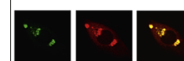


Available online at [www.sciencedirect.com](http://www.sciencedirect.com)

ScienceDirect

[www.elsevier.com/locate/brainres](http://www.elsevier.com/locate/brainres)

Brain Research



## Research Report

Tracking the consequences of morpho-orthographic decomposition using ERPs<sup>☆</sup>Joanna Morris<sup>a,\*</sup>, Jonathan Grainger<sup>b</sup>, Phillip J. Holcomb<sup>c</sup><sup>a</sup>School of Cognitive Science, Hampshire College, Amherst, MA 01002, USA<sup>b</sup>CNRS & Aix-Marseille University, Marseille, France<sup>c</sup>Tufts University, Medford, MA, USA

## ARTICLE INFO

## Article history:

Accepted 10 July 2013

Available online 18 July 2013

## Keywords:

Morphological processing

Masked priming

Event related potentials

Visual word recognition

Morpho-orthographic segmentation

## ABSTRACT

We examined the effects of morpho-orthographic decomposition on complex word processing using a combination of masked priming and ERP recordings. The process of morpho-orthographic decomposition was primed by the prior presentation of complex non-words (formed by the combination a legal stem and legal affix, e.g. huntity, cornity, scanity) as prime stimuli. Targets were semantically transparent complex words (e.g., hunter), semantically opaque pseudocomplex words (e.g., corner), and simplex words (e.g., scandal) that contained the same stem as primes or a different stem (e.g., huntity-hunter vs. farmity-hunter). We found a large early (150–200 ms) priming effect for transparent complex words only, followed by widely distributed priming effects between 200 and 300 ms and more spatially focused N400 priming effects for all types of target. Furthermore, in the 150–200 ms time-window, the ERP waveforms generated by pseudo-complex words patterned with those of complex words, both of which generated less negative-going waveforms compared with simplex words. In the N400 time-window, on the other hand, complex words differed from both pseudocomplex and simplex words. The results provide further support for early morpho-orthographic segmentation processes that operate independently of semantic transparency, and suggest that the output of these processes only affects the subsequent processing of truly complex words.

© 2013 Elsevier B.V. All rights reserved.

## 1. Introduction

Morphology is the aspect of language concerned with the structure and formation of words, and a morpheme is often described as the smallest linguistic unit that serves a grammatical function (Aronoff and Fudeman, 2011). Recognizing a word involves determining what the word means on the basis of its form—its orthographic form in the case of reading, or phonological form in the case of listening. Words can be

similar in meaning (synonyms) or form (homophones or homographs), but morphologically related words share both form and meaning. Morphologically related words therefore serve as an excellent model for examining the ways in which the representation of form and meaning interact during language comprehension. One standard paradigm used to investigate such issues is the priming paradigm, where a given target word is preceded by different types of prime stimuli, and the influence of these different primes on target

<sup>☆</sup>This research was supported by NIH Grant HD25889 to P. J. Holcomb., by ERC grant #230313 to J. Grainger.

\*Corresponding author. Fax: +1 413 559 5438.

E-mail addresses: [jmorris@hampshire.edu](mailto:jmorris@hampshire.edu), [joanna.morris@gmail.com](mailto:joanna.morris@gmail.com) (J. Morris).

word processing is evaluated. Morphologically related primes that share both semantic and orthographic features with targets (hunter-hunt) facilitate processing relative to unrelated primes (singer-hunt). Morphological priming is not simply the product of semantic and orthographic priming, since it can be distinguished from purely orthographic priming on the one hand, and from purely semantic priming on the other. That is, morphological priming produces stronger and longer lasting facilitation than does semantic priming, while orthographic priming tends to result in inhibition (e.g. Drews and Zwitserlood, 1995; Grainger et al., 1991). Thus, morphological priming effects are typically interpreted as reflecting access to some form of explicit representation of morphological structure during the recognition of morphologically complex words.

One key manipulation in prior research involves the distinction between morphologically related words that are also semantically related (semantically transparent morphological relationship: e.g., farmer, farm), compared with morphologically related words that have no obvious semantic relation (semantically opaque morphological relationship: e.g., apartment, apart) and words that only share a superficial morphological relation (pseudo-morphological relationship: e.g., corner, corn). A clear influence of semantic transparency in driving morphological priming effects has been found in studies using paradigms in which the primes are consciously available to participants, for example, delayed repetition priming (Marslen-Wilson and Zhou, 1999), cross-modal priming (Marslen-Wilson et al., 1994), and long stimulus onset asynchrony (SOA) priming (Rastle et al., 2000). However, results from many studies using the masked priming paradigm show statistically equivalent priming effects for semantically transparent primes (hunter-hunt) and semantically opaque or pseudocomplex primes (corner-corn) (e.g. Gold and Rastle, 2007; Lavric et al., 2007; Longtin et al., 2003; McCormick et al., 2008; Rastle et al., 2000, 2004). In the masked priming paradigm, a prime is visually presented for a very brief period, around 30–50 ms, and followed by clearly visible target stimuli. Prime visibility is further reduced by presenting a masking stimulus immediately before primes and, in certain studies, with a brief masking stimulus appearing between prime and target. The short prime duration, as well as the presence of masking stimuli, prevents the subject from consciously perceiving the prime, thereby eliminating any predictive strategies that subjects might use upon noticing the relations between primes and targets. Moreover, the close temporal proximity of the prime and target allow little time for the prime to be processed in isolation from the target, and thus any effects of the prime on responses to the target are presumed to reflect early automatic lexical processes (Forster and Davis, 1984). Therefore the emergence of priming for semantically opaque morphological primes in the masked priming paradigm has been interpreted as reflecting a morphological decomposition procedure that segments visually presented complex words into an orthographically defined stem and affix irrespective of semantic transparency.

Questions nonetheless remain about the point at which semantic information influences the processes involved in complex word recognition. Although many individual studies show statistically non-significant differences in the magnitude

of facilitation for semantically transparent and semantically opaque prime-target targets, some studies have reported significant differences (Diependaele et al., 2005, 2009; Feldman et al., 2009; Morris et al., 2007, for a review see Rastle and Davis, 2008). Moreover, (Crepaldi et al., 2010) have recently shown that irregularly inflected words, which cannot be orthographically decomposed in any simple fashion, prime their base forms more than orthographically matched and semantically unrelated control words. Crepaldi et al. selected nouns and verbs with irregular past tense and plural forms (e.g. fall, mouse) as targets. Each target word was paired with three different primes: the irregular past tense or plural form of the target (e.g. fell-FALL), monomorphemic words that were morphologically unrelated but orthographically related to the targets (e.g. fill-FALL), and finally monomorphemic words that were unrelated either semantically or orthographically to the targets (e.g., hope-FALL). Participants performed a lexical decision task on the targets. Results showed that response times were faster for targets that were preceded by an irregular inflection (fell-FALL) than for those preceded by an orthographically related (fill-FALL) or completely unrelated prime (hope-FALL). Because ‘fell’ and ‘fall’ do not share any orthographically defined morphological units, the ‘fell-FALL’ priming effect cannot be attributed to morpho-orthographic overlap between prime and target, but rather must be attributed to their semantic overlap, providing further support for models of complex word recognition that include a level of morpho-semantic representation located “at some intermediate level between the morpho-orthographic segmentation stage and the semantic system” (Crepaldi et al., 2010, p. 91).

Grainger and colleagues have proposed a dual-route model of morphological processing (Diependaele et al., 2005, *in press*; Morris et al., 2011), in which morphologically complex words activate, in parallel, units at two distinct representational levels: morpho-orthographic and morpho-semantic, the latter via whole-word form representations. Words that share a semantically opaque morphological or pseudo-morphological relationship (e.g. department-depart; corner-corn) prime each other only through shared representations at the morpho-orthographic level. In contrast, words that share a semantically transparent morphological relationship (e.g. hunter-hunt) will not only prime each other via morpho-orthographic representations, but will also be able to prime each other via the shared representations at the morpho-semantic level. This model therefore correctly predicted the pattern of results reported by Crepaldi et al. (2010). The present study provides a further test of this dual-route model, by examining the processing of semantically transparent derived words and semantically opaque pseudo-derived words as targets rather than prime stimuli. Following-up on our previous work we will combine the masked priming procedure with the recording of event-related potentials (ERPs), in order to provide a fine-grained temporal analysis of fast-acting automatic processes involved in the recognition of morphologically complex words

### 1.1. ERPs and masked morphological priming

Prior research using ERPs and masked priming has provided mixed evidence for an early influence of semantic transparency

on masked morphological priming. In a study conducted in our lab [Morris et al. \(2007\)](#) using the lexical decision task and a masked priming paradigm, examined ERPs and reaction times to targets primed by morphologically related words that were semantically transparent (e.g., hunter-hunt), morphologically related words that were semantically opaque (e.g., corner-corn) or orthographically, but not morphologically, related primes (e.g., scandal-scan). In this study both the N250 ERP component, thought to reflect the mapping of sublexical form representations onto whole-word form representations, and the N400 component, thought to reflect the mapping of whole-word form representations onto semantics, were found to be sensitive to semantic transparency (see [Grainger and Holcomb, 2009](#), for a review of N250 and N400 priming effects). Semantically transparent primes generated the largest priming effects, orthographic primes the smallest effects, and semantically opaque primes intermediate effects. However, in an ERP masked priming study testing the same conditions as [Morris et al. \(2007\)](#), [Lavric et al. \(2007\)](#) found no evidence for an influence of semantic transparency on priming effects on the N400 component. Priming effects were the same for targets preceded by semantically transparent and semantically opaque primes, and there was no priming effect for primes that shared an orthographic but no morphological relation. Finally, in a follow-up study ([Morris et al., 2008](#)), we found evidence for equivalent priming from semantically transparent (complex) and semantically opaque (pseudocomplex) primes between 200 and 250 ms post-target onset, with opaque priming effects disappearing between 250 and 300 ms, and priming effects emerging for all types of prime (transparent, opaque, orthographic) on the N400 component. Therefore, as with the evidence obtained from behavioral measures, we can only conclude from the ERP findings that there is evidence that pseudocomplex words (e.g., corner) undergo some form of morpho-orthographic decomposition. However, for the moment it is still not at all clear when semantic transparency begins to influence processing.

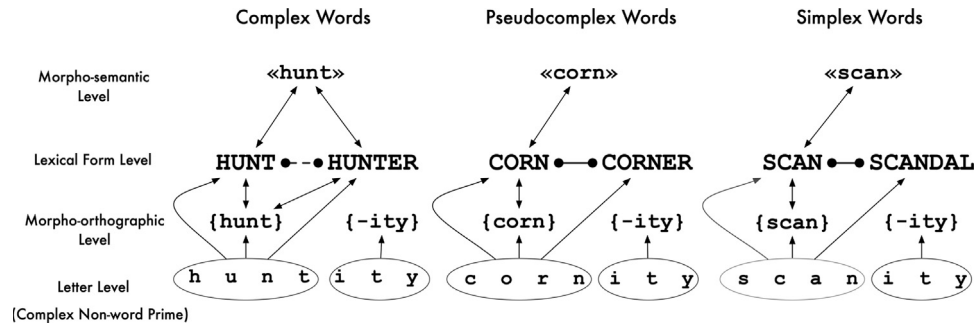
This mixed evidence in the ERP signal for an early phase of morpho-orthographic processing might be due to the fact that target stimuli in our prior studies, following the tradition of current research using masked priming to study morphological processing, were all monomorphemic. In these studies, morphological processes operating on complex prime stimuli are inferred as a result of the subsequent effects of such processes during the recognition of simplex targets (see [Diependaele et al., 2009](#); [Giraudo and Grainger, 2001](#), for exceptions). Countering this tradition, however, in a recent study [Lavric et al. \(2012\)](#) directly compared processing of complex, pseudocomplex, and simplex words in an unprimed lexical decision task with ERP recordings. They found that the ERP waveforms patterned together for complex and pseudocomplex words in an early time-window (190–220 ms post-stimulus onset), both differing from the simplex word ERPs, followed by complex and simplex words patterning together and differing from the pseudocomplex word ERPs in a slightly later time window (246–276 ms post-stimulus onset). This was taken as evidence for early morpho-orthographic decomposition operating on complex and pseudocomplex words, but not simplex words. In the present study we go one critical step further than in the

[Lavric et al. \(2012\)](#) study, by priming the process of morpho-orthographic decomposition prior to the presentation of the different types of target word.

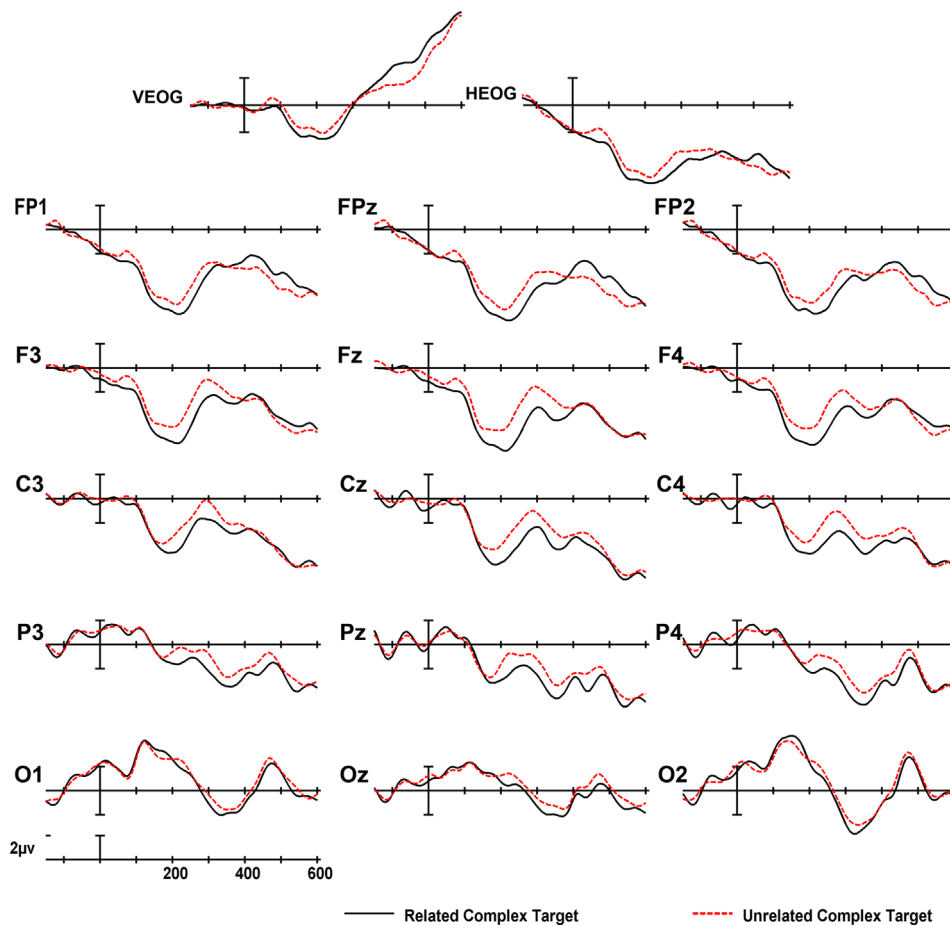
## 1.2. The present study

In the present study we examine responses to complex target words (i.e., words with a true morphological structure such as “hunter”), pseudocomplex target words (i.e., words with a superficial morphological structure such as “corner”), and simplex target words (i.e., words with no morphological structure such as “scandal”) preceded by complex nonword primes that can be related or unrelated to target words. This will allow us to directly examine ERP responses to morpho-orthographic decomposition as it occurs, or not, during the processing of target words of varying levels of complexity. Rather than priming complex targets with primes that are the stems of the corresponding target, as in the [Diependaele et al. \(2009\)](#) and [Giraudo and Grainger \(2001\)](#) studies, here the complex targets were primed by complex nonword primes formed of a combination of the stem of the target and a suffix that did not generate an existing word (e.g., farmity-farmer). The rationale here is that these nonword primes will induce morpho-orthographic decomposition ([Longtin and Meunier, 2005](#)), and we should be able to trace the influence of this morpho-orthographic decomposition during the processing of the complex target in the absence of any morpho-semantic priming. More precisely, in the present study we examined the effects produced by the brief masked presentation of complex nonword primes (e.g., huntity, cornity, scantity) on the recognition of semantically transparent complex words (e.g. hunter), semantically opaque pseudocomplex words (e.g., corner), and simplex words (e.g., scandal). We hypothesize that for each of the three types of target, the complex nonword primes will undergo morpho-orthographic decomposition. We therefore ought to be able to observe the influence of such morpho-orthographic priming during the processing of target words of different levels of complexity.

[Fig. 1](#) describes how this hypothesized decomposition of prime stimuli might differentially affect processing of the three types of target tested in the present study. This figure is adapted from [Morris et al. \(2011\)](#) for the specific conditions tested in the present study. First of all, related primes will generate equivalent orthographic priming effects across all target types (arrow linking letters and lexical form representations in [Fig. 1](#)). Second, related primes will activate embedded stems in the same way for each target type (arrow between letters and morpho-orthographic representations in [Fig. 1](#)). According to our dual-route model, the earliest possible difference in priming effects across target type will arise at the level of morpho-orthographic representations and the mapping of these representations onto lexical form representations. If, as hypothesized in our model, morpho-orthographic stem representations only send activation to whole-word representations having a semantically transparent morphological relationship with the stem (arrow between morpho-orthographic stem representation and whole-word representation of complex words in [Fig. 1](#)), then we should see an early difference in the priming effects to complex targets compared with both pseudocomplex and simplex targets.



**Fig. 1 – Processing of the different types of target word and corresponding related nonword primes tested in the present study (semantically transparent complex words – hunter, semantically opaque pseudocomplex words – corner, simplex words – scandal) according to our dual-route account of morphological processing. The figure shows how each combination of prime stimulus and target word (e.g., huntity-hunter) is processed at the letter, morpho-orthographic, lexical form, and morpho-semantic levels of representation. Excitatory connections are represented by arrows, and inhibitory connections by filled circles. The dashed inhibitory connection between “hunter” and “hunt” represents the modulation of lateral inhibition by semantic transparency.**



**Fig. 2 – ERP waveforms showing the responses to targets related (solid line), and unrelated (dashed line) primes for complex words. Negative voltages are plotted upward. Prime onset is marked by the vertical calibrating bar, and each tick mark on the x-axis represents 100 msec.**

Furthermore, activation of the lexical form representation of the embedded stem should generate inhibitory effects on the processing of both pseudocomplex and simplex targets (lines with filled circles in Fig. 1), and morpho-semantic representations should further facilitate processing of complex targets

(arrows connecting lexical form and morpho-semantic representations in Fig. 1) while having no influence on the processing of pseudocomplex and simplex targets.

Summing up the key predictions of the model presented in Fig. 1 to be tested in the present study: (1) in the earliest



priming effects (voltage differences between related and unrelated prime conditions) we expect to observe a pattern showing the strongest priming for complex targets compared with both pseudocomplex and simplex targets; (2) comparisons across the different types of target should reveal early morpho-orthographic decomposition for complex and pseudocomplex words, which should both differ from simplex words, whereas later differences should reveal effects of semantic transparency (Lavric et al., 2012).

## 2. Results

### 2.1. Visual inspection of ERPs

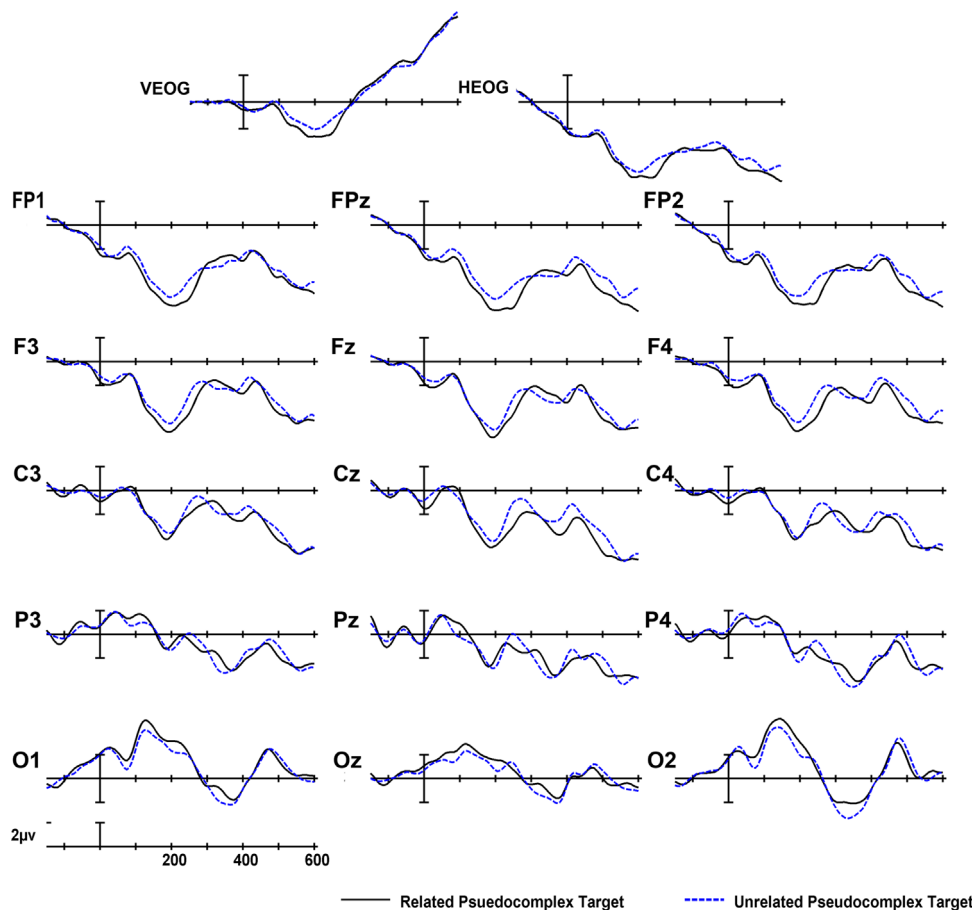
The ERPs for the complex, pseudocomplex and simplex prime conditions from 29 electrode sites are plotted against the ERPs for the corresponding unrelated prime conditions in Figs. 2–4. Fig. 5 shows the topography of ERP priming effects formed by subtracting ERPs to targets following related primes from ERPs to targets following unrelated primes, separately for the complex, pseudocomplex, and simplex target words. Fig. 6 shows the scalp distribution of the effects of target type for the unrelated prime conditions only, in order to remove confounding effects of differences in priming effects across target type. This figure plots the difference in amplitude between ERPs to targets in the (a) complex and

simplex conditions, (b) pseudocomplex and simplex conditions, and (c) pseudocomplex and complex conditions.

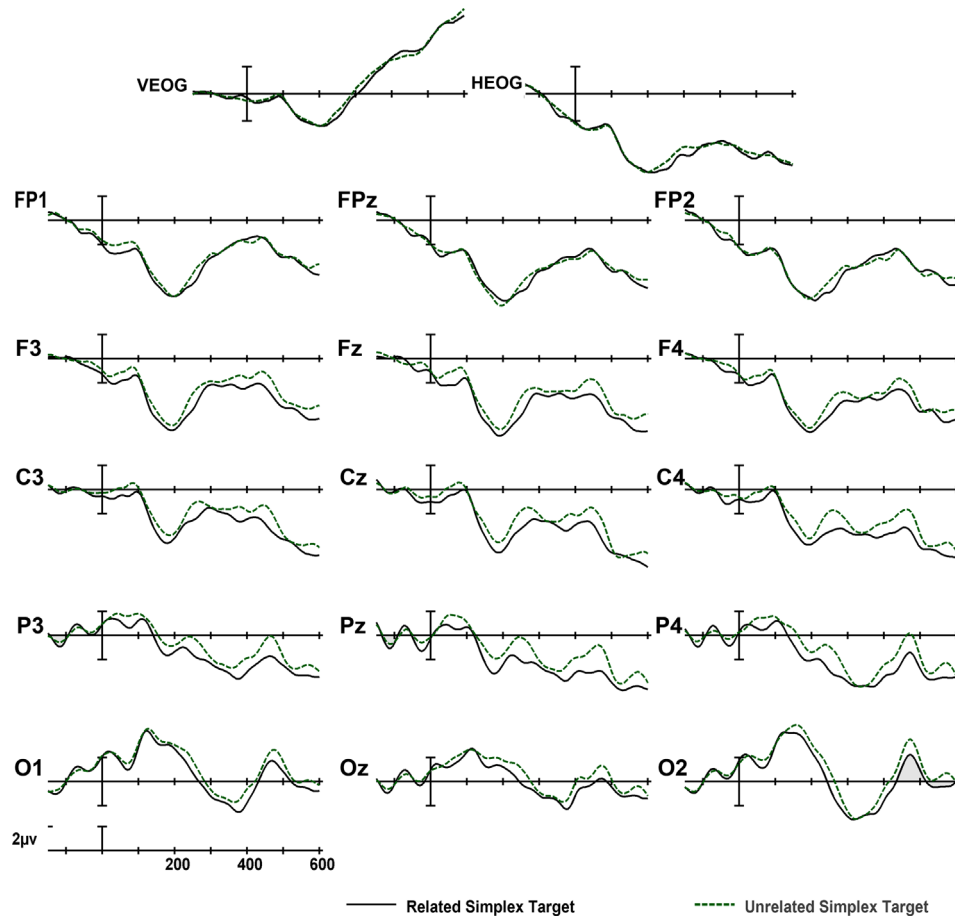
ERPs in the epoch from 0 to 800 ms after the onset of the target produced an initial small negative-going potential peaking between 50 and 100 ms immediately followed by a larger positivity peaking at around 200 ms. Following these early potentials, we observed a series of two negative deflections, the first peaking around 250 ms post-target (N250) and the second at around 400 ms post-target (N400). These two negativities were followed by a large positive deflection (the late positive component, or LPC).

### 2.2. The 150–200 ms time-window

Between 150 and 200 ms after target onset we found a significant RELATEDNESS by TARGET TYPE by ANTERIORITY interaction ( $F(8,176)=5.9$ ,  $p=0.001$ ) and a significant RELATEDNESS by TARGET TYPE by LATERALITY interaction ( $F(4,88)=3.1$ ,  $p=0.03$ ). Follow-up analyses revealed a marginally significant interaction between RELATEDNESS and TARGET TYPE at frontal pole sites ( $\alpha=0.01$ ,  $F(2,44)=4.23$ ,  $p=0.022$ ), where responses to unrelated targets were significantly more negative than to related targets for complex items only ( $\alpha=0.017$ ,  $F(1,22)=7.45$ ,  $p=0.012$ ). Given the spatial distribution and timing of this pattern it would appear to reflect an earlier onset of N250 priming for complex words.



**Fig. 3** – ERP waveforms showing the responses to targets related (solid line), and unrelated (dashed line) primes for pseudocomplex words. Negative voltages are plotted upward. Prime onset is marked by the vertical calibrating bar, and each tick mark on the x-axis represents 100 ms.



**Fig. 4 – ERP waveforms showing the responses to targets related (solid line), and unrelated (dashed line) primes for simplex words. Negative voltages are plotted upward. Prime onset is marked by the vertical calibrating bar, and each tick mark on the x-axis represents 100 ms.**

Further follow-up analyses examined the effects of TARGET TYPE separately for the related and unrelated prime conditions<sup>1</sup>. In the unrelated condition there was a significant TARGET TYPE by ANTERIORITY interaction ( $F(8,176)=3.09$ ,  $p=0.029$ ). When examining each level of anteriority separately, we found an effect of TARGET TYPE at occipital sites ( $\alpha=0.01$ ,  $F(2,46)=5.54$ ,  $p=0.008$ ). Pairwise comparisons on the occipital data ( $\alpha=0.017$ ) showed a significant difference between complex and simplex targets ( $p=0.002$ ), a marginally significant difference between pseudocomplex and simplex targets ( $p=0.037$ ), and no significant difference between complex and pseudocomplex targets ( $p>0.5$ ). The pattern of effects of TARGET TYPE with unrelated primes is shown in panel A of Fig. 6.

### 2.3. The 200–300 ms time-window

Between 200 and 300 ms after target onset, in the time-window typically associated with the N250, we found a

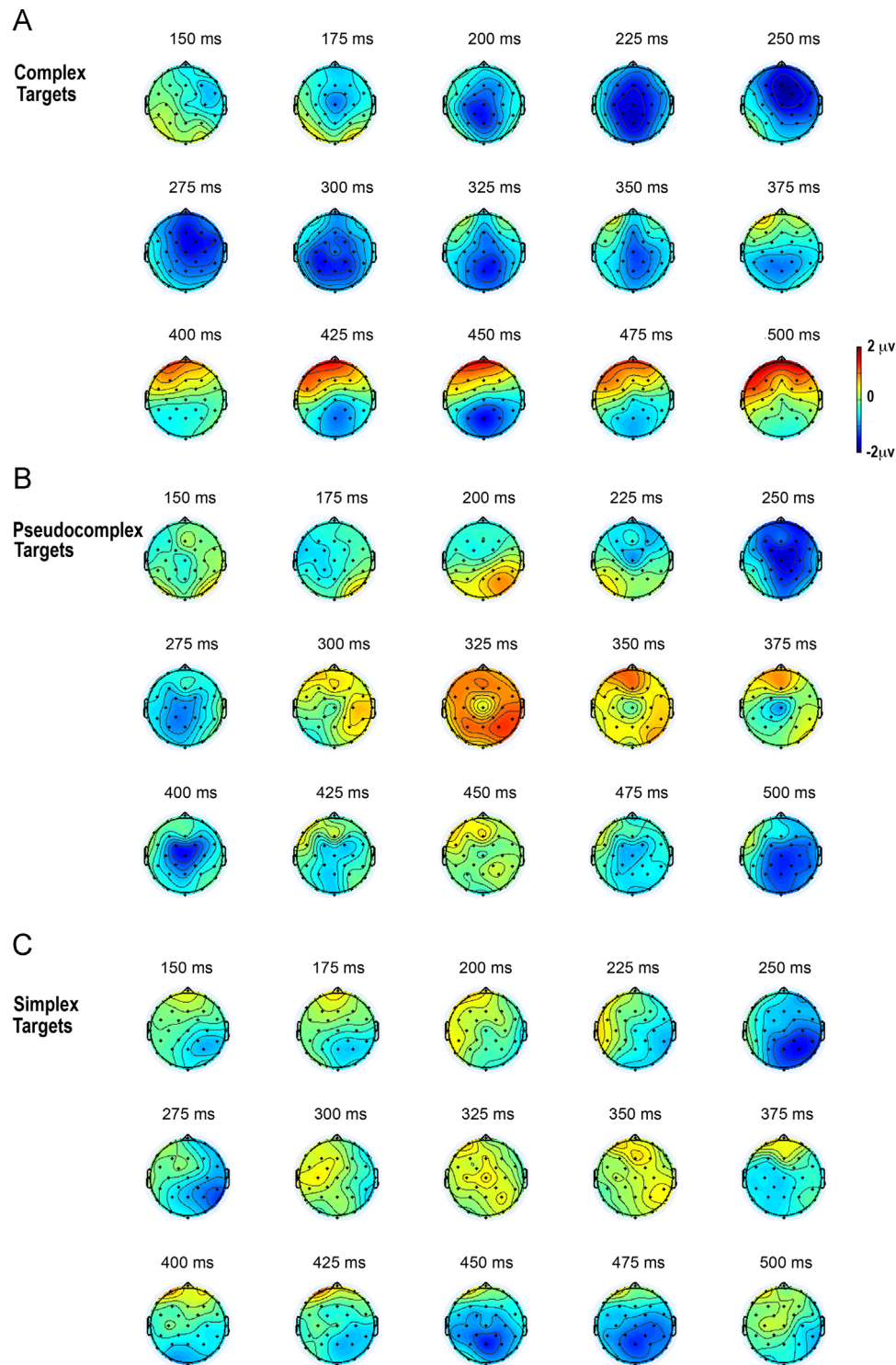
significant main effect of RELATEDNESS ( $F(1,22)=9.43$ ,  $p=0.006$ ), with unrelated primes generating more negative-going waveforms compared to related primes. In addition to the main effect of RELATEDNESS we also found a significant RELATEDNESS by TARGET TYPE by ANTERIORITY interaction ( $F(8,176)=3.16$ ,  $p=0.026$ ).

In a series of follow-up analyses we examined the TARGET TYPE by ANTERIORITY interaction ( $\alpha=0.017$ ) for related and unrelated targets separately. We found that this interaction was not significant for either related or unrelated targets (all  $ps>0.05$ ). Inspection of Fig. 5 suggests that the significant RELATEDNESS by TARGET TYPE by ANTERIORITY interaction likely reflects the slightly more anterior distribution of priming effects for complex targets plus the clearly posterior distribution of priming effects for simplex targets, even if such distributional differences are not significant when examined separately for each Target Type.

### 2.4. The 350–500 ms time-window

In this time-window we found a significant main effect of TARGET TYPE ( $F(2,44)=6.41$ ,  $p=0.005$ ). Follow-up analyses revealed that responses to transparent complex targets were less negative than to both pseudocomplex targets ( $\alpha=0.017$ ,  $F(1,22)=7.05$ ,  $p=0.014$ ) and simplex targets ( $\alpha=0.017$ ,  $F(1,22)=17.04$ ,  $p<0.001$ ), whereas responses to pseudocomplex and

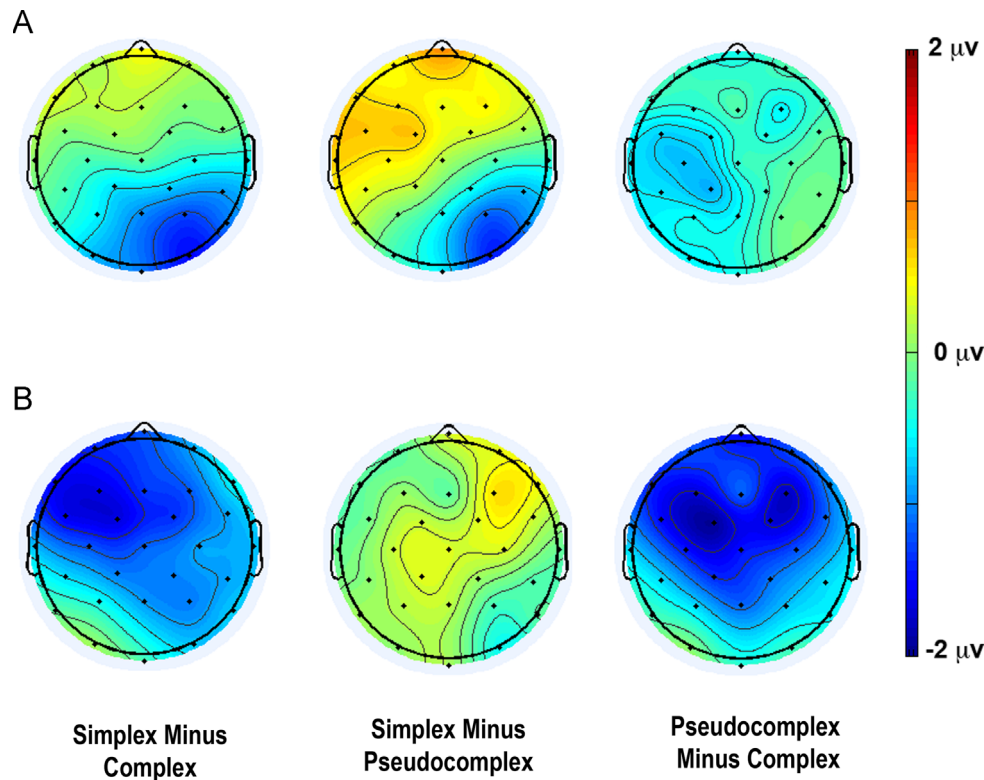
<sup>1</sup>Although we generally avoid direct comparisons of ERPs generated by different items, since this runs the risk of confounding condition and item effects which can both influence early ERPs, here this comparison is motivated by the hypotheses under test and it also provides a comparison point with the Lavric et al. (2012) study. To this end we very carefully matched the different target types on a number of variables (see Table 1).



**Fig. 5 – The spatial distribution and time-course of priming effects for the complex (top), pseudocomplex (middle), and simplex (bottom) target words. These voltage maps were formed from difference waves calculated by subtracting voltages in the related prime condition from voltages in the unrelated prime condition.**

simplex targets did not differ ( $p=0.6$ ). In addition to the effects of TARGET TYPE, in this time window we also found significant interactions between RELATEDNESS and ANTERIORITY ( $F(4,88)=4.61$ ,  $p=0.025$ ) and TARGET TYPE and ANTERIORITY ( $F(8,176)=3.22$ ,  $p=0.027$ ). Follow-up analyses ( $\alpha=0.01$ ) revealed that the RELATEDNESS by ANTERIORITY interaction was driven by

greater priming effects at posterior sites—a topographic distribution typical of the N400 (central:  $F(1,22)=4.7$ ,  $p=0.041$ ; parietal:  $F(1,22)=10.16$ ,  $p=0.004$ ; occipital:  $F(1,22)=5.7$ ,  $p=0.026$ ). In contrast, the effects of TARGET TYPE on the N400 were greater at anterior than at posterior sites. Responses to complex targets were less negative than to those of both



**Fig. 6 – The spatial distribution of the effects of target type found in the 150–200 ms time-window (panel A) and the 350–500 ms time window (panel B). For each time-window effects of the three possible pairwise comparisons enabling the calculation of voltage differences are shown: simplex vs. complex, simplex vs. pseudocomplex, and pseudocomplex vs. complex. Only data from unrelated prime conditions were used in these calculations.**

pseudocomplex targets (frontal:  $F(1,22)=7.22$ ,  $p=0.013$ ; central:  $F(1,22)=11.32$ ,  $p=0.003$ ; parietal:  $F(1,22)=7.72$ ,  $p=0.011$ ) and simplex targets (frontal pole:  $F(1,22)=8.72$ ,  $p=0.007$ ; frontal:  $F(1,22)=14.14$ ,  $p=0.001$ ; central:  $F(1,22)=31.48$ ,  $p<0.001$ ; parietal:  $F(1,22)=6.47$ ,  $p=0.019$ ).

Although we did not find a significant interaction between TARGET TYPE and RELATEDNESS in this time window time-window, we examined the effects of TARGET TYPE for the unrelated prime condition only in order to remove the possibility that the more positive responses of complex targets were simply caused by priming from related primes (see Fig. 6 panel B). We again found a significant main effect of TARGET TYPE ( $F(2,44)=6.0$ ,  $p=0.007$ ). The responses to transparent complex targets ( $\alpha=0.017$ ) were less negative than to simplex targets ( $F(1,22)=16.58$ ,  $p=0.001$ ). The difference between complex and pseudocomplex targets was also marginally significant ( $F(1,22)=5.02$ ,  $p=0.036$ ), and responses to pseudocomplex and simplex targets did not differ ( $p=0.35$ ).

### 3. Discussion

Semantically transparent complex words (e.g., hunter), semantically opaque pseudocomplex words (e.g., corner), and simplex words (e.g., scandal) served as targets in a masked priming study where primes were formed of the embedded stem/pseudostem/word combined with a suffix to form a non-existing word (huntity, cornity, scanity).

Effects of these related primes were measured relative to unrelated primes from the same stimulus category, using EEG recordings while participants monitored target words for animal names. We found priming effects for complex words, but not for pseudocomplex or simplex words in an early time window spanning 150–200 ms post-target onset. Following that, all types of target showed widely distributed priming effects in the time window typically associated with the N250 ERP component, and following that, more spatially focused priming effects in the N400 time window. In all cases, related primes produced less negative-going waveforms than unrelated primes. Effects of target word complexity, independently of priming, were found in the 150–200 ms epoch as well as in the N400 epoch. In the early time window, complex target words in the unrelated prime condition patterned with pseudocomplex targets, which both generated less negative-going waveforms than simplex words. In the N400 epoch, on the other hand, pseudocomplex words patterned with simplex words which both generated more negative-going waveforms than complex words (see Fig. 6 panel A).

The first important finding of the present study is the significant priming effect that was only seen for semantically transparent complex target words in the earliest time-window, between 150 and 200 ms post-target onset. The ERP waveforms generated by semantically transparent complex targets (e.g., hunter) were significantly more positive-going following a related nonword prime (huntity) compared with unrelated nonword primes (farmity) in frontal electrode sites.



Given the timing and the spatial distribution of this effect, we think that it likely reflects an early onset of priming on the N250 component (see Grainger et al., 2012, for a similar early onset of N250 priming with an anterior distribution). Given that there were no significant priming effects for either semantically opaque pseudocomplex targets or simplex targets in this time-window, the early priming effect would appear to be driven by processes that are specific to semantically transparent complex words. Within the framework of our dual-route model presented in Fig. 1, processing specific to semantically transparent complex words first arises during the mapping of sublexical morpho-orthographic representations onto the whole-word orthographic representations of morphologically complex words. That is, although pseudocomplex words might automatically undergo morpho-orthographic segmentation, the activation of the morpho-orthographic representation of the embedded stem (e.g., “corn” activated upon presentation of “corner”) would not directly influence activation of the whole-word orthographic representation of “corner”. In other words, although the process of morpho-orthographic segmentation might be blind to whether or not the stimulus is truly morphologically complex, the subsequent use of the segments extracted by this process is determined by whether or not the stimulus is truly complex. In line with this proposal is the fact that Diependaele et al. (2009) reported behavioral priming effects with stem primes and complex targets only when targets were transparent complex words.

Following the early priming effect for semantically transparent complex words, we found priming effects for all three types of target in the time window typically associated with the N250 ERP component. This general priming effect could simply be driven by the increased orthographic overlap between primes and targets in the related prime condition. N250 orthographic priming effects with nonword primes have been reported in several prior studies (e.g., Grainger et al., 2006; Holcomb and Grainger, 2006; see Grainger and Holcomb, 2009, for a review). Orthographically related primes are thought to facilitate the mapping of sublexical form representations onto whole-word form representations, and the efficiency of this mapping is thought to be reflected in a modulation of N250 amplitude (Grainger and Holcomb, 2009). Our results suggest that when targets are not embedded in prime stimuli, as is the case in standard morphological priming (e.g., hunter-hunt; corner-corn; scandal-scan), then N250 priming effects are mainly determined by non-morphological orthographic overlap (i.e., number of letters shared by prime and target). We would nevertheless argue that the early priming effect (150–200 ms post-target onset) seen only for complex targets most likely reflects an earlier onset of N250 priming in this condition, rather than priming of a distinct ERP component. This early morphological priming would be driven by the combined influence of letter-level information and morpho-orthographic representations in determining the onset of N250 priming effects (see Fig. 1).

In the N400 time-window, we again observed robust priming effects for all target types, which were albeit more spatially focused than the N250 priming effects. Related primes generated a reduced negativity compared with unrelated primes, and the size of this priming effect was

comparable for the three types of target word. This can be interpreted as orthographic priming, generated by differences in prime-target orthographic overlap (which was the same across target types), affecting the build-up in activation of whole-word orthographic representations for target words, and the subsequent mapping between such whole-word representations and semantics (Holcomb and Grainger, 2006; Grainger and Holcomb, 2009).

However, independently of priming effects, the nature of the target item differentially affected ERPs in the present study. In the early 150–200 ms time window, we found that in the unrelated prime condition ERPs to complex and pseudocomplex targets patterned together, and both were significantly less negative-going than the ERPs to simplex targets (although only marginally so for the pseudocomplex targets). This result is in line with the pattern reported by Lavric et al. (2012) in an ERP study comparing the processing of these three types of target word without any priming manipulation. Like Lavric et al. (2012), we take this early pattern as reflecting automatic morpho-orthographic decomposition operating on complex and pseudocomplex target words alike. In line with these findings are the results from MEG studies (Lewis et al., 2011; Solomyak and Marantz, 2010) showing that activity originating in the left fusiform gyrus and peaking between 140 and 190 ms after the onset of a visually presented word (the M170 component) increases as a function of the decomposability of words. Morphological decomposability was measured as the ratio of the stem frequency to the whole word frequency, and the effect of decomposability was found for both transparent complex words and opaque pseudocomplex words.

In our N400 analysis, on the other hand, there was clear evidence that transparent complex targets generated reduced N400 amplitudes compared with both pseudocomplex and simplex targets (see Fig. 6 panel B). One way to explain these effects of target type on later processing is in terms of an inhibitory influence of embedded words during the processing of target words containing the embedded item. According to the model presented in Fig. 1, these inhibitory effects will be strongest for pseudocomplex and simplex targets, and should be smaller and possibly even absent for transparent complex targets (the dashed line connecting morphologically related whole-word representations in Fig. 1). That is, related primes in the pseudocomplex (corner) and simplex (scandal) target conditions provide bottom-up support not only for the target word but also for the embedded word (corn, scan) that will compete with the target for identification, and this competition is thought to be reduced or even absent for complex targets. The fact that this specific pattern of effects of Target Type emerged in the N400 time-window is in line with the hypothesized role of lexical inhibition in driving these effects (see Massol et al., 2010, for evidence for the time-course of lexical inhibition during priming). Alternatively, it could be the morpho-semantic representation that is shared by a complex word and its embedded stem that is the source of the reduced N400 amplitude for these target words compared with both pseudocomplex and simplex words. This shared morpho-semantic representation would be activated by the complex word target and its embedded stem, and send excitatory feedback to both of these representations. Future research should help clarify how mechanisms such as

lateral inhibition and top-down feedback might underlie processing differences between complex, pseudocomplex, and simplex target words, and computational modeling will certainly play an important role here.

Summing up, the results of the present study are in line with the predictions of the model shown in Fig. 1, according to which sublexical morpho-orthographic decomposition, once it has occurred, will only influence the subsequent processing of transparent complex words. First of all, in the line with the results of Lavric et al. (2012), we found evidence for early morpho-orthographic decomposition in the form of ERP differences between the different types of target in the 150–200 ms epoch. Here both complex and pseudocomplex words patterned together, and differed from simplex words. In the same early time-window, however, ERP priming effects were only evident for complex words. According to the model shown in Fig. 1, this arises because the extraction of the stem from complex prime stimuli, which occurs for all target types, can only influence processing of related complex targets (e.g., huntity-hunter). Neither the stem “corn” extracted from “cornity”, nor the stem “scan” extracted from “scanity” can influence processing of related targets words (cornity-corner; scanity-scandal), given the hypothesized absence of connectivity between the relevant representations in the model (see Fig. 1). Therefore, although we provide further evidence in favor of rapid morpho-orthographic decomposition that is blind to semantic information (operating on pseudocomplex words like “corner”) and even to lexical status (operating on pseudowords like “cornity”), we also show that subsequent processing can only capitalize on such decomposition when reading truly complex words.

## 4. Experimental procedures

### 4.1. Participants

The participants for this study were 27 adults (16 women). The data from three participants were excluded from analysis, two because of excessive heartbeat artifact and one for excessive alpha activity. All participants were recruited from the Tufts University community and paid for their participation. The participants ranged in age from 18 to 22 years (mean 19.3 years). All were right-handed native English speakers with normal or corrected-to-normal vision, and none reported any linguistic or neurological impairment.

### 4.2. Stimuli

The stimuli for this study were 306 prime-target pairs in which the prime consisted of an existing English root combined with a bound suffix to form a morphologically complex nonword (e.g., ‘hunt’+‘-ity’  $\Rightarrow$  ‘huntity’), and targets were either semantically transparent complex words containing the same root as the prime (e.g. huntity-HUNTER), semantically opaque pseudocomplex words containing the same root as the prime (cornity-CORNER), or simplex words that had the prime word embedded at initial position, but for which the letters following the embedded prime word did not form a common English suffix (scanity-SCANDAL). We constructed

two lists each consisting of 306 prime-target pairs. In each list there were 102 semantically transparent complex targets, 102 semantically opaque pseudocomplex targets and 102 simplex targets. Primes were formed by dividing each target into either a stem and affix (in the case of the complex and pseudo-complex words, e.g., hunt-er, corn-er ) or into a possible stem and non-affix letter sequence (in the case of simplex words, e.g., scan-dal). Each stem was then combined with one of 23 possible affixes, taken from the CELEX English database (Baayen et al., 1995) to form a complex non-word prime. The list of affixes and their frequency of use in primes across the three conditions is given in Table 1.

For each type of target, 51 pairs were related in that the stem of the complex nonword prime was the same as the stem of the target, and 51 were unrelated. Related primes in one list were used as unrelated primes in the other list. Therefore, across all participants, each target appeared with both a related and unrelated prime, and the same primes appeared in the related and unrelated conditions, but each participant saw each item in only one condition.

Although we held the targets constant across the related and unrelated conditions, it was obviously not possible to hold the targets constant across the different target type conditions, where the categories in question contain different lexical items. As a result, across target types, targets were not matched on all variables known to affect lexical processing. In particular, affix and non-affix endings varied in frequency. In addition, affix tokens were not equally represented across the morphologically complex and pseudocomplex targets.

We measured semantic transparency using Latent Semantic Analysis (LSA; Landauer and Dumais, 1997). A one-way

**Table 1 – The list of affixes and the frequency with which each was used to construct primes for each of the three types of target.**

	Frequency		
	Complex	Pseudo complex	Simplex
-al	2	2	2
-able	4	4	4
-age	4	5	4
-ance	6	6	6
-ant	5	6	6
-ard	6	5	5
-ary	7	6	6
-ate	3	3	4
-ence	3	3	3
-er	4	3	4
-ern	3	3	3
-ery	4	4	4
-ible	2	3	3
-ic	5	5	5
-ify	4	4	4
-ish	4	5	4
-ist	3	3	2
-ity	5	5	5
-ize	6	7	6
-ment	6	6	6
-ness	6	5	6
-or	4	4	4
-ous	6	5	6

**Table 2 – Length and frequency characteristics of the related primes and targets. Frequencies (log10 scale) were obtained from the CELEX English lexical database (Baayen et al., 1995). Token frequencies were computed per million.**

	Complex		Pseudocomplex		Simplex	
	Mean	SE	Mean	SE	Mean	SE
LSA	0.32	0.02	0.06	0.01	0.06	0.01
Target length	7.08	0.12	6.83	0.13	6.88	0.11
Target CBF	14310.48	492.03	14132.79	498.45	13779.02	570.82
Target NS	0.77	0.12	0.97	0.14	1.02	0.17
Target LF	0.69	0.05	0.81	0.07	0.74	0.06
Nonword prime length	7.61	0.10	7.46	0.11	7.40	0.11
Stem length	4.49	0.09	4.34	0.09	4.26	0.07
Stem CBF	6836.33	383.92	7115.82	356.47	7481.60	373.00
Stem NS	5.62	0.48	6.53	0.56	6.94	0.52
Stem LF	1.40	0.07	1.37	0.08	1.31	0.08
Suffix length	2.59	0.07	2.49	0.08	2.62	0.09

ANOVA comparing LSA scores for the three types of prime/target pairs revealed a significant difference ( $F(2,285)=107.5$ ,  $p<0.001$ ). Post-hoc tests showed that the LSA scores of transparent pairs differed from those of both pseudocomplex ( $M_{\text{diff}}=0.26$  ( $SE=0.02$ ),  $p<0.001$ ), and orthographic pairs ( $M_{\text{diff}}=0.26$  ( $SE=0.02$ ),  $p<0.001$ ) but the scores of the latter two types of prime/target pairs did not differ ( $M_{\text{diff}}=0.002$  ( $SE=0.02$ ),  $p=1.0$ ). Across the three conditions, targets were equated for length, cumulative bigram frequency (CBF), neighborhood size (NS), log frequency (LF), stem length, stem cumulative bigram frequency, stem neighborhood size, stem log frequency, and suffix length<sup>2</sup> (see Table 2). Oneway ANOVAs comparing scores on each of these variables for the three types of prime/target pairs revealed no significant differences (all  $ps>0.1$ ).

As the task was semantic categorization with animal names as the target category, we added 124 filler items consisting of 62 animal names each in either the target or the prime position. When the animal names were in the target position, the primes were nonwords similar to those of the experimental items in that they consisted of part of the animal name acting as a stem followed by an affix (e.g., cougance-COUGAR). These primes could be either related or unrelated to the animal name. Animal names were also included in the prime position in order to monitor prime visibility. In these cases, targets resembled those of the experimental pairs in that they were either real complex words, pseudocomplex words or simplex words. Participants detected 89% of the animal targets but only 3% of the animal primes suggesting that the primes were effectively masked.

#### 4.3. Procedure

Participants were seated in a comfortable chair in a darkened room at a distance of 59 in. from the computer monitor. Each testing session began with a short practice block, followed by the experimental block. Participants were told that they would see a list of words appear on the computer monitor and were instructed to monitor the list of words for animal name and press a button on a game controller when an

animal name appeared. There were 31 such words in the prime position and 31 in the target position. Participants were told to read all other words passively (i.e., critical stimuli did not require an overt response). Visual stimuli were presented on a 19 in. monitor, with a diagonal viewable screen size of 18 in., and a width of approximately 14.5 in., with a screen resolution of 800 by 600 pixels set to a refresh rate of 100 Hz (which allows 10-ms resolution of stimulus control). Stimuli were displayed at high contrast as white letters (Verdana font) on a black background. Each letter was 40 pixels tall by 20 pixels wide. Targets which ranged from 5 to 11 characters measured 1.8–3.9 in. and subtended a visual angle of 1.7–3.8°. Primes were presented in lower case letters for 50 ms preceded by a 500 ms random consonant mask. The mask shared no letters in common with the target or with the prime. The target was presented immediately after the prime in upper case letters for 300 ms followed by a 900 ms ISI.

#### 4.4. Recording procedure

The electroencephalogram (EEG) was recorded from 29 active tin electrodes held in place on the scalp by an elastic cap (Electrode-Cap International). In addition to the 29 scalp sites, additional electrodes were attached to below the left eye (to monitor for vertical eye movement/blinks), to the right of the right eye (to monitor for horizontal eye movements), over the left mastoid bone (reference) and over the right mastoid bone (recorded actively to monitor for differential mastoid activity). All EEG electrode impedances were maintained below 5 k $\Omega$  (impedance for eye electrodes was less than 10 k $\Omega$ ). The EEG was amplified by an SA Bioamplifier with a bandpass of 0.01 and 40 Hz and the EEG was continuously sampled at a rate of 200 Hz throughout the experiment.

#### 4.5. Data analysis

ERPs time locked to the onset of target words in each category were formed off-line from trials free of excessive artifact or response error. Trials characterized by EOG artifact in excess of 70 mV were rejected, resulting in 7.9% of trials being discarded. We calculated the mean voltage in each of four time-windows centered at 175, 250, and 425 ms (i.e. 150–200 ms, 200–300 ms, and 350–500 ms after target onset),

<sup>2</sup>For simplex targets this was defined as the number of characters in the target that were not also contained in the embedded stem.



relative to a 100 ms baseline (50 ms pre-stimulus and 50 ms post-stimulus). These time epochs were chosen to minimize temporal overlap between measured components and because they correspond roughly to peaks in the waveforms elicited by the stimuli that were identified by visual inspection, as well as to the time windows associated with a series of electrophysiological effects (N250, N400) that have been hypothesized to correspond to the component processes involved in word recognition (Grainger and Holcomb, 2009). The early 150–200 ms window corresponds to a pre-N250 period that might capture an earlier onset of N250 priming effects in certain conditions. The 50 ms pre and 50 post stimulus baseline was chosen in order to minimize the effect of waveforms elicited by the prime stimulus given the close temporal proximity of the prime and target in this paradigm.

We analyzed the mean amplitude scores by selecting 15 representative sites distributed across the scalp (FP1, FPz, FP2, F3, Fz F4, C3, Cz, C4, P3, Pz, P4, O1, Oz, O2). This method of analyzing ERP data has been widely used in previous studies (Chauncey et al., 2008; Midgley et al., 2009) and provides a thorough analysis of the entire head while reducing the number of sites included in the analysis and hence the complexity of the interpretation. Our model included four within-subjects factors-RELATEDNESS, TARGET TYPE, ANTERIORITY and LATERALITY. The RELATEDNESS factor contrasted mean ERP amplitude scores for the related and unrelated word pairs, while the TARGET TYPE factor contrasted mean ERP amplitude scores for complex, pseudocomplex and simplex targets. To analyze the scalp distribution of the ERP effects, we included two factors, ANTERIORITY and LATERALITY. The ANTERIORITY factor represented the anterior–posterior distribution of effects and included five levels contrasting electrode locations from the back to the front of the head, while the LATERALITY factor represented the left–right distribution and included three levels contrasting electrode locations at left hemisphere, midline and right hemisphere locations.

LIST was included as a dummy variable. As there were no significant main effects of LIST (all  $p$ s > 0.1) we only report results concerning the main effects of the experimentally manipulated factors RELATEDNESS and TARGET TYPE, and the interaction of these factors with the topographic factors ANTERIORITY and HEMISPHERE. The alpha level was set at .05. The Greenhouse-Geisser correction was applied when evaluating effects with more than one degree of freedom in the numerator. In these cases, we report the uncorrected degrees of freedom and the corrected  $p$ -value. For interactions involving the experimental variables RELATEDNESS and TARGET TYPE, we conducted follow-up simple effects tests. As there is no consensual standard for the level of familywise error to use when testing simple effects, we chose to adjust the alpha level by dividing by the number of tests performed (Bonferroni correction).

## REFERENCES

- Aronoff, M., Fudeman, K.A., 2011. What is Morphology? Chichester, West Sussex, UK. Wiley-Blackwell, Malden, MA.
- Baayen, R.H., Piepenbrock, R., Gulikers, L., 1995. The CELEX Lexical Database (CD-ROM). Linguistic Data Consortium, University of Pennsylvania, Philadelphia: PA.
- Chauncey, K., Holcomb, P., Grainger, J., 2008. Effects of stimulus font and size on masked repetition priming: an event-related potentials (ERP) investigation. *Lang. Cognitive Processes* 23 (1), 183–200.
- Crepaldi, D., Rastle, K., Coltheart, M., Nickels, L., 2010. 'Fell' primes 'fall', but does 'bell' prime 'ball'? Masked priming with irregularly inflected primes. *J. Mem. Lang.* 63 (1), 83–99.
- Diependaele, K., Morris, J., Serota, R.M., Bertrand, D., Grainger, J., 2013. Breaking boundaries: letter transpositions and morphological processing. *Lang. Cognitive. Proc.* 28 (7), 988–1003, <http://dx.doi.org/10.1080/01690965.2012.719082>.
- Diependaele, K., Sandra, D., Grainger, J., 2005. Masked cross-modal morphological priming: unravelling morpho-orthographic and morpho-semantic influences in early word recognition. *Lang. Cognitive Processes* 20 (1), 75–114.
- Diependaele, K., Sandra, D., Grainger, J., 2009. Semantic transparency and masked morphological priming: the case of prefixed words. *Mem. Cognit.* 37 (6), 895–908.
- Drews, E., Zwitserlood, P., 1995. Morphological and orthographic similarity in visual word recognition. *J. Exp. Psychol.: Hum. Percept. Perform.* 21 (5), 1098–1116.
- Feldman, L.B., O'Connor, P.A., Del Prado Martín, F.M., 2009. Early morphological processing is morpho-semantic and not simply morpho-orthographic: a violation of form-then-meaning accounts of word recognition. *Psychon. Bull. Rev.* 16 (4), 684–691.
- Forster, K.I., Davis, C., 1984. Repetition priming and frequency attenuation in lexical access. *J. Exp. Psychol.: Learn., Mem. Cognit.* 10 (4), 680–698.
- Giraudo, H., Grainger, J., 2001. Priming complex words: evidence for supralexical representation of morphology. *Psychon. Bull. Rev.* 8 (1), 127–131.
- Gold, B.T., Rastle, K., 2007. Neural correlates of morphological decomposition during visual word recognition. *J. Cognitive Neurosci.* 19 (12), 1983–1993.
- Grainger, J., Holcomb, P.J., 2009. Watching the word go by: on the time-course of component processes in visual word recognition. *Lang. Linguist. Compass* 3 (1), 128–156.
- Grainger, J., Colé, P., Segui, J., 1991. Masked morphological priming in visual word recognition. *J. Mem. Lang.* 30 (3), 370–384.
- Grainger, J., Kiyonaga, K., Holcomb, P.J., 2006. The time course of orthographic and phonological code activation. *Psychol. Sci.* 17 (12), 1021–1026.
- Grainger, J., Lopez, D., Eddy, M., Dufau, S., Holcomb, P.J., 2012. How word frequency modulates masked repetition priming: an ERP investigation. *Psychophysiology* 49, 604–616.
- Holcomb, Grainger, J., 2006. On the time-course of visual word recognition: an ERP investigation using masked repetition priming. *J. Cognitive Neurosci.* 18 (10), 1631–1643.
- Landauer, T.K., Dumais, S.T., 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.
- Lavric, A., Clapp, A., Rastle, K., 2007. ERP evidence of morphological analysis from orthography: a masked priming study. *J. Cognitive Neurosci.* 19 (5), 866–877.
- Lavric, A., Elchlepp, H., Rastle, K., 2012. Tracking hierarchical processing in morphological decomposition with brain potentials. *J. Exp. Psychol.: Hum. Percept. Perform.* 38 (4), 811–816.
- Lewis, G., Solomyak, O., Marantz, A., 2011. The neural basis of obligatory decomposition of suffixed words. *Brain Lang* 118 (3), 118–127.
- Longtin, C.-M., Meunier, F., 2005. Morphological decomposition in early visual word processing. *J. Mem. Lang.* 53 (1), 26–41.
- Longtin, C.-M., Segui, J., Halle, P.A., 2003. Morphological priming without morphological relationship. *Lang. Cognitive Processes* 18 (3), 313–334.



- Marslen-Wilson, W., Zhou, X., 1999. Abstractness, allomorphy, and lexical architecture. *Lang. Cognitive Processes* 14 (4), 321–352.
- Marslen-Wilson, W., Tyler, L.K., Waksler, R., Older, L., 1994. Morphology and meaning in the english mental lexicon. *Psychol. Rev.* 101 (1), 3–33.
- Massol, S., Grainger, J., Dufau, S., Holcomb, P.J., 2010. Masked priming from orthographic neighbors: an ERP investigation. *J. Exp. Psychol.: Hum. Percept. Perform.* 36, 162–174.
- McCormick, S.F., Rastle, K., Davis, M.H., 2008. Is there a ‘fete’ in ‘fetish’? Effects of orthographic opacity on morpho-orthographic segmentation in visual word recognition. *J. Mem. Lang.* 58 (2), 307–326.
- Midgley, K.J., Holcomb, P.J., Grainger, J., 2009. Language effects in second language learners and proficient bilinguals investigated with event-related potentials. *J. Neurolinguistics* 22 (3), 281–300.
- Morris, J., Frank, T., Grainger, J., Holcom, P.J., 2007. Semantic transparency and masked morphological priming: an ERP investigation. *Psychophysiology* 44 (4), 506–521.
- Morris, J., Grainger, J., Holcomb, P.J., 2008. An electrophysiological investigation of early effects of masked morphological priming. *Lang. Cognitive Processes* 23 (7–8), 1021–1056.
- Morris, J., Porter, J.H., Grainger, J., Holcomb, P.J., 2011. Effects of lexical status and morphological complexity in masked priming: an ERP study. *Lang. Cognitive Processes* 26 (4–6), 558–599.
- Rastle, K., Davis, M.H., 2008. Morphological decomposition based on the analysis of orthography. *Lang. Cognitive Processes* 23 (7–8), 942–971.
- Rastle, K., Davis, M.H., New, B., 2004. The broth in my brother's brothel: morpho-orthographic segmentation in visual word recognition. *Psychon. Bull. Rev.* 11 (6), 1090–1098.
- Rastle, K., Davis, M.H., Marslen-Wilson, W.D., Tyler, L.K., 2000. Morphological and semantic effects in visual word recognition: a time-course study. *Lang. Cognitive Processes* 15 (4–5), 507–537.
- Solomyak, O., Marantz, A., 2010. Evidence for early morphological decomposition in visual word recognition. *J. Cognitive Neurosci.* 22 (9), 2042–2057.