaces from lexical cooccurrence. Behav Res Methods Instrum Comput 28(2):203-208.
- 73. Landauer TK, Dumais ST (1997) A solution to Plato's problem: The latent semantic analysis theory of acquisition, induction, and representation of knowledge. Psychol Rev 104(2):211-240.
- 74. Johnson-Laird PN (1983) Mental Models: Towards a Cognitive Science of Language, Inference, and Consciousness (Harvard Univ Press, Cambridge, MA).
- 75. van Dijk TA, Kintsch W (1983) Strategies in Discourse Comprehension (Academic, New York).
- 76. Gernsbacher MA (1990) Language Comprehension as Structure Building (Erlbaum, Hillsdale, NJ).
- 77. Zwaan RA, Radvansky GA (1998) Situation models in language comprehension and memory. Psychol Bull 123(2):162-185.
- 78. Johnson-Laird PN (1989) Mental models. Foundations of Cognitive Science, ed Posner MI (MIT Press, Cambridge, MA), pp 469-499.
- 79. Perfetti CA (1989) There are generalized abilities and one of them is reading. Knowing, Learning, and Instruction: Essays in Honor of Robert Glaser, ed Resnick LB (Erlbaum, Hillsdale, NJ), pp 307-335.
- 80. Graesser AC, Singer M, Trabasso T (1994) Constructing inferences during narrative text comprehension. Psychol Rev 101(3):371-395.
- 81. Zwaan RA, Langston MC, Graesser AC (1995) The construction of situation models in narrative comprehension: An event-indexing model. Psychol Sci 6:292-297.
- 82. Graesser AC, Millis KK, Zwaan RA (1997) Discourse comprehension. Annu Rev Psychol 48:163-189.
- 83. Schalk G, McFarland DJ, Hinterberger T, Birbaumer N, Wolpaw JR (2004) BCI2000: A general-purpose brain-computer interface (BCI) system. IEEE Trans Biomed Eng 51(6): 1034-1043.
- 84. Schalk G, Mellinger JA (2010) Practical Guide to Brain-Computer Interfacing with BCI2000 (Springer, London).