1	Hand gestures as visual prosody: BOLD responses to audio-
2	visual alignment are modulated by the communicative nature of
3	the stimuli
4	
5	
6	Emmanuel Biau ^a
7	Luis Moris Fernandez ^a
8	Henning Holle ^c
9	César Avila ^d
10	Salvador Soto-Faraco ^{a, b}
11	
12	^a Multisensory Research Group, Center for Brain and Cognition, Universitat
13	Pompeu Fabra, Barcelona, Spain.
14	^b Institució Catalana de Recerca i Estudis Avançats (ICREA), Barcelona, Spain.
15	^c Department of Psychology, University of Hull, UK.
16	^d Department of Psychology, Universitat Jaume I, Castelló de la Plana, Spain.
17	
18	Manuscript accepted for publication in Neuroimage
19	
20	Corresponding author: Emmanuel Biau
21	
22	Dept. de Tecnologies de la Informació i les Comunicacions
23	Universitat Pompeu Fabra
24	Roc Boronat, 138
25	08018 Barcelona
26	Spain
27	+34 691 752 040
28	emmanuel.biau@free.fr

29

30 ABSTRACT

31

public addresses. speakers accompany 32 During their discourse with spontaneous hand gestures (beats) that are tightly synchronized with the 33 prosodic contour of the discourse. It has been proposed that speech and beat 34 gestures originate from a common underlying linguistic process whereby both 35 speech prosody and beats serve to emphasize relevant information. We 36 hypothesized that breaking the consistency between beats and prosody by 37 38 temporal desynchronization, would modulate activity of brain areas sensitive to speech-gesture integration. To this aim, we measured BOLD responses as 39 40 participants watched a natural discourse where the speaker used beat gestures. In order to identify brain areas specifically involved in processing hand gestures 41 42 with communicative intention, beat synchrony was evaluated against arbitrary visual cues bearing equivalent rhythmic and spatial properties as the gestures. 43 44 Our results revealed that left MTG and IFG were specifically sensitive to speech 45 synchronized with beats, compared to the arbitrary vision-speech pairing. Our results suggest that listeners confer beats a function of visual prosody, 46 complementary to the prosodic structure of speech. We conclude that the 47 emphasizing function of beat gestures in speech perception is instantiated 48 through a specialized brain network sensitive to the communicative intent 49 conveyed by a speaker with his/her hands. 50

51

<u>Speech perception; Gestures;</u> Audiovisual speech; Multisensory Integration;
 MTG; fMRI.

- 54
- 55
- 56

57 **1. INTRODUCTION**

58

In everyday life, most communicative interactions between humans involve 59 auditory and visual information. Indeed, in addition to auditory speech, listeners 60 often have visual access to the speaker's lips, head, body posture and hand 61 gestures. Here we concentrate on the communicative impact of the cospeech 62 gestures that speakers produce with their hand movements while talking to 63 someone (McNeill, 1992). By combining behavioral and physiological measures 64 65 like event-related potentials (ERPs), prior studies have demonstrated that, for example, gestures describing an object or an action (i.e. iconic gestures) alter 66 67 semantic processing of the spoken message (Kelly et al., 2004; Kelly et al., 2009; Wu & Coulson, 2010) or help disambiguate semantically complex 68 69 sentences (Holle et al., 2007). These studies suggest that gestures provide 70 information not present in the verbal modality alone, and support the idea that 71 both streams of information are in fact components of a common integrated language system (McNeill, 1992; Kelly, Creigh & Bartolotti, 2009). 72

73

74 Many fMRI studies have investigated the degree to which gestures and speech recruit common brain areas. For example, a recent study by Dick et al. (2014) 75 established the implication of a fronto-temporal network of language-related 76 areas when iconic gestures provide complementary information to speech. The 77 Superior Temporal Sulcus (STS) and the Middle and Superior Temporal Gyri 78 (MTG/STG), which are well known to respond to audiovisual (AV) speech (Nath 79 and Beauchamp, 2012; Calvert et al., 2000; Callan et al., 2004; Macaluso et al., 80 2004; Meyer et al., 2004; Campbell, 2008), have been found to be sensitive to 81 the semantic relationship and congruency between gestures and the spoken 82 message (Marstaller & Burianova, 2014). Greater BOLD responses in the STS, 83 84 inferior parietal lobule and precentral sulcus were found for the perception of spoken sentences accompanied by semantically corresponding iconic gestures, 85 86 as compared to meaningless movements or auditory-only versions (Holle et al., 2010; Holle et al., 2008). Willems et al, (2009) also found greater activations in 87 88 the left STS/MTG when spoken sentences were presented with simultaneous pantomimes (i.e. speech-independent gestures) whose shape matched the verb 89 90 of the utterance in meaning, as compared to incongruent ones. Additionally, the

left Inferior Frontal Gyrus (IFG) has been often found to respond to the
manipulation of the semantic relationship between gesture and speech
(Marstaller & Burianova, 2014; Willems et al., 2009; Willems et al., 2007),
suggesting a role in the integration of both streams of information to support
sentence comprehension (Glaser et al., 2013; Uchiyama et al., 2008; Willems et al., 2007);
Hagoort, 2005).

Although very relevant, these past studies have focused mostly on the 97 neural correlates of hand gestures conveying semantic content, leaving aside 98 other important functions of gestures, like their role as prosodic markers of 99 speech (Guellaï, Langus & Nespor, 2014). Additionally, in these prior studies, 100 101 participants were typically presented with single sentences where gesture-102 speech interactions happen in an impoverished context (i.e., short speech 103 fragments containing an isolated gesture corresponding to a critical word). If 104 one considers gestures and speech as two complementary sides of a common 105 underlying language system, a natural continuous flow of visual (gestural) and audio (speech) streams might be essential for the system to remain fully 106 107 functional (Hubbard et al., 2009; Biau & Soto-Faraco, 2013; Biau et al., 2015). 108

109 In the present study, we address the neural correlates of spontaneous beat 110 gestures. As compared to the more commonly studied iconic gestures, beats 111 are much less sophisticated in semantic content. Generally, beats are rapid biphasic flicks of the hand with no semantic content, serving to highlight 112 relevant information and structure the narrative discourse (McNeill, 1992; So et 113 al., 2012). These kinds of gestures are, by far, the most frequent class of co-114 115 speech gesture, and their use is very evident in public addresses, such as 116 political discourses. Based on several evidences, it is now widely hypothesized 117 that beat gestures may also play a role in prosodic processing (Guellaï, Langus 118 & Nespor, 2014). First, beats seem to be very precisely aligned with speech 119 envelope. The functional phase of beats - the moment of maximum extension of 120 the movement, called the "apex" - is temporally aligned with the pitch accent of 121 its affiliate spoken word, increasing its prominence by modulating the acoustic properties of the accentuated syllable (Yasinnik, Renwick & Shattuck-Hufnagel, 122 2004; Krahmer & Swerts, 2007; Treffner and al., 2008; Leonard & Cummins, 123 124 2010). Second, the speakers use the timing of gesture's apexes to pack related

125 information together, possibly playing a role in the syntactic organization of sentences supported by prosody (Holle et al., 2012; Guellaï, Langus & Nespor, 126 127 2014). The few studies that have investigated the neural correlates of beat gestures support the prosodic hypothesis too. For instance, Biau & Soto-Faraco 128 (2013) found that beats modulate early ERPs time-locked to the affiliate words 129 130 onset, within the latency window corresponding to phonological processing. Holle et al. (2012) also found that beats in complex sentences modulated the 131 P600 ERP component, associated to syntactic analysis. Finally, in an fMRI 132 133 study, observers watched a speaker producing beats while spontaneously 134 speaking (Hubbard et al., 2009). The authors reported greater activations in the 135 left STG/S in response to speech when it was accompanied by beats as 136 compared to unrelated sign language gestures. They also reported greater 137 BOLD responses in the bilateral posterior STG/S, including the Planum Temporale (PT) for speech accompanied by beats compared to a still body. 138 139 Using beats from an actual fragment of continuous discourse ensured that 140 gestures were produced in a legitimate context and frequency. In addition, 141 spontaneous speech production ensured that the temporal relationship between the continuous beats stream and the rhythm of speech was maintained as in 142 natural language conversation (Biau et al., 2015). 143

144

145 Scope of the present study

146

147 We hypothesize that beat gestures are produced as an integral part of 148 the language system, providing the listener with visual prosodic information that 149 is aligned with the prosodic contour of the speech message. For this reason, we 150 advance that precise temporal alignment is essential to engage brain processes 151 related to the integration of beats and speech. If this is true, brain activations in 152 relevant integration areas may be sensitive to a breach in the temporal 153 synchrony of beats with respect to their speech affiliates (Marstaller & Burianova, 2014; Hubbard et al., 2009). To test this hypothesis, we used fMRI 154 155 while participants were presented with video clips in which the video was either synchronized with the audio track or lagged behind 800 milliseconds. With this 156 157 manipulation, we assumed that when beat's apexes fall out of synchrony with 158 their affiliated speech accentuations, their highlighting function would falter. Yet, 159 please note that desynchronization between beats and speech involves 160 temporal misalignment at many levels, from mere spatio-temporal correlations 161 of low level features to the misalignment in linguistic functions. Therefore, an 162 integral question in this framework is whether the putative prosodic function of beats relates to a generic mechanism of visual emphasis or, alternatively, 163 164 whether beats engage a specialized mechanism. Revealing such specialization is essential to attribute any beat-speech interaction effects to a common 165 underlying language system. For instance, it is relevant that in the study by 166 167 Holle et al. (2012), mentioned above, the authors did not find the same effects 168 on the P600 ERP component when speaker's moving hands (producing the 169 beats) were replaced with discs following equivalent spatio-temporal trajectories 170 in the visual display. The authors concluded that beats bear additional 171 communicative intentions above and beyond simple visual emphasis (e.g. intentions and postures that come along with the prosodic variations, which 172 173 might not be the case for an isolated disc).

Following Holle et al.'s logic, we wanted to single out brain areas that 174 175 play a relevant and specific role in beat-speech integration by looking at the 176 effect of beats-speech (de)synchronization, compared to the same effect when the speaker's hands are replaced by arbitrary visual cues (i.e., moving discs). 177 178 We hypothesized that the visual emphasis from arbitrary stimuli may differ from 179 the linguistic function that gestures have when combined with speech (i.e. when beat emphasis is synchronized with the speech prosody). If beat gestures 180 181 effectively confer a special communicative value to the spoken message, then 182 one should expect disparate effects of audio-visual synchrony for beat gestures as compare to visual cues. We set up a 2x2 design with the factors AV 183 184 synchrony (synchronous or asynchronous) and visual information (beats or discs) to test how the temporal alignment affects the integration of speech with 185 186 either type of visual information. The interaction between synchrony and visual 187 information is of essential interest because it allows isolating brain areas in which the impact of synchrony depends on which kind of visual information 188 189 (beats or discs) accompanies audio speech prosody. Please note that a simple comparison between synchronous-asynchronous would conflate brain areas 190 that are sensitive to generic, low level features as well as more specific 191 192 linguistic related attributes of the stimuli. Thus, in this study we will mainly

concentrate on brain areas where such an interaction arises. According to prior
literature, these areas might (though not exclusively) correspond to the ones
previously shown to be sensitive to gesture-speech integration, such as the left
STS/G but also the left IFG (Holle et al., 2007; Willems et al., 2007; Hubbard et
al., 2009; Holle et al., 2010; Marstaller & Burianova, 2014).

198

199 2. MATERIAL AND METHODS

200

201 2.1 Participants

202

203 Nineteen native speakers of Spanish (12 female, age range 19-29) took part in 204 the current study. All participants were right-handed with normal auditory acuity 205 as well as normal or corrected-to-normal vision. Participants gave informed 206 consent prior to participation in the experiment and the study was approved by 207 the University's ethics committee. Due to a technical problem, two participants 208 could not listen to the speech stream during fMRI data acquisition and were 209 therefore excluded from the statistical analysis. Thus, data from 17 participants 210 (12 females, age range: 22.4 ± 2.4 years old) were included in the imaging 211 analysis.

212

213 2.2 Material and stimuli

214

We extracted 44 video clips (18 s duration each) from a political discourse of 215 the former Spanish President Luis Rodríguez Zapatero, recorded at the palace 216 of La Moncloa and available on the official website (Balance de la acción de 217 218 Gobierno en 2010, 12-30-2010; http://www.lamoncloa.gob.es). During the whole public address, the speaker stood behind a lectern, with the upper part of the 219 220 body in full sight. The video clips were edited using Adobe Premiere Pro CS3. We visually inspected the entire discourse to select relevant segments of 221 speech, containing only beats and cohesive gestures (series of beats that link 222 successive points to a common concept) according to McNeill's definition. Clear 223 iconic gestures were not found but as gesture categories sit along a continuum 224 225 with fuzzy boundaries, some gestures may fall into multiple categories. Therefore 226 one cannot be absolutely certain that our stimuli never included a minimum of

semantic content in the hand shape. However, hand movements always conformed 227 to McNeill's definition of beat gestures. To avoid abrupt onsets and offsets, we 228 229 introduced 1 second audio-visual fade-in and -out at the beginning and end of each clip (respectively). In all the AV clips, the head of the speaker was masked 230 with a superimposed ellipse-shaped patch in order to remove any facial 231 232 information, such as lips or evebrow movements, as well as head movements. After editing, videos were exported using the following parameters: video 233 resolution 960x720, 25 fps compressor Indeo video 5.10, AVI format; audio 234 sample rate 48 kHz 16 bits Mono. As explained below, we created four different 235 versions for each video, corresponding to the four conditions of our 236 237 experimental design: Beat Synchronous (Bs), Beat Asynchronous (Ba), Disc Synchronous (Ds) and Disc Asynchronous (Ds) (Fig. 1). 238



240

239

Figure 1. Screenshots from (i) Beat and (ii) Disc conditions. Audio and video streams were either synchronized (Bs and Ds conditions) or desynchronized (audio lagged video by 32 frames, corresponding to 800 ms) with respect to audio in the Ba and Da conditions). Green arrow illustrates the trajectory of a beat gesture and the corresponding disc. The apex of the movement coincided in this case with the Spanish word 'crisis'.

Beat conditions: We selected 44 segments (18s each, 450 frames) of the 246 247 discourse in which the speaker naturally produced spontaneous beats (McNeill, 1992). For each clip, the speaker produced a minimum of 8 beats within the 18 248 s (mean number of gestures per clip: 12.8 ± 4.2). To create the Beat-249 250 Synchronous condition, audio and visual information remained synchronized as in the original discourse, with the speaker's hands fully visible (beat synchrony, 251 Bs). For the beat asynchrony (Ba) condition, audio and visual information were 252 desynchronized by inserting a lag of 800 ms (32 frames), leading to speech 253 254 preceding beat gestures.

Disc conditions: To create the disc conditions, the video was removed and the 256 257 hands were replaced by two discs that followed the hand trajectories of the 258 original clips. We defined the junction between the index and the thumb as the 259 reference point of both hands. We used Skin Color Estimation Application and ELAN software to detect pixel coordinates of hands frame-by-frame in each 260 Beat video (http://tla.mpi.nl/tools/tla-tools/elan; Max Planck Institute for 261 Psycholinguistics, The Language Archive, Nijmegen, The Netherlands; 262 263 Wittenburg et al., 2006). Reference point coordinates were reviewed and 264 corrected were necessary for both hands using custom-made scripts for Matlab 265 (MATLAB Release 2012b, The MathWorks, Inc., Natick, Massachusetts, United 266 States). The two discs representing the hands had a 40 pixel diameter size and 267 were flesh-colored (Red, Green, Blue color values: 246, 187 and 146) at their corresponding reference point. The background color was set to the average 268 269 value of a still frame of the speaker (Red Green Blue Value: 110, 114, and 104). We then created a synchronized (Disc Synchrony, Ds) and a desynchronized 270 271 (Disc Asynchrony, Da) condition following the same process as in the beat 272 condition.

273

Target videos: To ensure that stimuli were attended, participants performed an auditory detection task. For this, we used two clips from each experimental condition to create 8 targets. For each target video, the fundamental pitch of the original audio tracks was artificially shifted up three semitones (high pitch) for one syllable using Adobe's PitchShift filter while the intensity remained the same. In total, each participant was presented with 36 experimental and 8 target videos.

- 281
- 282 2.3 Procedure and Instructions
- 283

Participants were presented with 44 trials using E-Prime2 software. The order of trials was pseudo-randomized to avoid direct repetition of experimental conditions. Each trial consisted of a fixation cross with variable duration (from 7.5 to 8.5 seconds in steps of 0.25 seconds, uniformly distributed) followed by a video clip. The next trial began automatically after the end of the preceding

255

video. A total of four experimental lists were created, counterbalanced for thefour experimental conditions. Each participant saw one of the four lists.

291

292 Participants were instructed to perform an auditory detection task and press a button of the fMRI-compatible controller as soon as they detected an artificial 293 294 pitch change in the voice of the speaker. The hand holding the controller (left or right hand) was counterbalanced across participants (even though target trials 295 were not included in the statistical analysis). Participants were also instructed to 296 297 always look at the screen during the whole experiment as if they were watching television. Before the fMRI acquisition, participants performed a rapid training 298 299 with an extra target video presented in both Bs and Ds conditions as an 300 example of artificial pitch change. After the scanning session, participants were 301 given a questionnaire, asking 1) Did you perceive any asynchrony between 302 video and speech during the experiment? 2) What could the moving discs 303 represent? This questionnaire served to ensure that participants correctly attended to all videos. More importantly, it allowed us to evaluate if they could 304 305 perceive the asynchrony between video and speech.

306

307 2.4 fMRI acquisition

308

309 Imaging was performed in a single session on a 1.5 T Siemens scanner. We first acquired a high-resolution T1-weigthed structural image (GR\IR 310 TR=2200ms, TE=3.79ms, FA=15°, 256 x 256 x 160, 1mm isotropic voxel size). 311 312 Functional data was acquired in a single run consisting of 610 Gradient Echo EPI functional volumes (TE = 50 ms, TR = 2000 ms) not specifically co-planar 313 314 with the Anterior Commisure - Posterior Commisure line, acquired in an interleaved ascending order using a 64x 64 acquisition matrix with a FOV = 315 316 224. Voxel size was 3.5 x 3.5 x 3.5 mm with a 0.6 mm gap between slices, covering 94.3 mm in the Z axis.. The functional volumes were placed attempting 317 to cover the whole brain in 23 axial slices. The first four volumes were discarded 318 to allow for stabilization of longitudinal magnetization. 319

320

321 2.5 Imaging data analysing

322

FMRI data were analyzed using SPM12b (www.fil.ion.ucl.ac.uk/spm) and
Matlab R2013b (MathWorks).

325

326 2.5.1. Preprocessing

327

Standard spatial preprocessing was performed for all participants using the 328 following steps: Horizontal AC-PC reorientation; realignment and unwarp using 329 the first functional volume as reference, a least squares cost function, a rigid 330 body transformation (6 degrees of freedom) and a 2nd degree B-spline for 331 interpolation, creating in the process the estimated translations and rotations 332 333 occurred during the acquisition; slice timing correction using the middle slice as 334 reference using SPM8's Fourier phase shift interpolation; coregistration of the 335 structural image to the mean functional image using a normalized mutual information cost function and a rigid body transformation. The image was then 336 337 normalized into the Montreal Neurological Institute (MNI) space (Voxel size was changed during normalization to isotropic $3.5 \times 3.5 \times 3.5$ mm and interpolation 338 339 was done using a 4th B-spline degree). Functional data was smoothed using an 340 8-mm full width half-maximum Gaussian kernel to increase signal to noise ratio and reduce inter subject localization variability. To add an extra quality control 341 to the movement in participants, we used the Artifact Detection tools (ART) 342 (http://www.nitrc.org/projects/artifact_detect/) with 343 which the composite movement was calculated. This provides a single measure that comprises the 344 movement due to rotation and translation between volumes. All volumes with a 345 composite movement of more than 0.5 mm or more than 9 standard deviations 346 away from the global mean signal of the session were considered as outliers 347 348 (On average, 1.4% of the volumes per participant were detected as outliers). One regressor per outlier was added at the first level to discard any possible 349 350 influence of these volumes in the final analysis.

351

352 2.5.2. fMRI analysis

353

The time series for each participant were high-pass filtered at 128 s and prewhitened by means of an autoregressive model AR(1). At the first level (subjectspecific) analysis, box-car regressors modelling the occurrence of the four

conditions of interest (Bs, Ba, Ds and Da) and a fifth regressor for trials 357 358 containing a target, all modelled as 18s blocks, were convolved with the standard SPM12b hemodynamic response function. Additionally, several 359 regressors of no interest were included, including the six movement regressors 360 provided by SPM during the realign process, the extra composite movement 361 regressor calculated with ART and one regressor for each of the volumes 362 considered as outliers. The resulting general linear model produced an image 363 estimating the effect size of the response induced by each of the conditions of 364 365 interest. The images from the first level were used for the planned critical 366 contrasts in a second level analysis (inter-subject). At the second (inter-subject) 367 level, these images were entered into a random effects factorial design with five levels, corresponding to the four critical conditions, plus an additional subject 368 369 constant to account for non-condition-specific inter-subject variance. Correction for non-sphericity (Friston et al., 2002) was used to account for possible 370 371 differences in error variance across conditions and any non-independent error 372 terms for the repeated measures. Statistical images were assessed for cluster-373 wise significance using a cluster-defining threshold of p<0.001. The 0.05 374 Family-wise error correction critical cluster size was 31 voxels and was determined using random field theory (Data smoothing FWHM: 11.4mm, 375 11.2mm, 11.3 mm. Resel Count: 749.2), considering the whole brain as a 376 377 volume of interest. Contrasts vectors assessing the two main effects and the interaction were used. Although the whole interaction statistical parametric map 378 is presented, the discussion is limited to the clusters that showed an effect of 379 380 Beat gestures compared to Discs (Bs+Ba > Ds+Da), as our main interest is 381 focused on the parts of the brain that are involved in beat processing (for 382 unmasked results and additional contrasts, please see supplementary online 383 materials). To achieve this, we masked the interaction contrast, corrected as 384 explained above, with the Beat > Discs contrast (p-threshold (unc.) <0.05). MNI coordinates were classified as belonging to a particular anatomical region using 385 386 the SPM Anatomy Toolbox (Eickhoff et al., 2005).

387

388 3. RESULTS

389

390 3.1 Behavioral results

391

Participants correctly detected pitch deviation targets on $65.4\% \pm 31.7\%$ of the target trials and gave False Alarm (FA) responses only on $7.0\% \pm 13.6\%$ of the non-target trials.

395

396 3.2 Post-scanning questionnaire

397

When asked, after the scanning session, whether they perceived any 398 399 asynchrony between video and speech during the experiment, 12 participants 400 responded "yes"; 3 participants responded "yes, but not in the disc condition" and 2 participants responded "no". With respect to the second question ("What 401 402 could the moving discs represent?"), all participants responded "the hand of the 403 speaker. This suggests that the asynchrony between beats and speech was 404 noticeable, even though facial information was removed from videos. 405 Furthermore, this consistent response confirmed that the spatiotemporal characteristics of disc movements successfully mimicked the hand trajectories 406 407 in the Disc conditions. Both the behavioural and post-scanning questionnaire 408 results suggest that participants were attentive to the AV stimuli.

409

410 3.3 fMRI results

411

412 3.3.1 Differential effect of AV synchrony depending on visual information

413

414 The first contrast of interest concerns the interaction between synchrony and 415 visual information [(Bs-Ba) – (Ds-Da)]. This contrast is of particular interest as it 416 highlights the brain areas where the impact of synchrony depends on which kind of visual information (beats or discs) accompanies speech. We studied this 417 418 interaction in the areas that showed an effect of Beat > Disc (uncorrected mask p<0.05), as explained in the methods section (see Table 1). This restricts our 419 420 analysis to areas that are related to beat processing. The results revealed a significant interaction in BOLD responses in two different clusters of the left 421 Middle Temporal Gyrus and Superior Temporal Sulcus (MTG/STS), one more 422 posterior and one more anterior (respectively, pMTG and aMTG/STS). 423

424 Additionally, significant interactions in left IFG and left occipital cortex425 (Brodmann area 18) were observed.



Interaction

426 427

Figure 2. Interaction contrast [(Bs- Ba) – (Ds – Da) inclusively masked with the main effect of Beat (Bs+Da) compared to Disc (Ds+Da) using a p<0.05 cluster-corrected threshold with a minimum cluster size k = 31 and rendered on a 3D brain surface in MNI space (Left hemisphere). Error bars show 1 S.E.M of parameter estimates. IFG: Inferior frontal gyrus (-41 32 -11); Ant.MTG: anterior Middle temporal gyrus (-52 -7 -18); Post. MTG: posterior MTG (-59 -46 -4); Occipital (-20 -95 14).

434

435 These results suggest that synchrony differentially affects speech integration, 436 depending on the content of visual information. In particular, speech-gesture 437 synchrony seems to recruit left-hemisphere brain areas preferentially, as compared to other visual cues which share the same spatio temporal properties 438 but are arbitrary. Post-hoc analysis in the four significant clusters revealed that 439 activations were significantly greater when beats and audio were synchronized 440 (Bs) than asynchronous (Ba). Furthermore, the effect of synchrony on brain's 441 activations was exactly the opposite when beats were replaced by simple discs 442 (see Figure 2; see the significance of post-hoc simple main effects in the 443

Supplementary Material). It is worth noting that the areas which display this 444 445 pattern (MTG, IFG and Occipital cortex in the left hemisphere) and the directionality of the numerical effects of beat synchrony are well in line with 446 447 previous studies investigating gesture perception (Hubbard et al., 2009; Willems et al., 2009; Skipper et al., 2007; Holle et al., 2008, 2010), which further 448 449 reassures the interpretation of these activations. Yet, despite this is the pattern expected from prior results and support our hypothesis, one should be careful 450 from putting too much weight on it, given the post-hoc nature of the test. 451

- 452
- 453

454 3.3.2 Effect of type of visual information within temporal synchrony

455

456 Looking at the main effect of type of visual cue within the synchronous 457 conditions can reveal differences arising from the type of visual stimulus. The 458 contrast Beat Synchronous > Disc Synchronous revealed a greater BOLD 459 response in various brain areas when speech was accompanied by 460 synchronized beats (Bs), relative to synchronized discs (Ds) (see figure 3 and table 1). Not surprisingly, the greatest difference was observed in the occipital 461 cortex likely due to a pure difference in visual information between conditions. 462 The contrast also revealed differences in beyond visual brain areas, such as a 463 significantly greater BOLD activity in the left MTG/STS, as well as in the left 464 Inferior frontal Gyrus (left IFG) and left hippocampus. The contrast Ds>Bs 465 revealed greater BOLD activity when speech was accompanied by synchronous 466 467 discs rather than synchronous hand beats in the Superior Parietal areas bilaterally and right Angular Gyrus (see figure 3 and table 1). 468

469



470

© 2016. This manuscript version is made available under the CC-BY-NC-ND 4.0 license http://creativecommons.org/licenses/by-nc-nd/4.0/

471

Figure 3. Main effect of Beat Synchronous (Bs) compared to Disc Synchronous (Ds). Statistical
maps are thresholded at *P*-uncorrected <0.001 with a minimum cluster size k = 31 and rendered
on a 3D brain surface in MNI space. From left to right: left hemisphere, right hemisphere and an
axial cut at z=0. Hot colors indicate Bs > Ds. Cold colors indicate Ds> Bs.

476

477 3.3.3 Effect of synchrony between beat gestures and speech

478

The contrasts involving the comparisons Bs>Ba and Ba>Bs, restricted within 479 the beat gesture conditions, revealed no main effect of synchrony, when 480 performed at the whole brain level. Note that this particular result deviates from 481 Hubbard et al. (2009), who reported an effect of synchrony in the left STS/G 482 483 area. However, it must be mentioned that in Hubbard's study not only the actual 484 synchrony, but also the nature of the gestures themselves was substantially 485 changed between the synchronous and asynchronous condition (beats vs. ASL 486 gestures in the control condition, respectively). In any case, our result implies that despite the BOLD responses for synchronous gestures tend to be larger 487 488 than the BOLD responses for asynchronous gestures in the areas of significant 489 interaction (as revealed in the interaction analysis). However, as discussed in 490 the introduction, this effect cannot be fully interpreted without factoring in the responses of these areas to the disc synchrony/asynchrony conditions. This is 491 because several low-level generic, as well as language-specific responses to 492 493 synchrony are conflated in this contrast.

494 495

Hemisphere	Region	Corrected Cluster	Number of	Z - Score	Coordinates (mm) ^b		
		P-Value	Voxels ^a		X	У	z
Interaction [(Bs-Ba) -	- (Ds-Da)] masked with Beat > Dis	c (mask p-value	<0.05)				
L	Middle Temporal Gyrus	0,043	32	5,93	-59	-46	-4
L	Inferior frontal gyrus	0,048	31	4,36	-41	32	-11
L	Temporal Pole	·		4,35	-45	14	-18
L	Middle Temporal Gyrus	0,048	31	4,20	-52	-7	-18
L	Middle Temporal Gyrus			4,10	-59	-11	-14
L	Middle Temporal Gyrus			4,09	-59	-4	-21
L	Middle Occipital	0,039	33	4,04	-20	-95	14
L	Inferior Occipital			3,38	-31	-88	4
Beat Synchronous >	Disc Synchronous						

R	Lingual Gyrus	0.000	3080	Inf	8	-88	4
Ĺ	Cuneus	0,000	0000	Inf	-10	-98	18
L	Calcarine			Inf	-3	-88	-4
L	Middle Temporal Gyrus	0,000	151	5,22	-62	-11	-14
L	Temporal Pole			4,75	-48	18	-14
L	Inferior Frontal Gyrus			4,33	-41	28	-11
L	Thalamus	0,006	52	5,20	-24	-28	0
L	Middle Temporal Gyrus	0,001	75	4,90	-55	-46	0
L	Middle Temporal Gyrus			3,93	-48	-32	0
Disc Synchronous > Beat Synchronous							
L	Superior Parietal	0,006	50	4,75	-16	-70	56
R	Superior Parietal	0,009	47	3,73	22	-66	59
	Angular Gyrus			3,49	22	-56	49
	Superior Parietal			3,40	15	-59	63
Beat Synchrono	us > Beat Asynchronous						
No significantly activate regions							
Beat Asynchron	ous > Beat Synchronous						
No significantly activate regions							

496

497

Table 1.^a Number of voxels exceeding a voxel-height threshold of p < 0.001 using a p < 0.05
cluster-extend FWE correction. ^b First three maximum peaks more than 8 mm apart are reported
for each cluster.

501

502

503 4. DISCUSSION

504

In the present study, we investigated the neural correlates of spontaneous beat 505 506 gestures accompanying continuous, natural <u>spoken</u> discourses. Based on previous reports (McNeill, 1992; Yasinnik et al., 2004; Guellaï et al., 2014; Biau 507 508 et al., <u>2015</u>), we hypothesized that beats act as a visual counterpart of prosody. If this is the case, then breaking up the consistency between beat apexes and 509 510 speech prosody may affect speech processing. In terms of neural expression, we hypothesized that if beats are integrated as linguistically relevant 511 512 information, brain activity in relevant integration areas may be modulated by an 513 asynchrony between visual and audio streams. As an integral aspect of this 514 <u>question</u>, we addressed whether beats convey additional communicative aspects above and beyond arbitrary visual cues (discs) sharing the same 515 516 spatiotemporal properties (Holle et al., 2012). Beats are thought to translate speaker intentions, extending body posture accompanying speaker's prosody to 517 emphasize relevant segments of speech, which are available for listeners 518

519 during speech perception (So et al., 2012; Casasanto & Jasmin, 2009). If this is 520 the case, and beats play a linguistically relevant role above and beyond mere emphasis acting at low-level stages of stimulus processing, then the effect of 521 522 synchrony for beats should be different as compared to visual discs, in the relevant brain areas. Indeed, this guestion was answered with the interaction 523 524 term in our analysis, that indicates that the temporal synchrony of beats with 525 speech prosody has a differential impact on **BOLD** responses, as compared to other kinds of visual information (here, discs that replaced the speaker's hands). 526 527 The tendencies in the pattern of the interaction simple contrasts suggest greater 528 activations when beats and speech were presented in synchrony as compared 529 to asynchrony. Instead, the opposite pattern was observed when discs 530 accompanied speech. Based on this significant interaction pattern, we interpret 531 that, in addition to their emphasizing trajectory, beats also convey communicative aspects that simple discs are arguably lacking. 532

533

534 One surprising finding of our study is that the effect of synchrony for beats (i.e., 535 greater activity for synchronous as compared to asynchronous beats in left IFG and MTG) was not simply absent for the moving discs, but actually tended to be 536 reversed. When interpreting this cross-over interaction, it is also useful to take 537 into account whether the neural response in these areas represents an 538 activation or deactivation, relative to the implicit fixation cross baseline (see 539 parameter estimates in Fig. 2). Relative to this fixation cross baseline, only 540 speech accompanied by synchronous beats elicited activation in IFG, aMTG 541 542 and pMTG. This is consistent with the idea that IFG and posterior temporal lobe 543 are crucially involved in comprehending co-speech gestures (Holle et al., 2008, 544 2010, Willems et al., 2007, 2009). In contrast, a visual emphasis cue presented in asynchrony with speech (regardless of whether emphasis consisted of beats 545 546 or moving discs) did not activate these areas, which may reflect that temporally incongruent AV stimuli are less likely to be integrated and may even cause 547 suppression in multisensory areas (Noesselt et al., 2007). Interestingly, 548 processing speech accompanied by temporally congruent discs elicited a 549 reduction of activity in IFG, aMTG and pMTG, relative to fixation baseline. Such 550 a deactivation could possibly reflect a phasic inhibitory influence onto IFG, 551 552 aMTG and pMTG whenever speech is accompanied by temporally congruous 553 but unfamiliar visual emphasis cues, such as moving discs. An influence of 554 stimulus familiarity on AV integration in the temporal lobe has been 555 demonstrated before (Hein et al., 2007) and may extend to unfamiliar speech-556 accompanying visual emphasis cues, such as moving discs.

557

558 Our results are in line with previous fMRI studies that investigated neural correlates of iconic gestures (Holle et al., 2010; Holle et al., 2008; Willems et al., 559 560 2009; Willems et al., 2007). Particularly, one previous fMRI addressed natural hand beats co-occurring with continuous speech (Hubbard et al., 2009) and 561 reported a greater engagement of the STS compared to speech alone, an area 562 comparable to the one found in the present study. The authors also reported 563 564 greater BOLD activation in the left STS/G when speech was presented with the corresponding beat as compared to when presented with unrelated hand 565 movements. Please note that this comparison does not allow one to infer 566 whether the difference in left STS activation was produced by the lack of 567 568 synchrony between control gestures and speech, the lack of communicative value of control gestures, or an unknown combination of the two. When 569 570 Hubbard et al. compared speech accompanying beats to beats presented 571 without speech, no difference was observed, suggesting that the modulations in 572 the left STS/G reflect not only processing of biological movement but also integration of speech with the synchronized beat gestures. Indeed, the STS is 573 574 sensitive to various types of cross-modal correspondence including AV speech (sound-lip correspondence) in various previous studies (Nath and Beauchamp, 575 576 2012; Calvert et al., 2000; Callan et al., 2004; Macaluso et al., 2004; Meyer et 577 al., 2004).

578

579 In the present study, the interaction contrast suggests that BOLD response in 580 the left MTG was greater when speech was accompanied by beats as compared to discs (regardless of whether they were synchronized or not with 581 speech). At first glance, the greater response to stimuli containing beats in 582 occipital areas compared to those with discs may reflect a pure bottom-up effect 583 of richness of visual information (Figure 3). However, the interaction (Figure 2) 584 585 revealed also that the significant difference of BOLD activity in the visual areas 586 between beat and disc were dramatically reduced under asynchronous

presentations. This suggests that mere physical differences between beats and 587 588 discs conditions were not sufficient to explain their respective impact of synchrony in the indentified areas. The difference between beats and discs 589 590 might bring about more profound consequences. For example, in a previous ERP study, Holle et al. (2012) showed that a beat modulated the P600 591 592 component reflecting syntactic parsing, whereas a disc following the equivalent 593 trajectory did not. The authors suggested that the lack of communicative intention may explain the failure of simple discs to affect the neural correlates of 594 595 syntactic parsing. Here, the significant simple contrast Bs>Ds supports this 596 claim as it revealed greater activations not only in the occipital areas (although 597 certainly due to differences of visual information, the results are only 598 orientative), but also in the left MTG and left IFG areas. Indirectly, this result 599 also converges toward the idea a differential response to synchrony for using 600 discs that are not functionally associated with speech as part of a common 601 language system.

602

603 According to the effect of interaction on the neural activations, it seems that the 604 MTG responded to some additional language-related aspects associated with beat gestures during speech perception. Previous behavioral studies suggested 605 that some implicit pragmatic and intentional information from the speaker could 606 607 be extracted from beats, and influence speech encoding. For example, So et al, 608 (2012) showed that adult observers managed to remember more words from a spoken list when the words had previously been accompanied by a beat 609 610 gesture. As this memory improvement was not found in children, the authors 611 concluded that beat gestures conveyed communicative information but the 612 effect was functionally dependent on experiencing social interactions during development (McNeill, 1992). For example, listeners learn to interpret the 613 614 speaker's intention to underline relevant information with a beat through social experience. This association of communicative aspects between beats and 615 pitch accentuations was highlighted by Krahmer and Swerts (2007) who 616 showed that listeners perceived words as more salient when accompanied with 617 618 a beat gesture compared to same words presented in isolation. What is often 619 missing in these studies is whether the value of gestures and their integration of 620 speech simply depended on the general salience of the stimulus, or whether co-

speech gestures engaged a more specialized system. Although the listeners in 621 622 the present study could associate moving discs with movements of the hands and participants were able to detect an asynchrony between discs and speech, 623 synchronized gestures and synchronized discs elicited gualitatively distinct 624 patterns of brain activation (see contrast Bs>Ds). This suggests that during 625 626 perception listeners distinguished visual information functional related to some aspect of speech (beats) from arbitrary visual cues (discs). Here, this 627 information may require additional processes reflected by the differences of 628 629 activations in the MTG between beats and discs conditions.

630 In addition to the above explanation, the possible linguistic aspects engaged 631 when beats are present may be directly related to human movement 632 understanding and body postures, over and above to their interaction with 633 speech. The STS was found to respond to point-light representations of biological movements (Grossman et al., 2004; Pelphrey et al., 2004), actions 634 635 executed by humans (Thioux et al., 2008) and social visual cues (for reviews, see Nummenmaa & Calder, 2009; Allison, Puce & McCarthy, 2000). Herrington 636 637 et al, (2009) showed that the posterior STS was significantly more activated for trials in which participants perceived human point-light representations of 638 actions compared to non-human movements. In the present study, the discs did 639 640 not clearly represent a human form but clearly mimicked the trajectories described by hands during speech. In reference to the present study, listeners 641 could have associated discs trajectories with hands (as they identified in the 642 643 post-task guestionnaire). Yet, whatever aspect of biological motion engaged by 644 left MTG activations in the disc conditions, it was more strongly expressed 645 during beat conditions. Please note, however, that this possible perceptual 646 difference between beat gestures and discs in biological motion cannot explain 647 the whole pattern of results we found in the left MTG, because the interaction 648 term [(Bs – Ba) – (Ds – Da)] effectively controls for the different amounts of biological movement in the beat and disc conditions. 649

650

The present results also revealed an interaction between synchrony and visual information effects in the left IFG. Several fMRI studies have showed that the left IFG is sensitive to the semantic relationship between gesture and corresponding speech (Skipper et al., 2007; Willems et al., 2007; Willems et al.,

2009; Dick et al., 2009) and may be engaged in the unification of visual 655 656 (gestures) and audio (speech) complementary streams to facilitate comprehension (Willems et al., 2007; Hagoort, 2005). Recently, a meta-analysis 657 658 investigating the neural correlates shared between different types of gestures reported a common engagement of the left IFG during the perception of speech 659 660 accompanied with gestures as compared to a still body (Marstaller & Burianova, 2014). However, beat gestures do not convey semantic content, therefore the 661 IFG responses observed in the present study cannot be explained in terms of 662 663 semantic integration. Beyond meaning integration, the left IFG was also shown 664 to be involved in the process of syntactic analysis during sentence 665 comprehension (Glaser et al., 2013; Meyer et al., 2012; Obleser et al., 2011; 666 Uchiyama et al., 2008). As beats play a role in syntactic parsing (Holle et al., 667 2012), our results might correspond to an engagement of this area in the integration of beat information toward the parsing of the spoken stream, as 668 669 compared to moving discs. When beats were delayed (Ba condition), their apexes felt out from synchrony with pitch accents and likely out of the time 670 671 window of gesture-speech integration, potentially affecting the AV speech processing load (Habets et al., 2011; Obermeier et al., 2011; Obermeier & 672 Gunter, 2014). 673

674

675 It is worth noting that the simple main effect of synchrony for beat stimuli 676 (contrast Bs vs Ba) in left MTG, IFG and occipital cortex did not reach 677 significance in the whole brain analysis, but it is only revealed by the patterns of activations in the interaction contrasts following up on the interaction. Yet, the 678 post-hoc results obtained for the simple main effects restricted to the interaction 679 680 areas have to be often interpreted with caution (see Supplementary Materials). 681 In consequence, the interpretation of synchrony effects for beat gestures must 682 be linked to its effects relative to the disc condition. In other words, the disc synchrony manipulation can be seen as a baseline for the beat-synchrony 683 684 manipulation. However, this is indeed a theoretically relevant type of comparison as discussed Holle et al. (2012). In addition, if we go by the results 685 of previous studies, and extant knowledge the neural correlates of speech, we 686 687 feel safe in interpreting this pattern in line with the results of the interaction that 688 suggested a difference between synchronous and asynchronous beat

conditions (see Figure 2). Note, for example that a similar effect of AV 689 690 synchrony involving gestures in the left STG/S was reported in Hubbard et al. 691 (2009). In their study, however, as mentioned earlier, Hubbard et al. used unrelated sign language movements as a control condition, which not only 692 constitute a more dramatic asynchrony manipulation altogether (as speech and 693 694 gestures had completely different rhythms), but also changed the very nature of the visual stimuli from the synchronous to the asynchronous condition. Here, we 695 have looked at these two effects (confounded in Hubbard) separately, and 696 697 therefore it is not surprising that their individual neural correlates are more subtle. That is, in the present study, although delayed with respect to speech, 698 699 the rhythm of beats was maintained and might still be associable with the global 700 speech envelope. This may have diminished the detrimental impact of 701 desynchronized gestures on a listener's perception. This may also explain why 702 we did not observe any effect of synchrony in the right auditory cortex related to 703 auditory processing and prosody, as it was reported in Hubbard et al.'s results. A further relevant aspect in our study is that participants were asked to simply 704 705 focus on an auditory detection task. This is interesting because our results 706 cannot be attributed to an explicit monitoring of speech-gesture synchrony. On 707 the contrary, our auditory detection task may have decreased attention on visual information and effectively weakened the expression of beat synchrony 708 709 on speech processing networks.

710

Taken together, the present results provide new insights about the specificity of 711 left MTG and IFG in the processing of multimodal language (for a review, see 712 Campbell, 2008; Özürek, 2014). As participants were not explicitly asked to pay 713 714 attention to the speaker's hands, this suggests that the temporal correspondence between beats and speech prosody may be picked up 715 716 automatically. This is in line with previous proposals considering speech and gestures as two side of a same underlying language system (McNeill, 1992; 717 Kelly, Creigh and Bartolotti, 2009). Beats appear to convey additional 718 communicative value such as speakers' intentions, which are not available (or 719 at least, not extracted) from simple visual stimuli (Holle et al., 2012; So et al., 720 2012; Casasanto & Jasmin, 2009; McNeill, 1992). The access to concurrent 721 722 gestures during speech perception may engage the listeners and provide a better alignment between listener and speaker, improving speech processing
and information encoding. Finally, the fact that the speaker was a well-known
former Spanish president may have engaged some political sensitivity from
listeners. However, such a possible bias is unlikely to influence our results,
since participants viewed the same speaker across all four experimental
conditions.

729

730 5. CONCLUSION

731

We investigated the neural correlates of spontaneous beat gestures produced in continuous speech. Our results revealed that the synchrony affected <u>brain's</u> activations differently according to the visual information accompanying speech during perception. We concluded that beats <u>are linguistic</u> <u>information</u> by their trajectories aligned with speech prosody, but also communicative intentions of the speaker.

738

739 **AKNOWLEDGMENTS**

740

This research was supported by the Ministerio de Economia y Competitividad
(PSI2013-42626-P), AGAUR Generalitat de Catalunya (2014SGR856), and the
European Research Council (StG-2010 263145).

744

745 **REFERENCES**

746

- Allison, T., Puce, A., & McCarthy, G. (2000). Social perception from visual cues: role of the STS
 region. *Trends in Cognitive Sciences*, 4(7), 267–278.
- Biau, E., & Soto-Faraco, S. (2013). Beat gestures modulate auditory integration in speech
 perception. *Brain and Language*, 124(2), 143–52.
- Biau, E., Torralba , M., Fuentemilla, L., de Diego Balaguer, R., & Soto-Faraco, S. (2015).
 Speaker's hand gestures modulate speech perception through phase resetting of ongoing
 neural oscillations. *Cortex*, 68, 76-85.
- Brett, M., Anton, J-L., Valabregue, R., & Poline, J-B. Region of interest analysis using an SPM
 toolbox [abstract] Presented at the 8th International Conference on Functional Mapping of
 the Human Brain, June 2-6, 2002, Sendai, Japan. Available on CD-ROM in NeuroImage,
 Vol 16, No 2.

- Callan, D. E., Jones, J. A., Callan, A. M., & Akahane-Yamada, R. (2004). Phonetic perceptual
 identification by native- and second-language speakers differentially activates brain
 regions involved with acoustic phonetic processing and those involved with articulatoryauditory/orosensory internal models. *NeuroImage*, 22(3), 1182–94.
- Calvert, G. A., Campbell, R., & Brammer, M. J. (2000). Evidence from functional magnetic
 resonance imaging of crossmodal binding in the human heteromodal cortex. *Current Biology: CB*, 10(11), 649–57.
- Campbell, R. (2008). The processing of audio-visual speech: empirical and neural bases. *Philosophical Transactions of the Royal Society of London. Series B, Biological Sciences*,
 363(1493), 1001–10.
- Casasanto, D., & Jasmin, K. (2010). Good and bad in the hands of politicians: spontaneous
 gestures during positive and negative speech. *PloS One*, 5(7), e11805.
- Dick, A. S., Mok, E. H., Raja Beharelle, A., Goldin-Meadow, S., & Small, S. L. (2014). Frontal
 and temporal contributions to understanding the iconic co-speech gestures that
 accompany speech. *Human Brain Mapping*, *35*(3), 900–17.
- Dick, A. S., Goldin-Meadow, S., Hasson, U., Skipper, J. I., & Small, S. L. (2009). Co-speech
 gestures influence neural activity in brain regions associated with processing semantic
 information. *Human Brain Mapping*, 30(11), 3509–26.
- Friston, K. J., Glaser, D. E., Henson, R. N. A., Kiebel, S., Phillips, C., & Ashburner, J. (2002).
 Classical and Bayesian inference in neuroimaging: applications. *NeuroImage*, *16*(2), 484–
 <u>512.</u>
- Glaser, Y. G., Martin, R. C., Van Dyke, J. A., Hamilton, A. C., & Tan, Y. (2013). Neural basis of
 semantic and syntactic interference in sentence comprehension. *Brain and Language*, *126*(3), 314–26.
- Grossman, E. D., Blake, R., & Kim, C.-Y. (2004). Learning to see biological motion: brain activity
 parallels behavior. *Journal of Cognitive Neuroscience*, 16(9), 1669–79.
- Guellaï, B., Langus, A., & Nespor, M. (2014). Prosody in the hands of the speaker. *Frontiers in Psychology*, *5*, 700.
- Habets, B., Kita, S., Shao, Z., Ozyurek, A., & Hagoort, P. (2011). The role of synchrony and
 ambiguity in speech-gesture integration during comprehension. *Journal of Cognitive Neuroscience*, *23*(8), 1845–54.
- Hagoort, P. (2005). On Broca, brain, and binding: a new framework. *Trends in Cognitive Sciences*, 9(9), 416–23.
- Hein, G., Doehrmann, O., Müller, N. G., Kaiser, J., Muckli, L., & Naumer, M. J. (2007). Object
 familiarity and semantic congruency modulate responses in cortical audiovisual integration
 areas. *The Journal of Neuroscience: The Official Journal of the Society for Neuroscience*,
 27(30), 7881–7.
- Herrington, J. D., Nymberg, C., & Schultz, R. T. (2011). Biological motion task performance
 predicts superior temporal sulcus activity. *Brain and Cognition*, 77(3), 372–81.

- Holle, H., & Gunter, T. C. (2007). The role of iconic gestures in speech disambiguation: ERP
 evidence. *Journal of Cognitive Neuroscience*, 19(7), 1175–92.
- Holle, H., Gunter, T. C., Ruschemeyer, S. A., Hennenlotter, A., & Iacoboni, M. (2008). Neural
 correlates of the processing of co-speech gestures. *Neuroimage*, *39*(4), 2010-2024.
- Holle, H., Obermeier, C., Schmidt-Kassow, M., Friederici, A. D., Ward, J., & Gunter, T. C.
 (2012). Gesture facilitates the syntactic analysis of speech. *Frontiers in Psychology*, 3, 74.
- Holle, H., Obleser, J., Rueschemeyer, S.-A., & Gunter, T. C. (2010). Integration of iconic
 gestures and speech in left superior temporal areas boosts speech comprehension under
 adverse listening conditions. *NeuroImage*, 49(1), 875–84.
- Hubbard, A. L., Wilson, S. M., Callan, D. E., & Dapretto, M. (2009). Giving speech a hand:
 gesture modulates activity in auditory cortex during speech perception. *Human Brain Mapping*, 30(3), 1028–37.
- Kelly, S. D., Kravitz, C., & Hopkins, M. (2004). Neural correlates of bimodal speech and gesture
 comprehension. *Brain and Language*, 89(1), 253–60.
- Kelly, S. D., Ozyürek, A., & Maris, E. (2010). Two sides of the same coin: speech and gesture
 mutually interact to enhance comprehension. *Psychological Science*, 21(2), 260–7.
- Kelly, S. D., Ward, S., Creigh, P., & Bartolotti, J. (2007). An intentional stance modulates the
 integration of gesture and speech during comprehension. *Brain and Language*, 101(3),
 222–33.
- Krahmer, E., & Swerts, M. (2007). The effects of visual beats on prosodic prominence: Acoustic
 analyses, auditory perception and visual perception. *Journal of Memory and Language*,
 57(3), 396–414.
- Leonard, T., & Cummins, F. (2011). The temporal relation between beat gestures and speech.
 Language and Cognitive Processes, 26(10), 1457–1471.
- Macaluso, E., George, N., Dolan, R., Spence, C., & Driver, J. (2004). Spatial and temporal
 factors during processing of audiovisual speech: a PET study. *NeuroImage*, 21(2), 725–
 32.
- Marstaller, L., & Burianová, H. (2014). The multisensory perception of co-speech gestures A
 review and meta-analysis of neuroimaging studies. *Journal of Neurolinguistics*, 30, 69–77.
- Meyer, M., Steinhauer, K., Alter, K., Friederici, A. D., & von Cramon, D. Y. (2004). Brain activity
 varies with modulation of dynamic pitch variance in sentence melody. *Brain and Language*, 89(2), 277–89.
- Noesselt, T., Rieger, J. W., Schoenfeld, M. A., Kanowski, M., Hinrichs, H., Heinze, H.-J., &
 Driver, J. (2007). Audiovisual temporal correspondence modulates human multisensory
 superior temporal sulcus plus primary sensory cortices. *The Journal of Neuroscience: The*Official Journal of the Society for Neuroscience, 27(42), 11431–41.
- Nummenmaa, L., & Calder, A. J. (2009). Neural mechanisms of social attention. *Trends in Cognitive Sciences*, 13(3), 135–43.

- Obermeier, C., Holle, H., & Gunter, T. C. (2011). What iconic gesture fragments reveal about
 gesture-speech integration: when synchrony is lost, memory can help. *Journal of Cognitive Neuroscience*, *23*(7), 1648–63.
- Obermeier, C., & Gunter, T. C. (2014). Multisensory Integration: The Case of a Time Window of
 Gesture-Speech Integration. *Journal of Cognitive Neuroscience*, 1–16.
- Obleser, J., Meyer, L., & Friederici, A. D. (2011). Dynamic assignment of neural resources in
 auditory comprehension of complex sentences. *NeuroImage*, *56*(4), 2310–20.
- Pelphrey, K. A., Morris, J. P., & McCarthy, G. (2004). Grasping the intentions of others: the
 perceived intentionality of an action influences activity in the superior temporal sulcus
 during social perception. *Journal of Cognitive Neuroscience*, 16(10), 1706–16.
- 845
- Skipper, J. I., Goldin-Meadow, S., Nusbaum, H. C., & Small, S. L. (2007). Speech-associated
 gestures, Broca's area, and the human mirror system. *Brain and Language*, 101(3), 260–
 77.
- So, W. C., Sim Chen-Hui, C., & Low Wei-Shan, J. (2012). Mnemonic effect of iconic gesture and
 beat gesture in adults and children: Is meaning in gesture important for memory recall? *Language and Cognitive Processes*, 27(5), 665–681.
- Thioux, M., Gazzola, V., & Keysers, C. (2008). Action understanding: how, what and why.
 Current Biology : CB, 18(10), R431–4.
- Treffner, P., Peter, M., & Kleidon, M. (2008). Gestures and Phases: The Dynamics of SpeechHand Communication. *Ecological Psychology*, *20*(1), 32–64.
- Uchiyama, Y., Toyoda, H., Honda, M., Yoshida, H., Kochiyama, T., Ebe, K., & Sadato, N.
 (2008). Functional segregation of the inferior frontal gyrus for syntactic processes: a
 functional magnetic-resonance imaging study. *Neuroscience Research*, *61*(3), 309–18.
- Willems, R. M., Ozyürek, A., & Hagoort, P. (2007). When language meets action: the neural
 integration of gesture and speech. *Cerebral Cortex (New York, N.Y.: 1991)*, 17(10), 2322–
 33.
- Willems, R. M., Ozyürek, A., & Hagoort, P. (2009). Differential roles for left inferior frontal and
 superior temporal cortex in multimodal integration of action and language. *NeuroImage*,
 47(4), 1992–2004.
- Wu, Y. C., & Coulson, S. (2010). Gestures modulate speech processing early in utterances. *Neuroreport*, 21(7), 522–6.