Coding of multisensory temporal patterns in human superior temporal sulcus

Tömme Noesselt1,2*, Daniel Bergmann3,4†, Hans-Jochen Heinze3, Thomas Münte5 and Charles Spence6

1 Department of Biological Psychology, Otto-von-Guericke-Universität Magdeburg, Magdeburg, Germany
2 Center of Behavioral Brain Sciences, Otto-von-Guericke-Universität Magdeburg, Magdeburg, Germany
3 Department of Neurology, Otto-von-Guericke-Universität Magdeburg, Magdeburg, Germany
4 Psychosomatic Medicine, Asklepios Westklinikum Hamburg, Hamburg, Germany
5 Department of Neurology, Universität zu Lübeck, Lübeck, Germany
6 Crossmodal Research Laboratory, Department of Experimental Psychology, University of Oxford, Oxford, UK

Edited by: Zhuanghua Shi, Ludwig-Maximilians-Universität München, Germany
Reviewed by: Emiliana Macaluso, Fondazione Santa Lucia, Italy
Mark W. Greenlee, University of Regensburg, Germany
*Correspondence: Tömme Noesselt, Department of Biological Psychology, Otto-von-Guericke-Universität Magdeburg, Universitätsplatz 2, 39106 Magdeburg, Germany. E-mail: toemme@med.ovgu.de
†These authors equally contributed to this work.

INTRODUCTION

When observers are confronted with incongruent auditory and visual information, that information is often fused into a congruent multisensory percept. Spatial, semantic, and temporal factors have all been shown to contribute to this perceptual fusion (see e.g., Driver and Noesselt, 2008, for a review). The temporal relationship between inputs from different senses plays a particularly important role in multisensory integration (Köhler, 1947; Dennett, 1991; Spence and Squire, 2003; Kelly, 2005) and the perceived synchrony declines when the audio-visual asynchrony exceeds a certain temporal delay. When simple auditory beeps and visual flashes are being judged, subjects’ temporal synchrony window spans approximately 100 ms (Slutsky and Recanzone, 2001; Vatakis and Spence, 2006a) becoming broader/wider when stimuli are more complex (consisting of semantic content; Dixon and Spitz, 1980; McGrath and Summerfield, 1985; Spence and Squire, 2003; Miller and D’Esposito, 2005; Vatakis and Spence, 2006b, see also Vroomen and Keetels, 2010 for review).

Philosophers, psychologists, and neuroscientists have long been interested in how the temporal aspects of perception are represented in the brain. In the present study, we investigated the neural basis of the temporal perception of synchrony/asynchrony for audiovisual speech stimuli using functional magnetic resonance imaging (fMRI). Subjects judged the temporal relation of (a)synchronous audiovisual speech streams, and indicated any changes in their perception of the stimuli over time. Differential hemodynamic responses for synchronous versus asynchronous stimuli were observed in the multisensory superior temporal sulcus complex (mSTS-c) and prefrontal cortex. Within mSTS-c we found adjacent regions expressing an enhanced BOLD-response to the different physical (a)synchrony conditions. These regions were further modulated by the subjects’ perceptual state. By calculating the distances between the modulated regions within mSTS-c within single-subjects we demonstrate that the “auditory leading (AL)” and “visual leading (VL) areas” lie closer to “synchrony areas” than to each other. Moreover, analysis of interregional connectivity indicates a stronger functional connection between multisensory prefrontal cortex and mSTS-c during the perception of asynchrony. Taken together, these results therefore suggest the presence of distinct sub-regions within the human STS-c for the maintenance of temporal relations for audiovisual speech stimuli plus differential functional connectivity with prefrontal regions. The respective local activity in mSTS-c is dependent both upon the physical properties of the stimuli presented and upon the subjects’ perception of (a)synchrony.

Keywords: audiovisual, temporal perception, fMRI, speech, human

Several brain structures have been implicated in the multisensory integration of auditory and visual stimuli. Among them are the superior colliculi (Stein and Meredith, 1993), the superior temporal sulcus complex (STS-c), the intraparietal sulcus (IPS), the insular cortex, the claustrum and prefrontal areas (e.g., Calvert et al., 2000; Bushara et al., 2001; Calvert, 2001; Driver and Noesselt, 2008). Within the STS-c, areas within or close to the upper bank have been identified as key regions governing multisensory integration in both humans (Wright et al., 2003; Beauchamp, 2005a; Noesselt et al., 2007) and non-human primates (Benevento et al., 1977; Desimone and Gross, 1979; Bruce et al., 1981; Hikosaka et al., 1988; Barradough et al., 2005). Direct neuronal recordings from the superior temporal polysensory (STP) region in monkeys have revealed that neurons can respond to both visual and auditory stimuli in both the upper (Bruce et al., 1981; Hikosaka et al., 1988) and lower banks (Benevento et al., 1977). Barradough et al. (2005) reported neurons within the STS-c that respond to action-related congruent...
audiovisual stimulation. When focusing on complex, speech-related animal communication, results from studies in macaques suggest that temporal regions in the macaque brain (including in the STS-c) are activated by audiovisual species-specific vocalizations (Gil-da-Costa et al., 2004; Ghazanfar et al., 2008). In humans, using linguistic stimuli, van Atteveldt et al. (2004) found regions in the STS-c that responded to visually presented letters, spoken single letters, or their combination. As in the study by Wright et al. (2003) employing lip-movements plus spoken syllables, the STS-c response was greatest for audiovisual stimuli. van Atteveldt and colleagues (2004) reported that multisensory enhancement was seen for congruent but not for incongruent stimuli. However, other studies reported enhancements in functional magnetic resonance imaging (fMRI)-responses for incongruent stimuli within STS-c (e.g., van Atteveldt et al., 2007). These findings suggest that the STS-c is involved in the temporal binding of audiovisual stimuli. However, it still needs to be established whether congruent or incongruent audiovisual stimuli elicit a higher fMRI-signal in STS-c, or whether different subregions within the STS-c may differentially code multisensory temporal relations.

Hence, the aim of the present study was to investigate the functional neuroanatomy of the multisensory regions including STS-c and prefrontal cortex when perceiving a temporal (mis-)alignment of ecologically-valid long speech sequences; and to examine whether audiovisual temporal relationships may subdivide multisensory regions functionally. Subjects were shown videos of temporally aligned and misaligned video streams [either auditory leading (AL) or visual leading (VL)] and reported whether those were perceived as being synchronous or asynchronous. Importantly, they also reported changes of perceived timing during the presentation of each stimulus. This design enabled us to dissociate those neural processes that were related to perceptual switches and those related to stable perceptual states during the presentation of audiovisual speech sequences. To anticipate, we found differential BOLD-effects for the different temporal percepts (AL, VL, and synchrounous (AVS)) within adjacent subregions in human STS-c, plus differential interregional connectivity with prefrontal cortex.

METHODS

A temporal-threshold experiment was conducted prior to scanning, to account for any individual differences in temporal perception. By choosing bistable stimuli for each subject we maximized the number of trials per condition during the fMRI-experiment (see below). Subjects (n = 14, 7 female) were placed in a dark, sound-attenuated chamber after providing written informed consent in accord with local ethics. They were seated with eyes closed and were adapted individually to ensure easy comprehension of the auditory speech sequences despite the scanner head, TR 2 s, TE 30 ms, flip 80°, resolution 64 × 64 × 32 at 3.5 × 3.5 × 4 mm). The derived overlapping audio-visual activation map was then used to identify candidate multisensory areas (see below). Fourth, subjects wore earplugs; perceived loudness and balance were adapted individually to ensure easy comprehension of the auditory speech sequences despite the scanner.

FIGURE 1 | Overlap of visual and auditory BOLD-modulations for unisensory stimulus presentations (p < 0.005). This activation map was used as the search volume for the fMRI-analysis in the main experiment.

(NO)behavioral Systems, Inc., CA). Initially, 20 synchronous sequences plus 80 temporally shifted sequences were presented (−130 ms, −60 ms (AL) and 200 ms/400 ms (VL)), 20 video clips each, see Figure 2A). These asynchronies for threshold-determination were chosen in accord with previous reports (Dixon and Spitz, 1980). For the fMRI-experiment, those stimuli were chosen for each subject that had a similar number of synchrony and asynchrony judgments (called near-threshold below).

fMRI-DATA ACQUISITION

fMRI-data was acquired on a whole body Siemens 3 T Trio-scanner (Siemens, Erlangen, Germany) using a circular-polarized whole-head coil (BrukerBioSpin, Ettlingen, Germany). Subjects performed the same task as they had outside the scanner, reporting their responses with their right index, middle, and ring finger. Within the scanner subjects were presented three conditions: near-threshold VL, near-threshold AL plus the AVS condition. All other stimulus parameters were kept as in the behavioral experiment outside the scanner except for the following: first, a baseline period of 20 s was introduced after each video clip. Second, eye movements were monitored using an fMRI-compatible infrared recording system (Kanowski et al., 2007) plus evaluation software (PupilTracker, HumanScan, Erlangen, Germany). The eye movement data was analysed with MATLAB 6.5. Third, before the main fMRI-experiment, a functional localizer was run in which only unimodal auditory or unimodal visual stimuli from the videos were presented (331 volumes covering the whole head, TR 2 s, TE 30 ms, flip 80°, resolution 64 × 64 × 32 at 3.5 × 3.5 × 4 mm). The derived overlapping audio-visual activation map was then used to identify candidate multisensory areas (see below). Fourth, subjects wore earplugs; perceived loudness and balance were adapted individually to ensure easy comprehension of the auditory speech sequences despite the scanner.
FIGURE 2 | Experimental design and behavioral results. (A) Depicts an example of a video-clip presented in three conditions [i.e., auditory leading (top left, temporal lag from 60–120 ms), auditory and video synchronous (top middle), or visual leading (top right, temporal lag from 200–400 ms)]. Auditory and visual lags were determined in a preliminary threshold-determination-experiment. Stimuli were presented at 1° visual angle above fixation (lower boundary) up to 7° (upper boundary). The duration of all 20 video-clips was 23.7 s, the interstimulus interval was 20 s. Participants indicated whether they perceived the auditory stream leading, the visual stream leading, or the 2 streams as being synchronous. They were encouraged to report any changes in their perception during the presentation of each video. Note that the physical lag was fixed within each video clip near the individual’s synchrony/asyncrhony-threshold. (B) Radar graph depicts mean durations (time from one keypress to the next) of subjects’ (a)synchrony-percepts for each experimental condition during fMRI-scanning: perceptual states were longest when perception of (a)synchrony was congruent with physical stimulation. Therefore, in the fMRI-analysis, hemodynamic response functions (HRF) could be specifically modeled and extracted for each stable percept and perceptual switches using a mixed model (see below). (C) An example trial modeled with hemodynamic response functions for an auditory leading-stimulus (A_L). Gamma-curves depict perceptual switches/decisions, whereas box-car functions illustrate the sensory processing prior to the first decision and perceptual states. Purple curves stand for AV_S, red for A_L. Note that each box-car function was individually specified based on the trial-by-trial inter-button-press duration.
noise. The stimuli were presented using MR-compatible, electro-
dynamic headphones (MRconfon, Magdeburg, Germany).

During the main experiment functional volumes were col-
lected in four sessions (331 volumes each, covering the whole
head, TR 2s, TE 30 ms, flip 80°, resolution 64 × 64 × 32
at 3.5 × 3.5 × 4 mm). Additionally, for anatomical localization
an inversion-recovery EPI was acquired (TR 2s, TE 30 ms,
TI:1450 ms, resolution 64 × 64 × 32 at 3.5 × 3.5 × 4 mm, same
slice orientation and distortions as the functional volumes). The
first five volumes from each session were excluded from further
analysis. The remaining volumes were acquisition-corrected to
the first acquired slice of each volume, motion-corrected, nor-
malized at 2 mm³ voxel size and smoothed (6 mm), using SPM2
(Wellcome Department of Cognitive Neurology, London, UK).

GROUP-LEVEL STATISTICS

After pre-processing the data from a localizer run were modeled
with two box-car functions convolved with the hemodynamic
response function (HRF) for the auditory and visual trials. For
the localizer runs, blocks were compared to the baseline dur-
ing which no stimulus was present (p < 0.005; k > 10). An
audiovisual mask (i.e., overlap of unisensory visual and auditory
activations) was computed to identify candidate multisensory
structures (see Figure 1; cf. Beauchamp et al., 2004b; Beauchamp,
2005b; Noesselt et al., 2007; Szczyk et al., 2008).

Next, all experimental conditions were modeled with the HRF
with variable durations when appropriate (mixed model; see
Figure 2C). In particular, 21 conditions were defined in a mixed
model: three perceptual switches (subjects’ button press, event-
related), three perceptual states (time after button press, variable
block) and the initial stimulation (time before the first but-
ton press, variable block) for every stimulus condition (A_L, V_L,
and A_VS). To test condition effects, linear contrasts were used
for each subject and condition and masked inclusively with the
audiovisual overlap from the functional localizer. The result-
ing contrast images were applied to perform random effects
second-level analyses. The statistical parametric maps of the
t-statistics at each voxel were thresholded at p < 0.05 (small-
volume-corrected) and the spatial extent threshold was set at k >
5 voxels.

The following contrasts were computed: First, we identified
regions that responded to physical synchrony and asynchronous
conditions. Second, we identified regions that showed differen-
tial fMRI-signals for perceived synchrony vs. asynchrony con-
ditions. Finally, we computed interaction effects for differential
perceptual states with identical physical stimulation (i.e., asyn-
chronous vs. synchronous percepts separately for A_L, V_L, and A_VS
stimulation).

SINGLE-SUBJECT STATISTICS

We also analysed the data from individual subjects in order
to confirm our group-level results and to test the interaction
between stimulation and percepts formally. We identified for each
subject regions within STS-c using the identical contrasts as in
the group analysis above: for A_L stimulation: veridical A_L per-
cept > non-veridical synchronous percept; for A_VS stimulation:
veridical synchrony percept > both non-veridical percepts; for V_L
stimulation: veridical V_L percept > non-veridical synchronous
percept. Subject-specific regions of interest (ROI) were identi-
ﬁed by searching for signiﬁcant clusters of the three contrasts
of interest within subregions of the STS-c (anatomical crite-
rium) which expressed unisensory responses to both modalities
(additional functional criterion). We extracted the beta-weights
of all experimental conditions from these three local maxima for
each subject and tested whether these local maxima would express
signiﬁcantly different results across stimulations. Note that this
analysis is non-trivial and provides additional information, since
any BOLD-modulation of different perceptual states to the A_VS-
stimulation was left unspecified in the A_L and V_L stimulation
contrasts and vice versa.

ANALYSIS OF INTERREGIONAL CONNECTIVITY

Complementary to the analysis of local modulations of the
BOLD-response we investigated the effects of interregional con-
nectivity in the context of perception of A_VS, V_L, and A_L as
described above (Friston et al., 1997). We seeded our analyses
in the subject-specific local maxima in STS-c and analyzed
which other regions showed enhanced functional coupling in the
context of A_L percepts in the A_L condition (relative to non-
veridical synchronous percept in the A_L condition), in the
context of V_L percepts in the V_L condition (relative to the non-veridical synchronous percept in the A_L condition) and in the
context of synchronous percepts in the synchronous condi-
tion (relative to the non-veridical asynchronous percept in the
synchronous condition) using a model with 21 regressors (see
above) plus the physiological response and the psychophysio-
logical interaction as two additional regressors (see e.g., Noesselt
et al., 2007 for a similar approach) to reveal differential func-
tional interregional connections in the psychological context of
synchronous or asynchronous percepts. Three models were cal-
culated separately for each STS-local maximum (corresponding
to veridical A_L-percepts, veridical V_L-percepts, and veridical A_VS
percepts).

Differential group-level effects were calculated with an analysis
of variance (ANOVA) pertaining the three PPIs from the three
connectivity analysis.

ANALYSIS OF CONSISTENT PATTERNING OF SUBREGIONS

Finally, distances between single subject maxima in STS-c were
computed and analysed to reveal any systematic anatomical dis-
tribution of subjects’ local maxima for the A_L, V_L, and A_VS
representation. For this we used a three step approach: normaliza-
tion of MNI-coordinates, calculation of distances by subtracting
the normalized MNI-coordinates and calculation of Euclidian
distances in three-dimensional space. In particular, for the cal-
culation of distances, the MNI coordinates (in millimeters) of
the three contrasts and their respective local maxima were scaled
by adding the maximum negative value to all coordinates of
one dimension so that all values were positive. This procedure
was applied for the y and z extension/dimension; x coordinates
were converted into their absolute value. Second, coordinate val-
ues of the same dimension but different local maxima were sub-
tracted from each other (A_L/V_L minus synchrony and A_L
minus V_L). Finally, we computed Euclidean distances for the
difference measures: following Pythagoras’ Theorem, difference values of the x and y dimension (cathetuses) were squared and added together and the resulting value (hypotenuse) added to the squared z dimension difference. The square roots of the resulting values (again hypotenuse) represent the reported distances between voxels.

RESULTS

BEHAVIORAL RESULTS
The results of the behavioral experiment outside the scanner revealed that subjects’ judgments became more consistent with stimulation as the audiovisual delay increased. For the auditory stream leading condition, the mean delay for near-threshold stimuli was 105 ms (±35 ms) while for the visual stream leading condition it was 227 ms (±47 ms). Inside the scanner, subjects again judged temporal relations of the video clips while fMRI-data were acquired. The eye-movement data were analysed using both deviations from fixation and eye blinks (Kanowski et al., 2007). Three subjects who showed extensive eye movements or blinking were excluded from further analysis. In the remaining 11 subjects, neither “real” eye movements nor eye blinks showed any differential effect across the experimental conditions (i.e., eye movements < 1°).

During each video subjects (n = 11) switched 5.72 (2.34 SD) times toward a “congruent” perceptual state [i.e., one during which perception and the physical stimulus were identical] vs. 3.97 (2.0) times toward a non-veridical one. Moreover, subjects maintained veridical percepts for 9.13 (3.38) s on average, whereas non-veridical percepts lasted 6.04 (2.02) s (see Figure 2C). In contrast, subjects maintained veridical percepts for 9.13 (3.38) s on average, whereas non-veridical percepts lasted 6.04 (2.02) s (see Figure 2B) for length of stable durations as a function of the stimulus type). This allowed for an unbiased mixed model design (see Figure 2C and Kleinschmidt et al., 1998; Dosenbach et al., 2006 for similar approaches).

NEUROIMAGING RESULTS

Voxel-based group results
First, we computed candidate multisensory structures (i.e., the overlap of activation patterns found with unisensory visual and auditory stimuli before the main experiment; see Beauchamp et al., 2004b; Noesselt et al., 2007; Szycik et al., 2008, for similar approaches). These candidate multisensory structures comprised of bilateral superior temporal sulcus, bilateral anterior insula extending into prefrontal cortex plus bilateral premotor cortex.

When comparing stable perceptual states with switches we found stronger fMRI-responses in bilateral STS-c and lateral prefrontal cortex for the maintenance of perceptual states relative to switches whereas perceptual switches engaged posterior parietal regions plus anterior cingulate in accord with earlier studies (e.g., Heekeren et al., 2008). Since perceptual switches did not significantly modulate voxels within temporal regions, we then focused on the experimental effects of the different stimulus types and of stable perceptual states (i.e., inter-response intervals) within multisensory regions.

First, comparisons of AVS vs. \((V_L+A_L)\) perceptual states (collapsed over stimulus types) revealed modulations in adjacent subregions of bilateral multisensory STS-c, in right insular cortex, and in bilateral prefrontal areas (see Figure 3A and Tables 1A,B); note that both asynchronous and synchronous perceptual states modulated regions within STS-c, whereas only asynchronous perceptual states additionally modulated the anterior insula and prefrontal cortex (see Table 1). Second, comparisons of the physically AVS minus \((V_L+A_L)\) stimuli (regardless of perceptual states) revealed right-lateralised modulations in middle and posterior STS-c plus prefrontal cortex (see Figure 3B, purple spots). \(A_L\) and \(V_L\) stimuli (relative to synchronous stimuli; see Figure 3B, red and blue spots, respectively) showed enhanced BOLD-responses in bilateral STS-c, prefrontal cortex, and anterior insula (see Tables 2A–C for local maxima). Please note, that the time-related modulations are more widespread in the left hemisphere, which might be a reason for the left-sided dominance of synchronous representation reported in other studies (e.g., Miller and D’Esposito, 2005; Marchant et al., 2012).

Finally, we compared different perceptual states separately for each stimulus type (and not collapsed across stimulus type as above). Note that these stimulus-type-specific comparisons were designed to reveal perceptual effects for identical physical stimuli. Differential non-overlapping BOLD-modulations were again found in anterior insula, prefrontal cortex, and STS-c with only asynchronous perceptions expressing higher activations in the insula and prefrontal cortex (see Figure 4, plus Tables 3A–C). Within STS-c, distinct regions for synchronous and asynchronous perceptions were observed as a function of stimulus type. BOLD-modulations for AL and VL conditions (veridically perceived as asynchronous) revealed right-lateralised modulations in middle and posterior STS-c, whereas only asynchronous perceptions expressed higher activations in the insula and prefrontal cortex (see Figure 4, middle row and lower left panel). In the right hemisphere, regions within the STS-c responded to veridically perceived AVS and VL stimuli (see Figure 4, middle and bottom row). We also investigated whether we would find modulations in the fMRI-signal for the main effects of stimulus type, perception and perceptual states for each stimulus type outside the multisensory ROI. However no significant modulations were observed (PFWE-corrected < 0.05, since we did not have any a priori hypothesis).

Single-subject region-of-interest approach
Because of the possible anatomical differences between subjects within the STS-c (Ochiai et al., 2004), a ROI analysis was performed within single subjects to confirm and extend voxel-based group-level responses to physical and/or perceptual (a)synchrony.

For this ROI analysis, three differential temporal percepts were evaluated for each subject with the following contrasts: veridical (asynchronous) minus non-veridical synchronous perception for \(A_L\) and \(V_L\) speech stimuli; plus synchronous minus asynchronous perception for AVS stimulus trains. This analysis was applied again within subjects’ audiovisual masks. Mean beta weights responses (proportional to percent signal change) for the subjects’ perceptual states in every experimental condition were assessed for the three (a)synchrony areas and their respective local maxima. (Note, that these local maxima were identified by conducting comparisons of a limited number of perceptual states, regardless of any other differential effects between conditions.
FIGURE 3 | Voxel-based group BOLD-effect of subjects’ stable perceptual states (time from one keypress to the next, upper row) and the effects of the different stimulus types (lower row) within audiovisual activation maps (as defined by the overlap of unisensory stimuli) thresholded at $p < 0.05$ (small-volume-corrected). Note that the distribution of time-sensitive regions differed in the left and right hemisphere, with the left hemisphere showing a more widespread pattern than the right hemisphere (as evidenced by the formatting). (A) Comparison of synchrony $>$ asynchrony percepts collapsed over stimulus type (purple spots) highlights modulations reaching from posterior to middle STS-c. Adjacent regions within STS-c were also found to be relevant for stable asynchrony percepts $>$ synchronous ones (red spots; additionally, the asynchrony $>$ synchrony percepts-contrast produced significant modulations in prefrontal areas; not shown, see Table 1). (B) Differential BOLD-responses for the three stimulus types collapsed over perceptual state show significant effects at the right posterior STS-c (purple spots; plus premotor regions; not shown, but see Table 2) for synchronous relative to asynchronous stimulation; at both STS-c (blue spots) and prefrontal areas (not shown) for visual leading relative to synchronous stimulation, and at left posterior STS-c and right anterior/posterior STS-c (red spots plus modulations at precentral gyrus and prefrontal areas; not shown, see Table 2) for auditory leading relative to synchronous stimulation.

Thus, the analysis of BOLD-effects reported below, will provide additional information concerning the overall response patterns within the STS-c-subregions:

We extracted the beta weights for all perceptual states (3 states $\times$ 3 stimulus types) from the three local maxima within STS-c and conducted a $2 \times 3 \times 3 \times 3$ repeated measures ANOVA with the factors of hemisphere, type of (a)synchrony area, percept, and stimulus type (see Figure 5B). As no effect of hemisphere was found [$F(1, 10) < 1$; n.s.], beta weights averaged over hemispheres are displayed in Figure 5C. Interaction effects occurred between type of area, percept, and stimulus type [$F(8, 80) = 3.1; p < 0.01$] suggesting that, within each (a)synchrony area, beta weights change as a function of the subjects’ percept and stimulus type. Main effects were observed for type of (a)synchrony area [$F(2, 20) = 4.9; p < 0.05$] and percept [$F(13.33, 13.26) = 10.9; p < 0.01$]. Although post-hoc t-tests showed no significant effects, responses within the “V1 areas” were lower than in the other two areas. BOLD responses to synchronous
stimuli were significantly lower than to asynchronous stimuli \(t_{(20)} = -3.53; p < 0.01\). Interaction effects occurred between hemisphere and type of area \(F_{(2, 20)} = 8.04; p < 0.01\), type of area and percept \(F_{(4, 40)} = 3.48; p < 0.05\), type of area and stimulus type \(F_{(1,73, 17.3)} = 9.17; p < 0.01\), percept and stimulus type \(F_{(1,77, 10)} = 4.7; p < 0.05\).

Further analysis of the ANOVA-data (post-hoc t-tests) revealed that for each stimulus category, subjects’ BOLD responses were highest when a veridical judgment was made. Within the “AL area” (red), the mean BOLD response was highest when subjects perceived an AL stimulus as AL (veridical percept). The according beta weight differed statistically from the two other beta weights and their respective perceptual states \(t_{10} = 3.12; p < 0.05\), whereas the beta weights of the non-veridical percepts did not differ statistically from each other. The same pattern of results was also observed for the AVS region (yellow) \(t_{10} = 4.76; p < 0.001\) and VL percepts (blue) \(t_{10} = 2.72; p < 0.05\). Since, in the AVS area, veridical responses were not significantly different from BOLD-responses for other stimulus types, this region may serve additional sub-functions on top of the maintenance of synchrony perception. In general, these ROI-results reaffirm the functional micro-compartmentalization of the STS-c found in the voxel-based group results into areas specialized for the perception of distinct audiovisual temporal patterns.

**Interregional connectivity of STS-c-regions**

Moreover, we assessed whether the subregions within STS-c that consistently expressed differential local activity (see Figure 5) would also be functionally linked to other multisensory regions. We used the assumption-free “psychophysiological interaction” (PPI; Friston et al., 1997) and seeded our analysis in subject-specific STS-c maxima. We analysed whether the strength of functional coupling of these adjacent STS-c-regions with other multisensory regions would differ. We found that both AL and VL-regions in bilateral STS-c showed a significantly stronger coupling with right prefrontal regions than did the AVS-region (see Figure 6 and Table 4). Moreover, synchronous patches with the middle STS-c expressed a stronger functional connection with posterior STS-c-regions in the left hemisphere, whereas asynchronous patches showed a stronger coupling with posterior STS-c in the right hemisphere (see Table 4).

**Spatial configuration of STS-c-subregions**

In addition, we evaluated whether the spatial configuration of the identified sub-regions within bilateral STS-c showed a systematic spatial distribution across subjects: the analysis revealed that perceived asynchrony (AL or VL) and synchrony modulated distinct regions along the STS-c which were situated adjacent to one another (with asynchrony enclosing synchrony modulations). For every subject, this specific pattern differed in its position along STS-c but occurred regularly (see Figure 5A for illustrative subjects). Distances between the areas modulated by an interaction of stimulus type and perception were calculated. We found that, on average, the local maxima of the “AL” and “VL” areas were situated closer to “synchrony areas” (12.1 and 11.1 mm) than to each other (17.6 mm).

A 2 × 3 repeated measures ANOVA with the factors hemisphere and distance showed an effect of distance \(F_{(2, 20)} = 10.2; p < 0.001\). The distance between the “asynchrony areas” was statistically different from their respective distance to the “synchrony area” [AL: \(t_{10} = 3.77; p < 0.05\); VL: \(t_{10} = 3.40; p < 0.05\)]; the distances between the asynchrony areas and the “synchrony area” were similar \(t_{(10)} = 0.63; p = 0.55\). There was no effect of hemisphere \(F_{(1, 10)} < 1; \text{n.s.}\), nor any interaction between hemisphere and distance \(F_{(2, 20)} < 1; \text{n.s.}\).
Table 2 | Local maxima ($p < 0.05$, $k > 5$ small-volume-corrected) for (A) AVS minus (AL+VL) stimulation within multisensory regions (see Figure 3B, purple spots); (B) VL minus synchrony stimulation (see Figure 3B, blue spots); and (C) AL minus synchrony stimulation (see Figure 3B, red spots) collapsed across perceptual states.

| Anatomical structure | Hemisphere | Cluster size (voxels) | t-value | MNI coordinates |
|----------------------|------------|-----------------------|---------|-----------------|
| Anatomical structure |            |                       |         |                 |
| Anatomical structure | Cluster size (voxels) | t-value | MNI coordinates |
| Anatomical structure | x | y | z |

A. PHYSICAL SYNCHRONY > PHYSICAL ASYNCHRONY

Temporal cortex

| Anatomical structure | Hemisphere | Cluster size (voxels) | t-value | MNI coordinates |
|----------------------|------------|-----------------------|---------|-----------------|
| STS                  | R          | 9                     | 2.39 (0.03) | 54  | −46  | 14 |
| STS                  | R          | 8                     | 2.31 (0.04) | 62  | −50  | 10 |

Frontal lobe

| Anatomical structure | Hemisphere | Cluster size (voxels) | t-value | MNI coordinates |
|----------------------|------------|-----------------------|---------|-----------------|
| Prefrontal cortex    | R          | 14                    | 2.92 (0.008) | 50  | 36   | 12 |

B. PHYSICAL VISUAL LEADING ASYNCHRONY > PHYSICAL SYNCHRONY

Temporal lobe

| Anatomical structure | Hemisphere | Cluster size (voxels) | t-value | MNI coordinates |
|----------------------|------------|-----------------------|---------|-----------------|
| Anterior STS         | R          | 370                   | 3.81 (0.001) | 64  | −20  | −12 |
| Middle STS           | L          | 9                     | 3.05 (0.005) | −68 | −38  | 14 |
| Middle STS           | L          | 14                    | 2.95 (0.007) | −60 | −30  | 8  |

Frontal lobe

| Anatomical structure | Hemisphere | Cluster size (voxels) | t-value | MNI coordinates |
|----------------------|------------|-----------------------|---------|-----------------|
| Prefrontal cortex    | R          | 41                    | 2.78 (0.01) | 38  | 18   | 26 |
| Prefrontal cortex    | L          | 11                    | 2.73 (0.01) | −46 | 20   | 28 |
| Anterior insula/IFG  | L          | 8                     | 2.99 (0.006) | 50  | 42   | 2  |
| Anterior insula/IFG  | L          | 41                    | 3.88 (0.001) | −36 | 38   | −16|

C. PHYSICAL AUDITORY LEADING ASYNCHRONY > PHYSICAL SYNCHRONY

Temporal Lobe

| Anatomical structure | Hemisphere | Cluster size (voxels) | t-value | MNI coordinates |
|----------------------|------------|-----------------------|---------|-----------------|
| Anterior STS         | R          | 177                   | 3.55 (0.002) | 62  | −14  | −8  |
| Posterior/middle STS | R          | 122                   | 3.32 (0.002) | 54  | −46  | −2  |
| Middle STS           | L          | 12                    | 4.07 (0.001) | −68 | −38  | 14 |
| Posterior STS        | L          | 57                    | 2.86 (0.009) | −54 | −54  | 8  |

Frontal Lobe

| Anatomical structure | Hemisphere | Cluster size (voxels) | t-value | MNI coordinates |
|----------------------|------------|-----------------------|---------|-----------------|
| Precentral gyrus     | R          | 17                    | 3.27 (0.003) | 44  | 0    | 40 |
| Precentral gyrus     | R          | 7                     | 2.47 (0.02) | 48  | 6    | 44 |
| Anterior insula/IFG  | L          | 6                     | 3.31 (0.002) | −36 | 40   | −18 |
| Prefrontal cortex    | L          | 19                    | 2.46 (0.02) | −46 | 22   | 24 |

DISCUSSION

The present study investigated the neural basis of both the processing of physical properties and subjective perception of the temporal relationship between auditory and visual speech stimuli, thereby pinpointing the functional neuroanatomy of audiovisual temporal processing and perception in multisensory cortex in humans. We found that sub-regions within the superior temporal sulcus have a distinct response pattern during the maintenance of perceptual states and for the processing of physical stimulus differences regardless of subjects’ perceptual state. Within lateral prefrontal regions and anterior insula only the perception of asynchrony was consistently linked to an increase in BOLD-response. A ROI-based single-subject analysis corroborated and extended this pattern: three subregions within the STS-c showed a differential response for the different physical stimuli (AL, VL, and AVS). Responses were further enhanced if subjects’ perceptual states were congruent to the physical stimulus being presented. Further, analyses of interregional connectivity suggest that during the perception of asynchronous stimuli AL and VL regions within the STS-c are coupled more strongly to lateral prefrontal regions, whereas connectivity within posterior STS-c was lateralized with stronger connections of the middle with posterior STS-c in the left hemisphere for synchrony patches and with posterior STS-c in the right hemisphere for asynchronous patches. Finally, analysis of the anatomical patterning of these regions suggests that they are distributed regularly within the STS-c with a synchrony region being enclosed by asynchrony regions.

Previous neuroimaging studies have reported that the STS-c (among other structures) is involved in audiovisual temporal processing and synchrony perception (Calvert, 2001; Macaluso et al., 2004; Miller and D’Esposito, 2005; Dhamala et al., 2007; Noesselt et al., 2007; Stevenson et al., 2010; Marchant et al., 2012; see Driver and Noesselt, 2008, for a review). However, most of these studies investigating the crossmodal binding of semantically meaningful stimuli (Calvert et al., 2000; Calvert and Campbell, 2003; Macaluso et al., 2004) did not separate task- and perception-related effects; their reported modulations may therefore reflect a mixture of stimulus-, decision-, and perception-related processing.
multisensory areas, including STS-c, plus frontal regions. Noesselt et al. (2007) reported effects in contralateral STS-c for the processing of lateralized non-semantic synchronous audiovisual stimuli, but did not report effects for asynchronous audiovisual stimuli. In a related study, Marchant et al. (2012) observed left-sided synchrony representations in left STS-c. Meanwhile, van Atteveldt et al. (2004, 2007) identified lateral temporal areas (PT, STP, and STS-c) as major integration sites whenever audiovisual grapheme-morpheme pairs were being processed. While the intensity of modulations increased in auditory areas for semantically congruent conditions, the location of modulations within the STS-c changed as a function of the temporal distance/delay between vision and sound: asynchrony was predominantly processed at the eccentricity of the STS-c activation pattern, whereas smaller temporal delays were related to the activation’s core region. However, no effect of synchrony was reported for synchronous audiovisual letters in the STS-c and the reported activations for different audiovisual lags overlapped substantially.

In the present study, asynchronous percepts engaged the posterior STS-c, the anterior insula, and the prefrontal cortex bilaterally. Our results accord with previous imaging studies on temporal asynchrony which reported right-sided effects within the STS-c, supplementary motor areas (Miller and D’Esposito, 2005; Stevenson et al., 2007) and prefrontal (MFG, IFG) cortices (Bushara et al., 2001; Dhamala et al., 2007) in the perception of asynchrony. Our findings corroborate previous results and suggest that audiovisual prefrontal areas and the STS-c are functionally linked during the maintenance of the perception of audiovisual asynchrony. There is also corroborating anatomical evidence that the STS-c is reciprocally linked to prefrontal regions (see e.g., Yeterian et al., 2012). We speculate that the perception of asynchronous percepts may be more demanding than synchrony perception and requires the on-line updating of two separate working memory representations in prefrontal cortex with input from the STS-c. Alternatively, the separation of auditory and visual input may be processed by prefrontal cortical regions (in line with the notion of a hierarchical multisensory processing model, see e.g., Noppeney et al., 2010) and fed back into the STS-c. Future research in non-human primates or in humans using transcranial magnetic stimulation/transcranial direct current stimulation is needed to disentangle these two possibilities.

Most remarkably of all, our results indicate that the multisensory superior temporal sulcus complex (mSTS-c) can be further differentiated into subregions that process particular audiovisual temporal patterns. Anatomical studies in non-human primates that have investigated the anatomical texture of TPO (the likely homologue to the human STS-c; Beauchamp, 2005a) have provided evidence for three caudal-to-rostral subdivisions within this region (Cusick et al., 1995). Those subdivisions are distinct in terms of their chemoarchitecture. Seltzer and Pandya (1991) provided evidence that TPO consists of cytoarchitectonic subdivisions of which particularly the rostral part is directly connected to the insula. Further chemoarchitectonic results support the view that the upper bank of TPO in the rhesus monkey contains several different anatomical and functional zones.
Table 3 | Local maxima ($p < 0.05$, $k > 5$ small-volume-corrected) for (A) Auditory leading minus synchrony percepts during AL stimulation within multisensory regions (see Figure 4, red spots); (B) synchrony minus (AL + VL) percepts during AVS stimulation (see Figure 4, purple spots); and (C) VL minus synchrony percepts during VL stimulation (see Figure 4, blue spots).

| Anatomical structure | Hemisphere | Cluster size (voxels) | t-value  | MNI coordinates |
|----------------------|------------|-----------------------|----------|-----------------|
|                       |            |                       |          |                 |
| A. COHERENT AUDITORY LEADING PERCEPT > COHERENT SYNCHRONY PERCEPT |            |                       |          |                 |
| Temporal lobe         |            |                       |          |                 |
| Posterior/middle STS  | L          | 57                    | 2.62 (0.01) | −52 −42 4 |
| Posterior/middle STS  | L          | 18                    | 2.96 (0.01) | −64 −38 10 |
| Posterior/middle STS  | L          | 6                     | 2.85 (0.01) | −64 −36 −8 |
| Middle STS            | R          | 8                     | 3.43 (0.005) | −54 −30 −14 |
| Anterior/middle STS   | R          | 15                    | 2.95 (0.007) | −64 −42 12 |
| Frontal lobe          |            |                       |          |                 |
| Anterior insula       | L          | 20                    | 3.64 (0.001) | −32 28 −6 |
| anterior insula       | R          | 66                    | 6.52 (0.000) | 42 32 −6 |
| Precentral gyrus      | R          | 86                    | 3.98 (0.002) | 48 10 40 |
| precentral gyrus      | L          | 28                    | 4.67 (0.000) | −36 8 60 |
| Precentral gyrus      | L          | 11                    | 3.41 (0.006) | −40 8 38 |
| Prefrontal cortex     | R          | 191                   | 3.25 (0.003) | 54 28 12 |
| Prefrontal cortex     | L          | 12                    | 2.97 (0.01) | −50 6 44 |
| B. COHERENT SYNCHRONY PERCEPT > COHERENT ASYNCHRONY PERCEPT |            |                       |          |                 |
| Anterior STS          | R          | 447                   | 4.24 (0.001) | 62 −16 −4 |
| Anterior STS          | L          | 62                    | 4.05 (0.002) | −62 −24 4 |
| Posterior STS         | R          | 5                     | 2.66 (0.02) | 48 −52 10 |
| Posterior/middle STS  | L          | 29                    | 3.28 (0.007) | −50 −46 8 |
| Frontal lobe          |            |                       |          |                 |
| Anterior Insula/Prefrontal | L | 14                    | 2.63 (0.01) | −34 40 −14 |
| C. COHERENT VISUAL LEADING PERCEPT > COHERENT SYNCHRONY PERCEPT |            |                       |          |                 |
| Temporal lobe         |            |                       |          |                 |
| Posterior STS         | R          | 18                    | 3.5 (0.002) | 64 −50 2 |
| Posterior STS         | R          | 5                     | 2.41 (0.02) | 62 −50 14 |
| Posterior STS         | L          | 29                    | 3.47 (0.002) | −58 −50 8 |
| Frontal lobe          |            |                       |          |                 |
| Anterior insula       | R          | 72                    | 4.77 (0.000) | 44 40 −10 |
| anterior insula       | L          | 20                    | 2.71 (0.008) | −34 30 −2 |
| Precentral gyrus      | R          | 21                    | 3.42 (0.002) | 42 8 46 |
| Prefrontal cortex     | R          | 451                   | 3.76 (0.001) | 50 24 24 |
| Prefrontal cortex     | L          | 134                   | 3.56 (0.002) | −42 16 26 |

MNI, Montreal Neurological Institute; L, left; R, right.

(Padberg et al., 2003). They demonstrated that within those distinct neurochemical/connectional modules the STS-c shows a patchy organization of connections toward other cerebral regions. Those patches within the STS-c may have functional relevance. In a functional imaging study, Beauchamp et al. (2004a) reported that STS-c can be parcelled into unisensory auditory, visual, and multisensory patches. Our imaging analysis extends these findings and reveals distinct multisensory patches along the STS-c that encode separate audiovisual temporal patterns when the synchrony/asynchrony of continuous speech is being judged. Given that the identified synchrony patches lie in-between auditory- and visual-leading audiovisual patches, these modulations build up a chronological array that suggests the existence of a “time line.” Moreover, another publication (Fairhall and Macaluso, 2009) also reported a modulation of the fMRI-signal due to attention within middle but not posterior STS-c, when subjects processed congruent audiovisual speech, thereby suggesting a large-scale segregation of the STS-c along the anterior-posterior axis (though asynchronous representations seem to be more variable; see Tables 1–3). Moreover, Marchant et al. (2012) investigated the correspondence of an audiovisual behavioral benefit on BOLD-modulations in the cerebrum and found significant effects in middle but not posterior STS-c for synchronous stimulus trains. The results from our study—revealing an interaction effects in middle STS-c specific for temporal patterns and their perception plus an enhanced connectivity with more posterior regions—are in accord with this proposition (though note that our results did not reveal a clear anterior-posterior
FIGURE 5 | Panels showing the results of single-subject analyses. The contrasts displayed here represent subjective perceptions that were congruent with physical stimulation > incoherent perceptions for auditory/visual leading (AL, VL, red/blue spots) and synchronous stimuli (AVS, purple spots). (A) The colored spots indicate average local maxima (11 subjects) of areas that express higher activations for veridical percepts (see main text for contrast definitions) than for non-veridical ones within the STS-c region (region of interest). The white lines and their corresponding numbers display averaged distances in millimeters from one activation spot to the two others (see "Methods" section for details). Note that asynchrony spots are always more distant from each other than from synchrony activation. Colored lines show the individual anatomical curvatures of STS-c of the all subjects after normalization. (B) The middle row depicts the activation maps of three individual subjects for the above-described contrasts. Note that synchrony spots (purple) are enclosed by two asynchrony spots [blue and red spots; see also distances in panel (A)]. Such activation patterns were found in both left and right hemispheres. (C) Mean beta-weights (proportional to % signal change) for the local maxima in panel (A) were collapsed over hemispheres. Bars show the height of the BOLD-effect (y-axis) for each stable percept (auditory leading (red bars), synchronous (orange bars), and visual leading (blue bars)) for the three stimulus types (auditory leading, visual leading, and synchronous, x-axis) within each each of the local maxima shown in panel (A) (auditory leading percept maximum (left graph section), synchrony percept maximum (middle section of graph), and visual leading percept (right graph section)). BOLD-responses to asynchrony percepts within asynchrony percept maxima were always higher (outer left and right bars) than to any other percept for the different stimulus types. Within the synchrony percept maximum BOLD-responses to synchrony percepts were higher than asynchrony percepts whenever synchronous video clips were presented.
FIGURE 6 | Interregional connectivity of subjects’ stable veridical percepts (i.e., identical with the physical stimulus) during asynchronous stimulation thresholded at $p < 0.05, k > 5$ (small-volume-corrected). Left column: right prefrontal regions only expressed stronger coupling with temporal regions in the context of veridical asynchronous $>$ non-veridical synchronous perceptions during $A_L$ and $V_L$ stimulation (see Table 4 for all maxima). Right column shows the origin of the brain sections depicted on the left on a lateral group mean view.

Table 4 | Local Maxima ($p < 0.05, k > 5$ small-volume-corrected) of interregional connectivity in the context of (A) veridical synchrony percepts (relative to non-veridical asynchrony percepts) during $AV_S$ stimulation; (B) veridical auditory leading percepts (relative to non-veridical synchrony percepts) during $A_L$ stimulation; (C) veridical visual leading percepts (relative to non-veridical synchrony percepts) during $V_L$ stimulation.

| Anatomical structure       | Hemisphere | Cluster size (voxels) | $p$-value     | MNI coordinates |
|----------------------------|------------|-----------------------|---------------|-----------------|
|                            |            |                       | $p$-value     | $x$  | $y$  | $z$  |
| **A. PSYCHOPHYSICAL INTERACTION OF SYNCHRONY PERCEPTS** |            |                       |               | $x$  | $y$  | $z$  |
| Temporal regions           |            |                       |               | $x$  | $y$  | $z$  |
| Posterior STS              | L          | 31                    | 4.32 (0.001)  | −54  | −54  | 12   |
| **B. PSYCHOPHYSICAL INTERACTION OF AUDITORY LEADING PERCEPTS** |            |                       |               | $x$  | $y$  | $z$  |
| Temporal regions           |            |                       |               | $x$  | $y$  | $z$  |
| Anterior STS               | R          | 73                    | 3.57 (0.001)  | 64   | −12  | −8   |
| Posterior STS              | R          | 22                    | 3.32 (0.001)  | −50  | −46  | 16   |
| Frontal regions            |            |                       |               | $x$  | $y$  | $z$  |
| Middle/inferior frontal gyrus | R      | 168                   | 2.94 (0.002)  | 40   | 22   | 16   |
| Precentral gyrus           | L          | 54                    | 2.68 (0.005)  | −48  | −2   | 46   |
| **C. PSYCHOPHYSICAL INTERACTION OF VISUAL LEADING PERCEPTS** |            |                       |               | $x$  | $y$  | $z$  |
| Temporal regions           |            |                       |               | $x$  | $y$  | $z$  |
| Posterior STS/STG           | R          | 44                    | 2.42 (0.009)  | 52   | −46  | 0    |
| Frontal regions            |            |                       |               | $x$  | $y$  | $z$  |
| Inferior frontal gyrus     | R          | 21                    | 2.51 (0.008)  | 58   | 22   | 14   |

distinction for the main effects of physical vs. perceptual states). Finally, our results could be applied to nonhuman primates to enable more invasive measures [combined with fMRI (see Tsao et al., 2006)] to identify the pathways and neural mechanisms involved. A study in non-human primates on audiovisual face-voice integration (Ghazanfar et al., 2008) reported enhanced coupling of STS-c-neurons with auditory areas when processing audiovisual stimuli (Schroeder et al., 2008). Our results would predict the existence of distinct patches within mSTS-c that may differentially engage unisensory cortices via feedback connections (Driver and Noesselt, 2008).

In conclusion, we found a distinct pattern of modulations within mSTS-c reflecting an interaction between perceptual state and the physical properties of audiovisual speech stimuli. Our data therefore suggest that there is an aligned spatial representation of audiovisual temporal patterns parcellating the multisensory STS-c in humans, with differential functional connections to multisensory prefrontal regions.
ACKNOWLEDGMENTS
Tomme Noesselt, Daniel Bergmann, and Hans-Jochen Heinze were supported by DFG-SFB-TR31/TPA8; Thomas Münte by DFG-SFB-TR31/TPA7 and Charles Spence by the Alexander von Humboldt Foundation, Germany. Tomme Noesselt, Daniel Bergmann, and Charles Spence planned the experiment. Daniel Bergmann collected and analysed the data. All authors were involved in writing the MS.

REFERENCES
Barralough, N. E., Xiao, D., Baker, C. I., Oram, M. W., and Perrett, D. I. (2005). Integration of visual and auditory information by superior temporal sulcus neurons responsive to the sight of actions. J. Cogn. Neurosci. 17, 377–391.

Beauchamp, M. S. (2005a). See me, hear me: multisensory integration in lateral occipital-temporal cortex. Curr. Opin. Neurobiol. 15, 145–153.

Beauchamp, M. S. (2005b). Statistical criteria in fMRI studies of multisensory integration. Neuroinformatics 3, 93–113.

Beauchamp, M. S., Argall, B. D., Bodurka, J., Duyn, J. H., and Martin, A. (2004a). Unraveling multisensory integration: patchy organization within human STS-c multisensory cortex. Nat. Neurosci. 7, 1190–1192.

Benevento, L. A., Fallon, J., Davis, B. J., and Rezak, M. (1977). Auditory-visual interaction in single cells in the cortex of the superior temporal sulcus and the orbital frontal cortex of the macaque monkey. Exp. Neurol. 57, 489–472.

Bruce, C., Desimone, R., and Gross, C. G. (1981). Visual properties of neurons in a polysensory area in superior temporal sulcus of the macaque. J. Neurophysiol. 46, 369–384.

Bushara, K. O., Grafman, J., and Hallett, M. (2001). Neural correlates of auditory-visual stimuli onset asynchrony detection. J. Neurosci. 21, 300–304.

Calvert, G. A. (2001). Crossmodal processing in the human brain: insights from functional neuroimaging studies. Cereb. Cortex 11, 1110–1123.

Calvert, G. A., and Campbell, R. (2003). Reading speech from still and moving faces: the neural substrates of visible speech. J. Cogn. Neurosci. 15, 57–70.

Calvert, G. A., Campbell, R., and Brammer, M. J. (2000). Evidence from functional magnetic resonance imaging of crossmodal binding in the human heteromodal cortex. Curr. Biol. 10, 649–657.

Calvert, G. A., Hansen, P. C., Iversen, S. D., and Brammer, M. J. (2001). Detection of audio-visual integration sites in humans by application of electrophysiological criteria to the BOLD effect. Neuroimage 14, 427–438.

Cusick, C. G., Seltzer, B., Cola, M., and Griggs, E. (1995). Chemoarchitectonics and corticocortical terminations within the superior temporal sulcus of the rhesus monkey: evidence for subdivisions of superior temporal polysensory cortex. J. Comp. Neurol. 360, 513–535.

Dennett, D. (1991). Consciousness Explained. London: Penguin Press.

Desimone, R., and Gross, C. G. (1979). Visual areas in the temporal cortex of the macaque. Brain Res. 178, 363–380.

Dhamala, M., Assisi, C. G., Jirsa, V. K., Steinberg, F. L., and Keho, J. A. (2007). Multisensory integration for timing engages different brain networks. Neuroimage 34, 764–773.

Dixon, N. F. and Spitz, L. (1980). The detection of auditory visual desynchrony. Perception 9, 719–721.

Dosenbach, N. U., Visscher, K. M., Palmer, E. D., Miezin, F. M., Wenger, K. K., Kang, H. C., Burgund, E. D., Grimes, A. L., Schlaggar, B. L. and Petersen, S. E. (2006). A core system for the implementation of task sets. Neuro Image 50, 799–812.

Driver, J., and Noesselt, T. (2008). Multisensory interplay reveals crossmodal influences on ‘sensory-specific’ brain regions, neural responses, and judgments. Neuron 57, 11–23.

Fairhall, S., and Macaluso, E. (2009). Spatial attention can modulate audiovisual integration at multiple cortical and subcortical sites. Eur. J. Neurosci. 29, 1247–1257.

Friston, K. J., Buechel, C., Fink, G. R., Morris, J., Rolls, E., and Dolan, R. J. (1997). Psychophysiological and modulatory interactions in neuroimaging. Neuroimage 6, 218–229.

Ghazanfar, A. A., Chandrasekaran, C., and Logothetis, N. K. (2008). Interactions between the superior temporal sulcus and auditory cortex mediate dynamic face/voice integration in rhesus monkeys. J. Neurosci. 28, 4457–4469.

Gil-da-Costa, R., Braun, A., Lopes, M., Hauser, M. D., Carson, R. E., Herscovitch, P., and Martin, A. (2004). Toward an evolutionary perspective on conceptual representation: species-specific calls activate visual and affective processing systems in the macaque. Proc. Natl. Acad. Sci. U.S.A. 101, 17156–17152.

Heekeren, H. R., Marret, S., and Ungerleider, L. G. (2008). The neural systems that mediate human perceptual decision making. Nat. Rev. Neurosci. 9, 467–479.

Hikosaka, K., Iwai, E., Saito, H., and Tanaka, K. (1998). Polysensory properties of neurons in the anterior bank of the caudal superior temporal sulcus of the macaque monkey. J. Neurophysiol. 60, 1615–1637.

Kanowski, M., Rieger, J. W., Noesselt, T., Tempellmann, C., and Hinrichs, H. (2007). Endoscopic eye tracking system for fMRI. J. Neurosci. Methods 160, 10–15.

Kelly, S. D. (2005). “The puzzle of temporal experience,” in Cognition and the Brain: The Philosophy and Neuroscience movement, ed A. B. K. Akins (Cambridge: Cambridge University Press), 208–240.

Klein schmidt, A., Buchel, C., Zeki, S., and Frackowiak, R. S. (1998). Human brain activity during spontaneously reversing perception of ambiguous figures. Proc. Biol. Sci. 265, 2427–2433.

Köhler, W. (1947). Gestalt Psychology: An Introduction to New Concepts in Modern Psychology. New York, NY: Liveright Publication Corporation.

Macaluso, E., George, N., Dolan, R., Spence, C., and Driver, J. (2004). Spatial and temporal factors during processing of audiovisual speech: a PET study. Neuroimage 21, 725–732.

Marchant, J. L., Ruff, C. C., and Driver, J. (2012). Audiovisual synchonry enhances BOLD responses in a brain network including multisensory STS while also enhancing target-detection performance for both modalities. Hum. Brain Mapp. 33, 1212–1224.

McGrath, M., and Summerfield, Q. (1985). Intermodal timing relations and audio-visual speech recognition by normal-hearing adults. J. Acoust. Soc. Am. 77, 678–685.

Miller, L. M., and D’Esposito, M. (2005). Perceptual fusion and stimulus coincidence in the cross-modal integration of speech. J. Neurosci. 25, 5884–5893.

Noesselt, T., Rieser, J. W., Schoenfeld, M. A., Kanowski, M., Hinrichs, H., Heinze, H. J., and Driver, J. (2007). Audiovisual temporal correspondence modulates human multisensory superior temporal sulcus plus primary sensory cortices. J. Neurosci. 27, 11431–11441.

Noppeney, U., Ostwald, D., and Werner, S. (2010). Perceptual decisions formed by accumulation of audiovisual evidence in prefrontal cortex. J. Neurosci. 30, 7434–7446.

Ochiai, T., Grimali, S., Scardava, D., Roch, G., Horii, T., Riviere, D., Mangin, J. F., and Regis, J. (2004). Sulcal pattern and morphology of the superior temporal sulcus. Neuroimage 22, 706–719.

Padberg, J., Seltzer, B., and Cusick, C. G. (2003). Architecture and cortical connections of the upper bank of the superior temporal sulcus in the rhesus monkey: an analysis in the tangential plane. J. Comp. Neurol. 467, 418–434.

Schroeder, C. E., Lakatos, P., Kajikawa, Y., Partan, S., and Puce, A. (2006). Neuronal oscillations and visual amplification of speech. Trends Cogn. Sci. 12, 106–113.

Seltzer, B., and Pandya, D. N. (1991). Post-rolandoic cortical projections of the superior temporal sulcus in the rhesus monkey. J. Comp. Neurol. 312, 625–640.

Slutzky, D. A., and Recanzone, G. H. (2001). Temporal and spatial dependency of the ventralisquism effect. Neuronetwork 12, 7–10.

Spence, C., and Squire, S. (2003). Multisensory integration: maintaining the perception of synchrony. Curr. Biol. 13, R519–R521.

Stein, B. E., and Meredith, M. A. (1993). The Merging of the Senses. Cambridge, MA: MIT Press.

Stevenson, R. A., Altieri, N. A., Kim, S., Pisoni, D. B., and James, T. W. (2010). Neural processing of asynchronous audiovisual speech perception. Neuroimage 49, 33–38.

Szczylik, G. R., Tausche, P., and Münte, T. F. (2008). A novel approach
to study audiovisual integration in speech perception: localizer fMRI and sparse sampling. *Brain Res.* 1220, 142–149.
Tsao, D. Y., Freiwald, W. A., Tootell, R. B., and Livingstone, M. S. (2006). A cortical region consisting entirely of face-selective cells. *Science* 311, 670–674.
van Atteveldt, N., Formisano, E., Goebel, R., and Blomert, L. (2004). Integration of letters and speech sounds in the human brain. *Neuron* 43, 271–282.
van Atteveldt, N. M., Formisano, E., Blomert, L., and Goebel, R. (2007). The effect of temporal asynchrony on the multisensory integration of letters and speech sounds. *Cereb. Cortex* 17, 962–974.
Vatakis, A., and Spence, C. (2006a). Audiovisual synchrony perception for music, speech, and object actions. *Brain Res.* 1111, 134–142.
Vatakis, A., and Spence, C. (2006b). Audiovisual synchrony perception for speech and music assessed using a temporal order judgment task. *Neurosci. Lett.* 393, 40–44.
Vroomen, J., and Keetels, M. (2010). Perception of intersensory synchrony: a tutorial review. *Atten. Percept. Psychophys.* 72, 871–884.
Wright, T. M., Pelprey, K. A., Allison, T., McKeown, M. J., and McCarthy, G. (2003). Polysensory interactions along lateral temporal regions evoked by audiovisual speech. *Cereb. Cortex* 13, 1034–1043.
Yeterian, E. H., Pandya, D. N., Tomaiuolo, F., and Petrides, M. (2012). The cortical connectivity of the prefrontal cortex in the monkey brain. *Cortex* 48, 58–81.

**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 06 June 2012; paper pending published: 20 June 2012; accepted: 07 August 2012; published online: 28 August 2012.

Citation: Noesselt T, Bergmann D, Heinze H-J, Münte T and Spence C (2012) Coding of multisensory temporal patterns in human superior temporal sulcus. *Front. Integr. Neurosci.* 6:64. doi: 10.3389/fnint.2012.00064
Copyright © 2012 Noesselt, Bergmann, Heinze, Münte and Spence. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in other forums, provided the original authors and source are credited and subject to any copyright notices concerning any third-party graphics etc.