NeuroImage xxx (2014) xxx-xxx



Contents lists available at ScienceDirect

### NeuroImage



journal homepage: www.elsevier.com/locate/ynimg

# Supplementary motor area activations predict individual differences in temporal-change sensitivity and its illusory distortions

### Q1 Björn Herrmann \*, Molly J. Henry, Mathias Scharinger, Jonas Obleser

4 Max Planck Research Group "Auditory Cognition" at the Max Planck Institute for Human Cognitive and Brain Sciences, Stephanstraße 1A, 04103 Leipzig, Germany

#### 5 ARTICLE INFO

Article history:
 Accepted 16 July 2014
 Available online xxxx

9 Kevwords:

10 Temporal rate change

11 Time-pitch interdependence

12 Functional magnetic resonance imaging

13 Feature-selective attention

14 Perceptual illusion

#### ABSTRACT

Perception of time and temporal change is critical for human cognition. Yet, perception of temporal change is susceptible to contextual influences such as changes of a sound's pitch. Using functional magnetic resonance imag- 16 ing (fMRI), the current study aimed to investigate perception of temporal rate change and pitch-induced illusory 17 distortions. In a  $6 \times 6$  design, human participants (N = 19) listened to frequency-modulated sounds (~4 Hz) that 18 varied over time in both modulation rate and pitch. Participants judged the direction of rate change ('speeding 19 up' vs. 'slowing down'), while ignoring changes in pitch. Behaviorally, rate judgments were strongly biased by 20 pitch changes: Participants perceived rate to slow down when pitch decreased and to speed up when pitch in- 21 creased ('rate-change illusion'). The fMRI data revealed activation increases with increasing task difficulty in 22 pre-SMA, left putamen, and right IFG/insula. Importantly, activation in pre-SMA was linked to the perceptual sen- 23 sitivity to discriminate rate changes and, together with the left putamen, to relative reductions in susceptibility to 24 pitch-induced illusory distortions. Right IFG/insula activations, however, only scaled with task difficulty. These 25 data offer a distinction between regions whose activations scale with perceptual sensitivity to features of time 26 (pre-SMA) and those that more generally support behaving in difficult listening conditions (IFG/insula). 27 Hence, the data underscore that individual differences in time perception can be related to different patterns 28 of neurofunctional activation. 29

© 2014 Elsevier Inc. All rights reserved.

#### **32** 33

30

#### 35 Introduction

Almost all environmental events inherently carry temporal informa-36 tion. The most prominent examples are speech and music, which con-37 tain variations in item duration and fluctuations in temporal rate. 38 Critically, perceived time does not necessarily precisely reflect physical 39 40 stimulus features (Eagleman, 2008) but can be influenced by contextual factors such as sound intensity (Alards-Tomalin et al., 2013), visual 41 space (Huang and Jones, 1982; Jones and Huang, 1982) or changes in 42a sound's pitch (Boltz, 1998, 2011; Henry and McAuley, 2009, 2013; 4344Herrmann et al., 2013; Shigeno, 1986). For example, participants tend to overestimate the rate of a modulated sound when the sound's pitch 45 increases and underestimate the rate of a sound when the sound's 46 47 pitch decreases (Herrmann et al., 2013).

In order to investigate the neural underpinnings of time perception,
functional magnetic resonance imaging (fMRI) studies have been conducted to reveal where in the brain temporal information is processed.
Most fMRI studies thus far investigated interval or duration perception
(e.g., Coull et al., 2004; Lewis and Miall, 2003; Pouthas et al., 2005;
Rao et al., 2001; Tregellas et al., 2006), while a few other studies
examined the perception of rate (e.g., Bengtsson et al., 2009; Grahn

\* Corresponding author. *E-mail address:* bjoern.herrmann@outlook.com (B. Herrmann).

http://dx.doi.org/10.1016/j.neuroimage.2014.07.026 1053-8119/© 2014 Elsevier Inc. All rights reserved. and McAuley, 2009; Grahn and Rowe, 2009; Henry et al., in press; 55 McAuley et al., 2012). The most prominent regions associated with pro-56 cessing temporal information are the supplementary motor area (SMA), 57 pre-SMA, insular cortex, inferior frontal gyrus (IFG), inferior parietal cortex, cerebellum, and basal ganglia including the caudate and puta-59 men (Bengtsson et al., 2009; Coull et al., 2004, 2008; Grahn and 60 McAuley, 2009; Harrington et al., 1998, 2010; Lewis and Miall, 2003; 61 McAuley et al., 2012; Morillon et al., 2009; Nenadic et al., 2003; Rao et al., 2001; Schwartze et al., 2012; Teki et al., 2011; Tipples et al., 63 2013; Wiener et al., 2014; for a recent meta-analysis, see Wiener et al., 64 2010).

Another line of studies focused on the influences of task perfor- 66 mance and attentional dynamics in modulating the brain activity in 67 those regions implicated in timing functions (Coull and Nobre, 1998; 68 Coull et al., 2004; Henry et al., in press; Tregellas et al., 2006). For 69 example, increasing the difficulty of discriminating the duration or 70 rate of auditory or visual stimuli leads to increased activation in the 71 pre-SMA, right inferior frontal cortex, basal ganglia, and inferior parietal 72 cortex (Ferrandez et al., 2003; Henry et al., in press; Livesey et al., 2007; 73 Tregellas et al., 2006; Wencil et al., 2010; Wiener et al., 2014). Further- 74 more, selective attention to a specific temporal feature (e.g., duration) 75 while ignoring a different temporal feature (e.g., modulation rate) 76 leads to a reversal in the pattern of brain activity as a function of task 77 difficulty (Henry et al., in press). In this study, for a to-be-attended 78

### **ARTICLE IN PRESS**

B. Herrmann et al. / NeuroImage xxx (2014) xxx-xxx

temporal feature, the largest neural response was observed for small 79 80 (i.e., difficult to discriminate) stimulus changes, and the smallest neural response occurred for large (i.e., easy to discriminate) stimulus changes. 81 82 Critically, this differential effect was inverted for the to-be-ignored temporal feature, for which the largest neural response was observed when 83 stimulus change was large (i.e., maximally distracting), while the 84 smallest neural response was observed when stimulus change was 85 86 small (i.e., easy to ignore; Henry et al., in press). Taking these studies 87 together, it has been concluded that brain activation in regions asso-88 ciated with timing functions is also strongly affected by task perfor-89 mance and attentional dynamics in a timing task (Coull et al., 2004; Ferrandez et al., 2003; Henry et al., in press; Livesey et al., 2007; 90 Tregellas et al., 2006). 91

92Critically, individuals vary in their ability to discriminate, for example, small changes in duration or rate (Erb et al., 2012; Fitzgerald 93 and Wright, 2011; Grahn and McAuley, 2009; Moore et al., 1991), 94 and in their susceptibility to illusory distortions of duration or rate 95 96 (Dirnberger et al., 2012; Harrington et al., 2004; Herrmann et al., 2013; Tipples et al., 2013). Furthermore, individuals differ in the degree 97 to which brain activation in regions implicated in timing functions is 98 modulated by performance in a timing task (Coull et al., 2008; Wiener 99 et al., 2014), induction of a beat (Grahn and McAuley, 2009; Grahn 100 101 and Rowe, 2009), emotion-induced time distortions (Dirnberger et al., 2012), and general misestimation of time (Harrington et al., 2004; 102 103 Tipples et al., 2013).

Regarding distortions in perceived time, there have been a num-104 ber of previous attempts to relate brain activation to illusory time 105106 percepts (using within-participant and across-participant correlations). Several studies have observed a relation between brain activations and 107distortions of perceived time originating from global context effects 108(Harrington et al., 2004; Tipples et al., 2013), while others observed a re-109110 lation between brain activity and illusory percepts originating from ma-111 nipulations of nontemporal stimulus features such as visual motion, 112 visual looming, audio-visual interaction, or emotional content (Bueti and Macaluso, 2011; Dirnberger et al., 2012; Harrington et al., 2011; 113 Wittmann et al., 2010). With respect to the auditory domain, Bueti and 114 Macaluso (2011) were unable to observe a link between brain activa-115 116 tions and temporal illusions, although they observed such a link for the visual domain; this null effect in the auditory domain might be due to 117 the weak perceptual distortions elicited by the non-modulated tone 118 stimuli used in this study (as discussed also in Bueti and Macaluso, 119 120 2011). Pitch changes in sounds, on the other hand, are known to strongly influence percepts of time (Boltz, 1998, 2011; Henry and 121 McAuley, 2009, 2013; Herrmann et al., 2013; Shigeno, 1986), and 122123 are therefore well suitable for investigating the underlying neural functional activations. 124

125Thus, the current fMRI study focused on time distortions induced by auditory stimulation and followed previous research by focusing on 126modulations of brain activity by attentional dynamics and task perfor-127mance in brain regions associated with timing functions. In detail, 128we used frequency-modulated sounds that changed over time in mod-129130ulation rate (speeding up vs. slowing down) and pitch (decrease vs. in-131crease). We aimed to investigate (1) which brain regions are modulated by difficulty in a discrimination task when attending to modulation-rate 132changes while simultaneously ignoring changes in pitch; and (2) whether 133the degree of modulation in brain activity is linked to individual differ-134135ences in perceptual sensitivity (to modulation rate) and the magnitude of pitch-induced illusory distortions. 136

#### 137 Methods and materials

#### 138 Participants

Nineteen healthy adults aged 21–32 years (median: 24 years; 10
 female) participated in the current study. Participants were right handed and had no self-reported hearing problems or history of

neurological diseases. They gave written informed consent and were 142 paid 8 Euro per hour. The study was in accordance with the Declaration 143 of Helsinki and approved by the local ethics committee of the University 144 of Leipzig. 145

#### Acoustic stimulation

Stimuli consisted of frequency-modulated sounds with a duration of 147 4 s, in which modulation rate and pitch were manipulated in a  $6 \times 6$  148 design (see Fig. 1A). Stimuli were created in MATLAB (v7.11; The 149 MathWorks Inc.) using an adapted version of the *vco.m* function, and 150 were sampled at 44.1 kHz with 24-bit resolution. In brief, the *vco.m* 151 function generates a sine wave with time-sensitive frequency modulations by manipulating the phase angles of the waveform. Levels for 153 rate and pitch changes were chosen based on pilot testing, which 154 ensured that rate-pitch combinations would elicit robust illusory ratechange percepts (see also Herrmann et al., 2013).

For the factor modulation rate, the carrier frequency of the sounds 157 was modulated by a sinusoidal function, where the modulation rate 158 linearly changed from 4 Hz to one of six levels (4 Hz  $\pm 2.6\%, \pm 7.8\%, 159$  and  $\pm 13\%$ ) over the duration of the sound (Fig. 1A). The modulation 160 depth was fixed at  $\pm 20\%$  of the mean carrier frequency, which took 161 on one of six values ranging from 1200 Hz to 1500 Hz. Note that we 162 selected a modulation rate of 4 Hz (i.e., a period of 250 ms) which, 163 together with unpredictable starting phases of the modulation, deems 164 using any counting strategy in order to perform the task unlikely.

For the factor Pitch, the carrier frequency itself was manipulated 166 such that it linearly changed around the mean carrier frequency by 167  $\pm$  8%,  $\pm$  24%, and  $\pm$  40% over the duration of the sound. For carrier frequency manipulations, modulation depth was scaled according to the 169 instantaneous mean carrier frequency and thus remained consistent 170 with the logarithmic frequency scale critical for auditory perception 171 (Attneave and Olson, 1971; Burns, 1999; Fig. 1A). 172

Note that the stimulus-final phase (and consequently the starting 173 phase) of the frequency modulation was manipulated such that half of 174 the trials ended in the rising phase of the frequency modulation and 175 the other half in the falling phase (uniformly distributed across condi-176 tions). Hence, any perceptual differences potentially arising from differ-177 ent final trajectories of the sounds' frequency were controlled for across 178 trials.

#### Procedure

Sounds were presented at 55 dB sensation level (i.e., above the 181 participants' individual hearing threshold), which was determined 182 for a 1350-Hz pure tone at the beginning of the experiment. Then, 183 participants underwent a short familiarization session prior to scanning (including extreme examples of modulation-rate changes without changes in pitch) in order to familiarize them with the task. 186

A schematic outline of a trial is shown in Fig. 1B. Each trial in the 187 experiment comprised the presentation of a sound (jittered randomly 188 between 1.45 and 1.95 s following trial onset; Fig. 1B), followed by a 189 visual response prompt showing the letters "S" and "L" next to each 190 other (S – "schneller" and L – "langsamer", German for "faster" and 191 "slower", respectively). The response prompt always occurred along 192 with the onset of the 5th fMRI volume (TR) within a trial at 6.4 s. Partic- 193 ipants were asked to press the button for "S" whenever the modulation 194 rate increased and the button for "L" whenever the modulation rate 195 decreased. Participants were instructed to ignore changes in pitch. The 196 positions (left vs. right on the screen) of the letters "S" and "L" randomly 197 changed from trial to trial (uniformly distributed across conditions), and 198 the participant was given 2.3 s to indicate his/her response using the 199 index finger (left screen position) and ring finger (right screen position) 200 of the right hand. Subsequently, an additional visual response prompt 201 was presented (2.3 s after the first prompt) which asked participants 202 to judge on a three-point scale how confident they were with their 203

Please cite this article as: Herrmann, B., et al., Supplementary motor area activations predict individual differences in temporal-change sensitivity and its illusory distortions, NeuroImage (2014), http://dx.doi.org/10.1016/j.neuroimage.2014.07.026

180

B. Herrmann et al. / NeuroImage xxx (2014) xxx-xxx



**Fig. 1.** Stimulus design. A) Schematic of the 6 × 6 rate–pitch stimulus design (only four stimuli examples are shown). Stimuli were frequency-modulated sounds varying in modulation rate and pitch over the duration of the sound. Here, the frequency modulation of example sounds is depicted, with the sounds' frequency displayed on the y-axis and time on the x-axis. Note that changes in modulation rate are schematic and were smaller in the actual stimuli. B) Trial structure of the current fMRI experiment, including sound presentation, modulation rate judgment (S – "schneller" and L – "langsamer", German for "faster" and "slower", respectively), and confidence ratings. The position of "S" and "L" on the screen (left versus right) varied from trial to trial, and thus prohibited specific motor preparation during sound presentation.

decision in the modulation-rate judgment (Fig. 1B; see also Wilsch et al.,in press).

The experiment was divided into six blocks. Within each block, each of the  $6 \times 6$  unique modulation rate  $\times$  pitch combinations was presented once, resulting in 36 stimulus presentations per block. In addition, four silent trials were presented per block, where no acoustic stimulation occurred.

Auditory stimuli were presented via MRI-compatible headphones (Commander XG, Resonance Technology, Inc.). Visual response prompts were projected through an LCD projector onto a mirror screen attached to the head coil. An IBM-compatible computer running with the presentation software (Neurobehavioral Systems, Albany, CA) controlled the stimulation.

#### 217 Behavioral data analysis

Behavioral data were analyzed in two ways. First, general task diffi-218culty was investigated. In detail, each trial's response was given a binary 219code (correct; incorrect) and then weighted by the corresponding 220 confidence rating, such that the single-trial response coding could take 221on one of six values [0 0.2 0.4 0.6 0.8 1]. For example, a response code 222of 0.6 refers to a correct response for which the participant was 223unconfident (confidence rating of 1). Single-trial responses were 224averaged over trials. We refer to the resulting dependent measure as 225hit rate. In order to test whether rate manipulations affected listeners' 226rate-change discrimination performance, hit rates were averaged across 227pitch conditions (rate marginal) and a one-way repeated measures 228 229analysis of variance (rmANOVA) was calculated to compare rate conditions. In order to test whether pitch manipulations affected lis-230teners' rate-change discrimination performance, hit rates were averaged 231across rate conditions (pitch marginal) and a one-way rmANOVA was 232 calculated to compare pitch conditions. Note that here the rate  $\times$  pitch 233234interaction was not of interest because pitch-induced illusory influences 235on rate-change discrimination were investigated using proportion of 'speeding up' responses (see below). Whenever the assumption of 236237sphericity was violated (according to a significant Mauchly's test at  $P \le 0.05$ ), Greenhouse–Geisser correction was applied (Greenhouse 238239and Geisser, 1959).

For the second analysis, each trial's response was again given a 240binary code (slowing down; speeding up) and then weighted by the 241 corresponding confidence rating, such that the single-trial response 242 coding could take on one of six values [0 0.2 0.4 0.6 0.8 1]. For example, 243a response code of 0.6 refers to a 'speeding up' response for which the 244 participant was unconfident (confidence rating of 1). Single-trial re-245sponses were averaged over trials. We refer to the resulting dependent 246measure as proportion of 'speeding up' responses. From the proportion 247248of 'speeding up' responses two measures were extracted. First, perceptual sensitivity to discrimination rate changes was estimated 249 as the slope of the linear fit to weighted proportions of 'speeding 250 up' responses as a function of z-normalized rate levels (average 251 across pitch conditions). We henceforth refer to the slope measure 252 as 'discrimination index'. A larger (i.e., steeper) slope indicates better 253 rate-discrimination sensitivity (Macmillan and Creelman, 2005, 254 p. 121). Second, strength of illusory rate-change percepts was estimated 255 as the slope of the linear fit to the proportion of 'speeding up' responses 256 as a function of z-normalized pitch levels (average across rate condi-257 tions). We henceforth refer to this measure as 'illusion index'. A larger 258 slope indicates stronger illusory rate-change percepts. In order to test 259 whether participants were able to discriminate rate changes and 260 whether participants were biased in their rate-change percept, the dis- 261 crimination index and the illusion index were separately tested against 262 zero using one-sample t-tests. Subsequently, the two indices were test- 263 ed against each other using a paired-sample t-test. Importantly, the dis- 264 crimination and illusion indices were used as covariates in the fMRI 265 analyses (see below). 266

#### fMRI recordings and preprocessing

Data were recorded on a 3-T MedSpec 30/100 scanner (Bruker, 268 Ettlingen, Germany) using a birdcage head coil.  $T_2^*$ -weighted images 269 were acquired using an interleaved silent steady state (ISSS) sequence 270 (Mueller et al., 2011; Schwarzbauer et al., 2006), with a TR of 1.6 s, TE 271 of 30.36 ms, flip angle of 73° (Ernst angle), acquisition bandwidth of 272 101 kHz, matrix =  $64 \times 64$  pixels, and FOV = 19.2 cm<sup>2</sup>, resulting in 273 an in-plane resolution of  $3 \times 3$  mm. Slice thickness was 4 mm, with a 274 1-mm interslice gap. During auditory stimulation, the magnetization 275 was kept in a silent steady state (4 volumes) followed by the subse- 276 quent acquisition of 3 volumes during which the visual response 277 prompts and corresponding button presses were delivered (Fig. 1B). 278 Following the experimental blocks, a Fieldmap image was acquired 279 for use during realignment. Existing high-resolution T<sub>1</sub>-weighted 280 magnetization-prepared rapid gradient-echo images were taken from 281 the database of the Max Planck Institute for Human Cognitive and 282 Brain Sciences. T1 images were acquired on a 3-T MAGNETOM TIM 283 Trio scanner (Siemens, Erlangen, Germany) according to the following 284 parameters: TR = 1.3 s, TA = 10 ms, TE = 3.93 ms, matrix =  $256 \times 285$ 240, FOV =  $256 \times 240$ , resulting in a resolution of 1 mm  $\times$  1 mm  $\times$  286 1.5 mm (interpolated to 1-mm isotropic voxel size during spatial 287 normalization). 288

Data were analyzed using SPM8 (Welcome Trust Centre for Neu- 289 roimaging, London, UK) and custom MATLAB scripts. Preprocessing 290 comprised rigid-body spatial realignment and unwarping using the 291 Fieldmap image, segmentation of the  $T_1$  image, coregistration to 292 the  $T_1$  image according to spatial normalization parameters from 293

### **ARTICLE IN PRESS**

B. Herrmann et al. / NeuroImage xxx (2014) xxx-xxx

segmentation, normalization to Montreal Neurological Institute (MNI) space, interpolation to  $3 \times 3 \times 3$ -mm voxel size, and smoothing with an 8-mm FWHM isotropic Gaussian kernel. First-level analyses were modeled using a finite impulse response (FIR) function, and a high-pass filter of 0.1 Hz was applied to remove slow drifts in the data.

### fMRI analysis: overall brain activation to acoustic stimulation and task performance

In order to test for general brain activations elicited during acoustic 302 303 stimulation and task performance, one design matrix was constructed 304 for each participant and a general linear model (GLM) was calculated 305 (first-level analysis). The design matrix included one regressor per block indexing sound trials and a second regressor per block indexing 306 silent trials. Blocks were modeled as regressors of no interest. Effects 307 of overall brain activation were calculated as sound trials vs. silent trials 308 (contrast vector: [1–1]). On the second level, the resulting contrast 309 values were compared to zero using a one-sample t-test. t-Values 310 were transformed to z-scores, and activations were cluster-extent 311 threshold corrected for multiple comparisons ( $P \le 0.05$ ) based on 312 Monte Carlo simulations as described in Slotnick and Schacter (2004) 313 314 and Slotnick et al. (2003). Voxels with z-scores equal or greater than 3153.719 ( $P \le 0.0001$ ) and a cluster extent of nine voxels were considered statistically significant. Anatomical labels were determined using the 316 xjView toolbox (http://www.alivelearn.net/xjview) implemented in 317 SPM8. 318

#### 319 fMRI analysis: neural activation as a function of increasing task difficulty

Next, two GLMs were calculated for each participant (first-level analysis), one for the six rate conditions and one for the six pitch conditions. Each design matrix included six regressors per block (six rate or pitch conditions) and blocks were modeled as regressors of no interest. The resulting beta values for rate conditions and the beta values for pitch conditions were separately subjected to the analyses described in the following.

In order to investigate neural responses related to task performance 327 (i.e., increasing task difficulty), hit rates were correlated with neural ac-328 tivation (Fig. 4). In detail, for each participant and separately for each 329voxel, a linear function was fitted to the estimated coefficients (beta 330 331 values) from each GLM as a function of hit rate. In order to allow for a possible separation of task-difficulty increases due to rate versus pitch 332 333 changes, linear fits were separately calculated for the rate conditions 334 and for the pitch conditions (see Fig. 4B). On the second level, voxels that significantly scaled with increasing task difficulty (decreasing hit 335 336 rate) were determined by testing the slopes of the linear fits against zero using a one-sample t-test. t-Values were transformed to z-scores 337 and activations were cluster-extent threshold corrected for multiple 338 comparisons ( $P \le 0.05$ ; Slotnick et al., 2003; Slotnick and Schacter, 339 2004). Voxels with z-scores equal or greater than 2.576 ( $P \le 0.005$ ) 340 341 and a cluster extent of twenty-four voxels were considered statisti-342 cally significant. Anatomical labels were determined using the xjView toolbox (http://www.alivelearn.net/xjview) implemented 343 in SPM8. 344

Further correlations were carried out using the slopes of the above-345 346 described linear fits to investigate whether the modulation in neural activation as a function of increased task difficulty (decreasing hit rate) 347 has perceptual consequences related to either (a) sensitivity to discrim-348 inate changes in rate or (b) pitch-induced distortions of rate changes. 349 For each significant cluster, individual slopes from the linear fit to beta 350values as a function of hit rate (reflecting the degree of modulation in 351 brain activation with increasing task difficulty) were extracted from 352 the voxel yielding the highest z-score within the cluster. Extracted 353 slopes were then correlated with the discrimination index and the illu-354 355 sion index estimated from the behavioral data.

#### Results

#### Behavioral results

Fig. 2A depicts hit rates (weighted proportion correct) for rate 358 conditions and pitch conditions. One-way rmANOVAs conducted 359 separately for rate conditions and for pitch conditions revealed 360 significant differences in hit rate for both rate and pitch condi- 361 tions (rate:  $F_{5.90} = 78.72$ , P < 0.001,  $\varepsilon = 0.485$ ; pitch:  $F_{5.90} = 25.21$ , 362 P < 0.001,  $\varepsilon = 0.596$ ). These differences in performance were best cap- 363 tured by subtracting hit rate averaged across the two conditions with 364 the largest stimulus change from hit rate averaged across the two con-365 ditions with the smallest stimulus change. For rate conditions, this dif- 366 ference was significantly smaller than zero ( $t_{18} = -13.52$ , P < 0.001), 367 indicating that discrimination of rate change was most difficult for 368 small changes in modulation rate. For pitch conditions, the difference 369 between small and large stimulus changes was significantly greater 370 than zero ( $t_{18} = 4.49$ , P < 0.001), indicating that discrimination of 371 rate change was most difficult for large changes in pitch (Fig. 2A, top 372 right). 373

The analyses reported in the previous paragraph for hit rate are in- 374 formative regarding general task difficulty. The sensitivity to discrimi- 375 nate rate changes and the degree to which changes in pitch induced 376 illusory rate-change percepts are more clearly investigated using pro- 377 portion of 'speeding up' responses (Fig. 2B). The slope of a linear fit 378 to proportions of 'speeding up' responses as a function of rate levels 379 (z-normalized) provides an individual measure of rate-discrimination 380 sensitivity (discrimination index). The slope of a linear fit to proportions 381 of 'speeding up' responses as a function of pitch levels (z-normalized) 382 provides an individual measure of the strength of rate-change illusion 383 (illusion index). Both indices (slopes) were significantly larger than 384 zero (rate:  $t_{18} = 11.96$ , P < 0.001; pitch:  $t_{18} = 7.41$ , P < 0.001; Fig. 2B, 385 top right), thus indicating that participants were able to discriminate 386 rate changes, but at the same time were biased in their rate-change per-387 cepts by changes in pitch. Noteworthy, rate slopes (discrimination 388 index) were larger than pitch slopes (illusion index), indicating that 389 percepts of rate change were, as expected, more strongly driven by 390 rate changes than by pitch changes ( $t_{18} = 2.98$ , P = 0.008). 391

In addition, we tested whether perceptual sensitivity for rate chang-  $_{392}$  es was linked to pitch-induced illusions by correlating the discrimina-  $_{393}$  tion index with the illusion index. We observed a significant negative  $_{394}$  correlation (r = -0.638, t<sub>18</sub> = 3.42, P = 0.003), indicating that partic-  $_{395}$  ipants who were more sensitive to rate changes tended to perceive  $_{396}$  weaker illusions. This correlation reflects the necessity of reduced illu-  $_{397}$  sions with increased perceptual sensitivity to rate changes, while re-  $_{398}$  duced perceptual sensitivity could in principle be independent from  $_{399}$  changes in pitch.

#### Overall brain activation related to sound processing and task performance 401

Fig. 3 depicts brain regions that were more strongly activated during402sound presentation and general task performance than during silent tri-403als. Activated brain regions included the bilateral superior temporal404gyri, left inferior frontal gyrus (pars opercularis), left precentral gyrus405and pre-SMA. Table 1 shows MNI peak voxel coordinates and the correstion406sponding statistical results.407

#### Neural activation as a function of task performance (increasing task 408 difficulty) 409

In order to investigate changes in neural activation coupled to task 410 performance, for each participant a linear function was fitted to neural 411 activation (beta values from separate GLMs using rate and pitch condi-412 tions as covariates) as a function of hit rate, separately for rate and for 413 pitch conditions. The resulting slopes reflect a measure of brain activa-414 tion modulation by increasing task difficulty, which we refer to as 415

Please cite this article as: Herrmann, B., et al., Supplementary motor area activations predict individual differences in temporal-change sensitivity and its illusory distortions, NeuroImage (2014), http://dx.doi.org/10.1016/j.neuroimage.2014.07.026

B. Herrmann et al. / NeuroImage xxx (2014) xxx-xxx



**Fig. 2.** Behavioral performance. A) Bottom left: Mean hit rate (proportion correct weighted by confidence ratings) for the  $6 \times 6$  rate–pitch conditions. Rate and pitch marginal means are provided at the bottom right and top left, respectively. Top right: Hit rates for large changes in rate or pitch subtracted from hit rates for small changes ( $\Delta$ hit rate). For both rate and pitch conditions, this difference is significantly different from zero. B) Bottom left: Mean proportion of 'speeding up' responses (weighted by confidence ratings) for the  $6 \times 6$  rate–pitch conditions. Rate and pitch marginal means are provided at the bottom right and top left, respectively. Top right: Discrimination index (slope of linear fit to rate marginal) and illusion index (slope of linear fit to pitch marginal). Both indices are significantly larger than zero, and also significantly different from each other. \*P < 0.05.

416 'neural task-difficulty index'. Fig. 4 describes the analysis using the data417 of one participant.

Individual slopes were then tested against zero on the group (sec-418 ond) level (Fig. 5A). Significant activation clusters for rate conditions 419 were observed in pre-SMA and in the right IFG (pars opercularis)/insula. 420 421 That is, activation strength (beta values) in pre-SMA and right IFG/ insula increased with decreasing hit rate (increasing task difficulty). 422For pitch conditions, a similar modulation of activation strength was ob-423 served in the left putamen, such that activation strength (beta values) 424 increased with decreasing hit rate (increasing task difficulty). For illus-425426 tration purposes, the predicted neural activation as a function of hit rate (calculated from the linear coefficients) for each participant is 427 depicted in Fig. 5A (middle panels). Table 2 summarizes the results. 428

429 Neural activation linked to rate-change discrimination and pitch-induced430 illusions

Figs. 4C and 5A middle panel) was extracted for each participant. Slopes435were subsequently correlated with the discrimination index (slope of436linear fit to behavioral proportions of 'speeding up' responses as a func-437tion of rate levels) and the illusion index (slope of linear fit to behavioral438proportions of 'speeding up' responses as a function of pitch levels).439Significant correlations of the neural task-difficulty index (for rate440

activation with increasing task difficulty; neural task-difficulty index; 434

conditions) with the discrimination index and illusion index were do- 441 served in pre-SMA (r = -0.536,  $t_{18} = -2.618$ , P = 0.018; r = 0.456, 442  $t_{18} = 2.115$ , P = 0.049; respectively), but not in the right IFG/insula 443 (r = -0.136,  $t_{18} = -0.566$ , P = 0.579; r = 0.001,  $t_{18} = 0.006$ , 444 P = 0.996; respectively). In other words, participants showing stron- 445 ger modulation of pre-SMA activation with increasing task difficulty 446 were more sensitive to discriminate rate changes and showed smaller 447 pitch-induced rate-change illusions. Activations in the right IFG/ insula 448 cortex, on the other hand, only scaled with task difficulty, but were un- 449 related to individual differences in perceptual sensitivity or illusion 450 strength (Fig. 5B).

Further correlations were conducted for maximally-activated voxels
within each significant region. The slope of the linear fit to beta values as
a function of hit rate (reflecting the degree of modulation of brain

In left putamen, neural task-difficulty index (for pitch conditions) was 452 correlated with the illusion index (r = 0.483,  $t_{18} = 2.275$ , P = 0.036), 453



Fig. 3. Brain regions more strongly activated during sound presentation than during silence.

Table 1

#### B. Herrmann et al. / NeuroImage xxx (2014) xxx-xxx

t1.1 t1.2

MNI coordinates and statistics for peak voxels in significant clusters arising from contrasting neural activation to sound trials with neural activation to silent trials SMA - supplet1.3 mentary motor area, STG - superior temporal gyrus, IFG - inferior frontal gyrus. t1.4

-						
t1.5	Region	MNI peak coordinate (mm)	z-Score/number of voxels/volume (cm <sup>3</sup> )			
t1.6	Right STG	51 - 4 - 8	6.62/634/17.12			
t1.7	Left SIG	-57 - 4 - 2	6.68/6/4/18.20			
t1.8	Left IFG	- 57 8 25	4.96/65/1.76			
t1.9	Left precentral gyrus	-54 - 743	4.95/49/1.32			
t1.10	Pre-SMA	-6855	4.73/27/0.73			

but not with the discrimination index (r = -0.054, t<sub>18</sub> = -0.222, 454455P = 0.827). In other words, participants showing stronger modulation of left putamen activation with increasing task difficulty showed small-456er pitch-induced rate-change illusions, while activity modulations in 457left putamen were not related to the sensitivity to discriminate rate 458changes (Fig. 5B). Note that although the discrimination index and the 459illusion index share some variance (as indicated by the significant cor-460 relation reported above), they dissociate on the neural level such that 461 the degree of brain activity modulation in left putamen is correlated 462 463 only with the illusion index, but not with the discrimination index.

464 In order to test whether overall activations are related to behavioral performance, we also correlated the discrimination index and the illu-465 sion index with the activity from the baseline contrast. For this analysis, 466 we used the sound vs. silence contrast images (baseline, see above) and 467 extracted values from the same voxels in pre-SMA, right IFG/insula, and 468 469left putamen as identified by the linear fit analysis (modulation of brain activity by hit rate). Overall brain activity in these regions was correlat-470 471 ed neither with the discrimination index nor with the illusion index (for 472all; P > 0.25), indicating that overall brain activity is unrelated to being sensitive to modulation rate changes and to the magnitude of illusory 473474distortions.

Finally, in order to confirm that the observed across-participant cor-475relation effects are not due to the specific selection of brain regions 476 using all data in the linear fit analysis (Kriegeskorte et al., 2009), we 477

also split the six data blocks into two independent data sets (three 478 blocks each) at the first level. One half of the blocks (1, 3, and 5) was 479 used to test for brain regions that were modulated by increasing task 480 difficulty (linear fits to betas as a function of hit rate; neural task- 481 difficulty index). Given the reduced number of data points, only the 482 pre-SMA out of the three regions (pre-SMA, right IFG/insula, left 483 putamen) survived the whole-brain significance threshold (data not 484 shown). Critically, using the other half of the data (block 2, 4, and 6), 485 modulation of brain activity by increasing task difficulty (linear fits; 486 neural task-difficulty index) was calculated for the identified pre-SMA 487 region. The magnitude of brain activity modulation (neural task- 488 difficulty index for rate conditions) was then correlated with the 489 discrimination index and the illusion index (also derived from 490 blocks 2, 4, and 6). In accordance with the analysis using all blocks, 491 both correlations were significant (discrimination index: r = -0.555, 492  $t_{18} = -2.751$ , P = 0.014; illusion index: r = 0.469,  $t_{18} = 2.190$ , 493 P = 0.043), confirming that participants showing stronger modulation 494 of pre-SMA activation with increasing task difficulty were more sensi- 495 tive to discriminate rate changes and showed smaller pitch-induced 496 rate-change illusions. 497

#### Discussion

This functional imaging study investigated individual patterns of 499 neural activations in response to temporal rate change and illusory dis- 500 tortions thereof. We observed the following: (1) Perception of temporal 501 rate change is biased by changes in pitch; (2) Neural activations in pre- 502 SMA, right IFG/insula and the left putamen were modulated by the dif- 503 ficulty of rate-change discrimination; (3) However, only activation 504 changes in pre-SMA and the left putamen were linked to the perceptual 505 sensitivity to discriminate rate changes and to relative reductions of il- 506 lusory distortions. The present data provide evidence that individual 507 differences in time perception can be related to different patterns of 508 neurofunctional activation. In particular the results offer a dissociation 509 of pre-SMA and left putamen versus the right IFG/insula. In the follow- 510 ing sections, these findings are discussed in more detail. 511

498



Fig. 4. Single-participant (first-level) analysis of brain activation-hit rate correlation. A) Hit rates (proportion correct weighted by confidence ratings) for one participant, illustrated similar to those of Fig. 2A. On the right, mean hit rates are depicted for pitch conditions (averaged over rate) and rate conditions (averaged over pitch). B) Brain activations (beta values; red; calculated for all voxels) for the six rate and the six pitch conditions (top and bottom, respectively) and the corresponding hit rates (from A; blue). Single-participant brain activation is shown for the peak voxel from the group analysis (marked in red) in left putamen (pitch; MNI coordinate: [-18 8 4]) and in pre-SMA (rate; MNI coordinate: [3 17 46]). C) Linear fits to brain activation data (beta values) as a function of hit rate from the same peak voxel (here marked in white). This analysis is separately conducted for rate and pitch conditions. Linear fits were calculated for each voxel. The slope (linear coefficient) provides a measure of modulation of brain activation as a function of hit rate (task difficulty). The participantspecific brain map of slopes (neural task-difficulty index) is subsequently subjected to the second-level analysis.

B. Herrmann et al. / NeuroImage xxx (2014) xxx-xxx



**Fig. 5.** Brain activation as a function of perceptual performance. A) Activation increase with task difficulty (decrease in hit rate) for rate conditions (cyan) and pitch conditions (red). The middle panels show the predicted betas (zero-centered) from linear fits for the peak voxel of each significant cluster (individuals – gray; mean – colored). The slope of the linear fit reflects an index of neural task difficulty. B) The degree of brain activity modulation by hit rate (neural task-difficulty index in A) correlated with the discrimination index and the illusion index. n.s. – not significant, SMA – supplementary motor area, IFG – inferior frontal gyrus.

#### 512 Perceptual interdependence of temporal rate and pitch

In the current study, participants were presented with stimuli that 513 changed simultaneously in modulation rate and pitch over the duration 514of the sound. The two measures derived from the proportion of 'speed-515ing up' responses (i.e., the 'discrimination index' and the 'illusion index') 516indicate that participants judged the modulation rate but at the same 517time were biased in their judgments by changes in pitch. That is, pitch 518increases led to overestimations of the sound's modulation rate and 519pitch decreases led to underestimations of the sound's modulation 520521rate. This finding is in line with a previous magnetoencephalography (MEG) study using similar stimulation (Herrmann et al., 2013) and 522 agrees with reports of pitch-induced biases in perceived musical 523tempo (Boltz, 1998, 2011), interval length (Crowder and Neath, 5241994; Henry and McAuley, 2009; Pfeuty and Peretz, 2010; Shigeno, 5251986), and sound duration (Henry and McAuley, 2013; Yoblick and 526Salvendy, 1970). 527

It has been proposed that pitch-induced illusory distortions of temporal rate percepts are related to learned structural covariations of time 529 and pitch in natural sounds, which coincide with expectations about 530 how stimuli in the acoustic environment will unfold (Boltz, 2011; 531 Jones et al., 1978). For example, the speeding up of rate in music and 532 speech is often accompanied by increases in pitch and vice versa. 533 When such structural covariations are violated perception can become 534 biased in a direction that would fulfill the violated expectation (Boltz, 535 2011; Henry and McAuley, 2013) — in the current study, participants 536 were biased towards perceiving stimuli with increasing pitch as speeding up and stimuli with decreasing pitch as slowing down. 538

Modulation of brain activation by increasing task difficulty during temporal 539 rate-change discrimination 540

Our first analysis aimed to investigate activation changes related to 541 task difficulty during temporal rate-change discrimination. Analyses 542

#### t2.1 Table 2

t2.2 MNI coordinates and statistics for peak voxels in significant clusters arising from linear fits to individual neural activation (beta values) as a function of hit rate (task difficulty). Right-most column: correlation of brain activation modulation with discrimination index and illusion index. n.s. – not significant, SMA – supplementary motor area, IFG – inferior frontal gyrus.

t2.4	Region	Contrast	MNI peak coordinate (mm)	z-Score/number of voxels/volume (cm <sup>3</sup> )	Correlation (r): discrimination index/illusion index
t2.5	Pre-SMA	Rate	3 17 46	-4.17/154/4.16	$-0.536^{*}/0.456^{*}$
t2.6 t2.7	Left putamen	Pitch	-1884	- 3.03/28/0.76	$-0.054^{\text{n.s.}}/0.483^*$
t2.8	* P < 0.05.				

### **ARTICLE IN PRESS**

B. Herrmann et al. / NeuroImage xxx (2014) xxx-xxx

were separately carried out for rate and for pitch conditions, given that
the behavioral data indicated opposite task performance effects for rate
conditions (hit rate decreased with smaller changes in the sound's
modulation rate) compared to pitch conditions (hit rate decreased
with larger changes in the sound's pitch).

For rate conditions, brain activation linearly increased with increas-548ing task difficulty (decreasing hit rate) in the pre-supplementary motor 549area (pre-SMA) and the right-hemispheric inferior frontal gyrus (IFG, 550551pars opercularis), extending into the insula. For pitch conditions, brain 552activation linearly increased with increasing task difficulty (decreasing 553hit rate) in the left putamen (Fig. 5 and Table 2). These observations are in line with previous studies showing the strongest activation in 554the pre-SMA, basal ganglia and right IFG for time discriminations that 555556were most challenging (Ferrandez et al., 2003; Henry et al., in press; Livesey et al., 2007; Tregellas et al., 2006; Wencil et al., 2010; Wiener 557 et al., 2014). 558

Although a number of studies have failed to control for systematic 559 differences in task difficulty between timing and control conditions 560(e.g., Hinton et al., 2004; Kudo et al., 2004; Pouthas et al., 2005), those 561that have highlight the role of the pre-SMA, IFG, and putamen in pro-562cessing specifically temporal information (Coull et al., 2004, 2008; 563Harrington et al., 2010; Morillon et al., 2009; Rao et al., 2001). In these 564565previous studies, however, it has proven challenging to separate brain 566 activations to timing functions per se from processes related to attention to time (Henry et al., in press). In fact, following research on 567feature-selective attention (e.g., Corbetta et al., 1990; Saenz et al., 5682002; Schoenfeld et al., 2007; Treue and Martinez Trujillo, 1999), 569570Coull et al. (2004) reasoned that attending to the temporal features of a stimulus leads to modulations of brain activation in regions specifical-571ly tuned to temporal information (see also Coull, 2004; Macar et al., 5722006). Under the assumption that participants in the current study 573574might have focused more strongly on the modulation rate when rate 575changes were small and more strongly ignored the pitch when pitch 576changes were large (both of which led to decreasing hit rate), stronger activations for more difficult stimulus conditions might have been 577observed in regions tuned specifically to temporal information. 578

However, another possibility comes from a recent MEG study, which 579 580 used similar stimulation as employed in the current study and emphasized the role of modality-specific auditory sensory encoding of rate-581 pitch interdependence (Herrmann et al., 2013; see also Shigeno, 5821986). In this study, perception of temporal rate change and illusory dis-583584tortions thereof was linked to neural phase patterns of low-frequency oscillatory responses in auditory cortex. While acknowledging the fun-585damental differences between MEG versus fMRI recordings (Bowman 586587 et al., 2007; D'Esposito et al., 1999; Hämäläinen et al., 1993; Horwitz et al., 2000), these MEG findings raise the possibility that temporal 588589information per se is encoded already early in the sensory system (Kanai et al., 2011), while activity modulations in pre-SMA, right 590IFG/insula and putamen are indeed linked to changes in task diffi-591culty and/or attentional dynamics during temporal rate-change 592discrimination. 593

594 Perceptual sensitivity and illusory distortions are related to modulation of 595 neural activation in pre-SMA and putamen

In the current study, two measures were extracted from behavioral performance patterns that relate to the ability with which a participant discriminates temporal rate changes. The 'discrimination index', on the one hand, is a measure of the participant's sensitivity to discriminate modulation rate changes, while the 'illusion index' captures the degree to which a participant is biased in his/her modulation rate judgment by changes in pitch.

While previous timing studies investigated modulations of brain
activity due to task difficulty and attentional dynamics (Coull et al.,
2004; Ferrandez et al., 2003; Henry et al., in press; Livesey et al., 2007;
Tregellas et al., 2006), the current study extended this research by

examining the link between brain activity changes and individual perceptual abilities. We observed that the degree of brain activity modula-608 tion by task difficulty in pre-SMA correlated with the individuals' 609 sensitivity to and illusory distortions of perceived temporal rate. That 610 is, individuals whose brain activity was strongly modulated by task difficulty showed higher perceptual sensitivity to modulation rate changes 612 and were relatively less influenced by changes in pitch. Furthermore, 613 similar to pre-SMA, the degree to which brain activity was modulated by task difficulty in left putamen correlated with the illusion index. In 615 contrast, activity in the right IFG/insula only scaled with task difficulty 616 but was unrelated to perceptual sensitivity or illusion strength. 617

Functionally, activity modulations in pre-SMA and left putamen 618 might be related to complementary aspects of attending to versus 619 ignoring features of temporally varying sounds. In particular, pre-SMA 620 activity modulations were related to the difficulty in discriminating/ 621 attending to rate changes (decreasing hit rate with small rate changes), 622 while left putamen activity modulations were related to the difficulty in 623 ignoring pitch changes (decreasing hit rate with large pitch changes). 624 Thus, enhanced activity modulation in pre-SMA might reflect the de-625 gree of attention to rate changes, which in turn led to higher sensitivity 626 to discriminate rate changes and to a reduction in illusory distortions. 627 On the other hand, enhanced modulation of brain activity in putamen 628 might reflect the degree to which pitch changes could be ignored, 629 which in turn led to reduced illusory rate-change distortions, but did not necessarily result in better perceptual sensitivity. 631

Previous fMRI studies reported individual differences in brain activa- 632 tion in regions implicated in temporal processing (pre-SMA, insula, 633 basal ganglia, inferior parietal cortex) as a function of task performance 634 (Coull et al., 2008; Wiener et al., 2014), general time distortions 635 (Harrington et al., 2004; Tipples et al., 2013), and emotion-induced 636 time illusions (Dirnberger et al., 2012). The current results are generally 637 in line with these previous observations, but are also critically different 638 with respect to the previously observed relation between brain activity in 639 IFG/insula (among other areas) and temporal illusions (Bueti and 640 Macaluso, 2011; Dirnberger et al., 2012; Tipples et al., 2013; Wittmann 641 et al., 2010) — such a relation was notably absent in the current study. 642 This discrepancy could be due to differences in experimental stimuli 643 (visual versus auditory), features inducing illusory percepts (global ver- 644 sus local; emotional versus non-emotional) as well as the specific fMRI 645 and behavioral contrasts chosen for between-participant correlations. In 646 fact, in order to examine task performance and attentional dynamics in 647 a temporal discrimination task, the current study focused explicitly on 648 behavior-related modulations of brain activity which in turn were related 649 to individual differences in perceptual sensitivity and the degree of illuso- 650 ry distortions. 651

Consequently, the current study reveals a distinction between brain 652 areas in which difficulty-related activation is correlated with individual 653 differences in perceptual sensitivity and the magnitude of illusory 654 distortions (pre-SMA, left putamen), and those that are not (right 655 IFG/insula). The absence of a relationship between perceptual sensitiv-656 ity and activity modulation in the right IFG/insula suggests that these 657 areas support difficult listening tasks rather unspecifically. Consistent 658 with this suggestion, activity modulations in the right IFG/insula have 659 been reported for a variety of challenging task conditions (Dosenbach 660 et al., 2008; Duncan and Owen, 2000; Nelson et al., 2010) including 661 but not limited to temporal discriminations (e.g., Binder et al., 2004; 662 Erb et al., 2013; Pastor et al., 2006; Wencil et al., 2010). For a detailed 663 discussion of the right IFG/insula see Eckert et al. (2009). 664

In contrast, activations in pre-SMA were related to the individuals' 665 sensitivity to discriminate temporal rate changes and, together with 666 the left putamen, to relative reductions in pitch-induced illusory distor-667 tions. Two alternative interpretations are viable. On the one hand, these 668 findings support the role of the pre-SMA and basal ganglia in general 669 timing functions as suggested previously (Grahn and Rowe, 2009; 670 Harrington et al., 2010; Meck et al., 2008; Pouthas et al., 2005; 671 Schwartze et al., 2012). That is, activations in the respective regions 672

B. Herrmann et al. / NeuroImage xxx (2014) xxx-xxx

are important for temporal rate-change discriminations and for re-673 ductions of the illusion magnitude via sharpening the rate-change 674 675 representation. On the other hand, the current data also support a 676 feature-selective attention interpretation (Coull et al., 2004; Henry et al., in press; Livesey et al., 2007; Tregellas et al., 2006). That is, 677 brain activation might have been more strongly modulated for par-678 ticipants who intensively focused on the modulation rate changes 679 and at the same time intensively ignored changes in pitch. In turn, 680 681 perceptual sensitivity increased in these participants relative to the ones who less intensively focused on rate changes and were more sus-682 683 ceptible to pitch changes. Although it is unlikely that perceptual sensitivity is solely dependent on attentional factors, the current results 684 685 critically demonstrate that activation changes in pre-SMA support the 686 behavioral outcome of time perception, and allow a functional dissociation from the IFG/insula. 687

#### 688 Conclusions

The current study investigated perception of temporal rate change 689 and illusory distortions thereof using functional imaging. Neural activa-690 tions in the pre-SMA, right IFG/insula and left putamen increased with in-691 creasing difficulty to discriminate rate changes. Critically, the data offer a 692 distinction between regions whose activations scale with individuals' per-693 ceptual sensitivity to features of time (pre-SMA, putamen) and those that 694 more generally support difficult listening conditions (IFG/insula). As such, 695 696 the data underscore that individual differences in time perception can be 697 related to different patterns of neurofunctional activation.

#### 698 Acknowledgments

Q3 Q2 This research is supported by the Max Planck Society. B.H., M.J.H.,
700 M.S. and J.O. are supported by a Max Planck Research Group grant to
701 J.O. We thank Toralf Mildner for his help in preparing the ISSS702 protocol as well as for his support during data collection. Two anony703 mous reviewers provided constructive comments on an earlier version
704 of this manuscript.

#### 705 References

738

739

- Alards-Tomalin, D., Leboe-McGowan, L.C., Mondor, T.A., 2013. Examining auditory kappa effects through manipulating intensity differences between sequential tones. Psychol. Res. 77, 480–491.
- Attneave, F., Olson, R.K., 1971. Pitch as a medium: a new approach to psychophysical scal ing. Am. J. Psychol. 84, 147–166.
- Bengtsson, S.L., Ullén, F., Ehrsson, H.H., Hashimoto, T., Kito, T., Naito, E., Forssberg, H., Sadato,
   N., 2009. Listening to rhythms activates motor and premotor cortices. Cortex 45,
   62–71.
- Binder, J.R., Liebenthal, E., Possing, E.T., Medler, D.A., Ward, B.D., 2004. Neural correlates of sensory and decision processes in auditory object identification. Nat. Neurosci. 7, 295–301.
- Boltz, M.G., 1998. Tempo discrimination of musical patterns: effects due to pitch and rhythmic structure. Percept. Psychophys. 60, 1357–1373.
- Boltz, M.G., 2011. Illusory tempo changes due to musical characteristics. Music Percept.
   28, 367–386.
- Bowman, F.D., Guo, Y., Derado, G., 2007. Statistical approaches to functional neuroimaging data. Neuroimaging Clin. N. Am. 17, 441–458.
- 723 Bueti, D., Macaluso, E., 2011. Physiological correlates of subjective time: evidence for the 724 temporal accumulator hypothesis. Neuroimage 57, 1251–1263.
- Burns, E.M., 1999. Intervals, scales, and tuning. In: Deutsch, D. (Ed.), The Psychology of Music, vol. 2. Academic Press, San Diego, CA, pp. 215–264.
- Corbetta, M., Miezin, F.M., Dobmeyer, S., Shulman, G.L., Petersen, S.E., 1990. Attentional modulation of neural processing of shape, color, and velocity in humans. Science 248, 1556–1559.
- Coull, J.T., 2004. fMRI studies of temporal attention: allocating attention within, or
   towards, time. Cogn. Brain Res. 21, 216–226.
- Coull, J.T., Nobre, A.C., 1998. Where and when to pay attention: the neural systems for
   directing attention to spatial locations and to time intervals as revealed by both
   PET and fMRI. J. Neurosci. 18, 7426–7435.
- Coull, J.T., Vidal, F., Nazarian, B., Macar, F., 2004. Functional anatomy of the attentional modulation of time estimation. Science 303, 1506–1508.
   Coull, I.T., Nazarian, B., Vidal, F., 2008. Timing, storage, and comparison of stimulus
  - Coull, J.T., Nazarian, B., Vidal, F., 2008. Timing, storage, and comparison of stimulus duration engage discrete anatomical components of a perceptual timing network. J. Cogn. Neurosci. 20, 2185–2197.

Crowder, R.G., Neath, I., 1994. The influence of pitch on time perception in short melodies.						
Music Percept. 12, 379–386.	741					
D'Esposito, M., Zarahn, E., Aguirre, G.K., 1999. Event-related functional MRI: implications	742					

- for cognitive psychology. Psychol. Bull. 125, 155–164. 743 Dirnberger, G., Hesselmann, G., Roiser, J.P., Preminger, S., Jahanshahi, M., Paz, R., 2012. Give 744 it time: neural evidence for distorted time perception and enhanced memory 745
- it time: neural evidence for distorted time perception and enhanced memory 745 encoding in emotional situations. Neuroimage 63, 591–599. 746 Dosenbach, N.U., Fair, D.A., Cohen, A.L., Schlaggar, B.L., Petersen, S.E., 2008. A dual-networks 747
- architecture of top-down control. Trends Cogn. Sci. 12, 99–105. 748 Duncan, J., Owen, A.M., 2000. Common regions of the human frontal lobe recruited by 749
- diverse cognitive demands. Trends Neurosci. 23, 475:483–475:483, 750 Eagleman, D.M., 2008. Human time perception and its illusions. Curr. Opin. Neurobiol. 18, 751
- 131–136. 752 Eckert, M.A., Menon, V., Walczak, A., Ahlstrom, J., Denslow, S., Horwitz, A.R., Dubno, J.R., 753 2009. At the heart of the ventral attention system: the right anterior insula. Hum. 754 Brain Mapp. 30, 2530–2541. 755
- Erb, J., Henry, M.J., Eisner, F., Obleser, 2012. Auditory skills and brain morphology predict Q4 individual differences in adaptation to degraded speech. Neuropsychologia 50, 757 2154–2164. 758
- Erb, J., Henry, M.J., Eisner, F., Obleser, 2013. The brain dynamics of rapid perceptual adaptation to adverse listening conditions. J. Neurosci. 33, 10688–10697. 760
- Ferrandez, A.M., Hugueville, L., Lehéricy, S., Poline, J.B., Marsault, C., Pouthas, V., 2003. Basal 761 ganglia and supplementary motor area subtend duration perception: an fMRI study. 762 Neuroimage 19, 1532–1544.
- Fitzgerald, M.B., Wright, B.A., 2011. Perceptual learning and generalization resulting from 764 training on an auditory amplitude-modulation detection task. J. Acoust. Soc. Am. 129, 765 898–906. 766
- Grahn, J.A., McAuley, J.D., 2009. Neural bases of individual differences in beat perception. 767 Neuroimage 47, 1894–1903. 768
- Grahn, J.A., Rowe, J.B., 2009. Feeling the beat: premotor and striatal interactions in musicians and nonmusicians during beat perception. J. Neurosci. 29, 7540–7548. 770
- Greenhouse, S.W., Geisser, S., 1959. On methods in the analysis of profile data. 771 Psychometrika 24, 95–112. 772
- Hämäläinen, M.S., Hari, R., Ilmoniemi, R.J., Knuutila, J., Lounasmaa, O.V., 1993. Magnetoen-773
   cephalography theory, instrumentation, and applications to noninvasive studies of 774
   the working human brain. Rev. Mod. Phys. 65, 413–497. 775
- Harrington, D.L., Haaland, K.Y., Knight, R.T., 1998. Cortical networks underlying mechanisms of time perception. J. Neurosci. 18, 1085–1095.
   777
- Harrington, D.L.,Boyd, L.A., Mayer, A.R., Sheltraw, D.M., Lee, R.R., Huang, M., Rao, S.M., 2004. 778 Neural representation of interval encoding and decision making. Cogn. Brain Res. 21, 779 193–205. 780
- Harrington, D.L.,Zimbelman, J.L.,Hinton, S.C.,Rao, S.M., 2010. Neural modulation of temporal encoding, maintenance, and decision processes. Cereb. Cortex 20, 1274–1285. 782
- Harrington, D.L., Castillo, G.N., Fong, C.H., Reed, J.D., 2011. Neural underpinnings of distortions in the experience of time across senses. Front. Integr. Neurosci. 5 (Articles 32). 784
- Henry, M.J., McAuley, J.D., 2009. Evaluation of an imputed pitch velocity model of the auditory kappa effect. J. Exp. Psychol. 35, 551–564.
   Yang M.M. Machine D. 2001.
- Henry, M.J., McAuley, J.D., 2013. Perceptual distortions in pitch and time reveal active 787 prediction and support for an auditory pitch-motion hypothesis. PLoS One 8, e70646. 788
   Henry, M.J., Herrmann, B., Obleser, J., 2014. Selective attention to temporal features on Q6
- neura, n.g., riermann, p., objeser, J., 2014. selective attention to temporal features on Q6 nested time scales. Cereb. Cortex. http://dx.doi.org/10.1093/cercor/bht240 (in press). 790 Herrmann, B., Henry, M.J., Grigutsch, M., Obleser, J., 2013. Oscillatory phase dynamics in 791
- neural entrainment underpin illusory percepts of time. J. Neurosci, 33, 15799-15809. 792 Hinton, S.C., Harrington, D.L., Binder, J.R., Durgerian, S., Rao, S.M., 2004. Neural systems 793
- Supporting timing and chronometric counting: an fMRI study. Cogn. Brain Res. 21, 794 183-192.
   Horwitz, B., Friston, K.J., Taylor, J.G., 2000. Neural modeling and functional brain imaging: 796
- an overview. Neural Netw. 13, 829–846. 797
- Huang, Y.L., Jones, B., 1982. On the interdependence of temporal and spatial judgments. 798 Percept. Psychophys. 32, 7–14. 799
- Jones, B., Huang, Y.L., 1982. Space-time dependencies in psychophysical judgment of 800 extent and duration: algebraic models of the tau and kappa effects. Psychol. Bull. 801 91, 128–142. 802
- Jones, M.R., Maser, D.J., Kidd, G.R., 1978. Rate and structure in memory for auditory 803 patterns. Mem. Cognit. 6, 246–258. 804
- Kanai, R., Lloyd, H., Bueti, D., Walsh, V., 2011. Modality-independent role of the primary auditory cortex in time estimation. Exp. Brain Res. 209, 465–671. 806
- Kriegeskorte, N., Simmons, W.K., Bellgowan, P.S.F., Baker, C.I., 2009. Circular analysis in systems neuroscience: the dangers of double dipping. Nat. Neurosci. 12, 535–540. 808
- Kudo, K., Miyazaki, M., Kimura, T., Yamanaka, K., Kadota, H., Hirashima, M., Nakajima, Y., 809
   Nakazawa, K., Ohtsuki, T., 2004. Selective activation and deactivation of the human 810
   brain structures between speeded and precisely timed tapping responses to identical 811
   visual stimuli. Neuroimage 22, 1291–1301.
- Lewis, P.A., Miall, R.C., 2003. Brain activation patterns during measurement of sub- and supra-second intervals. Neuropsychologia 41, 1583–1592. 814
- Livesey, A.C., Wall, M.B., Smith, A.T., 2007. Time perception: manipulation of task difficulty 815 dissociates clock functions from other cognitive demands. Neuropsychologia 45, 816 321–331. 817
- Macar, F., Coull, J., Vidal, F., 2006. The supplementary motor area in motor and perceptual 818 time processing: fMRI studies. Cogn. Process. 7, 89–94. 819
- Macmillan, N.A., Creelman, C.D., 2005. Detection Theory: A User's Guide. Lawrence 820 Erlbaum Associates, Inc., Mahwah, NJ. 821
- McAuley, J.D., Henry, M.J., Tkach, J., 2012. Tempo mediates the involvement of motor areas 822 in beat perception. Ann. N. Y. Acad. Sci. 1252, 77–84. 823
- Meck, W.H., Penney, T.B., Pouthas, V., 2008. Cortico-striatal representation of time in 824 animals and humans. Curr. Opin. Neurobiol. 18, 145–152. 825

#### B. Herrmann et al. / NeuroImage xxx (2014) xxx-xxx

- Moore, B.C.J., Glasberg, B.R., Gaunt, T., Child, T., 1991. Across-channel masking of changes in modulation depth for amplitude- and frequency-modulated signals. Q. J. Exp. Psychol.
   A 43, 327–347.
- Morillon, B., Kell, C.A., Giraud, A.-L., 2009. Three stages and four neural systems in time estimation. J. Neurosci. 29, 14803–14811.
- Mueller, K., Mildner, T., Fritz, T., Lepsien, J., Schwarzbauer, C., Schroeter, M.L., Möller, H.E.,
   2011. Investigating brain response to music: a comparison of different fMRI acquisition schemes. Neuroimage 54, 337–343.
- Nelson, S.M., Dosenbach, N.U., Cohen, A.L., Wheeler, M.E., Schlaggar, B.L., Petersen, S.E.,
   2010. Role of the anterior insula in task-level control and focal attention. Brain Struct.
   Funct. 214, 669–680.
- Nenadic, I., Gaser, C., Volz, H.-P., Rammsayer, T., Häger, F., Sauer, H., 2003. Processing of temporal information and the basal ganglia: new evidence from fMRI. Exp. Brain Res. 148, 238–246.
- Pastor, M.A., Macaluso, E., Day, B.L., Frackowiak, R.S.J., 2006. The neural basis of temporal auditory discrimination. Neuroimage 30, 512–520.
- Pfeuty, M., Peretz, I., 2010. Abnormal pitch-time interference in congenital amusia:
  evidence from an implicit test. Atten. Percept. Psychophys. 72, 763–774.
- Pouthas, V., George, N., Poline, J.-B., Pfeuty, M., VandeMoorteele, P.-F., Hugueville, L.,
   Ferrandez, A.-M., Lehéricy, S., LeBihan, D., Renault, B., 2005. Neural network involved
   in time perception: an fMRI study comparing long and short interval estimation.
- Hum. Brain Mapp. 25, 433–441.
  Rao, S.M.,Mayer, A.R.,Harrington, D.L., 2001. The evolution of brain activation during temporal processing. Nat. Neurosci. 4, 317–323.
- Saenz, M., Buracas, G.T., Boynton, G.M., 2002. Global effects of feature based attention in human visual cortex. Nat. Neurosci. 5, 631–632.
- Schoenfeld, M.A., Hopf, J.-M., Martinez, A., Mai, H.M., Sattler, C., Gasde, A., Heinze, H.-J.,
   Hillyard, S.A., 2007. Spatio-temporal analysis of feature-based attention. Cereb.
   Cortex 17, 2468–2477.
- Schwartze, M., Rothermich, K., Kotz, S.A., 2012. Functional dissociation of pre-SMA and SMA-proper in temporal processing. Neuroimage 60, 290–298.

- Schwarzbauer, C., Davis, M.H., Rodd, J.M., Johnsrude, I., 2006. Interleaved silent steady state857(ISSS) imaging: a new sparse imaging method applied to auditory fMRI. Neuroimage85829, 774–782.859
- Shigeno, S., 1986. The auditory tau and kappa effects for speech and nonspeech stimuli. 860 Percept. Psychophys. 4, 9–19. 861
- Slotnick, S.D., Schacter, D.L., 2004. A sensory signature that distinguishes true from false 862 memories. Nat. Neurosci. 7, 664–672. 863

Slotnick, S.D., Moo, L.R., Segal, J.B., Hart Jr., J., 2003. Distinct prefrontal cortex activity associated 864 with item memory and source memory for visual shapes. Cogn. Brain Res. 17, 75–82.
 Teki, S., Grube, M., Kumar, S., Griffiths, T.D., 2011. Distinct neural substrates of duration-866

Source, and the set of the set

- subjective experience of short durations (less than 2 seconds). PLoS One 8, e54669. 869 Tregellas, J.R., Davalos, D.B., Rojas, D.C., 2006. Effect of task difficulty on the functional 870
- ing gain in macaque visual cortex. Nature 399, 575–579. 873 Wencil, E.B., Coslett, H.B., Aguirre, G.K., Chatterjee, A., 2010, Carving the clock at its compo-874
- nent joints: neural bases for interval timing. J. Neurophysiol. 104, 160–168. 875 Wiener, M., Turkeltaub, P.E., Coslett, H.B., 2010. The image of time: a voxel-wise metaanalysis. Neuroimage 49, 1728–1740. 877
- Wiener, M., Lee, Y.-S., Lohoff, F.W., Coslett, H.B., 2014. Individual differences in the mor-878 phometry and activation of time perception networks are influenced by dopamine genotype. Neuroimage 89, 10–22. 880
- Wilsch, A., Henry, M.J., Herrmann, B., Maess, B., Obleser, J., 2014. Alpha oscillatory dynamics Q7 index temporal expectation benefits in working memory. Cereb. Cortex. http:// dx.doi.org/10.1093/cercor/bhu004 (in press).
- Wittmann, M., van Wassenhove, V., Craig, A.D.B., Paulus, M.P., 2010. The neural substrates 884 of subjective time dilation. Front. Hum. Neurosci. 4 (Article 2).
- Yoblick, D.A., Salvendy, G., 1970. Influence of frequency on the estimation of time for auditory, visual and tactile modalities: the kappa effect. J. Exp. Psychol. 86, 157–164.