Distinct neural substrates of individual differences in components of reading comprehension in adults with or without dyslexia

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Abstract

Reading comprehension is a complex task that depends on multiple cognitive and linguistic processes. According to the updated Simple View of Reading (Gough and Tunmer, 1986; Hoover and Gough, 1990), individual differences in reading comprehension are explained by two main skills: word decoding (i.e., accuracy of identifying words in print) and language comprehension (i.e., ability to understand spoken language). Although Decoding Accuracy and Fluency were not differentiated in the original model in children, the theory has since been expanded to include Fluency, measured by the rate of decoding words or naming stimuli, as an additional contributor, particularly in adulthood (Silverman et al., 2013; Tighe and Schatschneider, 2016; Tilstra et al., 2009). In the current manuscript, we differentiate between the two constructs and refer to accuracy as Decoding and fluency as Fluency. Support for the Simple View of Reading comes from numerous studies demonstrating that decoding and language comprehension are correlated but separable skills, together accounting for a large variance in reading comprehension performance across development (Aaron et al., 1999; Catts et al., 2003; de Jong and van der Leij, 2002; Hoover and Gough, 1990; Singer and Crouse, 1981). The current study examined the neural correlates of decoding, fluency, and language comprehension and examined whether those correlates were dissociable in the brain during naturalistic reading, as posited by the Simple View of Reading framework.

Introduction

Reading a connected text involves multiple cognitive and linguistic processes. According to the Simple View of Reading (Gough and Tunmer, 1986; Hoover and Gough, 1990), individual differences in reading comprehension are explained by two main skills: word decoding (i.e., accuracy of identifying words in print) and language comprehension (i.e., ability to understand spoken language). Although Decoding Accuracy and Fluency were not differentiated in the original model in children, the theory has since been expanded to include Fluency, measured by the rate of decoding words or naming stimuli, as an additional contributor, particularly in adulthood (Silverman et al., 2013; Tighe and Schatschneider, 2016; Tilstra et al., 2009). In the current manuscript, we differentiate between the two constructs and refer to accuracy as Decoding and fluency as Fluency. Support for the Simple View of Reading comes from numerous studies demonstrating that decoding and language comprehension are correlated but separable skills, together accounting for a large variance in reading comprehension performance across development (Aaron et al., 1999; Catts et al., 2003; de Jong and van der Leij, 2002; Hoover and Gough, 1990; Singer and Crouse, 1981). The current study examined the neural correlates of decoding, fluency, and language comprehension and examined whether those correlates were dissociable in the brain during naturalistic reading, as posited by the Simple View of Reading framework.

An important prediction of the Simple View model is that the independence of word decoding and language comprehension will be particularly evident at the lower end of comprehension performance. Indeed, it has been shown that despite the positive and reciprocal association
between decoding and language comprehension in typical readers (i.e., better decoding is associated with better language comprehension and vice versa), the correlation between the two skills weakens with lower decoding skills (Hoover and Gough, 1990; Singer and Crouse, 1981; Stanovich et al., 1984). Additionally, several studies have demonstrated that the relative contribution of language comprehension to reading comprehension increases with better decoding skills, both developmentally across grades (Catts et al., 1999, 2003, 2005; Hoover and Gough, 1990; Storch and Whitehurst, 2002; Vellutino et al., 1994) and across skill levels within one age group (Hoover and Gough, 1990).

An outstanding question exists, however, as to whether the relative contribution of the components to reading comprehension also differs in individuals with reading impairment. Developmental dyslexia (hereafter “dyslexia”) is an unexplained difficulty in learning to read, affecting approximately 10–12% of all individuals (Lyon, 1995). It is an outcome of multiple risk and protective factors (Ozernov-Palchik et al., 2016; Pennington, 2006; van Bergen et al., 2014), but frequently originates from a deficit in phonological awareness (the ability to identify and manipulate speech sounds) (Stanovich and Siegel, 1994; Vellutino et al., 1996; Wagner et al., 1997). This difficulty results in slower and less accurate decoding. It has been suggested that slow fluency and labored decoding constrain comprehension during reading in individuals with dyslexia (Crain and Shankweiler, 1990), but since behavioral evidence is limited in differentiating the underlying mechanisms of performance, there has been insufficient evidence in support of this hypothesis. Given that the associations among the Simple View constructs differ based on decoding skill levels, it is plausible that the neurocognitive mechanisms underlying these skills, and the relations among them, will differ in individuals with dyslexia. For example, individuals with dyslexia have demonstrated greater reliance on language comprehension skills during reading, suggesting the use of linguistic context to bootstrap the decoding deficits that accompany this disorder (Nation and Snowling, 1998; Perfetti and Roth, 1981; Stanovich and Siegel, 1994). A compensatory account of dyslexia has been proposed that suggests increased reliance on language skills for reading comprehension in individuals with dyslexia to compensate for decoding difficulty (Snowling, 2005; Stanovich, 1980). Accordingly, reading comprehension will likely suffer in individuals with dyslexia who do not have these cognitive and linguistic resources available to them. Since similar reading comprehension performance and similar patterns of association among variables may have vastly distinct underlying neurocognitive mechanisms, using neuroimaging measures to study the Simple View components separately in adults with and without dyslexia can reveal these differences.

Although much is known about the neurocognitive basis of impaired single-word decoding in dyslexia, very little is known about the neurocognitive basis of impaired reading comprehension in dyslexia. This represents a major gap in knowledge because it is the comprehension impairment that places the greatest burden on people with dyslexia throughout their education. The current study evaluated whether the proportional contributions of decoding, fluency, and language comprehension to reading comprehension performance, as well as the neural correlates of these components, differ in adults with dyslexia as compared to typical readers. The neural correlates of variation in the three skills have yet to be directly compared within the same individuals, but there are separate literatures examining which neural systems are related to decoding, fluency, and language comprehension performance. Better decoding skills have been associated with increased activation in the left temporoparietal, occipitotemporal, and inferior frontal regions (Martin et al., 2015), and reduced activation in these regions has been consistently demonstrated in individuals with dyslexia (Pugh et al., 2008; Richlan et al., 2011, 2013).

Language comprehension is supported by a distributed network of cortical regions including areas along the middle and superior temporal gyri and inferior frontal cortex that are known to play a role in language processing (Binder et al., 2009; Chai et al., 2016; Ferstl and von Cramon, 2001; Horowitz-Kraus et al., 2013; Huettner, 1989; Price, 2012), in addition to parietal, prefrontal, and posterior medial regions (e.g., cingulate cortex and precuneus) associated with attention and updating (Gernsbacher and Kaschak, 2003; Price, 2012; L. C. Robertson and Ivry, 2000; Roe et al., 2018; St George et al., 1999; Yarkoni, et al., 2008b). Increased reading fluency has been associated with increased recruitment of the left-hemispheric occipitotemporal (i.e., visual word form) region (Benjamin and Gaab, 2012; Langer et al., 2015). To dissociate the neural substrates of individual variation in each skill, we directly compared the neural correlates of these skills within the same individuals.

To evoke reading comprehension processes, we designed a novel paragraph-reading task during imaging, which is a departure from the many prior studies employing single words or, more rarely, single sentences in isolation (Aboud et al., 2016; Richlan et al., 2011; Roe et al., 2018; Ryherd et al., 2018). Most of these tasks involved meta-cognitive judgments about words or sentences, such as making judgments about rhyming, spelling, or semantic plausibility (e.g., Price et al., 1997; Rimrodt et al., 2009). Despite the substantial contribution of these studies to understanding how the neural mechanisms that support reading differ in relation to specific skills, tasks requiring overt decisions or responses raise the likelihood that the neural responses observed reflect secondary, task-specific executive and motor demands in addition to reading processes. Such tasks, therefore, may be of limited utility in elucidating how the semantic, syntactic, and phonological processes interact in a naturalistic context to support reading comprehension (Rayner, 1998; Weber et al., 2014; Yarkoni, et al., 2008). Additionally, reading text sequentially, one word or a few words at a time, creates unnatural breaks and can hinder comprehension as it occurs during normal ecological conditions of reading connected text. Thus, a naturalistic paragraph reading design is ideal for investigating regions involved during comprehension in relation to individual differences in reading comprehension skills.

The current study applied a self-paced naturalistic fMRI task of reading text in 37 adults with and without dyslexia to examine the following: 1) whether decoding, fluency, and language comprehension make similar contributions to variance in reading comprehension in adults with or without dyslexia; 2) whether individual differences in decoding, fluency, and language comprehension are associated with distinct neural patterns of correlation during reading; 3) whether these neural correlates differ based on dyslexia status; 4) which of the neural correlates contributes to unique variance in reading comprehension skills across the groups and within each group.

This study is the first to investigate in-scanner connected-text reading in relation to individual differences in decoding, fluency, and comprehension skills. Using an unconstrained reading-aloud paradigm provides ecological validity as well as the ability to probe multiple aspects of reading as they unfold implicitly in natural reading. Based on the behavioral evidence, we hypothesized that decoding and fluency would be associated with distinct neural substrates of reading compared with language comprehension. We also predicted that decoding and fluency would involve the ventral reading systems and comprehension would involve brain regions implicated in semantic knowledge and cognitive control. Because of evidence of increased reliance on linguistic and cognitive systems in dyslexia, we expect that the neural correlates of the individual Simple View of Reading components would be different in individuals with dyslexia. Finally, we predicted that the neural substrates of each of the components would contribute to unique variance in predicting individual differences in reading comprehension.

Methods

Participants

Adults (n = 18 with dyslexia, n = 19 typical readers, age 18–41 years, \( M = 26.6 \), \( SD = 6.3 \)) participated in the study. The original sam-
ple included 60 participants, but 12 participants were determined ineligible during the behavioral session, 3 participants were subsequently excluded due to incidental findings in MRI, and 8 participants (5 with dyslexia, 3 typical readers) were excluded due to an unacceptable number of scans with motion outliers (see Neuroimaging Processing). The demographic information for all participants is reported in Supplemental Table 1. All participants met eligibility criteria including: being a native speaker of American English; right-handed; born after at least 36 weeks gestation; no sensory or perceptual difficulties other than corrected vision; no history of head or brain injury or trauma; no neurological, neuropsychological, or developmental disorder diagnoses; no medications affecting the nervous system; nonverbal IQ standard score > 85 (Wechsler Adult Intelligence Scale Matrices subtest, Wechsler, 1981). Hearing tests were completed for all participants, and one participant with atypical hearing was excluded. The study was approved by the Committee on the Use of Humans as Experimental Subjects (COUHES) at the Massachusetts Institute of Technology. All participants provided informed written consent in order to participate.

Behavioral measures

All participants completed a comprehensive battery of standardized reading, language, and cognitive assessments, as well as a background questionnaire. In the current analyses, we included four standardized tests that capture the separate components of reading comprehension: decoding, fluency, and language comprehension. Reading comprehension comprised the outcome variable for the behavioral analyses. The decoding construct was measured by the Word Attack subtest of the Woodcock Reading Mastery Test - Revised/Normative Update (Woodcock, 2011); the language comprehension construct was measured by the Listening Comprehension subtest of WRMT-R/NU; and reading comprehension was measured by the reading comprehension subtest of the Gray Oral Reading Test-4 (GORT; Wiederholt et al., 2001). The fluency construct was measured by the Letters subtest of Rapid Automatized Naming (RANL; Wolf and Denckla, 2005) because letter naming does not involve semantic processing or word decoding. Additional standardized and background measures, including measures used to characterize participants as having dyslexia, are reported in Table 1. Participants were included in the dyslexia group (Dys) based on a report of life-long reading difficulty and/or clinical diagnosis and based on a performance below the 25th percentile on at least two out of four standardized subtests of timed or untimed word or nonword reading (Test of Word Reading Efficiency’s Sight Word Efficiency and Phonemic Decoding Efficiency (TOWRE; Torgesen et al., 2012); WRMT’s Word ID and Word Attack). Participants were included in the typical reader group (Typ) based on performance at or above the 25th percentile on all four of the above subtests.

Neuroimaging acquisition

Imaging was performed using a Siemens 3T MAGNETOM Trio, a Tim System (Siemens Medical Solutions, Erlangen, Germany), and a commercial Siemens 32 channel head coil. Structural data were collected using MPRAGE with 2530 ms TR, FOV = 256, 1 × 1 × 1 mm resolution. Functional data were collected with 3 × 3 × 3.6 mm resolution, 2000 ms TR, 30 ms TE, 90° flip angle, 64 × 64 base resolution, and 32 slices approximately parallel to the AC/PC line with coverage of the entire cortex. Prior to each scan, four images were acquired and discarded to allow longitudinal magnetization to reach equilibrium. PACE, an online prospective motion correction algorithm (Thesen et al., 2000), was implemented to reduce the effect of motion artifacts on functional data.

Naturalistic reading task

The task consisted of three conditions, with seven 16-second blocks for each condition. Participants read expository (i.e., texts written to convey factual information on a topic) paragraphs out loud in their normal reading voice and rate inside the MRI scanner. Words in each paragraph were developed and matched based on the age of acquisition, frequency, and imageability. For the control blocks, participants verbally indicated whether arrows on the screen were pointing up or down (i.e., by saying “up” or “down”) to control for motor artifacts related to speech production. Participants’ speech was recorded with an MRI-compatible
microphone. Participants were also presented with a fixation cross and were asked to keep still and relax during the fixation condition.

**Behavioral analyses**

We used multiple regression to examine whether the three Simple View constructs (Language Comprehension, Word Attack, RAN Letters) contribute unique variance to reading comprehension. The models also included sex and age as control variables. The initial model included all participants who completed behavioral data (n = 60). The models in dyslexia and typical participants included only participants who were also included in the neuroimaging analysis (n = 37). The Shapiro test was used to evaluate whether variables violated the normality assumptions, and if the test was significant, permuted linear models were implemented using lmPerm (Wheeler et al., 2016). Multicollinearity among the variables in the regression models was evaluated using the Variance Inflation Factor (VIF) from the oslr package in R (Hebbali, 2018). In order to select the most parsimonious model in explaining reading comprehension from the three constructs, stepwise regression and variable relative importance analyses were conducted with 1,000 permutations.

**In-scanner performance**

To ensure the validity of the in-scanner task, audio recordings of participants reading the stories inside the scanner were analyzed for all participants with adequate in-scanner recordings (n=36). Two measures were calculated: 1) task-fluency: a total score was derived from the total number of words read for each of the stories; 2) task-accuracy: a mean score of number of words read accurately (using GORT scoring procedures) across the stories by a researcher blind to group assignment. To validate the in-scanner measure, pairwise Pearson correlation analyses were conducted relating the scores to the GORT accuracy and fluency measures.

**Neuroimaging processing and data analysis**

We preprocessed the fMRI data using FEAT (FMRI Expert Analysis Tool) Version 6.00, part of FSL version 5.0.2 (FMRIB's Software Library, www.fmrib.ox.ac.uk/fsl) (Smith et al., 2004). High-resolution T1-weighted structural images were skull stripped. The functional data was then registered to the high-resolution structural image using Boundary Based Registration (BBR) algorithms (Greve and Fischl, 2009). We used FMRIB’s Linear Image Registration Tool (FLIRT; Jenkinson et al., 2002) to register the structural data to standard space (2 mm MN152). The following pre-modeling processing included the following steps: spatial smoothing with a Gaussian kernel of FWHM 5 mm; grand-mean intensity normalization by a single multiplicative factor; high pass temporal filtering. Statistical analysis was carried out with FILM using a double-gamma HRF model. The model included 6 motion regressors, their temporal derivatives, and nuisance regressors that modeled out single TR’s identified to have excessive motion according to a framewise displacement (FD) > 0.9 mm (Siegel et al., 2014). Participants (N = 8) who lost more than 40% of frames (i.e., 70) due to FD censorship were excluded from the analysis. There were no significant group differences in the total number of outliers (t(31.7) = 1.42, p = 0.17) and the correlation between number of outliers and out-of-scanner GORT reading comprehension was not significant (r(34) = 0.07, p = 0.68). We carried out second-level analyses using a fixed effects model (Woolrich, 2008; Woolrich et al., 2004) with a cluster-forming threshold of z > 3.1 and a cluster probability of p < 0.05, using Gaussian random field theory (Worsley, 2001). Brain regions are reported in MNI coordinates and identified using the Harvard-Oxford atlas in the FMRIB software. For visualization of the statistical maps, we projected the data onto a standard volume using Mango software (http://tti.utahsc.sa.edu/mango). Coordinates and voxel size of group analysis results were obtained using the FSL version 5.0.2 Cluster tool.

**Whole-brain analyses**

Across all participants, we ran a voxel-wise regression to examine the relation between Decoding (WA), Language Comprehension (LC), and Fluency (RAN), and whole-brain activation during passage reading versus arrows contrast. These scores were entered separately as covariates to a third-level group analysis using FLAME stage 1 (z > 3.1, P’s < 0.05; Woolrich, 2008; Woolrich et al., 2004) with Sex and Age as nuisance covariates. In order to identify the activation overlap at a group level among the three constructs, we performed a pair-wise conjunction analysis to determine the regions that were significantly active (z = 3.1) (Nichols et al., 2005).

**Whole-brain ROI regression analyses**

We evaluated whether the regions associated with individual differences in each of the three constructs contribute uniquely and significantly to differences in reading comprehension. We extracted mean BOLD percent signal change for passages > arrows contrast from the clusters identified in the whole-brain regression models with each of the three constructs. The significant cluster values across participants were included in multivariate regression models using the same procedures as in the behavioral analyses.

**Literature-based Functional Regions of Interest (ROIs) analyses- by group**

Based on the findings from the whole-brain analyses, regions of interest were selected from masks designed to delin-

| Table 1 | Participant characteristics for Typ and Dys groups. Bolded constructs indicate the measures used in the brain analyses. For all measures standard scores are reported. |
|---------|-----------------------------------|---------|---------|---------|
| Construct | Typical | Dyslexia | Significance |
| Age (years) | 26.14 ± 6.15 | 27.07 ± 6.53 | n.s. |
| Nonverbal IQ | 114.28 ± 13.41 | 107.96 ± 15.18 | n.s. |
| Word ID | 108.75 ± 7.55 | 89.92 ± 9.96 | < 0.001 |
| **Word Attack** | 102 ± 8.37 | 77.25 ± 9.61 | < 0.001 |
| Sight Word Efficiency | 109.1 ± 13.99 | 89.15 ± 9.38 | < 0.001 |
| Phonemic Decoding | 104.33 ± 7.62 | 83.29 ± 6.81 | < 0.001 |
| Vocabulary | 114.44 ± 7.77 | 107.71 ± 8.05 | < 0.001 |
| Reading Comprehension | 10.26 ± 1.97 | 7.77 ± 1.48 | < 0.001 |
| Blending Words | 12.17 ± 2.62 | 10.60 ± 2.96 | 0.045 |
| Elision | 9.9 ± 1.88 | 8.12 ± 2.51 | 0.005 |
| Memory for Digits | 10.65 ± 2.11 | 9.76 ± 3.05 | n.s. |
| Language Comprehension | 107.42 ± 7.24 | 101.39 ± 7.41 | 0.052 |
| RAN-Letters | 109.52 ± 5.44 | 101.61 ± 5.93 | < 0.001 |
| RAN 2-Set | 114.21 ± 8.91 | 103 ± 7.25 | < 0.001 |
eate language and domain-general multi-demand systems (Table 2). The masks were downloaded from a publicly available website (https://evlab.mit.edu/funcloc). We used a set of 5 out of the 8 left-hemispheric language masks created using sentences > nonwords contrast (Fedorenko et al., 2010). We chose left-hemispheric masks corresponding to the four clusters identified in the whole-brain analyses for the three constructs: superior temporal (L_STG), angular gyrus (L_AngG), supramarginal gyrus (L_SMG), and middle temporal gyrus (L_MTG). We also selected two prefrontal regions that, although they did not show a significant association in our whole-brain analysis, have been consistently reported in previous studies in relation to decoding and language comprehension skills: the opercular part of the inferior frontal gyrus (IFG) and inferior frontal gyrus orbitalis and its orbital part (L_IFGorb) (Martin et al., 2015; Price, 2012). Three masks from the right hemisphere homologues of the left-hemispheric language-selective regions were also used, including right angular gyrus (R_AngG), inferior frontal gyrus (R_IFG) and inferior frontal gyrus OB (R_IFGorb). It is still unclear whether the prefrontal regions recruited serve broader functional targets or whether they are part of the language network (Hancock et al., 2017). To delineate these possibilities, we included two prefrontal ROIs from the multiple demand (MD) system: posterior parietal (PostParietal) and the insula (Insula). The masks were derived using a hard > easy spatial working memory contrast (Blank et al., 2014). Finally, we selected an additional reading-specific mask from the Neurosynth database for the left putative visual form area variable (VWFA). Mean activation values for the significant voxels for the stories > arrows contrast were extracted for each participant using the masks. We fit a permuted linear regression model using the lmPerm package (Wheeler et al., 2016) predicting each of the three skills from the literature-based ROIs across both Typ and Dys groups as follows:

\[
W_A \sim \text{Age} + \text{Sex} + \text{PostParietal} + L_1 \text{FG} + R_1 \text{FG} + L_1 \text{FGorb} + R_1 \text{FGorb} \\
L_C \sim \text{Age} + \text{Sex} + \text{PostParietal} + \text{Insula} + VWF A + R_2 \text{FG} + L_2 \text{VFG} + L_2 \text{Occ} + R_2 \text{VFG} + L_2 \text{FGorb} + R_2 \text{FGorb} \]

For each of the six models, we implemented a stepwise regression analysis to identify the optimal set of predictors for each construct separately in the Typ and the Dys groups. Predictors with high VIF were excluded from each model on a model-by-model basis.

**Results**

**Behavioral**

The two groups were similar in terms of sex (10 females in Typ and 11 females in Dys) and age (p’s > 0.98). There were significant differences in age between male and female participants (t(40.57) = −2.48, p = 0.02), with older male participants. Additionally, across both groups, males performed above females on many of the reading behavioral measures. Sex differences on measures of language and reading, as well as in the brain, have been reported across multiple studies (e.g., Lietz, 2006; Reilly, Neumann, & Andrews, 2019; Rutter et al., 2004; Shaywitz et al., 1995). This topic is beyond the scope of the current study but is discussed elsewhere (e.g., Krafnick & Evans, 2019). We therefore included Age and Sex in the models.

Age, Sex, and the three constructs of Language Comprehension (LC), Fluency (RANL), and Decoding (WA), were combined in a stepwise regression model with 1,000 permutations to identify whether, in accordance with the Simple View of Reading, each of the three constructs contributed independent and significant variance in explaining reading comprehension. The R² for the entire model was 0.49 (p < 0.001) and the best set of predictors included the three constructs in order of their relative importance: (1) Language Comprehension (contributing 41% of R², \( \eta^2 = 0.151 \)), (2) Decoding (contributing 38% of R², \( \eta^2 = 0.139 \)), and (3) Fluency (contributing 19% of R², \( \eta^2 = 0.177 \)). Correlations among the three constructs revealed a significant positive association between Fluency and Decoding (r(52) = 0.6, p < 0.001), a nonsignificant association between Fluency and Language Comprehension (r(52) = 0.14, p = 0.3), and a borderline significant association between Language Comprehension and Decoding (r(52) = 0.26, p = 0.06). Although the three constructs were moderately correlated, the VIF values for all variables ranged from 1.09 to 1.64, suggesting that multicollinearity is not an issue of concern for these regression models (O’Brien, 2007).

To test whether the constructs of reading comprehension are different based on dyslexia status, the models described above were repeated in the Dys and Typ groups in the final neuroimaging sample (n = 18 and 19, respectively, excluding participants who were not eligible to be part of either of the two groups based on their reading performance). For Typ, the overall model explained 36.3% of total variance in reading comprehension, with Language Comprehension accounting for 82% (\( \eta^2 = 0.26 \)) of total variance and making the only significant contribution to the model. For Dys, the model accounted for 28.3% of total variance with Language Comprehension contributing 28% (\( \eta^2 = 0.189 \)), Decoding, 8% (\( \eta^2 = 0.04 \)); and Fluency, 5% (\( \eta^2 = 0.08 \)) of total variance.

**In-scanner performance**

Correlational analyses confirmed the construct validity of the in-scanner task. Task-fluency scores were significantly associated with GORT fluency scores (r(36) = 0.73, p < 0.001) and task-accuracy scores...
Table 2
Regions of significant activation in stories > arrows whole-brain contrasts.

<table>
<thead>
<tr>
<th>Area</th>
<th>No. of voxels</th>
<th>t value</th>
<th>Coordinates (X, Y, Z)</th>
<th>fROI labels</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Group-level (stories &gt; arrows)</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Left occipitotemporal cortex (including cerebellum)</td>
<td>17654</td>
<td>8.43</td>
<td>(−20, −96, −6)</td>
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<tr>
<td>Left inferior frontal gyrus</td>
<td>9995</td>
<td>7.5</td>
<td>(−56, −8, 0)</td>
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<tr>
<td>Left superior/middle frontal</td>
<td>7098</td>
<td>7.16</td>
<td>(60, −4, 0)</td>
<td></td>
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<tr>
<td>Left paracingulate</td>
<td>1491</td>
<td>6.34</td>
<td>(−8, 12, 66)</td>
<td></td>
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<tr>
<td>Left caudate/putamen</td>
<td>1172</td>
<td>5.57</td>
<td>(−2, 14, 0)</td>
<td></td>
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<tr>
<td>Left medial frontal cortex</td>
<td>417</td>
<td>4.77</td>
<td>(−4, 50, −10)</td>
<td></td>
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<tr>
<td><strong>(arrows &gt; stories)</strong></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right angular gyrus</td>
<td>4110</td>
<td>7.23</td>
<td>(62, −40, 38)</td>
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<tr>
<td>Posterior cingulate</td>
<td>2680</td>
<td>5.85</td>
<td>(−6, −24, 28)</td>
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<tr>
<td>Left supramarginal gyrus</td>
<td>1401</td>
<td>6.99</td>
<td>(−58, −48, 42)</td>
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<tr>
<td>Right lingual gyrus</td>
<td>1084</td>
<td>5.75</td>
<td>(6, −80, −4)</td>
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<tr>
<td>Left lateral occipital/middle temporal gyrus</td>
<td>415</td>
<td>5.18</td>
<td>(−52, −74, 10)</td>
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<td>Left lingual gyrus</td>
<td>402</td>
<td>4.84</td>
<td>(−20, −56, −8)</td>
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<td>Left precuneus</td>
<td>230</td>
<td>4.55</td>
<td>(−12, −66, 30)</td>
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<td>Right anterior parahippocampal cortex</td>
<td>225</td>
<td>6.06</td>
<td>(20, −2, −36)</td>
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<tr>
<td><strong>Typ vs. Dys</strong></td>
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<tr>
<td>Fluency</td>
<td></td>
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<tr>
<td>Left occipitotemporal</td>
<td>135</td>
<td>4.51</td>
<td>(−42, −54, −6)</td>
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<tr>
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<tr>
<td>Left supramarginal</td>
<td>160</td>
<td>4.32</td>
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<td>L_SMG</td>
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<td>147</td>
<td>4.29</td>
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<td>138</td>
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<tr>
<td>Language Comprehension</td>
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<td>Right precuneus</td>
<td>156</td>
<td>4.78</td>
<td>(10, −80, 38)</td>
<td>PostParietal</td>
</tr>
<tr>
<td>Left precuneus</td>
<td>139</td>
<td>5.13</td>
<td>(−2, −72, 56)</td>
<td></td>
</tr>
<tr>
<td>Language Comprehension</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Left precuneus/cingulate</td>
<td>2336</td>
<td>5.32</td>
<td>(−4, −16, 30)</td>
<td>PostParietal</td>
</tr>
<tr>
<td>Right insula</td>
<td>956</td>
<td>4.88</td>
<td>(28, 0, 8)</td>
<td>Insula</td>
</tr>
<tr>
<td>Left occipitotemporal cortex</td>
<td>257</td>
<td>4.37</td>
<td>(−22, −52, −2)</td>
<td>VWFA</td>
</tr>
<tr>
<td>Right posterior temporal gyrus</td>
<td>223</td>
<td>5.07</td>
<td>(68, −34, −2)</td>
<td>R_PostTemp</td>
</tr>
<tr>
<td>Left angular</td>
<td>217</td>
<td>4.22</td>
<td>(−54, −46, 38)</td>
<td>L_Ang</td>
</tr>
<tr>
<td>Left lateral occipital/middle temporal gyrus</td>
<td>216</td>
<td>4.08</td>
<td>(−44, −78, 14)</td>
<td>L_MTG</td>
</tr>
<tr>
<td>Left putamen</td>
<td>198</td>
<td>4.41</td>
<td>(−28, −2, 12)</td>
<td></td>
</tr>
<tr>
<td>Right angular gyrus</td>
<td>172</td>
<td>4.08</td>
<td>(46, −50, 40)</td>
<td>R_Ang</td>
</tr>
<tr>
<td><strong>Decoding &amp; Language Comprehension Conjunction</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right cuneus</td>
<td>150</td>
<td>4.49</td>
<td>(6, −78, 40)</td>
<td></td>
</tr>
<tr>
<td>Left superior parietal lobe</td>
<td>52</td>
<td>4.76</td>
<td>(−4, −70, 58)</td>
<td></td>
</tr>
<tr>
<td>Right precuneus</td>
<td>17</td>
<td>3.6</td>
<td>(4, −62, 60)</td>
<td></td>
</tr>
<tr>
<td><strong>Language Comprehension &amp; Fluency Conjunction</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Left angular/supramarginal gyrus</td>
<td>46</td>
<td>4.08</td>
<td>(−56, −46, 38)</td>
<td></td>
</tr>
<tr>
<td>Left middle temporal gyrus</td>
<td>4</td>
<td>3.19</td>
<td>(−52, −70, 4)</td>
<td></td>
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</tbody>
</table>

were significantly associated with GORT accuracy ($r(36) = 0.65, p < 0.001$). Additionally, both in-scanner measures significantly correlated with GORT comprehension scores (accuracy: $r(36) = 0.52, p < 0.001$; fluency: $r(36) = 0.65, p < 0.001$).

Whole-brain

To examine the activation induced by the reading task, we conducted a one-sample $t$-test for stories > arrows for the entire group (Fig. 2A). This contrast yielded significant activation in the left-hemisphere ventral occipitotemporal and inferior frontal regions (occipital fusiform gyrus, inferior temporal gyrus, inferior frontal gyrus). Additional activations included right middle/superior frontal regions, basal ganglia (caudate, putamen), and paracingulate and cingulate regions. The Typ group, as compared to Dys group, showed significantly greater activation in left occipitotemporal region, which includes the putative VWFA (Fig. 2B). There were no significant clusters with greater activation in Dys as compared to Typ. For the whole-brain correlation with each of the three constructs, there were significant negative correlations with Language Comprehension and Decoding, and a significant positive correlation with Fluency (Table 2, Fig. 3). There were no significant clusters for the opposite relationships. Better Language Comprehension was significantly associated with less activation in bilateral posterior temporal and inferior parietal (including angular gyrus) and posterior parietal regions, insula, precuneus, and putamen. Better Decoding was significantly associated with less activation in medial and lateral posterior parietal cortex, including in the cuneus and precuneus. Finally, better Fluency was significantly associated with greater activation in left dorsal temporoparietal and ventral occipitotemporal regions, including the visual word form area.

The conjunction analysis for each pair of the constructs revealed overlapping regions of significant correlation between Language Comprehension and Decoding in posterior parietal regions. There were overlapping regions of correlated activation for Language Comprehension and Fluency in the left supramarginal/angular gyrus cluster and in a
small middle temporal cluster. There were no significantly overlapping clusters between Fluency and Decoding.

Whole-brain ROIs

Overall, 11 ROIs were extracted for the significant clusters identified in the whole-brain regression analyses (3 clusters for Fluency, 2 clusters for Decoding, and 6 clusters for Language Comprehension). All clusters were added into a regression model with Reading Comprehension as the predicted variable and Sex and Age as the control variables (Fig. 3-D). For all variables, the variance inflation factor (VIF) values were below 10 except one of three Fluency clusters. This cluster, which was in the left supramarginal gyrus, was excluded from the regression models. The Shapiro–Wilk test revealed that some of the ROI values were not normally distributed (W’s < 0.95, p’s < .02). Accordingly, the lmPerm package (Wheeler, et al., 2016) in R was used to calculate permuted linear regression for all analyses. The R² for the entire model was 0.21 (p < .001) and the best set of predictors included two of the Language Comprehension ROIs (lc3-left middle temporal gyrus: 21.25% of R²; lc4-left angular gyrus: 42.74% of R²).

Literature-based fROIs by group

The Shapiro–Wilk test revealed that several of the fROI’s were not normally distributed and permuted linear regressions were calculated for all analyses with a stepwise approach to select the best model fit. VIF was evaluated for each model and clusters that exceeded the value of 10 were excluded from the particular model. For the Typ group, Fluency was significantly predicted by VWFA activation only, accounting for 10.37% of the variance (η² = 0.12), with higher Fluency associated with increased activation in VWFA. Three fROIs had to be included in the Language Comprehension model due to high VIF: Insula, L_IOcc, and R_IFG. Language Comprehension was significantly predicted by Age (r² = 0.13, η² = .11), Sex (r² = 0.28, ηp² = 0.19), PostParietal (r² = 0.28, ηp² = 0.05), VWFA (r² = 0.11, ηp² = .06), R_Ang (r² = 0.12, ηp² = 0.19), and R_IFGorb (r² = 0.04, ηp² = 0.08). Increased recruitment of these regions was associated with lower Language Comprehension scores. For the Decoding model, R_IFGorb had to be excluded due to a high VIF. Decoding was significantly predicted by Age only (r² = 0.17, ηp² = 0.18).

For the Dys group, Fluency was significantly predicted by Age (r² = 0.07, ηp² = 0.004), Sex (r² = 0.19, ηp² = 0.25), and L_SMG (r² = 0.1, ηp² = 0.1). Better Fluency was associated with increased activation of this region. Several fROIs had to be removed from the Language Comprehension model due to high VIF: PostParietal, Insula, and R_SMG. Language Comprehension was significantly predicted by L_IFG (r² = 0.1, ηp² = 0.04) and L_IFGorb (r² = 0.18, ηp² = 0.2). Better Language Comprehension was associated with increased activation of L_IFGorb and decreased activation of L_IFG. Due to high VIF, R_IFGob was removed from the Decoding model. Decoding was predicted by Post-Parietal (r² = 0.09, ηp² = 0.05), L_IFG (r² = 0.04, ηp² = 0.01), L_IFGob (r² = 0.4, ηp² = 0.48), and R_IFG (r² = 0.16, ηp² = 0.14). Increased activation of these regions was associated with lower Decoding.

To test which fROIs explained significant variance in reading comprehension performance in Dys and Typ groups, we included clusters that were significant in the six Decoding, Language Comprehension, and Fluency models in a stepwise regression model predicting Reading Comprehension separately in each group. In the Typ group, after removing several clusters due to high VIF (Insula, R_SMG, R_IFG), the overall model explained 49.58% of variance in Reading Comprehension scores with Sex explaining 30% of total variance (ηp² = 0.03), with higher scores for males. The significant fROIs included: PostParietal explaining
14% ($\eta^2 = 0.21$), L_IFGorb explaining 6% ($\eta^2 = 0.001$), and L_IFG explaining 14% ($\eta^2 = 0.14$). For Dys, after the high VIF variables were removed (L_SMG, Insula, R_SMG), the model accounted for 55.83% of total variance with the only significant predictor PostParietal explaining 25% of total variance ($\eta^2 = 0.13$).

Discussion

The current study investigated, in adults with and without dyslexia, how brain activation during a naturalistic passage reading task was related to out-of-scanner components of reading comprehension as proposed by the Simple View of Reading: decoding accuracy, decoding fluency, and language comprehension. Consistent with the Simple View framework, the three constructs explained unique and significant variance in reading comprehension. We found distinctive patterns of activation in relation to individual differences in fluency and language comprehension. The activation related to decoding accuracy, however, was entirely overlapping with some of the regions associated with language comprehension. Activation in the inferior parietal and middle temporal language-comprehension ROIs made the largest contribution to reading comprehension. Importantly, confirming the validity of our fMRI paradigm, the stories > arrows contrast involved the expected dorsal and ventral reading regions. When examining the neural correlates of the three constructs in each of the groups using literature-based regions of interest, we found evidence that the neural substrates of comprehension differ in adults with and without dyslexia and the relative behavioral contribution of the three components to reading comprehension was different as well. Our results suggest that during naturalistic reading, individuals with dyslexia recruit cognitive and linguistic systems to support word decoding, thereby diverting resources from, and potentially impeding, comprehension processes. These findings reveal the underlying mechanisms of reading comprehension deficits of adults with dyslexia.

Behavioral findings

The behavioral results confirmed the basic tenants of the Simple View of Reading and demonstrated a unique and significant contribu-

Fig. 4. A) Correlations between Language Comprehension, Fluency, and Decoding scores and the fROIs and among the fROIs separately by group. B) fROIs that demonstrated a significant association in the stepwise regression model for each of the subskills for the Typ group (red), Dys group (blue).

Fig. 5. Association with reading comprehension for the best set of fROI predictors for Dys – Posterior Parietal (right panel) and for Typ – Posterior Parietal, Left IFG (left panel), and Left IFGorb (not shown).
dyslexia is due to the slow and impaired word reading that creates a bottleneck for comprehension of texts because of additional cognitive demands (Crain and Shankweiler, 1990).

Associations and dissociations among neural correlates of the Simple View of Reading

Decoding correlates

Across all participants, we found a negative correlation between decoding accuracy (Word Attack scores) and activation in bilateral posterior parietal and precuneus regions. Separate analyses by group using a literature-based posterior parietal multi-domain fROI, revealed that the negative association between posterior parietal regions and decoding was driven by the dyslexic group only. Additional correlates of decoding in the dyslexic group included bilateral IFG and right IFG orbitals. Posterior parietal/precuneus regions have been implicated in attentional modulation of complex tasks (Cavanna and Trimble, 2006). The precuneus has been frequently invoked in studies of language comprehension (Binder et al., 2009; Price, 2012) and reading (Chyl et al., 2018; Rimrodt et al., 2009; Roe et al., 2018; Ryherd et al., 2018; Schulz et al., 2008, 2009; Shaywitz et al., 2002), and there is evidence to suggest it represents the extent to which attentional resources are allocated to these tasks (Kuperberg et al., 2003; Schulz et al., 2008, 2009). Consistent with the current findings, several studies have demonstrated negative associations between activation in the precuneus and out-of-scanner reading (Chyl et al., 2018; Rimrodt et al., 2009; Ryherd et al., 2018). Cumulatively, these findings suggest that adults with poorer decoding skills invoked the attentional regions during reading to a greater extent than adults with better decoding skills. Importantly, the current study is the first study employing a connected-text paradigm in relation to individual differences in the subcomponents of reading. Previous studies that have identified the neural systems underlying reading in relation to decoding skills used single-word or sentence-reading tasks. The current findings, therefore, provide initial evidence of a relation between activation in the precuneus and individual differences in decoding skill as applied to reading connected text.

The increased importance of prefrontal regions for reading in individuals with dyslexia has been highlighted by previous studies (see Hancock et al., 2017). Several meta-analyses have documented increased activation in these regions in individuals with dyslexia relative to controls during phonological tasks (Maisog et al., 2008; Richlan et al., 2013). Hyperactivation in these regions has been interpreted as compensatory. In order to establish a compensatory role for these systems, however, it is important to demonstrate their engagement in relation to behavioral performance (Hancock et al., 2017). The few studies that have done this have demonstrated both positive and negative associations between bilateral inferior frontal regions and reading-related skills (Bach et al., 2010; Horowitz-Kraus et al., 2013; Ingvar et al., 2002; Patael et al., 2018; Rumsey et al., 1994; Ryherd et al., 2018). We found that individuals with worse Word Attack scores in the dyslexic group recruited the bilateral inferior frontal regions to a greater extent. The inferior frontal gyrus, bilaterally, represents the anterior language system involved in language comprehension and production, but has also been associated with domain-general cognitive and perceptual functions including attention, working memory, inhibitory control, planning/goal-directed behaviors, fluid intelligence, and consciousness (Hancock et al., 2017). Inferior frontal regions have been implicated in the resolution of phonological ambiguity via top-down modulation of activity in posterior phonological systems (Burton et al., 2000; Gow et al., 2008; Myers, 2007; Zatorre et al., 1996). These regions were also associated with better Language Comprehension skills in the dyslexic group. Together, the current findings provide novel evidence for the increased involvement of language comprehension and cognitive systems during reading to compensate for poor decoding skills in dyslexia.
Language comprehension correlates

We found increased activation of regions that are part of the cingulo-opercular networks (i.e. cuneus, cingulate, insula) as well as basal ganglia regions to be associated with worse language comprehension performance in the whole-brain analysis. These regions have been shown to support executive functions (e.g., inhibitory control, attentional selection, conflict resolution, maintenance and manipulation of task sets) for both linguistic and non-linguistic tasks (e.g., Duncan and Owen, 2000; Fedorenko et al., 2013; Hugdahl et al., 2015; see Fedorenko, 2014). The few previous studies investigating the neural correlates of individual differences in language comprehension also found increased engagement of these attentional and cognitive control systems during reading (Roe et al., 2018; Ryherd et al., 2018). Furthermore, cingulate and prefrontal regions have been associated with updates in and integration of the reader’s mental model during discourse comprehension (Gernsbacher and Kaschak, 2003; Robertson et al., 2000; St George et al., 1999; Whitney et al., 2009; Yarkoni et al., 2008).

Additionally, there was a negative correlation with clusters of activation in the bilateral middle temporo-polar gyrus and angular gyrus. These regions are thought to be part of a distributed semantic system across modalities (e.g., speech and text) and stimuli (language, faces) (Hartwigs et al., 2015; Price, 2012; Price et al., 1997). The engagement of the semantic network for reading as a function of comprehension skills is in accordance with the behavioral literature linking poor comprehension to deficits in semantic processing (see Landi and Ryherd, 2017). The current findings, however, indicate a negative association between comprehension skills and the engagement of this network, showing increased recruitment of these regions in participants who had poorer language comprehension. These findings are consistent with several previous studies that found a negative association between word reading skill and the engagement of the semantic network (Rimrodt et al., 2009; Welcome and Joanisse, 2012), but not with other studies that found a positive association (Aboud et al., 2016; Ettinger-Veenstra et al., 2016; Ryherd et al., 2018). For example, one study found decreased activation for skilled comprehenders in regions including angular gyrus and middle temporo-polar gyrus during word reading (Welcome and Joanisse, 2012). In another study, there was a positive association between comprehension skills and activation in middle temporo-polar regions and a negative association in inferior parietal regions during sentence-by-sentence passage reading (Ryherd et al., 2018). Therefore, our findings extend the previous literature in demonstrating that during naturalistic reading adults with lower comprehension skills engage both the domain-general executive systems and the language-specific semantic system to a greater extent.

When examining the Language Comprehension correlates by group, however, the semantic system identified in the whole-brain analysis was found to be negatively associated with comprehension skills in the typically reading group only. Furthermore, the posterior parietal area was also recruited to a greater extent in the typical adults with poor comprehension skills, but not in the dyslexic adults. In the dyslexic group, better Language Comprehension skills were associated with increased activation of the left IFG only. Thus, the domain-general posterior cognitive system and the semantic system were deployed to support reading in adults with poor comprehension but adequate decoding.

Fluency correlates

Across all participants, better Fluency was positively associated with increased activation of the left ventral occipital-temporal region and left temporo-parietal regions. In the typically reading group, better Fluency was associated with increased activation of the putative visual word form area (VWFA), a region that develops specialization for automatic word recognition with increased reading experience (Centanni et al., 2018; Kronbichler et al., 2004; McCandliss et al., 2003). Accordingly, previous studies have demonstrated an increased recruitment of VWFA with increased reading fluency demands (Benjamin and Gaab, 2012; Langer et al., 2015). In the Dys group, better Fluency skills were associated with increased recruitment of the left temporoparietal systems, comprising the dorsal phonological reading network as well as the semantic systems. The posterior middle temporal region has been implicated in previous studies as an additional region of difference in fluency performance (Meyler et al., 2007; Roe et al., 2018).

The differentiation between the two groups in the neural correlates of fluency is consistent with developmental theories of neural specialization for reading (Pugh et al., 2000; Sandak et al., 2004; Younger et al., 2017). Less skilled readers engage the dorsal temporo-parietal network for phonological analysis and recoding, whereas skilled readers rely on the rapid, ventral stream for orthographic computations. The automaticity of the ventral system facilitates reading fluency which in turn is thought to lead to better reading comprehension. This theory is supported by metanalytic findings showing a more consistent recruitment of the ventral system during reading in adults as compared to children (Martin et al., 2015). Together, the current results support the notion that RAN, as a measure of fluency, represents the automaticity of integration across the individual components of the reading circuit: the orthographic, phonological, and semantic systems (Norton and Wolf, 2011). In individuals with dyslexia, fluency depends on more efficient utilization of the dorsal system, but in typical readers, increased fluency depends on the automaticity of the ventral system.

Decoding & language comprehension

We examined whether there were commonly activated voxels for decoding and language comprehension skills. We found that the regions associated with decoding completely overlapped with the regions associated with language comprehension. Specifically, poor decoding and poor comprehension across all participants were associated with increased engagement of the domain-general posterior cognitive system region previously demonstrated to be involved in cognitive control during reading (Horowitz-Kraus et al., 2013; Roe et al., 2018). Interestingly, in the dyslexic group the posterior parietal cognitive-control ROI was more activated in reading with decreasing decoding skills, but in the typically reading group the region was more involved with decreased language comprehension skills. The overlap in neural correlates of Language Comprehension and Decoding were also evident in left inferior frontal regions, which supported both decoding and comprehension in the Dys group. These results extend behavioral findings in support of the idea that both cognitive and linguistic resources are deployed at different stages during reading based on dyslexia status: in individuals with dyslexia they support lower-level word identification and in typical readers they support higher-level comprehension.

Semantic regions explaining reading comprehension

We found that even though behaviorally the three constructs made significant and unique contributions to reading comprehension, only the contribution of the two semantic regions, left angular gyrus and middle temporal gyrus, was significant. Deficits in the semantic domain have been most consistently implicated in poor oral comprehension both in neuroimaging and in behavioral literature (see Landi and Ryherd, 2017). Furthermore, the developmental behavioral literature suggests that the unique contribution of decoding skills to variance in reading comprehension decreases across development and with increased mastery of reading (Catts et al., 2005; Hoover and Gough, 1990; Schatschneider et al., 2004). Instead, in these studies, the greatest contribution to reading comprehension was made by language comprehension skills as well as the shared variance between decoding and comprehension. Our neuroimaging findings in adults, therefore, mirror these behavioral results with the caveat that due to our enrollment criteria for dyslexia, the decoding skills in the current sample are oversampled on the lower end of the skill distribution. Therefore, the significant behavioral contribution of decoding to reading comprehension in our models could be due to the severity of the decoding deficits
in our sample. Nevertheless, this is the first study to directly compare the relative contribution of the different neural components of reading to out-of-scanner reading comprehension. The findings provide the first neural evidence for the crucial role of the semantic network, above phonological and orthographic systems, for reading comprehension in adults.

Differences in neural correlates of the Simple View of Reading in dyslexia

An important implication of the Simple View theory is that the interactive relationship between language comprehension and word reading influences reading comprehension. This interaction implies that the relative contribution of the individual components to reading comprehension varies based on the levels of each of the component skills. Based on this formulation, in individuals with dyslexia, a greater relative contribution of decoding to reading comprehension than in typical readers would be expected. Our behavioral findings support this prediction. Furthermore, our neural findings provide important insights into the underlying mechanisms of the Simple View of Reading in dyslexia. In typical readers, consistent with the behavioral results, there were no significant differences in activation patterns in relation to decoding skills. In adults with dyslexia, both linguistic and cognitive systems were deployed to support decoding. The inferior frontal language system was recruited in individuals with lower decoding and higher comprehension skills, suggestive of increased linguistic top-down modulation of word identification in dyslexia, in order to compensate for the impairments in the posterior left-hemispheric reading regions.

In addition to the significant contribution of the semantic system to reading comprehension across all participants, the patterns of neural contributions to explaining variance in reading comprehension skills differed between the two groups. In the typically reading group, increased recruitment of the posterior parietal and left inferior frontal regions was associated with worse reading comprehension skills. In the dyslexic group, the multi-domain posterior parietal region made the only significant contribution to explaining differences in reading comprehension. Increased recruitment of this system was associated with better reading comprehension, but it was also recruited in relation to poor decoding skills in the dyslexic group. This suggests that the contribution of the cognitive mechanisms to explaining reading comprehension in adults with dyslexia is modulated through the recruitment of this system for decoding.

Reading comprehension occurs when lower-level processes, such as word decoding, are integrated with higher-level comprehension systems (Perfetti et al., 2008; Perfetti and Roth, 1981). These processes take place within a domain-general cognitive system with limited processing resources. Strong decoding skills free up cognitive capacity for higher-level comprehension of text and for processes related to forming inferences while reading (Thurlow and Broek, 1997). Our findings offer insight into neural mechanisms of poor reading comprehension in adults with dyslexia, possibly indicating that this weakness could be due to insufficient cognitive resources to support linguistic comprehension when these resources are instead recruited to support accurate reading. The crucial role of these domain-general cognitive processes for comprehension is buttressed by extensive behavioral evidence of cognitive impairments in individuals with poor comprehension across multiple domains (e.g., planning, working memory, response inhibition, task switching) (Cutting et al., 2009; Locascio et al., 2010; Potocki et al., 2017; Protocass Pappas et al., 2007; Sesma et al., 2009). Furthermore, evidence from poor readers supports the specificity of cognitive impairments for tasks involving reading, but not for other tasks (Roe et al., 2018). Our neuroimaging results therefore lend support to the proposal that in poor readers under naturalistic reading conditions, cognitive control resources are strained by word decoding, creating a bottleneck and weakening the contribution of these resources towards integration and comprehension (Hudson et al., 2005; Pikulski & Chard, 2005).

Conclusions and implications

Our findings provide a snapshot of the neural underpinnings of naturalistic reading and show how these mechanisms are related to individual abilities across different reading comprehension components based on the Simple View of Reading. It is possible that the central role of domain-general networks during reading in the current study is the result of the expository nature of the text used for the fMRI task. There has been some evidence showing that expository texts are more difficult to process (Best et al., 2004; Graesser and McNamara, 2011; McNamara et al., 2004) and place higher demands on executive functions networks than narrative texts, particularly in terms of inferencing and planning/organizing information (Aboud et al., 2019; Barenta et al., 2009; Eason et al., 2012; Miller et al., 2013). The Simple View of Reading does not account for the interaction between reader differences and text characteristics and therefore these were not examined in the current study. Future studies are needed to delineate neural systems supporting expository versus narrative text types in relation to individual reader skills.

Additionally, as discussed above, the nature of the interaction among the three components changes across development, with greater independence of decoding and language comprehension found in older readers. The contributions of vocabulary and background knowledge to comprehension also increase with age and text difficulty. Furthermore, although there is convincing evidence for the robustness of the Simple View model across orthographies, the relative influence of decoding and language comprehension on reading comprehension varies across orthographies (Florit and Cain, 2011). In more transparent orthographies, fluency plays a greater role than in English. This suggests that our findings are limited to inference about English-speaking adults and that developmental studies across orthographies are needed to examine whether similar patterns of neural correlations are evident in children. Further, future studies may benefit from larger samples that support additional analyses and that may reveal more about variance within and between typically developing and dyslexic readers.

Despite the need for further investigations, the current study is the first to identify the neural correlates of the Simple View framework, as they unfold during naturalistic reading. The Simple View has been a predominant framework of reading comprehension, and the current findings support the application of the model to dyslexia and reveal different behavioral and neural contributions to reading comprehension in this disorder. Our findings shed light on the mechanisms of comprehension deficits in dyslexia by showing that individuals with dyslexia recruit linguistic and cognitive brain regions to support word identification at the potential expense of comprehension.

Declaration of Competing Interest

None declared

Data and Code Availability

The behavioral and fROI data and the analysis code used to support the findings of this study have been deposited in a Github repository (https://github.com/oozernov/neural_correlates_svr)

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