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Sequencing at the syllabic and supra-syllabic levels during speech perception: an fMRI study

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12	Keywords: syllabic information, supra-syllabic information, supratemporal plane, speech processing,
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42 Abstract

The processing of fluent speech involves complex computational steps that begin with the 43 segmentation of the continuous flow of speech sounds into syllables and words. One 44 question that naturally arises pertains to the type of syllabic information that speech 45 46 processes act upon. Here, we used functional magnetic resonance imaging to profile regions, using a combination of whole-brain and exploratory anatomical region-of-47 interest (ROI) approaches, that were sensitive to syllabic information during speech 48 perception by parametrically manipulating syllabic complexity along two dimensions: (1) 49 individual syllable complexity, and (2) sequence complexity (supra-syllabic). We 50 manipulated the complexity of the syllable by using the simplest syllable template—a 51 52 consonant and vowel (CV)-and inserting an additional consonant to create a complex onset (CCV). The supra-syllabic complexity was manipulated by creating sequences 53 composed of the same syllable repeated 6 times (e.g. /pa-pa-pa-pa-pa-pa/) and sequences 54 55 of 3 different syllables each repeated twice (e.g. /pa-ta-ka-pa-ta-ka/). This parametrical design allowed us to identify brain regions sensitive to (1) syllabic complexity 56 independent of supra-syllabic complexity, (2) supra-syllabic complexity independent of 57 syllabic complexity and, (3) both syllabic and supra-syllabic complexity. High-resolution 58 59 scans were acquired for 15 healthy adults. An exploratory anatomical ROI analysis of the supratemporal plane (STP) identified bilateral regions within the anterior two-third of the 60 61 planum temporale, the primary auditory cortices as well as the anterior two-third of the superior temporal gyrus that showed different patterns of sensitivity to syllabic and supra-62 syllabic information. These findings demonstrate that during passive listening of syllable 63 sequences, sublexical information is processed automatically, and sensitivity to syllabic 64 65 and supra-syllabic information is localized almost exclusively within the STP.

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68 1. Introduction

The speech signal is undoubtedly one of the most complex auditory signals that humans are exposed to, requiring multiple computational steps to parse and convert acoustic waves into discrete linguistic units from which meaning can be extracted. Unsurprisingly, given such complexity, the manner in which the human brain accomplishes the complex computational steps leading to the comprehension of speech remains far from understood.

Functional neuroimaging studies of speech perception offer converging evidence 75 suggesting that the supratemporal plane (STP), and superior temporal sulcus (STS) play a 76 critical role in the processing of speech sounds. This finding is quite robust having been 77 78 observed under different types of speech perception tasks (i.e. passive listening, monitoring and discrimination tasks as well as neural adaptation paradigms) and with 79 80 different types of speech stimuli (words, pseudo-words, syllables, phonemes). For instance, neuroimaging studies contrasting the neural activity evoked by speech stimuli to 81 82 the neural activity associated with the processing of acoustically complex non-speech sounds or silence have reliably reported clusters of activation within the STP and/or STS 83 (Zatorre et al., 1992;Binder et al., 1996;Binder et al., 1997;Dhankhar et al., 1997;Celsis et 84 85 al., 1999;Burton et al., 2000;Scott et al., 2000;Benson et al., 2001;Vouloumanos et al., 2001; Joanisse and Gati, 2003; Wilson et al., 2004; Liebenthal et al., 2005; Rimol et al., 86 2005; Wilson and Iacoboni, 2006; Obleser et al., 2007; Okada et al., 2010; Zhang et al., 87 2011;Tremblay et al., 2012). In addition, neuropsychological evidence demonstrate that 88 bilateral lesions to the superior temporal lobes can result in pure word deafness, a deficit 89 associated with impaired word comprehension but relatively intact ability to process non-90 speech sounds (Buchman et al., 1986; Tanaka et al., 1987; Poeppel, 1996). While both 91 functional and neuropsychological studies provide strong evidence regarding the 92 importance of the STP and STS for the perception of speech sounds, the specific 93 94 contribution of each of the sub-regions that form this large cortical area to speech perception is still uncertain; whether it is related to the processing of acoustical, 95 sublexical or lexical information. 96

97 Several neuroimaging studies have contrasted the neural activity evoked by 98 different sublexical units (e.g. consonant clusters, phonemes, syllables) to the processing of nonspeech or unintelligible speech sounds (e.g. sinewave analogs, tones, 99 environmental sounds, noise, spectrally rotated syllables, silence) to isolate speech 100 specific processes. These studies reported reliable activation within supratemporal 101 regions (e.g. the superior temporal gyrus (STG), the transverse temporal gyrus (TTG) and 102 planum temporal (PT)), the (STS), the middle temporal gyrus (MTG) and, in some 103 104 instances, in the inferior parietal lobule (IPL), and the inferior frontal gyrus (IFG) (Demonet et al., 1992;Zatorre et al., 1992;Binder et al., 1994;Dhankhar et al., 105 1997; Giraud and Price, 2001; Vouloumanos et al., 2001; Liebenthal et al., 2005; Rimol et 106 al., 2005; Pulvermuller et al., 2006; Obleser et al., 2007; Tremblay et al., 2012). The 107 consistency of the STP and STS results in studies using words or sublexical units suggest 108 109 that these regions might be involved in the conversion of acoustical information into

phonological representations. However, because these studies have contrasted different
 types of sublexical units to non-speech or unintelligible speech-sounds, the level of
 processing (e.g. acoustical/phonetic, phonemic, syllabic, supra-syllabic) at which
 mechanisms implemented within the STP and STS operate remains unclear.

Neuroimaging studies in which phonological mechanisms are engaged by the use 114 of an explicit task (discrimination, rhyming) can more readily target specific mechanisms 115 operating at different sublexical levels (phonemic, syllabic, supra-syllabic) and offer 116 valuable insights into the functional contribution of STP regions to the perception of 117 speech sounds. For instance, STP and STS activation have been reported in studies using 118 a variety of auditory tasks: phonetic discrimination (Burton et al., 2000), rhyming (Booth 119 et al., 2002), syllable identification (Liebenthal et al., 2013), monitoring (Rimol et al., 120 2005) and phonemic judgments (Jacquemot et al., 2003). Other studies using a neural 121 adaptation paradigm to target phonetic processing have also identified regions within the 122 STP that responded more strongly to stimuli that were part of different phonemic 123 categories than those that felt within the same phonemic category (Dehaene-Lambertz et 124 125 al., 2005; Joanisse et al., 2007). Taken together, these studies support the notion of a key involvement of the STP and STS in processing sounds at different levels (phonemic, 126 syllabic). However, despite their importance, studies using explicit speech perception 127 tasks requiring judgments on speech sounds probably recruit to greater extent 128 phonological processes than do more naturalistic speech tasks. It is therefore unclear 129 whether similar regions would be recruited in the absence of a task. It is also unclear 130 131 whether phonological mechanisms operating at different levels (phonemic, syllabic, supra-syllabic) engage the same or different neural networks. Despite the scarcity of 132 133 studies addressing this issue, in a recent functional magnetic resonance imaging (fMRI) 134 study, McGettigan and colleagues (2011) manipulated both the complexity of syllabic and supra-syllabic information in pseudo-words during a passive listening task. Syllabic 135 complexity was manipulated by varying the number of consonant clusters (0 vs. 2) and 136 137 supra-syllabic complexity was manipulated by varying the number of syllables (2 vs. 4). An effect of supra-syllabic complexity was observed in the bilateral PT. However, no 138 139 positive¹ effect of syllabic complexity was reported. In contrast, Tremblay and colleague (2011), also using fMRI, varied syllabic complexity as indexed by the presence or 140 141 absence of consonant clusters during the passive listening of words and found that the right PT was sensitive to the syllabic complexity manipulation, supporting the idea that 142 143 the supratemporal cortex plays a role in processing syllabic information (Grabski et al., 2013). 144

One question that arises from this literature is whether specific sublexical processes can be localized to specific regions within the STP and STS. In the current experiment, we were interested in investigating the distinct and shared effects of syllabic and supra-syllabic complexity on brain activity during the processing of auditory sequences. To this aim, we parametrically manipulated phonological complexity along two dimensions (1) individual syllable complexity (presence or absence of a consonant

¹ The authors reported several brain areas in which blood-oxygen-level dependent (BOLD) signal magnitude was higher for pseudowords without consonant clusters than for pseudowords containing consonant clusters.

151 cluster in the syllable onset) and (2) sequence-level complexity (the ordering of syllables 152 within a sequence). Given the importance of the STP and STS in the processing of auditory information, we conducted an exploratory anatomical ROI analysis focusing on 153 154 a fine-grain parcellation of the supratemporal cortex and STS based on our previous work (Tremblay et al., 2012; Tremblay et al., 2013) to determine whether sub-regions within 155 the STP and STS process similar or different kind of sublexical information during 156 passive speech perception (i.e. syllabic or supra-syllabic). In these prior studies, we 157 158 demonstrated that sub-regions within the STP exhibited different patterns of sensitivity to speech sounds during speech perception and production, suggesting that the STP contains 159 a mosaic of functionally distinct areas. It is therefore possible that sub-regions within the 160 STP are processing the speech signal in different manners and at different levels, with 161 some focusing on spectral information, while others on syllable- or sequence-level 162 information. Based on the results from our previous studies, we hypothesized that some 163 sub-regions within the STP (in particular the PT) and STS would show similar patterns of 164 activation for both manipulations while others would show a preference for one 165 manipulation. For example, we expected the primary auditory cortex to be sensitive to 166 both manipulations, as both syllabic and supra-syllabic complexity increase acoustic 167 complexity. We also expected the PT to be sensitive to the syllabic manipulation based 168 on previous results (Tremblay and Small, 2011). 169

170 2. Materials and methods

171 2.1 Participants

172 The participants were 15 healthy right-handed (Oldfield, 1971) native French speakers (9 females; 26.8±4.8 years; range 21-34, education 17.3±1.9 years), with normal 173 hearing and no history of language or neurological/neuropsychological disorders. Hearing 174 was assessed using a pure tone audiometry (clinical audiometer, AC40, Interacoustic) for 175 each ear separately for the following frequencies: 0.25, 0.5, 1, 2, 3, 4, 5, 8, 12 and 16 176 177 kHZ. Then for each participant, a standard pure tone average (PTA: average of threshold at 0.5, 1 and 2 kHz) was computed for the left $(17.13\pm3.78 \text{ dB})$ and right ear (18.68 ± 3.17) 178 dB), since most of the speech sounds fall within this range (Stach, 2010). All participants 179 were screened for depression (Yesavage et al., 1982) and their cognitive functioning was 180 evaluated using the Montreal Cognitive Assessment scale (MOCA) (Nasreddine et al., 181 2005). All participants were within normal range on the MOCA and none of the 182 183 participants were depressive. The study was approved by the committee on research ethics of the Institut Universitaire en santé mentale de Québec (#280-2012). 184

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186 **2.2 Stimuli and task**

187 The experimental task consisted in listening passively (i.e. without performing a task) to sequences of syllables. To investigate sublexical phonological processing, we 188 189 used sequences of syllables instead of pseudowords to avoid lexical effects. Prior research has demonstrated that pseudowords, given their close resemblance to words, 190 191 activate regions involved in lexical access and in some cases they do so to an even greater extent than words (Newman and Twieg, 2001;Burton et al., 2005). Thus the use of 192 pseudowords renders the dissociation between lexical and sublexical phonological 193 processing extremely hard. For this reason, we decided to used syllable strings rather than 194 195 words to alleviate potential lexical effects. The degree of complexity of each sequence was manipulated along two phonological dimensions: syllabic and supra-syllabic 196

complexity. Each factor had two levels (simple or complex), resulting in a 2x2
experimental design matrix (See Table 1).

199 Syllabic complexity refers to the presence or absence of a consonant cluster (e.g. /gr/): simple syllables were composed of a single consonant and vowel (CV) and complex 200 syllables were composed of a consonant cluster and a vowel (CCV). Supra-syllabic 201 complexity refers to the number of different syllables in a sequence: simple sequences 202 were composed of the same syllable repeated 6 times (e.g. /ba-ba-ba-ba-ba/) and 203 complex sequences were composed of three different syllables each repeated twice (e.g. 204 205 /ba-da-ga-ba-da-ga/). While these two manipulations increase phonological complexity, they target different levels of processing; syllabic (individual unit) and supra-syllabic 206 (sequence of units). 207

All syllables were created by selecting among five frequent French vowels, which 208 included two front vowels $(/i/, /\epsilon/)$, two back vowels (/o/, /u/), and one central vowel (a), 209 and twelve frequent French consonants, which included four labial consonants (/b/, /p/,210 (v/, f/), four coronal consonants (/d/, /n/, /t/, /l/) and four dorsal consonants (/g/, /n/, /k/, /l/)211 / μ /). These vowels and consonants were combined to form sixty simple syllables (CV) 212 and sixty complex syllables (CCV). Each syllable was repeated a total of three times (i.e. 213 in three different sequences). Six-syllable sequences were created by producing 214 sequences of three different syllables twice (/pa-ta-ka-pa-ta-ka), or by repeating one 215 216 syllable six times (/pa-pa-pa-pa-pa/). A native young adult male French speaker from Ouebec City pronounced all syllable sequences naturally in a sound attenuated booth. 217 Each sequence was recorded five times and the best exemplar was selected to use in the 218 experiment. The syllable sequences were recorded at 44.1 KH using a unidirectional 219 microphone connected to a sound card (Fast Track C-400, M-audio), saved directly to 220 disk using Sound Studio 4.5.4 (Felt Tip Software, NY, USA), and edited offline using 221 222 Wave Pad Sound Editor 4.53 (NHC Software, Canberra, Australia). Each syllable sequence was edited to have an average duration of 2400ms. The duration of the syllable 223 224 sequences was the same across all experimental conditions (i.e. 2400ms). The root mean square (RMS) intensity was then normalized across all sound files. Individual sequences 225 226 were not repeated during the course of the experiment.

227										
	Table 1: Experimental conditions									
228	Code	Syllable type	Sequence type	Examples						
220	SS	Simple	Simple	/ba-ba-ba-ba-ba/						
225	SC	Simple	Complex	/fo-de-ro-fo-de-ro/						
230	CS	Complex	Simple	/kli-kli-kli-kli-kli/						
	CC	Complex	Complex	/bri-dre-klou-bri-dre-klou/						

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232 **2.3 Procedure**

This experimental paradigm resulted in four conditions of 30 trials each, for a total of 120 trials. Each trial lasted 6.5 s. A resting baseline condition was interleaved with the experimental conditions (60 trials). The randomization of the experimental and baseline conditions was optimized using Optseq2

237 (<u>http://surfer.nmr.mgh.harvard.edu/optseq/</u>). The four conditions were equally divided

- into two runs. A passive listening experimental paradigm was used; participants were not
- required to produce any overt response. All stimuli were presented during the delay in
- 240 acquisition (see Image acquisition section below) using Presentation Software
- 241 (Neurobehavioral System, CA, USA) through high-quality MRI-compatible stereo
- electrostatic earplugs (Nordic Neurolab, Norway), which provide 30 dB of soundattenuation.
- 244

245 **2.4 Image acquisition**

A 3T Philips Achieva TX MRI scanner was used to acquire anatomical and 246 functional data for each participant. Structural MR images were acquired with a T1-247 weighted MPRAGE sequence (TR/TE=8.2/3.7 ms, flip angle = 8° , isotropic voxel size = 248 1mm³, 256x256 matrix, 180 slices/volume, no gap). Single-shot EPI BOLD functional 249 images were acquired using a SENSE reduction factor of 2 to reduce the number of phase 250 251 encoding steps. In order to ensure that syllables were intelligible, a sparse image acquisition technique (Eden et al., 1999;Edmister et al., 1999;Hall et al., 1999;Gracco et 252 al., 2005) was used. A silent period of 4360 ms was interleaved between each volume 253 254 acquisition. The syllable sequences were presented 360 ms after the onset of the silent period. One hundred and eighty functional volumes were acquired across 2 runs 255 (TR/TE=6500/30 ms, volume acquisition = 2140 ms; delay in TR 4360 ms, 40 axial)256 257 slices parallel to AC/PC, voxel size= 3x3x3, no gap; matrix = 80x80; FoV= 240x240mm). This study was part of a larger project, which also included a speech production 258 task and a speech perception in noise $task^2$. Those two tasks will not be discussed as part 259 of this manuscript. The speech perception task that is the focus of the present manuscript 260 was always presented first to participants, followed by the speech production task and the 261 speech perception in noise task. Participants were not told until the production task that 262 they would be required to produce speech. This was done in order to avoid priming 263 264 subvocal rehearsal during the speech perception task. The speech perception in noise task has been reported elsewhere (Bilodeau-Mercure et al., 2014). 265

266 2.5 Data Analysis

267 2.5.1 fMRI time-series analyses

All functional time-series were motion-corrected, time-shifted, de-spiked and mean-normalized using AFNI (version 10.7, intel 64; Cox, 1996). All time points that occurred during excessive motion (i.e. > 1 mm) (Johnstone et al., 2006) were censored. The anatomical scan of each participant was aligned to their registered EPI time series using local Pearson correlations (Saad et al., 2009). The alignment was verified and

 $^{^{2}}$ Not all participants took part in all three tasks. Here we report the data from 15 young adults, whereas Bilodeau-Mercure et al., (2014) reported the data for a subset (11) of these participants, who performed the speech perception in noise task.

273 manually adjusted when necessary. For each participant and for each run a finite impulse 274 response ordinary least squares model was used to fit each time point of the hemodynamic response function for each of the four experimental conditions using 275 276 AFNI's tent basis function (SS, SC, CS, CC). Additional regressors for the mean, the linear and quadratic trend components as well as the six motion parameters were also 277 included. This model-free deconvolution method allows the shape of the hemodynamic 278 279 response to vary for each condition rather than assuming a single response profile for all 280 conditions (Meltzer et al., 2008). The interval modeled covered the entire volume acquisition (2.14 sec), starting with stimulus onset and continuing at intervals of 6.5 sec 281 282 (i.e. silent period and volume acquisition) for 13 sec (i.e. 2 TR). All analyses (wholebrain and ROIs) focused on the first interval (i.e. the first TR). The resulting time-series 283 were projected onto the 2-dimensional surfaces where all subsequent processing took 284 place. 285

For each participant, FreeSurfer was used to create a surface representation of the 286 participant's MRI (Dale et al., 1999; Fischl et al., 1999) by inflating each hemisphere of 287 the anatomical volumes to a surface representation and aligning it to a template of 288 average curvature. SUMA was used to import the surface representations into the AFNI 289 3D space and to project the pre-processed time-series from the 3-dimensional volumes 290 onto the 2-dimensional surfaces. Both the surface representations and the pre-processed 291 time-series were standardized to a common mesh reference system (Saad et al., 2004). 292 The time-series were smoothed on the surface to achieve a target smoothing value of 293 294 6mm using a Gaussian full width half maximum (FWHM) filter. Smoothing on the surface as opposed to the volume ensures that white matter values are not included, and 295 296 that functional data located in anatomically distant locations on the cortical surface are 297 not averaged across sulci (Argall et al., 2006).

298 2.5.2 Group-level node-wise analyses

299 Whole-brain group analyses were performed using SUMA on the participants' beta values resulting from the first level analysis (Saad et al., 2004). The group level 300 analyses focused on (1) the effect of passive auditory sequence perception on the Blood 301 oxygenation level dependent (BOLD) signal (2) the effect of syllabic and supra-syllabic 302 complexity on the BOLD signal during auditory sequence perception, (3) the contrast 303 between the effect of syllabic and supra-syllabic complexity, and (4) the conjunction of 304 305 the syllabic and supra-syllabic complexity effects. To identify regions recruited during the perception of auditory sequences, a node-wise linear regression was conducted 306 (perception > 0, one sample t-test option in the AFNI 3dttest++ program). To investigate 307 the effect of syllabic and supra-syllabic complexity, a 2-way repeated measure ANOVA 308 (rANOVA) was conducted (3dANOVA program) with syllabic complexity (simple, 309 310 complex) and supra-syllabic complexity (simple and complex) as within-subjects factors. To identify regions that exhibited a stronger response to one of the manipulations (i.e. 311 312 syllabic or supra-syllabic), we computed, at the individual subject level, the effect of syllabic complexity (complex syllables - simple syllables) and the effect of supra-313 syllabic complexity (complex sequences – simple sequences). At the group level, the 314 resulting t-maps were submitted to a paired sample t-test, to determine whether the two 315 contrasts (i.e. syllable and sequence contrast) differed (AFNI 3dttest++ program). For the 316 conjunction, we computed a map of the joint activation, for each subject, for syllabic and 317

supra-syllabic complexity (syllabic \cap supra-syllabic). Only voxels that were significant at 318 319 p = .05 (uncorrected) in both individual maps were included in the conjunction map. A group level average of the conjunction maps was then generated. All resulting group 320 321 maps were corrected for multiple comparisons using the Monte Carlo procedure implemented in Freesurfer. This correction implements a cluster-size threshold procedure 322 to protect against Type I error. For the first three analysis, based on the simulation results, 323 324 it was determined that a family-wise error (FWE) rate of p < 0.001 is achieved with a 325 minimum cluster size of 157 contiguous surface nodes, each significant at p < 0.01. For the conjunction analysis, we adopted a more lenient correction (a FWE rate of p 0.05 was 326 327 achieved with a minimum cluster size of 202 contiguous surface nodes, each significant at p < 0.05. 328

329 2.5.3 Exploratory anatomical ROI analysis

To examine the role of supratemporal regions in the processing of syllabic and 330 supra-syllabic information, we conducted an exploratory anatomical ROI analysis 331 focusing on a set of 16 a priori selected anatomical regions. In a previous study, using a 332 similar fine-grained parcellation, we demonstrated that several STP regions exhibited 333 differential sensitivity pattern to auditory categories (i.e. syllables or bird songs) and 334 sequence regularity (Tremblay et al., 2012). Here we used a similar parcellation scheme 335 with the addition of the superior temporal sulcus to investigate the sensitivity of these 336 regions to syllabic and supra-syllabic information. These bilateral ROIs included the 337 planum polare (PP), the superior temporal gyrus (STG), the superior temporal sulcus 338 339 (STS), the transverse temporal gyrus (TTG), the transverse temporal sulcus (TTS), the planum temporale (PT), the caudal segment of the Sylvia fissure (SF). These ROIs were 340 anatomically defined on the participant's individual cortical surface representation using 341 342 an automated parcellation scheme (Fischl et al., 2004;Desikan et al., 2006). This 343 parcellation scheme relies on a probabilistic labeling algorithm based on the wellestablished anatomical convention of Duvernov (1991). The anatomical accuracy of this 344 345 method is high, approaching that of manual parcellation (Fischl et al., 2002;Fischl et al., 2004; Desikan et al., 2006). The advantage of using anatomical (as opposed to functional) 346 347 ROIs based on individual micro-anatomical landmarks is that it can capture inter-subject anatomical variability, something that is loss when using normalized templates (i.e. 348 349 functional ROIs based on group level data or cytoarchitectonic maps). It is also more anatomically precise. Thus, given that we were specifically interested in exploring the 350 351 functional anatomy of the STP/STS, the choice of an anatomical ROI approach was logical. 352

To augment the spatial resolution of the FreeSurfer anatomical parcellation, we manually 353 subdivided the initial parcellation of each participant's inflated surface in the following 354 manner: the STS, the STG, the PT were subdivided into equal thirds whereas the SF, the 355 TTG and the TTS were subdivided into equal halves, resulting in 16 ROIs (refer to Fig. 1 356 357 and Table 2 for details). The use of thus modified FreeSurfer parcellation scheme is advantageous for several reasons: (1) it is based on a well-recognized anatomical 358 parcellation scheme, (2) it is systematic, (3) it is easily replicable across participants and 359 studies, and (4) it has been shown to reveal functional subdivisions within the STP. 360

361 For each participant, we extracted the mean percentage of BOLD signal change in each

- of the 16 bilateral ROIs. First, we determined which ROIs were significantly active
- 363 during the auditory perception of the sequences by testing the following hypothesis using
- FDR-corrected t-tests (Benjamini and Hochberg, 1995;Genovese et al., 2002) (q = .05):

365 (i) perception > 0, (n = 32, one-sample t-tests).

- For each ROI that was significantly active, we conducted a 3-way ANOVA with repeated
- 367 measurements on the magnitude of the BOLD signal as a function of hemisphere, syllabic 368 and supra-syllabic complexity. Within each ROI, all main effects as well as 2-way and 3-
- way interactions were examined using Bonferroni corrected paired-sample t-tests
- way interactions were examined using Bomerroin conected parted-sample t-($\alpha = 0.05$). A directed p values are reported
- 370 (α =0.05). Adjusted p-values are reported.

Table 2: Surface des	cription of the ROI parcellation
Regions	Description
Superior temporal	The FreeSurfer STS ROI is bounded anteriorly by the temporal
sulcus (STS)	pole, medially by the STG, laterally by the MTG and posteriorly by
	the IPL. We divided this region into roughly equal thirds along the
	rostro-caudal axis (STGp, STGm, STGa).
Superior temporal	The FreeSurfer STG ROI runs from the rostral edge of the STS to
gyrus (STG)	the supramarginal gyrus. It is bounded medially by the SF. We
	divided this region into roughly equal thirds along the rostro-caudal
	axis (STGp, STGm, STGp).
Planum temporale	The FreeSurfer PT ROI is bounded anteriorly by the TTS, medially
(PT)	by the SF, laterally by the STG and posteriorly by the
	supramarginal gyrus. We divided this region into roughly equal
	thirds along the rostro-caudal axis (PTp, PTm, PTa).
Transverse	The FreeSurfer TTS ROI is bounded posteriorly by the PT and
temporal sulcus	anteriorly by the TTG. We divided this region into two halves along
(TTS)	the medial-lateral axis.
Transverse	The FreeSurfer TTG ROI is bounded rostrally by the rostral extent
temporal gyrus	of the TTS, caudally by the caudal portion of the insular cortex,
(TTG)	laterally by the STG and medially by the SF. We divided this
	region in roughly equal halves along a medial-lateral axis.
Caudal segment of	The FreeSurfer posterior SF ROI runs from the lower end of the
the Sylvian Fissure	central sulcus to the end of the posterior ascending ramus (Dahl et
(SF)	al., 2006). We divided this region in roughly equal halves.
Planum polare	Unedited version of FreeSurfer. It is bounded rostrally by the
(PP)	temporal pole, caudally by the TTG and medially by the
	parahippocampal gyrus.

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373 3. Results

374 **3.1 Whole brain results**

The first whole-brain analysis focused on identifying brain regions that were significantly recruited during the perception of auditory sequences regardless of syllabic and supra-syllabic complexity. The node-wise linear regression identified regions within
the bilateral precentral gyrus, inferior frontal gyrus, medial superior frontal gyrus and
supratemporal cortex, as well as the left cingulate gyrus and right superior frontal gyrus
that were more active than during the perception of auditory sequences than the baseline
(i.e. rest) (for details, refer to Fig. 2 and Table 3).

The second analysis sought to identify brain regions that were sensitive to syllabic 382 complexity, supra-syllabic complexity. The node-wise rANOVA showed significant main 383 effects of syllabic complexity and supra-syllabic complexity within the STP (for details, 384 refer to Table 4 and Fig. 3A,B). As illustrated in Figure 3A, for the syllabic complexity 385 manipulation, significant clusters of activation were observed within the left TTGI 386 extending posteriorly into the SFp, and medially into the inferior sulcus of the insula as 387 well as the right TTGl extending posteriorly into the SFa, laterally into the STGm and 388 389 medially into the inferior circular sulcus of the insula (for details, refer to Table 4A). 390 These two regions were significantly more active for the complex syllables than the simple syllables. As illustrated in Fig. 3B, an effect of supra-syllabic complexity was 391 found within the left STGm extending medially into the STSm, and TTSI as well as the 392 right STGa/m, the right central sulcus and the right superior frontal gyrus. Only the 393 clusters within the STP were significantly more active for the complex sequences (see 394 395 Table 4B). No significant two-way interaction between syllabic complexity and suprasyllabic complexity was found. 396

The third analysis sought to determine whether brain regions responded more to one complexity manipulation than the other. The node-wise t-test showed that the effect of supra-syllabic complexity was stronger than the effect of syllabic complexity within supratemporal plane regions in the left STSp, STGp, and STGa, whereas the effect of syllabic complexity was stronger than the effect of supra-syllabic complexity in the left TTGl (for details, refer to Table 5 and Fig. 3C).

The last analysis focused on identifying regions that were sensitive to both 403 404 experimental manipulations. As illustrated in Fig. 3D, the conjunction between the syllabic complexity contrast and the supra-syllabic contrast revealed overlapping 405 406 activation for both experimental manipulation within left supratemporal plane regions (TTSm, TTSl, PTa, STGm), the cuneus as well as right supratemporal plane regions 407 (TTSm, TTSl, SFp), the right supramarginal gyrus, and the right subparietal sulcus. For 408 each area that responded to both manipulations, we quantified the number of participants 409 for which the two effects overlapped. As can be seen in Fig. 3D, less than 5 participants 410 411 shared common overlapping regions.

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Table 3: FWE-corrected whole-brain for the	speech per	ceptio	n netw	vork.				
Auditory sequences > rest	<u> </u>	1						
Anatomical location	Hemi	X	У	Z	F-value	p-value	Cluster size (nodes)	Area (mm)
STGa extending into the MTG, STGp,	Left						· ·	
SMG and circular sulcus of the insula		-54	3	-7	9.29	p<0.00001	14323	4808
(multiple clusters)								
Precentral gyrus extending into the central		58	2	10	5 01	n<0.00001	2300	855
sulcus, and the inferior frontal gyrus		-38	2	19	5.91	p<0.00001	2300	855
Medial superior frontal gyrus		-7	-4	56	7.29	p<0.00001	1589	439
Precentral gyrus		-51	-6	45	5.74	0.00004	863	252
Cingulate gyrus and sulcus		-4	-12	38	4.59	0.00035	543	155
Central sulcus		-36	-27	48	4.29	0.00064	442	149
Lateral-occipito-temporal sulcus		-43	-50	-10	5.31	0.00009	397	121
Parieto-occipital sulcus		-15	-58	14	3.84	0.002	191	53
Supramarginal gyrus		-50	-44	47	4.74	0.00026	180	41
Cingulate gyrus		-12	-41	1	3.52	0.0031	158	24
STGm extending into the TTGl, STGa, MTG and STSp	Right	61	-22	2	11.28	p<0.00001	11839	4200
Cingulate gyrus and sulcus (multiple clusters)		15	-23	45	6.94	p<0.00001	791	177
Subcentral gyrus and sulcus		58	-4	11	4.67	0.0003	660	171
Medial superior frontal gyrus (multiple clusters)		8	-24	53	4.38	0.0005	524	143
Central sulcus		45	-9	38	7.74	p<0.00001	270	102
Inferior circular sulcus of the insula and PP (2 clusters)		42	0	-20	4.62	0.0003	363	100
Middle frontal gyrus and precentral sulcus (2 clusters)		41	1	47	5.34	0.00008	359	86
Superior circular sulcus of the insula		38	-16	22	5.29	0.00009	171	61
Central sulcus		31	-28	50	3.87	0.0015	196	51
Superior frontal sulcus		25	0	47	4.33	0.00059	176	50
Precentral sulcus		19	-9	62	6.10	0.00002	204	47
Superior temporal sulcus		48	-54	21	-6.21	0.00002	297	41
Superior parietal gyrus		13	-74	44	4.94	0.0002	205	40

All coordinates are in MNI space and represent the peak surface node for each of the cluster (FWE: p=0.001, minimum cluster size: 157 contiguous surface nodes, each significant at p < .01). 2 clusters indicate that the activation cluster is not continuous.

19 -31

-8

4.32

0.0006

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Parahippocampal gyrus

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Table 4: FWE-corrected whole-brain BOLD res	sults.							
A. Syllabic complexity								
Anatomical location	Hemi	X	У	Z	t-value	p-value	Cluster size (nodes)	Area (mm)
TTGl extending into the SFp and the inferior circular sulcus of the insula	Left	-55	-14	2	5.84	0.00005	1618	596
TTGl extending into the SFa, STGm and inferior circular sulcus of the insula	Right	63	-14	4	5.26	0.0001	1106	370
B. Supra-syllabic complexity								
STGm extending into the TTSI and STSm (multiple clusters)	Left	-60	-12	-5	6.49	0.00002	1277	443
STGm and STGa		61	-4	-4	5.91	0.00004	240	83
Central sulcus	Right	37	-19	42	-4.71	0.0004	266	79
Superior frontal gyrus (multiple clusters)		8	-26	54	-6.82	0.000008	249	65

All coordinates are in MNI space and represent the peak surface node for each of the cluster (FWE: p=0.001, minimum cluster size: 157 contiguous surface nodes, each significant at p < .01). T-values are reported instead of F-values. T-values were obtained by contrasting the two levels of complexity for each experimental factor while collapsing across the other one. 2 clusters or multiple clusters indicate that the activation cluster is not continuous. 415

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Table 5: FWE-corrected whole-brain BOLD res	sults.							
A. Supra-syllabic complexity > Syllabic comp	plexity							
Anatomical location	Hemi	X	У	Z	t-value	p-value	Cluster size (nodes)	Area (mm)
STSp and STGp (two clusters)	Left	-50	-44	0	5.66	0.00005	600	196
STGa		-56	-2	-7	4.93	0.0002	172	78
TTGl		-37	-38	15	-4.63	0.0003	172	58
Central sulcus	Right	1	-11	16	-4.25	0.0007	211	60

All coordinates are in MNI space and represent the peak surface node for each of the cluster (FWE: p=0.001, minimum cluster size: 157 contiguous surface nodes, each significant at p < .01). T-values are reported instead of F-values. T-values were obtained by contrasting the two levels of complexity for each experimental factor while collapsing across the other one. 2 clusters or multiple clusters indicate that the activation cluster is not continuous

419 **3.2 Exploratory supra-temporal ROI analyses**

Only the ROIs that were significantly activated for speech perception were included in the subsequent analyses. Of the 32 ROIs, only the bilateral STSp was not significantly activated. For each remaining ROI (n=15), we investigated the main effects of hemisphere, syllabic complexity, supra-syllabic complexity as well as the two-way interactions between hemisphere*syllabic complexity, hemisphere*supra-syllabic complexity, syllabic complexity*supra-syllabic complexity and three-way interaction between hemisphere*syllabic complexity*supra-syllabic complexity.

426 Bonferroni adjusted p-values are reported.

As shown in Fig. 4, a main effect of syllabic complexity was observed in the TTGI 427 $(F_{(1,14)}=26.44, p=0.0002)$, the TTGm $(F_{(1,14)}=31.11, p=0.00007)$, the TTSl $(F_{(1,14)}=29.4, p=0.00009)$, 428 the TTSm ($F_{(1,14)}$ =17.13, p=0.001), the STGm ($F_{(1,14)}$ =8.71, p=0.011), the SFp ($F_{(1,14)}$ =5.90, p=0.029), 429 the SFa ($F_{(1,14)}$ =9.84, p=0.007), the PTa ($F_{(1,14)}$ =13.61, p=0.002) and the PTm ($F_{(1,14)}$ =4.84, p=0.045). 430 We then determined the type of stimuli driving the effect. For all nine regions, a stronger effect was 431 observed for complex than simple syllables (paired sample t-tests, Bonferroni corrected). For the 432 SFa, a significant hemisphere*syllabic complexity interaction was also observed ($F_{(1,14)}=8.39$, 433 p=0.012). Paired sample t-tests revealed that the source of the interaction was due to the presence of 434 an effect of syllabic complexity for the left SFa (t=4.39, p=0.003) but not the right SFa (t=1.358, 435 p=0.59) (for details, refer to Fig. 3). For the PTm, a significant syllabic complexity*supra-syllabic 436 complexity interaction was noted. Paired sample t-tests revealed that this interaction was due to the 437 presence of an effect of syllabic complexity for the complex (t=2.95, p=0.044) but not the simple 438 439 sequences (t=0.01, p=1) (for details, refer to Fig. 4). For the TTSm, a significant syllabic complexity*hemisphere interaction was observed. Paired sample t-tests revealed that this interaction 440 was due to a marginally significant difference when we computed a differential complexity score per 441 442 hemisphere [complex - simple syllable] and compared these scores across hemispheres (t=-2.51, p=0.06). A significant 3-way interaction was observed in the STSa. To investigate the source of the 443 3-way interaction, two-way interactions were computed. A two-way interaction between syllabic 444 complexity and hemisphere was found for complex sequences ($F_{(1,14)}=7.32$, p=0.018) but not for 445 simple sequences ($F_{(1,14)}$ =.413, p=0.531). Paired sample t-tests were computed. A marginally 446 significant difference (t=-2.67, p=0.054) was found when we computed a differential complexity 447 448 score per hemisphere [complex - simple syllable] and compared these scores across hemispheres. The overall pattern that emerges with regard to the syllabic manipulation is a significant 449 increase in sensitivity for complex syllables (i.e. CCV) relative to simple syllables (i.e. CV) in the 450

increase in sensitivity for complex syllables (i.e. CCV) relative to simple syllables (i.e. CV) in the
TTGl, TTGm, TTSm, TTSl, STGm, SFp, SFa, PTa and PTm. Furthermore, the SFa demonstrated a
lateralization effect during the processing of syllabic information (the left SFa was sensitive to the
syllabic manipulation but not the right SFa). Lastly, the PTm was the only region where an
interaction between the syllabic and supra-syllabic manipulations was observed. In this region, the
effect of syllabic complexity was restricted to complex sequences.

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As shown in Fig. 5, a main effect of supra-syllabic complexity was observed in the STSm ($F_{(1,14)}=5.89$, p=0.03), the STGa ($F_{(1,14)}=5.39$, p=0.036), the STGm ($F_{(1,14)}=27.38$, p=0.0001), the PTa ($F_{(1,14)}=8.64$, p=0.01), the TTSl ($F_{(1,14)}=10.95$, p=0.005), the TTSm ($F_{(1,14)}=11.67$, p=0.004), and the TTGm ($F_{(1,14)}=8.619$, p=0.011). We determined that for all seven regions, the complex sequences were driving the main effect of supra-syllabic complexity as they elicited higher levels of BOLD signal than simple sequences (paired sample t-tests, Bonferroni corrected). For the STSm and SFp, a hemisphere*supra-syllabic interaction was observed (STSm: $F_{(1,14)}=10.06$, p=0.007, 464 SFp: $F_{(1,14)}$ =11.84, p=0.004). For both regions, paired sample t-tests revealed that the source of the interaction was due to an effect of supra-syllabic complexity in the left hemisphere (STSm: t=3.851, 465 p=0.004, SFp: 2.55, p=0.046) but not the right hemisphere (STSm: t=0.64, p=1, SFp: t=.965, p=1). 466 The overall pattern that emerges with regard to the supra-syllabic manipulation is a 467 significant increase in sensitivity for complex sequences (i.e. three different syllables) relative to 468 simple sequences (i.e. same syllable repeated 3x) in the STSm, STGa, STGm, PTa, TTSl, TTSm and 469 470 TTGm. In addition, in two regions, the STSm and SFp an effect of hemisphere was observed. For both of these regions, the effect of supra-syllabic complexity was only observed in the left 471 hemisphere. 472 473 In sum, the pattern that emerges from the ROI analysis suggest that some ROIs (STGm, TTSl, TTGm, TTSm, PTa) are sensitive to both experimental manipulations while others are only sensitive 474 to one experimental manipulation (i.e. svllabic: left SFa, PTm, TTGI: supra-svllabic: left STSm, left 475 476 SFp; for details refer to Fig. 6). In addition, for ROIs that were sensitive to both manipulations, the

magnitude of the manipulations was equivalent given the absence of syllabic complexity*supra-

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480 4. Discussion

syllabic complexity interaction within these regions.

Neuroimaging studies have consistently documented the role of two large and functionally 481 heterogeneous cortical areas, the STP and STS, in the perception of speech sounds. However, a 482 detailed understanding of the role STP and STS in the processing of sublexical information has not 483 yet emerged. This is largely related to the intrinsic complexity of the speech signal. Indeed, 484 comprehending speech requires the interaction of complex sensory, perceptual and cognitive 485 mechanisms. The question, then, that naturally arises is whether these regions shows differential 486 patterns of activation as of function of the type of information being processed (syllabic versus supra-487 syllabic) (functional heterogeneity) and the specific sub-region (spatial heterogeneity). 488

The main objective of the current study was to examine, using fMRI, whether the processing 489 of syllabic and supra-syllabic information during a passive listening task involve similar or distinct 490 networks, with an emphasis on the STP and the STS. A passive listening paradigm was used in order 491 to minimize task-related cognitive/executive demands. Given the importance of the STP and STS in 492 speech processing, we conducted an exploratory ROI analysis focusing on 16 bilateral STP/STS sub-493 494 regions to determine whether differential patterns of activation would be observed as a function of the type of information processed (i.e. syllabic or supra-syllabic). To preface the discussion, the 495 results from the whole-brain analysis identified a network of regions involved in the perception of 496 speech sounds that is consistent with previous neuroimaging studies that contrasted the processing of 497 sublexical speech units to rest (Benson et al., 2001;Hugdahl et al., 2003;Wilson et al., 2004;Rimol et 498 al., 2005; Wilson and Iacoboni, 2006). In addition, the results clearly demonstrate that the processing 499 500 of auditory syllable sequences recruits both the left and right hemisphere, consistent with the notion that the processing of speech sounds is bilateral (Hickok and Poeppel, 2004;2007;Hickok, 2009). The 501 highly consistent results from the whole-brain and ROI analysis demonstrate that both syllabic and 502 supra-syllabic information are processed during passive listening. The anatomical specificity 503 afforded by the ROI analyses allowed us to go further in exploring the specific functional 504 contribution of sub-regions within the STP and STS during the perception of speech sounds. The 505 findings are discussed below. 506

Results from the whole-brain analyses demonstrate widespread bilateral supratemporal
 activation resulting from the syllabic manipulation. The widespread extent of this activation was not

expected based on previous fMRI results (McGettigan et al., 2011;Tremblay and Small, 2011). Of the 509 510 few studies that have investigated the effect of consonant clusters during passive speech perception, in one study, activation within the right PT was scaled to syllabic complexity (Tremblay and Small, 511 2011) and in the other, no positive effect was reported (McGettigan et al., 2011). Our finding of 512 widespread supratemporal effects may be related to the type of stimuli used. While in the present 513 study we used meaningless sequences of syllables, Tremblay and al. (2011) used whole words, for 514 which the mapping of sounds to linguistic representations may be more automatic, requiring less 515 resources for the processing of syllabic information. However, if the processing of syllabic 516 information interacts with lexical status, an effect of complexity should have been observed in the 517 McGettigan et al., (2011) study given that pseudo-words were used, which are not overlearned 518 stimuli with a stored lexical representation. It is possible that the absence of an effect of syllabic 519 complexity in the latter study is attributable to a less salient experimental manipulation. In the present 520 study, we contrasted sequences of syllables with either six or no consonant clusters, yielding a very 521 robust effect. Although the use of a passive listening paradigm minimized attention-directed 522 523 processes, mimicking more closely naturalistic speech perception situations, the use of syllable as experimental stimuli might have taxed to a greater extent phonological processes than the use of 524 pseudo-words and words. This line of reasoning is consistent with neuropsychological and 525 neurophysiological evidence suggesting that language comprehension does not depend on the 526 processing of sublexical units (i.e. units smaller than words, such as syllables, phonemes, and 527 phonetic features). For instance, it has been shown that patients with good word-level auditory 528 529 comprehension abilities can fail on syllable and phoneme discrimination tasks (Basso et al., 1977;Boatman et al., 1995). Similarly, electrocortical mapping studies have provided evidence that 530 phonological processes (e.g. syllable discrimination) and auditory word comprehension processes are 531 not entirely circumscribed to the same STP regions (for a review, refer to: Boatman, 2004). In sum, 532 while syllabic complexity effects are observed in sequences of syllables, further research need to 533 determine whether and how syllabic information contributes to the perception of speech sounds and 534 language comprehension. 535

Both whole-brain and exploratory ROI analyses identified a region that was sensitive to the 536 presence or absence of consonant clusters; the lateral part of the primary auditory cortex (TTGl). In 537 addition, the exploratory ROI analysis also identified the left SFa and PTm, as regions being 538 sensitive to the syllabic manipulation. These results tentatively suggest that this effect stems from the 539 addition of an extra consonant in the onset of the syllable and not from differences between adjacent 540 syllables (i.e. two different syllables). This pattern of response is consistent with the hypothesis that 541 these regions are sensitive to the structure of the syllable (i.e. whether it is phonologically complex or 542 not). Whether these regions respond to the complexity of the syllabic structure in general or to a 543 specific component of the syllable (i.e. onset, rhyme, nucleus, or coda) however remains to be 544 determined. Though the specific contribution of these three regions in the processing of syllabic 545 information is still awaits further specifications, these three regions are nonetheless robustly activated 546 during the perception of sublexical speech sounds (Benson et al., 2001;Hugdahl et al., 2003;Wilson 547 et al., 2004; Rimol et al., 2005; Wilson and Iacoboni, 2006). 548

An alternative hypothesis that could explain the complexity effect related to the addition of a consonant to form a cluster is that these regions are responding to an increase in phonological working memory due to an increase in sequence length. This is because the addition of a consonant cluster to increase syllabic complexity also increases the length of the sequence. However, previous studies that have manipulated item length to investigate phonological working memory have reported mixed results that seem dependent upon (1) how length was modulated (CV-CCV vs. number of syllables), (2) the type of stimuli used (words, pseudowords), and (3) task demands (passive

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556 listening, judgment or naming task) (Okada et al., 2003;Strand et al., 2008;McGettigan et al., 2011). The most consistent finding is that stimulus length as defined by the number of syllable yields more 557 reliable results than the addition of consonant clusters. Moreover, if our syllabic manipulation results 558 reflected an increase in phonological working memory, we would expect this contrast to vield 559 clusters of activation within the pre-motor cortex, the IFG, and the inferior parietal lobule, that is, 560 regions that are typically recruited during verbal working memory tasks (Paulesu et al., 1993;Honey 561 et al., 2000; Marvel and Desmond, 2012). However, none of these regions was found in any of our 562 contrasts. 563

Another alternative hypothesis is that the syllabic effect is due to an increase in 564 acoustic/phonemic complexity. Indeed, consonant clusters are more complex than single consonants 565 both acoustically and phonemically. Given that we parametrically varied both syllabic and supra-566 syllabic complexity, if this hypothesis were correct, we would expect the same regions to also exhibit 567 568 an effect of supra-syllabic complexity since the presentation of 3 different syllables as opposed to the same syllable presented 3 times also increases acoustical complexity. In addition, we would also 569 expect to see a syllabic complexity*supra-syllabic complexity interaction driven by a syllabic 570 complexity effect for both simple and complex sequences and a stronger effect of syllabic complexity 571 for the complex sequence. This pattern of result was not found in the SFa or the TTGl or the PTm. 572 However, in the PTm, a region identical to the one reported by Tremblay and colleagues (2011), 573 sensitivity to the syllabic manipulation was found only for the complex sequences. Combined with 574 the observation that this region is involved in speech production (Dhanjal et al., 2008; Tourville et al., 575 2008;Peschke et al., 2009;Zheng et al., 2010) and that its activation magnitude varies as a function of 576 577 syllabic complexity during both speech perception and production (Tremblay and Small, 2011), the result from the current study provides additional support to the hypothesis that the right PT is 578 involved in converting external auditory input into a phonological representation. Our results are in 579 agreement with this hypothesis because an effect of syllabic complexity only emerged in this region 580 when the sequences were composed of three different syllables (i.e. high supra-syllabic complexity). 581 In itself, the addition of a consonant cluster increases the complexity of the syllable template. The 582 additional complexity associated with processing three different sounds (high supra-syllabic 583 584 complexity) enhances the syllabic manipulation, as three different consonant clusters have to be mapped onto phonological representations as opposed to three single consonants. In sum, the current 585 results lend further support to the notion that regions within the posterior STP are important for the 586 processing of phonological information, perhaps through a template matching mechanisms that uses 587 spectrotemporal information to access stored syllabic representations (Griffiths and Warren, 588 2002; Warren et al., 2005). 589

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Both the whole-brain and exploratory ROI analyses identified two regions, the STSm and 591 592 STGa that were sensitive only to the supra-syllabic manipulation. This pattern of response suggests 593 that these regions are involved in tracking changes that affect the structure of the sequence. In the 594 present study, after having heard the second syllable of a sequence, participants could determine whether they would hear the same syllable again (i.e. in the case of simple sequences) or a different 595 596 syllable (i.e. in the case of complex sequences). Thus, after the second syllable, for simple sequences the continuation was completely deterministic and prediction about upcoming sounds could be made. 597 This pattern of response is also consistent with results from studies that have investigated the 598 599 perception of speech sounds using a neural adaptation and oddball paradigm. In these studies, cluster of activation were observed within these regions in response to the presentation of a deviant stimulus 600 (Vouloumanos et al., 2001; Joanisse et al., 2007). Overall, the results suggest that these regions are 601 involved in representing sequences overtime. Thus, speech perception mechanisms, even in the 602 absence of a task, are sensitive to changes that affect the structural properties of auditory sequences, 603

604 consistent with previous work (Tremblay et al., 2012).

Both whole-brain and exploratory ROI analyses also identified a group of regions that was 605 sensitive to both manipulations. These regions included the STGm, the TTGm, the TTSI, the TTSm, 606 the PTa and the SFp. Sensitivity to both manipulations suggests that these regions do not exhibit a 607 differentiation in processing syllabic or supra-syllabic information. In a previous neuroimaging study 608 using the same parcellation scheme of the supratemporal plane, both the TTSI and PTa responded to 609 speech and non-speech sounds, whereas the STGm, SFp and TTGm exhibited an absolute preference 610 for speech sounds (Tremblay et al., 2012), consistent with the idea that regions located anterior and 611 lateral the primary auditory cortex are involved in processing changes in spectro-temporal features 612 (Scott and Johnsrude, 2003). These results suggest that both syllabic and supra-syllabic information 613

614 recruits common mechanisms involved in processing acoustical information.

In the current study, we explored the neural mechanisms involved in the processing of syllabic and supra-syllabic information during passive speech perception. We demonstrated that both syllabic and supra-syllabic information are processed automatically during passive speech listening, a finding that is consistent with the finding of distinct neural representations for syllable and sequence-level information during speech production (Bohland and Guenther, 2006;Peeva et al., 2010). Importantly, these findings suggest that processing of sublexical information is automatic, at least during the processing of meaningless syllable sequences. Future studies need to examine whether the processing

- of sub-lexical information is automatic and necessary during language comprehension using more
- naturalistic stimuli such as words or connected speech. It is possible that the recruitment of
- 624 phonological mechanisms depends upon the context, or the kind or quality of auditory stimuli being
- processed. Degraded speech stimuli, for instance, could recruit sublexical phonological mechanisms
- to a greater extent than high-quality speech sounds. Nevertheless, the present study offers new insight
- 627 into the functional neuroanatomy of the system involved in sublexical phonological processing,
- highlighting the importance of the anterior two-thirds of the PT, the primary auditory cortices and the
- 629 middle part of the STS and STG in these processes.

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6. **Figure legends** 838

Figure 1: Panel A. Supratemporal and STS parcellation. Anatomical parcellation of the STP and STS 839

displayed on a flattened schematic representation. Panel B. Anatomical parcellation of the 840 supratemporal plane displayed on a lateral view of a left hemisphere smoothed white matter inflated 841 surface. 842

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Figure 2: Whole-brain analysis of BOLD response illustrating regions significantly active during 844 speech perception. Activation is shown on the group average smoothed flattened surfaces. All 845 analyses are controlled for multiple comparisons using a cluster extent of 157 vertices, and a single 846 node threshold of p < 0.01, to achieve a family-wise error rate of p < 0.001.

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Figure 3: Whole-brain analysis of BOLD response. Activation is shown on the group average 848

smoothed flattened surfaces. The first three analyses (Panel A,B,C) are controlled for multiple 849

comparisons using a cluster extent of 157 vertices, and a single node threshold of p < 0.01, to achieve 850

851 a family-wise error rate of p < 0.001. The last analysis (Panel D) is controlled for multiple

comparisons using a cluster extent of 202 vertices, and single node threshold of p < 0.05, to achieve a 852 family-wise error rate of p < 0.05. Panel A illustrates regions significantly active for the contrast 853

between levels of syllabic complexity (complex > simple). Panel B illustrates regions significantly 854

active for the contrast between levels of supra-syllabic complexity (complex > simple). Panel C 855

illustrates regions that were differently active for the two complexity contrasts ([complex sequence -856

simple sequence] - [complex syllable - simple syllable]). Panel D illustrates regions significantly 857

active for the conjunction of syllabic and supra-syllabic complexity (syllabic complexity \cap supra-858

syllabic complexity). The color scheme represents the number of participants in which an overlap 859 between the two manipulations was for (less than 5). 860

Figure 4: Patterns of syllabic complexity effects observed in exploratory STP and STS ROI analysis. 861

The results mapped onto a flattened schematic representation of STP and STS showing the 862

parcellation used in this study (different areas shown not to scale). Areas in dark purple represent a 863

main effect of complexity and areas in lighter purple represent areas were an interaction was 864

observed (hemisphere*syllabic complexity for the SFa and syllabic complexity*supra-syllabic 865 complexity for the PTm). Legend: PP = planum polare; TTG = transverse temporal gyrus (m =

866 medial, 1 =lateral); TTS = transverse temporal sulcus (m = medial, 1 = lateral); PT = planum

867 temporale (a = anterior, m = middle, p = posterior); SF = caudal Sylvian fissure (a = anterior, p = anterior)868

posterior); STG = superior temporal gyrus (a = anterior, m = middle, p = posterior); STS = superior 869

temporal sulcus (a = anterior, m = middle, p = posterior); * significant contrast at $p_{FWE}=0.05$, 870

Bonferonni corrected; n.s. non-significant contrast. Error bars represent standard error from the 871

872 mean.

873 Figure 5: Patterns of supra-syllabic complexity effects observed in exploratory STP and STS ROI

analysis. The results mapped onto a flattened schematic representation of STP and STS showing the 874

parcellation used in this study (different areas shown not to scale). Areas in dark blue represent a 875

main effect of complexity and areas in lighter blue represent areas were an interaction was observed 876

- (hemisphere*syllabic complexity for the SFp and STSm). Legend: PP = planum polare; TTG = 877
- transverse temporal gyrus (m = medial, l = lateral); TTS = transverse temporal sulcus (m = medial, l 878

- 879 =lateral); PT = planum temporale (a = anterior, m = middle, p = posterior); SF = caudal Sylvian 880 fissure (a = anterior, p = posterior); STG = superior temporal gyrus (a = anterior, m = middle, p = 881 posterior); STS = superior temporal sulcus (a = anterior, m = middle, p = posterior); * significant 882 contrast at p_{FWE} =0.05, Bonferonni corrected; n.s. non-significant contrast. Error bars represent 883 standard arror from the mean
- standard error from the mean.
- Figure 6: Patterns of main effects of syllabic and supra-syllabic complexity observed in exploratory
- 885 STP and STS ROI analysis. The results mapped onto a flattened schematic representation of STP and
- 886 STS showing the parcellation used in this study (different areas shown not to scale). Areas in dark
- 887 purple represent a main effect of syllabic complexity, areas in blue represent a main effect of supra-888 syllabic complexity, areas in orange represent areas where both an effect of syllabic and supra-
- syllable complexity, areas in orange represent areas where both an effect of synable and suprasyllable complexity were observed and areas in grey represent areas where no effect was observed.
- Legend: PP = planum polare; TTG = transverse temporal gyrus (m = medial, l = lateral); TTS =
- transverse temporal sulcus (m = medial, l =lateral); PT = planum temporale (a = anterior, m =
- middle, p = posterior); SF = caudal Sylvian fissure (a = anterior, p = posterior); STG = superior
- temporal gyrus (a = anterior, m = middle, p = posterior); STS = superior temporal sulcus (a =
- 894 anterior, m = middle, p = posterior).

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